

Health Risk and Exposure Assessment for Ozone

Second External Review Draft

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TABLE OF CONTENTS

	TABLE OF CONTENTS	v
	TABLE OF FIGURES	xi
	TABLE OF TABLES	xxv
1	INTRODUCTION	1-1
1.1	HISTORY	1-3
1.2	CURRENT RISK AND EXPOSURE ASSESSMENT: GOALS AND PLANNED	
	APPROACH	1-5
1.3	ORGANIZATION OF DOCUMENT	1-6
2	OVERVIEW OF EXPOSURE AND RISK ASSESSMENT DESIGN	2-1
2.1	POLICY-RELEVANT EXPOSURE AND RISK QUESTIONS	2-2
2.2	AIR QUALITY CHARACTERIZATION	2-4
	2.2.1 O ₃ Chemistry and Response to Changes in Precursor Emissions	2-5
	2.2.2 Sources of O_3 and O_3 Precursors	2-6
	2.2.3 Simulation of Meeting Existing and Alternative Standards	2-7
	2.2.5 Exposures of Concern	2-8
	2.2.6 Health Endpoints	2-9
	2.2.7 Exposure and Concentration-response Functions for Health Endpoints	2-13
	2.2.8 At-risk Populations	2-14
2.3	URBAN-SCALE MODELING OF INDIVIDUAL EXPOSURE	2-15
	2.3.1 Microenvironmental O ₃ Concentrations	2-16
	2.3.2 Human Activity Patterns	2-17
	2.5.5 Modeling of Exposures Associated with Simulating Just Meeting O ₃ Standards	2-19
	2.3.4 Considerations in Selecting Urban Case Study Areas for the Exposure	
	Analysis	2-19
2.4	RISK ASSESSMENT	2-19
	2.4.1 Attributable Risk	2-20
	2.4.2 Modeling of Risk for Total Exposure to O ₃	2-21
	2.4.3 Distributions of Risk Across O_3 concentrations	2-22
2.5	MODELING OF RISKS ASSOCIATED WITH SIMULATING JUST MEETING (STANDARDS	O ₃ 2-22
2.6	CONSIDERATIONS IN SELECTING URBAN CASE STUDY AREAS FOR THE	Ξ
	RISK ANALYSIS	2-23
2.7	RISK CHARACTERIZATION	2-23
2.8	REFERENCES	2-25

3	SCOPE
3.1	OVERVIEW OF EXPOSURE AND RISK ASSESSMENTS FROM LAST REVIEW 3-2
	3.1.1 Overview of Exposure Assessment from Last Review
	3.1.2 Overview of Risk Assessment from Last Review
3.2	PLAN FOR THE CURRENT EXPOSURE AND RISK ASSESSMENTS
3.3	CHARACTERIZATION OF UNCERTAINTY AND VARIABILITY IN THE
	CONTEXT OF THE O ₃ EXPOSURE AND RISK ASSESSMENT
3.4	AIR QUALITY CHARACTERIZATION
3.5	EXPOSURE ASSESSMENT
3.6	URBAN-SCALE LUNG FUNCTION RISK ANALYSES BASED ON APPLICATION
_	OF RESULTS FROM CONTROLLED HUMAN EXPOSURE STUDIES
3.7	URBAN CASE STUDY AREA EPIDEMIOLOGY-BASED RISK ASSESSMENT3-17
3.8	NATIONAL-SCALE MORTALITY RISK ASSESSMENT
3.9	PRESENTATION OF EXPOSURE AND RISK ESTIMATES TO INFORM THE O ₃ NAAQS POLICY ASSESSMENT
3.10	REFERENCES
4	AIR QUALITY CONSIDERATIONS
4.1	INTRODUCTION
4.2	OVERVIEW OF O ₃ MONITORING AND AIR QUALITY DATA
4.3	OVERVIEW OF URBAN-SCALE AIR QUALITY INPUTS TO RISK AND EXPOSURE ASSESSMENTS 4-4
	4 3 1 Urban Case Study Areas 4-5
	4.3.2 Recent Air Quality
	4.3.3 Air Quality Adjustments for "Just Meeting" Existing and Potential Alternative O ₃ Standards
4.4	OVERVIEW OF NATIONAL-SCALE AIR QUALITY INPUTS
4.5	UNCERTAINTIES IN MODELING OF RESPONSES TO EMISSION REDUCTIONS
	TO JUST MEET EXISTING AND POTENTIAL ALTERNATIVE STANDARDS 4-38
4.6	REFERENCES
5	CHARACTERIZATION OF HUMAN EXPOSURE TO O ₃ 5-1
5.0	OVERVIEW
5.1	SYNOPSIS OF O3 EXPOSURE AND EXPOSURE MODELING
	5.1.1 Human Exposure
	5.1.2 Estimating O ₃ Exposure
	5.1.3 Modeling O ₃ Exposure Using APEX
5.2	SCOPE OF THE EXPOSURE ASSESSMENT
	5.2.1 Urban Areas Selected
	5.2.2 Time Periods Simulated
	5.2.3 Ambient Concentrations Used

	5.2.4 Meteorological Data Used	5-10
	5.2.5 Populations Simulated	5-11
	5.2.6 Key Physiological Processes And Personal Attributes Modeled	5-15
	5.2.7 Microenvironments Modeled	5-16
	5.2.8 Model Output	5-17
5.3	EXPOSURE ASSESSMENT RESULTS	5-23
	5.3.1 Overview	5-23
	5.3.2 Exposure Modeling Results for Base Air Quality	5-24
	5.3.3 Exposure Modeling Results for Simulations of Just Meeting Existing and Alt O ₃ Standards	ernative 5-25
5.4	TARGETED EVALUATION OF EXPOSURE MODEL INPUT AND OUTPUT 5-35	Γ DATA
	5.4.1 Analysis of Time-Locaton-Activity Data	5-35
	5.4.1.1 General Evaluation of CHAD Study Data: Historical and Recently Acqui 5-36	red Data
	5.4.1.2 Exposure-Relevant Personal Attributes Included in CHAD and APEX Sin Individuals	mulated
	5.4.1.3 Evaluation of Afternoon Time Spent Outdoors for CHAD and Survey Par 5-37	rticipants
	5.4.1.4 Evaluation of Afternoon Time Spent Outdoors for ATUS Survey	
	Participants	5-38
	5.4.1.5 Evaluation of Outdoor Time and Exertion Level for Asthmatics and Non-Asthmatics in CHAD	5-39
	5.4.2 Characterization of Factors Influencing High Exposures	5-40
	5.4.3 Exposure Results for additional at-risk populations and Lifestages, Exposure scenarios, and Air Quality Input Data Used	5-41
	5.4.3.1 Exposures Estimated for All School-age Children During Summer Month Neither Attending School or Performing Paid Work	ns, 5-41
	5.4.3.2 Exposures Estimated for Outdoor Workers During Summer Months	5-43
	5.4.3.3 Exposures Estimated for All School-age Children When Accounting for A Behavior	Averting 5-45
	5.4.3.4 Comparison of APEX Estimated Exposures Using Three Different Base Quality Data Sets: AQS, VNA, and EVNA	Case Air 5-46
	5.4.3.5 Comparison of APEX Estimated Exposures Using Two Different Adjuste Quality Data Sets: Quadratic Rollback and HDDM	ed Air 5-48
	5.4.4 Limited Performance Evaluations	5-49
	5.4.4.1 Personal Exposure Comparisons	
	5.4.4.2 Ventilation Rate Comparisons	
	5.4.4.2 Ventilation Rate Comparisons5.4.4.3 Evaluation of Longitudinal Profile Methodology	5-51 5-55

	5.5.1 Treatment of Variability	. 5-56
	5.5.2 Characterization of Uncertainty	. 5-57
5.6	KEY OBSERVATIONS	. 5-65
5.7	REFERENCES	. 5-71
6	CHARACTERIZATION OF HEALTH RISKS BASED ON CONTROLLED	
	HUMAN EXPOSURE STUDIES	6-1
6.1	INTRODUCTION	6-1
	6.1.1 Development of Approach for Current Risk Assessment	6-2
	6.1.2 Comparison of Controlled Human Exposure- and Epidemiologic-based Risk	
	Assessments	6-3
6.2	SCOPE OF LUNG FUNCTION HEALTH RISK ASSESSMENT	6-3
	6.2.1 Selection of Health Endpoints	6-4
	Studies	6-5
	6.2.3 Controlled Human Exposure Studies	6-6
	6.2.4 The McDonnell-Stewart-Smith (MSS) Model	6-8
	6.2.5 The Exposure-Response Function Approach Used in Prior Reviews	.6-16
6.3	O ₃ RISK ESTIMATES	. 6-21
	6.3.1 Lung Function Risk Estimates Based on the McDonnell-Stewart-Smith Model	6-22
	6.3.2 Lung Function Risk Estimates Based on the Exposure-Response Functions	()
	Approach Used in Prior Reviews	. 6-28
	Approach	. 6-29
6.4	EVALUATION OF THE MSS MODEL	.6-35
	6.4.1 Summary of Published Evaluations	.6-35
	6.4.2 Children	.6-35
	6.4.3 Threshold vs. Non-Threshold Models	. 6-36
6.5	CHARACTERIZATION OF UNCERTAINTY	. 6-37
	6.5.1 Statistical Model Form	. 6-38
	6.5.2 Convergence of APEX Results	.6-41
	6.5.3 Application of Model for All Lifestages	. 6-42
	6.5.4 Application of Model for Asthmatic Children	.6-43
	6.5.6 Qualitative Assessment of Uncertainty	.6-43
6.6	DISCUSSION	6-46
67	REFERENCES	6-50
		.0.50
/	CHARACTERIZATION OF HEALTH RISK BASED ON EPIDEMIOLOGICAT STUDIES	_ 7-1
71	GENERAL APPROACH	7_1
/.1	7.1.1 Basic Structure of the Risk Assessment	,-1 7_1
	7.1.2 Calculating O ₃ -Related Health Effects Incidence	
7.2	AIR QUALITY CONSIDERATIONS	.7-13
		-

7.3	SELECTION OF MODEL INPUTS AND ASSUMPTIONS	.7-15
	7.3.1 Selection of Urban Study Areas	.7-15
	7.3.2 Selection of Epidemiological Studies and Specification of	
	Concentration-Response Functions	.7-17
	7.3.3 Baseline Health Effect Incidence and Prevalence Data	.7-30
- 4	7.3.4 Population (demographic) Data	. /-31
7.4	ADDRESSING VARIABILITY AND UNCERTAINTY	.7-31
	7.4.1 Treatment of Key Sources of Variability	.7-34
	7.4.2 Qualitative Assessment of Uncertainty	. /-38
75	LIDDAN STUDY ADEA DECLUTS	. 7-43 7 47
1.5	URBAN STUDY AREA RESULTS	. /-4/
	 7.5.1 Assessment of Health Risk After Just Meeting the Existing 75 ppb standard 7.5.2 Assessment of Health Risk Associated with Simulating Meeting Potential 	.7-65
	Alternative Standards of 70, 65, and 60 ppb	. /-6/
	Fstimates	7_72
7.6	KEY OBSERVATIONS REGARDING OVERALL CONFIDENCE IN THE RISK	7 83
77		- 1-05 7 06
1.1	REFERENCES	. /-80
8	NATIONAL SCALE MORTALITY RISK BURDEN BASED ON APPLICATION	N OF
	RESULTS FROM EPIDEMIOLOGICAL STUDIES	8-1
8.1	NATIONAL-SCALE ASSESSMENT OF MORTALITY RELATED TO O ₃ EXPOSURE	8-1
	8.1.1 Methods	8-2
	8.1.2 Results	8-6
	8.1.3 Sensitivity Analysis	.8-15
_	8.1.4 Discussion	. 8-19
8.2	EVALUATING THE REPRESENTATIVENESS OF THE URBAN STUDY AREA THE NATIONAL CONTEXT	IS IN .8-21
	8.2.1 Analysis Based on Consideration of National Distributions of Risk-Related Attributes	. 8-22
	8.2.2 Analysis Based on Consideration of National Distribution of O ₃ -Related	
	Mortality Risk	.8-45
	8.2.3 Analysis Based on Consideration of National Responsiveness of O ₃ Concentra	tions
	to Emissions Changes	.8-49
0.2	8.2.3 Discussion	0-70
8.3	REFERENCES	. 8-78
9 SY	YNTHESIS	9-1
9.1	INTRODUCTION	9-2
92		
1.2	SUMMARY OF KEY RESULTS	9-2
1.2	SUMMARY OF KEY RESULTS	9-2 9-2

	9.2.3 Health Risks Based on Controlled Human Exposure Studies (Chapter 6)9.2.4 Health Risks Based on Epidemiological Studies (Chapters 7 and 8)	9-13 9-18
9.3	COMPARISON OF RESULTS ACROSS EXPOSURE, LUNG FUNCTION RISK, AND EPIDMEIOLOGY-BASED MOTALITY AND MORBIDITY RISK	
	ANALYSES	9-27
	9.3.1 Evaluation of Exposures and Risks After Just Meeting the Existing Standard9.3.2 Reductions in Exposure and Risk Metrics after Just Meeting Alternative Standards	9-27
0.4		9-31
9.4	RISK RESULTS	9-36
	9.4.1 Representativeness of Selected Urban Case Study Areas in Reflecting Areas Across the Nation with Elevated Risk	9-36
	9.4.2 Representativeness of Selected Urban Case Study Areas in Reflecting Responsiveness of Risk to Just Meeting Existing and Alternative O ₃	0.07
		9-31
9.5	OVERALL ASSESSMENT OF CONFIDENCE IN EXPOSURE AND RISK	
	RESULTS	9-38
	9.5.1 Uncertainties in Modeling O ₃ Responses to Meeting Standards	9-39
	9.5.2 Uncertainties in Modeling Exposure and Lung-function Risk	9-40
	9.5.3 Uncertainties in Modeling Epidemiological-based Risk	9-40
9.6	OVERALL INTEGRATED CHARACTERIZATION OF RISK IN THE CONTEXT KEY POLICY RELEVANT QUESTIONS	OF 9-42
9.7	REFERENCES	9-47

TABLE OF FIGURES

Figure 2-1. Overview of Exposure and Risk Assessment Design.	2-2
Figure 2-2. Causal Determinations for O ₃ Health Effects.	2-11
Figure 3-1. Conceptual Diagram for Air Quality Characterization in the Health REA	3-10
Figure 3-2. Conceptual Diagram for Population Exposure Assessment	3-13
Figure 3-3. Conceptual Diagram of O ₃ Lung Function Health Risk Assessment Based on	
Controlled Human Exposure Studies	3-15
Figure 3-4. Conceptual Diagram of Urban Case Study Area Health Risk Assessment Based	l on
Results of Epidemiology Studies.	3-18
Figure 3-5. Conceptual Diagram of National O3 Mortality Risk Assessment Based on Resu	lts of
Epidemiology Studies	3-23
Figure 4-1. Map of Monitored 8-hour O ₃ Design Values for the 2006-2008 Period	4-3
Figure 4-2. Map of Monitored 8-hour O ₃ Design Values for the 2008-2010 Period	4-4
Figure 4-3. Flowchart of Air Quality Data Processing for Different Parts of the Urban-scal	e Risk
and Exposure Assessments	4-5
Figure 4-4. Trends in Annual 4th Highest 8-hour Daily Maximum O3 Concentrations in pr	b for
the 15 Urban Case Study Areas for 2006-2010. Urban areas are grouped int	to 3
regions: Eastern (top), Central (middle), and Western (bottom)	4-7
Figure 4-5a. Maps of the 5 Eastern U.S. Urban Case Study Areas Including O ₃ Monitor	
Locations	4-11
Figure 4-5b. Maps of the 5 Central U.S. Urban Case Study Areas Including O ₃ Monitor	
Locations	4-12
Figure 4-5c. Maps of the 5 Western U.S. Urban Case Study Areas Including O ₃ Monitor	
Locations	4-13
Figure 4-6. Flowchart of HDDM adjustment methodology to inform risk and exposure	
assessment	4-17
Figure 4-7. Distributions of composite monitor 8-hour daily maximum O ₃ concentrations fr	rom
ambient measurements (black), quadratic rollback (blue), and the HDDM	
adjustment methodology (red) for meeting the existing standard. Values are	based
on the Zanobetti & Schwartz study areas for April-October of 2006-2008	4-21

- Figure 4-8. Distributions of composite monitor 8-hour daily maximum O₃ concentrations from ambient measurements (black), quadratic rollback (blue), and the HDDM adjustment methodology (red) for meeting the existing standard. Values are based on the Zanobetti & Schwartz study areas for June-August of 2006-2008........4-22
- Figure 4-9. Distributions of composite monitor 8-hour daily maximum values for the 12 urban case study areas in the epidemiology-based risk assessment. Plots depict values based on ambient measurements (base), and values obtained with the HDDM adjustment methodology showing attainment of 75, 70, 65 and 60 ppb standards. Values shown are based on CBSAs for April-October of 2007. Note that the HDDM adjustment technique was not able to adjust air quality to show attainment of a 60 ppb standard in New York, so no boxplot is shown for that case...........4-25

Figure 4-13. Maps of 4th highest (top) and May-September average (bottom) daily maximum 8-
hour O_3 concentrations in Houston for 2006-2008 ambient measurements (left),
HDDM adjustment to meet the existing standard (center), and HDDM adjustment
to meet the alternative standard of 65 ppb (right). Squares represent measured
values at monitor locations; circles represent VNA estimates at census tract
centroids4-30
Figure 4-14. May-September average 8-hour daily maximum O ₃ concentrations in ppb, based on
a Downscaler fusion of 2006-2008 average monitored values with a 12km 2007
CMAQ model simulation
Figure 4-15. June-August average 8-hour daily 10am-6pm mean O ₃ concentrations in ppb, based
on a Downscaler fusion of 2006-2008 average monitored values with a 12km
2007 CMAQ model simulation
Figure 4-16. April-September average 1-hour daily maximum O3 concentrations in ppb, based on
a Downscaler fusion of 2006-2008 average monitored values with a 12km 2007
CMAQ model simulation4-34
Figure 4-17. Frequency and Cumulative Distributions of the Three Fused Seasonal Average O_3
Surfaces Based on all CMAQ 12 km Grid Cells
Figure 4-18. 2006-2008 O ₃ Design Values Versus 2006-2008 Fused Seasonal Average O ₃ Levels
for the CMAQ 12km Grid Cells Containing O3 Monitors
Figure 5-1. Conceptual Framework Used for Estimating Study Area Population O ₃ Exposure
Concentrations
Figure 5-2. Percent of asthmatic school-age children in all study areas with at least one O_3
exposure at or above 60 ppb-8hr while at moderate or greater exertion using base
air quality (2006-2010), stratified by year (top left panel) or by study area (bottom
left panel)
Figure 5-3. Percent of asthmatic school-age children in Atlanta with at least one O_3 exposure at
or above 60 ppb-8hr (left top panel), 70 ppb-8hr (middle top panel), and 80 ppb-
8hr (right top panel while at moderate or greater exertion, years 2006-2010 air
quality adjusted to just meet the existing and alternative O ₃ standard levels. The
multi-panel display (bottom) illustrates the same exposure results expanded to

reflect individual data points by year, standard averaging period, and benchmark Figure 5-4. Percent of asthmatic school-age children in Atlanta with multiple O₃ exposures at or above 60 ppb-8hr while at moderate or greater exertion, years 2006-2010 air quality adjusted to just meet the existing and alternative O₃ standard levels....5-23 Figure 5-5. Percent of all school-age children with at least one daily maximum 8-hr average O_3 exposure at or above 60, 70, and 80 ppb while at moderate or greater exertion, years 2006-2010, air quality adjusted to just meet the existing and potential alternative standards......5-30 Figure 5-6. Percent of asthmatic school-age children with at least one daily maximum 8-hr average O₃ exposure at or above 60, 70, and 80 ppb while at moderate or greater exertion, years 2006-2010, air quality adjusted to just meet the existing and Figure 5-7. Percent of all asthmatic adults with at least one daily maximum 8-hr average O_3 exposure at or above 60, 70, and 80 ppb-8hr while at moderate or greater exertion, years 2006-2010, air quality adjusted to just meet the existing and potential Figure 5-8. Percent of all older adults with at least one daily maximum 8-hr average O₃ exposure at or above 60, 70, and 80 ppb-8hr while at moderate or greater exertion, years 2006-2010, air quality adjusted to just meet the existing and potential alternative Figure 5-9. Percent of all school-age children with multiple daily maximum 8-hr average O₃ exposures at or above 60 ppb while at moderate or greater exertion, years 2006-2010, air quality adjusted to just meet the existing and potential alternative Figure 5-10. Comparison of the percent of all school-age children having daily maximum 8-hr average O₃ concentration at or above 60 ppb during June, July, and August in Detroit 2007: using any available CHAD diary ("All CHAD Diaries") or using CHAD diaries having no time spent in school or performing paid work ("No

- Figure 5-12. Percent of all school-age children (left panel) and asthmatic school-age children (right panel) having daily maximum 8-hr average O₃ concentration at or above benchmark levels during a 2-day simulation in Detroit, base air quality, August 1-2, 2007. Red bars indicate exposure results when considering effect of averting....
 5-46

- Figure 5-15. Distribution of daily average O₃ exposures (top panels) and daily afternoon outdoor time (bottom panels) and for DEARS study participants (left panels) and APEX simulated individuals (right panels) in Wayne County, MI, July-August 2006 5-50

- Figure 5-20. Incremental increases in percent of all school-age children exposed to O₃ at or above 60 ppb-8hr for each study area, year 2006-2010 air quality. Average

	percent (left panels), maximum percent (right panels), at least one exposure (top
	panels), at least two exposures (bottom panels) per year
Figure 6-1.	Two-Compartment Model
Figure 6-2.	Distribution of Responses (Lung Function Decrements in FEV1) Predicted by the
	MSS Model for 20-Year Old Individuals. Exposure to 100 ppb O_3 at Moderate
	Exercise (40 L/min, BSA=2 m2) Under the Conditions of a Typical 6.6-hour
	Clinical Study
Figure 6-3.	Median Response (Lung Function Decrements in FEV1) Predicted by the MSS
	Model for 20-Year Old Individuals. Exposure to 100 ppb O ₃ at Moderate Exercise
	(40 L/min, BSA=2 m2) Under the Conditions of a Typical 6.6-hour Clinical
	Study
Figure 6-4.	Median Response (FEV1 Decrements) Predicted by the MSS Threshold and Non-
	Threshold Models for 20-Year Old Individuals, Constant 100 ppb O ₃ Exposure, 2
	Hours Heavy Exercise (30 L/min-m2 BSA)6-15
Figure 6-5.	Probability of Response $\geq 10\%$ Predicted by the MSS Threshold and Non-Threshold
	Models for 20-Year Old Individuals, Constant 100 ppb O3 Exposure, 2 Hours
	Heavy Exercise (30 L/min-m2 BSA)6-15
Figure 6-6.	Probabilistic Exposure-Response Relationships for FEV1 Decrements $\geq 10\%$ for 8-
	Hour Exposures At Moderate Exertion, Ages 18-35. Values associated with data
	points are the number of subject-exposures at each exposure concentration 6-20
Figure 6-7.	Risk results for all school-aged children with ≥ 1 occurrences of FEV1 decrements \geq
	10, 15, 20% for all cities, year, and scenarios (y-axis is percent of children
	affected)
Figure 6-8.	Risk results for all school-aged children with ≥ 1 occurrences of FEV1 decrements \geq
	10% under the 0.07 ppm alternative standard showing variability across cities
	(horizontally) and years (vertically)
Figure 6-9.	Distribution of Daily FEV1 Decrements $\geq 10\%$ Across Ranges of 8-hour Average
	Ambient O ₃ Concentrations (Los Angeles, 2006 recent air quality)
Figure 6-10	. Comparison of E-R and MSS Model (restricted to 8-hour average EVR \geq 13)
	Response Functions (Atlanta 2006 base case, ages 18-35)

Figure 6-11. Distribution of Daily Maximum 8-hour Average EVR For Values of $EVR \ge 13$
(L/min-m2) (midpoints on vertical axis) (Atlanta 2006 base case, ages 18-35)
Figure 6-12. Sensitivity (Percent Change) of Population With One or More FEV1 Decrements \geq
10% to a 5% Increase in Individual MSS Model Parameter Estimates
Figure 6-13. Lung Function Risk Results, Incremental Increases In Risk For Increasing Standard
Levels: Percent of All School-aged Children With FEV1 Decrement $\geq 10\%$,
Highest Value For Each Study area Over Years
Figure 6-14. Lung Function Risk Results, Incremental Increases In Risk For Increasing Standard
Levels: Percent of All School-aged Children With FEV1 Decrement $\geq 10\%$, Mean
Value For Each Study Area Over Years
Figure 7-1. Flow Diagram of Risk Assessment for Short-term Exposure Studies
Figure 7-2. Heat Maps for Short Term O ₃ -attributable Mortality (Just meeting existing standard
and risk reductions from just meeting alternative standards) (2007) (Smith et al.,
2009 C-R functions)7-55
Figure 7-3. Heat Maps for Short Term O ₃ -attributable Mortality (Just meeting existing standard
and risk reductions from just meeting alternative standards) (2009) (Smith et al.,
2009 C-R functions) (see Key at bottom of figure)
Figure 7-4. Plots of Short-Term O ₃ -attributable All-Cause Mortality for Meeting Existing
standard and Alternative Standards (Smith et al., 2009) (Simulation year 2007 and
2009)
Figure 7-5. Plots of Short-Term O ₃ -attributable Respiratory HA for Meeting Existing standard
and Alternative Standards (Medina-Ramon, et al., 2006) (Simulation year 2007
and 2009)
Figure 7-6. Plots of Long-Term O ₃ -attributable Respiratory Mortality for Meeting Existing
standard and Alternative Standards (Jerrett et al., 2009) (Simulation year 2007
and 2009)
Figure 7-7. Sensitivity Analysis: Short-Term O3-attributable Mortality (air quality-related factors
including study area size and method used to simulate attainment of existing and
alternative standard levels) (2009) SA1-smaller (Smith-based) study area, SA2-
alternative method for simulating standards

Figure 7-8. Sensitivity Analysis: Short-Term O₃-attributable Mortality (C-R function specification) (2009) SA1-regional Bayes-based adjustment; SA2-copollutant model (PM₁₀); SA3-Zanobetti and Schwartz-based effect estimates7-80 Figure 8-2. Estimated annual non-accidental premature deaths (individuals) in 2007 associated with average 2006-2008 May-September average 8-hr daily maximum O₃ levels Figure 8-3. Estimated annual all-cause premature deaths (individuals) in 2007 associated with average 2006-2008 June-August average 8-hr daily mean (10am-6pm) O_3 levels Figure 8-4. Estimated annual adult (age 30+) respiratory premature deaths (individuals) in 2007 associated with average 2006-2008 April-September average 1-hr daily max O₃ Figure 8-5. Estimated percentage of May-September total non-accidental mortality (all ages) attributable to 2006-2008 average O₃ levels by county using Smith et al. (2009) Figure 8-6. Estimated percentage of June-August total all-cause mortality (all ages) attributable to 2006-2008 average O_3 levels by county using Zanobetti and Schwartz (2008) Figure 8-7. Estimated percentage of April-September respiratory mortality among adults age 30+ attributable to 2006-2008 average O_3 levels by county using Jerrett et al. (2009) Figure 8-8. Cumulative distribution of county-level percentage of all-cause, all-year, and all-age Figure 8-9. Cumulative percentage of total O_3 deaths by baseline O_3 concentration. O_3 concentrations are reported as May-September average 8-hr daily maximum for results based on Smith et al. (2009) effect estimates, June-August average 8-hr mean (10am to 6pm) for results based on Zanobetti and Schwartz (2008) effect estimates, and April-September average 1-hr daily maximum for results based on

Figure 8-10.Regions used in the sensitivity analysis based on the Smith et al. (2009) regional-
prior Bayes-shrunken city-specific and regional average effect estimates (Source:
Samet et al. 2000)
Figure 8-11. O ₃ -attributable premature deaths by region as calculated by applying Smith et al.
(2009) regional prior Bayes-shrunken and regional average effect estimates, as
compared with the national prior Bayes-shrunken and national average effect
estimates as in the main results
Figure 8-12. Comparison of county-level populations of urban case study area counties to the
frequency distribution of population in 3,143 U.S. counties
Figure 8-13. Comparison of county-level seasonal mean 8-hr daily maximum O ₃ concentrations
in urban case study area counties to the frequency distribution of seasonal mean
8-hr daily maximum O_3 concentrations in 671 U.S. counties with O_3 monitors
Figure 8-14. Comparison of 2007 county-level 4^{th} high 8-hr daily maximum O_3 concentrations in
urban case study area counties to the frequency distribution of 2007 4 th high 8-hr
daily maximum O ₃ concentrations in 725 U.S. counties with O ₃ monitors 8-34
Figure 8-15. Comparison of county-level all-cause mortality in urban case study area counties to
the frequency distribution of all-cause mortality in 3,137 U.S. counties8-35
Figure 8-16. Comparison of county-level non-accidental mortality in urban case study area
counties to the frequency distribution of non-accidental mortality in 3,135 U.S.
counties
Figure 8-17. Comparison of city-level all-cause mortality risk coefficients from Zanobetti and
Schwartz (2008) in urban case study areas to the frequency distribution of all-
cause mortality risk coefficients from Zanobetti and Schwartz (2008) in 48 U.S.
cities
Figure 8-18. Comparison of city-level national prior Bayes-shrunken non-accidental mortality
risk coefficients from Smith et al. (2009) in urban case study areas to the
frequency distribution of national prior Bayes-shrunken non-accidental mortality
risk coefficients from Smith et al. (2009) in 98 U.S. cities

- Figure 8-24. Comparison of city-level air conditioning prevalence in urban case study areas to the frequency distribution of air conditioning prevalence in 76 U.S. cities 8-44

- Figure 8-34. Map of O₃ trends at specific monitors in the New York area. All upward and downward facing triangles represent statistically significant trends from 1998-2011 (p < 0.05), circles represent locations with no significant trends. Sites used in Smith et al (2009) and the Zanobetti and Schwartz (2008) epidemiology studies are represented by colored dots. Only monitors with at least seven years of data are displayed. The pink star indicates the site with the higher design value in 2011. The MSA border as defined by the U.S. census bureau is delineated by the light blue line. Left panel shows trends in annual 4th highest 8-hr daily maximum O₃ values, center panel shows trends in annual median 8-hr daily maximum O₃ values.
- Figure 8-35. Map of O_3 trends at specific monitors in the Chicago area. All upward and downward facing triangles represent statistically significant trends from 1998-2011 (p < 0.05), circles represent locations with no significant trends. Sites used in Smith et al (2009) and the Zanobetti and Schwartz (2008) epidemiology studies are represented by colored dots. Only monitors with at least seven years of data are displayed. The pink star indicates the site with the higher design value in

- Figure 8-40. Histograms of U.S. population living in locations with increasing and decreasing mean O₃. Values on the x-axis represent change in mean O₃ (ppb) from the 2007 base CMAQ simulation to the 50% NOx cut CMAQ simulation. The percentages of the U.S. population living in areas that have changes less than -1 ppb, from -1 to +1 ppb, and greater than 1 ppb are shown on the y-axis. Left plots show population numbers in locations not included in one of the cases study areas while right plots show population numbers in locations included in one of the case study areas. Top plots show changes in January monthly mean O₃, middle plots show changes in seasonal mean June-August O₃, and bottom plots show changes in seasonal mean April-October O₃.
- Figure 8-41. Histograms of U.S. population living in locations with increasing and decreasing mean O₃. Values on the x-axis represent change in mean O₃ (ppb) from the 2007 base CMAQ simulation to the 90% NOx cut CMAQ simulation. The percentages of the U.S. population living in areas that have changes less than -1 ppb, from -1

- Figure 8-44. Histograms of U.S. population living in locations with increasing and decreasing mean O₃. Values on the x-axis represent the change in seasonal mean (April-October) O₃ from the 2007 base CMAQ simulation to the 50% NOx cut CMAQ simulation. The percentages of the U.S. population living in areas that have changes less than -1 ppb, from -1 to +1 ppb, and greater than 1 ppb are shown on the y-axis. Left plots show population numbers in locations not included in one of the cases study areas while right plots show population numbers in locations included in one of the urban case study areas. Bottom plots show histograms for low-mid population density areas while top plots show histograms for high population density areas.
- Figure 8-45. Histograms of U.S. population living in locations with increasing and decreasing mean O₃. Values on the x-axis represent the change in seasonal mean (April-October) O₃ from the 2007 base CMAQ simulation to the 90% NOx cut CMAQ simulation. The percentages of the U.S. population living in areas that have changes less than -1 ppb, from -1 to +1 ppb, and greater than 1 ppb are shown on the y-axis. Left plots show population numbers in locations not included in one of the cases study areas while right plots show population numbers in locations included in one of the urban case study areas. Bottom plots show histograms for

low-mid population density areas while top plots show histograms for high
population density areas
Figure 9-1. Distributions of composite monitor 8-hour daily maximum O ₃ concentrations from
ambient measurements (black), quadratic rollback (blue), and the HDDM
adjustment methodology (red) for meeting the existing standard9-
Figure 9-2. Effects of just meeting existing (columns 1 and 2) and alternative (columns 3 through
8) standards on percent of children (ages 5-18) with at least one O_3 exposure at or
above 60, 70, and 80 ppb-8hr while at moderate or greater exertion, years 2006-
2010
Figure 9-3. Effects of just meeting existing (75 ppb) and alternative standards on percent of
children (ages 5-18) exceeding 60 ppb exposure benchmark, highest value across
years for each urban case study area, 2006-2010
Figure 9-4. Effects of just meeting existing (column 1) and alternative (columns 2-4) standards
on percent of children (ages 5-18) with FEV1 decrement > 10 , 15, and 20%, years
2006-2010
Figure 9-5. Impact of just meeting existing (75 ppb) and alternative standards on percent of
children (ages 5-18) with FEV ₁ decrement $> 10\%$, highest value for each urban
case study area, 2006-20109-18
Figure 9-6. Impacts of just meeting existing (75 ppb) and alternative standard levels on mortality
risk per 100,000 population for 2007 and 2009
Figure 9-7. Impacts of just meeting existing and alternative standard levels on adult (ages 65 and
older) respiratory hospital admissions risk per 100,000 population for 2007 and
2009
Figure 9-8. Comparison of Exposure (Row 1) Lung Function Risk (Row 2) and Epidemiology-
Paged Dick (Down 2 and 4) Matrice after Just Masting the Existing 75 nph
Based Risk (Rows 5 and 4) Metrics after Just Meeting the Existing 75 ppb
Standard
Standard

TABLE OF TABLES

Table 3-1. Short-term O ₃ Exposure Health Endpoints Evaluated in Urban Case Study Areas
Table 4-1. Monitor and Area Information for the 15 Urban Case Study Areas in the Exposure
Modeling and Clinical Study Based Risk Assessment4-6
Table 4-2. Monitor and Area Information for the 12 Urban Case Study Areas in the
Epidemiology Based Risk Assessment4-9
Table 4-3. Summary Statistics Based on the Three Fused Seasonal Average O ₃ Surfaces Based
on all CMAQ 12 km Grid Cells4-36
Table 4-4. Correlation Coefficients Between the Three Fused Seasonal Average O ₃ Surfaces
Based on all CMAQ 12 km Grid Cells
Table 4-5. Correlation Coefficients and Ratios of the 2006-2008 O ₃ Design Values to the 2006-
2008 Fused Seasonal Average O3 Levels for the CMAQ 12km Grid Cells
Containing O ₃ Monitors
Table 4-6. Summary of Qualitative Uncertainty Analysis of Key Air Quality Elements in the O_3
NAAQS Risk Assessment4-42
Table 5-1. General Characteristics of the Population Exposure Modeling Domain Comprising
Each Study Area5-9
Table 5-2. Asthma Prevalence for Children and Adults Estimated by APEX in Each Simulated
Study Area5-12
Table 5-3. Consolidated Human Activity Database (CHAD) Study Information and Diary-days
Used by APEX
Table 5-4. Ventilation equation coefficient estimates (b_i) and residuals distributions (e_i)
Table 5-5. Microenvironments Modeled, Calculation Method Used, and Variables Included
Table 5-6. Characterization of Key Uncertainties in Historical and Current APEX Exposure
Assessments
Table 5-7. Mean and Maximum Percent of all School-age Children Estimated to Experience at
Least One Daily Maximum 8-hr Average Exposure to O ₃ at or Above Selected
Health Benchmark Levels5-68

Table 5-8.	Mean and Maximum Percent of All School-age Children Estimated to Experience at
	Least Two Daily Maximum 8-hr Average Exposures to O3 At or Above Selected
	Health Benchmark Levels5-70
Table 6-1.	Estimated Parameters in the MSS Models6-11
Table 6-2.	Age Term Parameters for Application of the 2012 MSS Threshold Model to All Ages
Table 6-3.	Study-specific Ozone Exposure-response Data for Lung Function Decrements Based
	on Correcting Individual Responses for the Effect on Lung Function of Exercise
	in Clean Air, Ages 18-356-16
Table 6-4.	Ranges of percents of population experiencing one or more days during the ozone
	season with lung function decrement (ΔFEV_1) more than 10%. The numbers in
	this table are the minimum and maximum percents estimated over all cities and
	years
Table 6-5.	Ranges of percents of population experiencing one or more days during the ozone
	season with lung function decrement (ΔFEV_1) more than 15%. The numbers in
	this table are the minimum and maximum percents estimated over all cities and
	years
Table 6-6.	Percents of the General Population and Outdoor Workers (ages 19-35) Experiencing
	1 or More and 6 or More FEV_1 Decrements $\geq 15\%$ (based on Atlanta 2006 APEX
	simulations)6-28
Table 6-7.	Ranges of percents of school-aged children experiencing one or more days during the
	ozone season with lung function decrement (ΔFEV_1) more than 10 and 15%. The
	numbers in this table are the minimum and maximum percents estimated over all
	cities and years
Table 6-8.	Comparison of responses from the MSS model with responses from the population
	exposure-response (E-R) method. 2006 existing standard, ages 5 to 186-30
Table 6-9.	Comparison of MSS Model and E-R Model of Previous Reviews for Atlanta, Mar 1-
	Oct 30, 2006, ages 18-35
Table 6-10	. Comparison of MSS Model and E-R Model of Previous Reviews for Los Angeles,
	Jan 1-Dec 31, 2006, ages 18-35

Table 6-11. Comparison of Responses from the MSS 2010 Model with Responses from
McDonnell et al. (1985)6-36
Table 6-12. Percents of the population by age group with one or more days during the ozone
season with lung function (FEV $_1$) decrements more than 10, 15, and 20% (Atlanta
2006 base case). MSS Threshold model, monitors air quality
Table 6-13. Percents of the population by age group with one or more days during the ozone
season with lung function (FEV $_1$) decrements more than 10, 15, and 20% (Atlanta
2006 base case). MSS No-Threshold model, monitors air quality
Table 6-14. MSS threshold model estimated parameters with confidence intervals
Table 6-15. Convergence results for the Atlanta 2006 base case with 200,000 simulated
individuals. Percents of the population by age group with one or more days (and
six or more days) during the ozone season with lung function (FEV $_1$) decrements
more than 10, 15, and 20%. Minimum and maximum values and ranges over 40
APEX runs6-42
Table 6-16. Summary of Qualitative Uncertainties of Key Modeling Elements in the O ₃ Lung
Function Risk Assessment6-44
Table 7-1. Information on the 12 Urban Case Study Areas in the Risk Assessment
Table 7-2. Overview of Epidemiological Studies Used in Specifying C-R Functions
Table 7-3. CBSA-based Study Areas with Multiple Effect Estimates from the Smith et al., 2009
Study
Table 7-4. Summary of Qualitative Uncertainty Analysis of Key Modeling Elements in the O ₃
NAAQS Risk Assessment7-41
Table 7-5. Specification of the Core and Sensitivity Analyses (air quality simulation)
Table 7-6. Specification of the Core and Sensitivity Analyses (alternative C-R function
specification)7-47
Table 7-7. Short-Term O ₃ -attributable All Cause Mortality Incidence (2007 and 2009) (Smith et
al., 2009 C-R Functions)7-53
Table 7-8. Percent of Total All-Cause Mortality Attributable to O_3 and Percent Change in O_{3-}
Attributable Risk (2007 and 2009) (Smith et al., 2009 C-R functions)7-54

Table 7-9. Short-Term O ₃ -attributable Morbidity Incidence, Percent of Baseline and Reduction
in O3-attributable Risk – Respiratory-Related Hospital Admissions (2007 and
2009)
Table 7-10. Short-Term O ₃ -attributable Morbidity Incidence, Percent of Baseline and Reduction
in Ozone-attributable Risk – Emergency Room Visits (2007 and 2009)7-59
Table 7-11. Short-Term O ₃ -attributable Morbidity Incidence, Percent of Baseline and Reduction
in Ozone-attributable Risk – Asthma Exacerbations (2007 and 2009)7-60
Table 7-12. Long-Term O ₃ -attributable Respiratory Mortality Incidence (2007 and 2009) (Jerrett
et al., 2009 C-R Functions)
Table 7-13. Long-Term O ₃ -attributable Respiratory Mortality Percent of Baseline Incidence and
Percent Reduction in O ₃ -attributable Risk (simulation years 2007 and 2009)
(Jerrett et al., 2009 C-R Functions)7-63
Table 7-14. Sensitivity Analysis for Long-Term O ₃ -attributable Respiratory Mortality –
Alternative C-R Function Specification (regional effect estimates) % of baseline
all-cause mortality and change in O ₃ -attribuable risk (2009) (Smith et al., 2009,
O ₃ season)
Table 7-15. Sensitivity Analysis for Long-Term O3-attributable Respiratory Mortality –
Alternative C-R Function Specification (national O ₃ -only effect estimates) % of
baseline all-cause mortality and change in O ₃ -attribuable risk (2009) (Smith et al.,
2009, O ₃ season)
Table 8-1. Estimated annual O ₃ -related premature mortality in 2007 associated with 2006-2008
average O ₃ concentrations (95th percentile confidence interval)
Table 8-2. Mean, median, 2.5 percentile, and 97.5 percentile of the estimated percentage of
mortality attributable to ambient O ₃ for all 3087 counties in the continental U.S
Table 8-3. Sensitivity of estimated O ₃ -attributable premature deaths to the application of the 5th
lowest and 5th highest city-specific risk estimates found by Smith et al. (2009)
and Zanobetti and Schwartz (2008) to the gridcells between the cities included in
those studies
Table 8-4. Sensitivity of estimated O ₃ -attributable premature deaths to the application of Smith et
al. (2009) regional prior Bayes-shrunken city-specific and regional average effect

estimates, as compared with the national prior Bayes-shrunken city-specific an	ıd
national average effect estimates as in the main results	8-18
Table 8-5. Data Sources for O ₃ risk-related Attributes	8-25
Table 8-6. Summary Statistics for Selected O ₃ Risk-related Attributes	8-29
Table 8-7. Broad Regional Annual Trends of Concurrent O3 Concentrations and Emissions of	f
NOx and VOCs over the 2000-2011 Time Period	8-61
Table 9-1. Area and Monitoring Information for the 15 Case Study Areas	.9-3
Table 9-2. General Patterns in Seasonal (May-Sept) Mean of Daily Maximum 8-hour O_3	
Concentrations after Adjusting to Meet Existing and Alternative Standards	.9-5

LIST OF ACRONYMS/ABBREVIATIONS

AER	air exchange rate
AHRQ	Agency for Healthcare Research and Quality
APEX	Air Pollution Exposure Model
AQI	Air Quality Index
AQS	Air Quality System
ATUS	American Time Use Survey
BenMAP	Benefits Mapping and Analysis Program
BRFSS	Behavioral Risk Factor Surveillance System
BSA	body surface area
CAA	Clean Air Act
CASAC	Clean Air Science Advisory Committee
CDC	Center for Disease Control and Prevention
CDF	cumulative distribution functions
CH ₄	methane
CHAD	Consolidated Human Activity Database
CI	confidence interval
CMAQ	Community Multi-scale Air Quality
CO ₂	carbon dioxide
C-R	Concentration Response (function)
ED	emergency department
EGU	electric generating unit
EPA	U.S. Environmental Protection Agency
ER	emergency room
eVNA	enhanced Voronoi Neighbor Averaging
EVR	equivalent ventilation rate
FEM	Federal Equivalent Method
FEV1	one-second forced expiratory volume
FRM	Federal Reference Method

FVC	forced vital capacity
HA	hospital admissions
HDDM	Higher-order Decoupled Direct Method
HNO ₃	nitric acid
HO ₂	hydro-peroxy radical
HUCP	Healthcare Cost and Utilization Program
IPCC	Intergovernmental Panel on Climate Change
IRP	Integrated Review Plan
ISA	Integrated Science Assessment
LML	lowest measured level
MATS	Modeled Attainment Test Software
METs	metabolic equivalents of work
MSA	Metropolitan Statistical Area
MT	metric ton
NAAQS	National Ambient Air Quality Standards
NCDC	National Climatic Data Center
NEI	National Emissions Inventory
NO	nitric oxide
NO ₂	nitrite
NO _x	nitrogen oxides
O ₃	Ozone
OAQPS	Office of Air Quality Planning and Standards
ОН	hydroxyl radical
PA	Policy Assessment
PDI	pain on deep inspiration
PI	posterior interval
PM	particulate matter
ppb	parts per billion
ppm	parts per million
PRB	Policy Relevant Background
REA	Risk and Exposure Assessment

RR	relative risk
SAB	Science Advisory Board
SEDD	State Emergency Department Databases
SES	socioeconomic status
SID	State Inpatient Databases
SO2	sulfur dioxide
STE	stratosphere-troposphere exchange
TRIM Expo	Total Risk Integrated Methodology Inhalation Exposure
VE	ventilation rate
VNA	Voronoi Neighbor Averaging
VOC	volatile organic carbon
WHO	World Health Organization

1 **INTRODUCTION**

2	The U.S. Environmental Protection Agency (EPA) is presently conducting a review of
3	the national ambient air quality standards (NAAQS) for ozone (O ₃), and related photochemical
4	oxidants. The NAAQS review process includes four key phases: planning, science assessment,
5	risk/exposure assessment, and policy assessment/rulemaking. ¹ This process and the overall plan
6	for this review of the O ₃ NAAQS is presented in the Integrated Review Plan for the Ozone
7	National Ambient Air Quality Standards (IRP, U.S. EPA, 2011a). The IRP additionally presents
8	the schedule for the review; identifies key policy-relevant issues; and discusses the key scientific,
9	technical, and policy documents. These documents include an Integrated Science Assessment
10	(ISA), Risk and Exposure Assessments (REAs), and a Policy Assessment (PA). This draft Health
11	REA is one of the two quantitative REAs developed for the review by the EPA's Office of Air
12	Quality Planning and Standards (OAQPS); the second is a Welfare REA. This draft Health REA
13	focuses on assessments to inform consideration of the review of the primary (health-based)
14	NAAQS for O ₃ .
15	The existing primary (health-based) NAAQS for O_3 is set at a level of 75 ppb (0.075
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¹ For more information on the NAAQS review process see http://www.epa.gov/ttn/naaqs/review.html. ² On March 30, 2009, EPA held a public consultation with the CASAC Ozone Panel on the draft IRP. The final IRP took into consideration comments received from CASAC and the public on the draft plan as well as input from senior Agency managers.

1 available since the previous review on the health effects of O_3 includes information on exposure,

2 physiological mechanisms by which O₃ might adversely impact human health, an evaluation of

3 the toxicological and controlled human exposure study evidence, and an evaluation of the

4 epidemiological evidence, including information on reported concentration-response (C-R)

5 relationships for O₃-related morbidity and mortality associations, and also includes information

6 on potentially at-risk populations and life-stages.³

7 This REA is a concise presentation of the conceptual model, scope, methods, key results, 8 observations, and related uncertainties associated with the quantitative analyses performed. This 9 REA builds upon the health effects evidence presented and assessed in the ISA, as well as 10 CASAC advice (Samet, 2011), and public comments on a scope and methods planning document 11 for the REA (here after, "Scope and Methods Plan," U.S. EPA, 2011). Preparation of this second 12 draft REA draws upon the final ISA and reflects consideration of CASAC and public comments on the first draft REA (Frey and Samet, 2012). This second draft health REA is being released, 13 14 concurrently with the second draft welfare REA and second draft PA for review by the CASAC 15 O₃ Panel at a public meeting scheduled for March 25-27, 2014, and for public comment.

16 The second draft PA presents a staff evaluation and preliminary staff conclusions of the 17 policy implications of the key scientific and technical information in the ISA, and second draft REAs. When final, the PA is intended to help "bridge the gap" between the Agency's scientific 18 19 assessments presented in the ISA and REAs, and the judgments required of the EPA 20 Administrator in determining whether it is appropriate to retain or revise the NAAQS. The PA 21 integrates and interprets the information from the ISA and REAs to frame policy options for 22 consideration by the Administrator. In so doing, the PA recognizes that the selection of a specific 23 approach to reaching final decisions on primary and secondary NAAQS will reflect the 24 judgments of the Administrator. The development of the various scientific, technical and policy 25 documents and their roles in informing this NAAQS review are described in more detail in the 26 second draft PA.

³ The ISA also evaluates scientific evidence for the effects of O₃ on public welfare which EPA will consider in its review of the secondary O₃ NAAQS. Building upon the effects evidence presented in the ISA, OAQPS has also developed a second draft of a second REA titled *Ozone Welfare Effects Risk and Exposure Assessment* (U.S. EPA, 2013).

1 **1.1 HISTORY**

2 As part of the last O₃ NAAQS review completed in 2008, EPA's OAQPS conducted 3 quantitative risk and exposure assessments to estimate exposures above health benchmarks and 4 risks of various health effects associated with exposure to ambient O_3 in a number of urban study 5 areas, selected to illustrate the public health impacts of this pollutant (U.S. EPA 2007a, U.S. 6 EPA, 2007b). The assessment scope and methodology were developed with considerable input 7 from CASAC and the public, with CASAC generally concluding that the exposure assessment 8 reflected generally accepted modeling approaches, and that the risk assessments were well done, 9 balanced and reasonably communicated (Henderson, 2006a). The final quantitative risk and 10 exposure assessments took into consideration CASAC advice (Henderson, 2006a; Henderson, 11 2006b), and public comments on two drafts of the risk and exposure assessments.

The exposure and health risk assessment conducted in the last review developed exposure and health risk estimates for 12 urban areas across the U.S., based on 2002 to 2004 air quality data. That assessment provided annual or O₃ season-specific exposure and risk estimates for these years of air quality and for air quality scenarios, simulating just meeting the then-existing 8-hour O₃ standard set in 1997 at a level of 0.08 ppm and several alternative 8-hour standards. The strengths and limitations in the assessment were characterized, and analyses of key uncertainties were presented.

19 Exposure estimates from the last assessment were used as an input to the risk assessment 20 for lung function responses (a health endpoint for which exposure-response functions were 21 available from controlled human exposure studies). Exposure estimates were developed for the 22 general population and population groups including school age children with asthma as well as 23 all school age children. The exposure estimates also provided information on exposures to 24 ambient O₃ concentrations at and above specified benchmark levels (referred to as "exposures of 25 concern"), to provide some perspective on the public health impacts of health effects associated 26 with O_3 exposures in controlled human exposure studies that could not be evaluated in the 27 quantitative risk assessment (e.g., lung inflammation, increased airway responsiveness, and 28 decreased resistance to infection). For several other health endpoints, O₃-related risk estimates 29 were generated using concentration-response relationships reported in epidemiological or field 30 studies, together with ambient air quality concentrations, baseline health incidence rates, and 31 population data for the various locations included in the assessment. Health endpoints included

1-3

in the assessment based on epidemiological or field studies included: hospital admissions for
 respiratory illness in four urban areas, premature mortality in 12 urban areas, and respiratory
 symptoms in asthmatic children in 1 urban area.

4 Based on the 2006 Air Quality Criteria for Ozone (U.S. EPA, 2006), the Staff Paper 5 (U.S. EPA, 2007), and related technical support documents (including the REAs), the proposed 6 decision was published in the Federal Register on July 11, 2007 (72 FR 37818). The EPA 7 proposed to revise the level of the primary standard to a level within the range of 0.075 to 0.070 8 ppm. Two options were proposed for the secondary standard: (1) replacing the current standard 9 with a cumulative seasonal standard, expressed as an index of the annual sum of weighted hourly 10 concentrations cumulated over 12 daylight hours during the consecutive 3-month period within 11 the O_3 season with the maximum index value (W126), set at a level within the range of 7 to 21 12 ppm-hours, and (2) setting the secondary standard identical to the revised primary standard. The 13 EPA completed the review with publication of a final decision on March 27, 2008 (73 FR 14 16436), revising the level of the 8-hour primary O_3 standard from 0.08 ppm to 0.075 ppm, as the 15 3-year average of the fourth highest daily maximum 8-hour average concentration, and revising 16 the secondary standard to be identical to the revised primary standard.

17 Following promulgation of the revised O₃ standard in March 2008, state, public health, 18 environmental, and industry petitioners filed suit against EPA regarding that final decision. 19 At EPA's request, the consolidated cases were held in abeyance pending EPA's 20 reconsideration of the 2008 decision. A notice of proposed rulemaking to reconsider the 21 2008 final decision was issued by the Administrator on January 6, 2010. Three public 22 hearings were held. The Agency solicited CASAC review of the proposed rule on January 23 25, 2010, and additional CASAC advice on January 26, 2011. On September 2, 2011, the 24 Office of Management and Budget returned the draft final rule on reconsideration to EPA for 25 further consideration. EPA decided to coordinate further proceedings on its voluntary 26 rulemaking on reconsideration with this ongoing periodic review, by deferring the 27 completion of its voluntary rulemaking on reconsideration until it completes its statutorily-28 required periodic review. In light of that, the litigation on the 2008 final decision proceeded. 29 On July 23, 2013, the Court ruled on the litigation of the 2008 decision, denying the 30 petitioners suit except with respect to the secondary standard, which was remanded to the 31 Agency for reconsideration. The second draft PA provides additional description of the court

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1 ruling with regard to the secondary standard.

CURRENT RISK AND EXPOSURE ASSESSMENT: GOALS AND PLANNED APPROACH

4 The goals of the current quantitative exposure and health risk assessments are to provide 5 information relevant to answering questions regarding the adequacy of the existing O_3 standard 6 and the potential improvements in public health from meeting alternative standards. To meet 7 these goals, this assessment provides results from several analyses, including (1) estimates of the 8 number of people in the general population and in at-risk populations and lifestages with O_3 9 exposures above benchmark levels, while at moderate or greater exertion levels; (2) estimates of 10 the number of people in the general population and in at-risk populations and lifestages with 11 impaired lung function resulting from exposures to O_3 ; and (3) estimates of the potential 12 magnitude of premature mortality and selected morbidity health effects in the population, 13 including at-risk populations and lifestages, where data are available to assess these groups. For 14 each of the analyses, we provide estimates for recent ambient levels of O_3 and for air quality 15 conditions simulated to just meet the existing O₃ standard and alternative standards.

16 In presenting these results, we evaluate the influence of various inputs and assumptions 17 on the exposure and risk estimates to more clearly differentiate alternative standards that might 18 be considered, including potential impacts on various at-risk populations and lifestages. We also 19 evaluate the distribution of risks and patterns of risk reduction and uncertainties in those risk 20 estimates. In addition, we have conducted an assessment to provide nationwide estimates of the 21 potential magnitude of premature mortality associated with recent ambient O₃ concentrations, to 22 more broadly characterize this risk on a national scale. This assessment includes an evaluation of 23 the distribution of risk across the U.S., to assess the extent to which we have captured the upper 24 end of the risk distribution with our urban study area analyses.

This current quantitative risk and exposure assessment builds on the approach used and lessons learned in the last O_3 risk and exposure assessment, and focuses on improving the characterization of the overall confidence in the exposure and risk estimates, including related uncertainties, by incorporating a number of enhancements, in terms of both the methods and data used in the analyses. This risk assessment considers a variety of health endpoints for which, in staff's judgment, there is adequate information to develop quantitative risk estimates that can meaningfully inform the review of the primary O_3 NAAQS.

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1 1.3 ORGANIZATION OF DOCUMENT

2 The remainder of this document is organized as follows. Chapter 2 provides a conceptual 3 framework for the risk and exposure assessment, including discussions of O_3 chemistry, sources 4 of O_3 precursors, exposure pathways and microenvironments where O_3 exposure can be high, at-5 risk populations and lifestages, and health endpoints associated with O_3 . This conceptual 6 framework sets the stage for the scope of the risk and exposure assessments. Chapter 3 provides 7 an overview of the scope of the quantitative risk and exposure assessments, including a summary 8 of the previous risk and exposure assessments, and an overview of the current risk and exposure 9 assessments. Chapter 4 discusses air quality considerations relevant to the exposure and risk 10 assessments, including available O₃ monitoring data, and important inputs to the risk and 11 exposure assessments. Chapter 5 describes the inputs, models, and results for the human 12 exposure assessment, and discusses the literature on exposure to O_3 , exposure modeling 13 approaches using the Air Pollution Exposure Model (APEX), the scope of the exposure 14 assessment, inputs to the exposure modeling, sensitivity and uncertainty evaluations, and 15 estimation of results. Chapter 6 describes the estimation of health risks based on application of 16 the results of controlled human exposure studies, including discussions of health endpoint 17 selection, approaches to calculating risk, and results. Chapter 7 describes the estimation of health 18 risks in selected urban areas based on application of the results of observational epidemiology 19 studies, including discussions of air quality characterizations, model inputs, variability and 20 uncertainty, and results. Chapter 8 describes the national scale risk characterization and urban 21 area representativeness analysis. Chapter 9 provides an integrative discussion of the exposure 22 and risk estimates generated in the analyses drawing on the results of the analyses based on both 23 clinical and epidemiology studies, and incorporating considerations from the national scale risk 24 characterization.

1

2 OVERVIEW OF EXPOSURE AND RISK ASSESSMENT DESIGN

2 In this chapter, we summarize our framework for assessing exposures to O_3 and the 3 associated risks to human populations. Figure 2-1 provides an overview of the general design of 4 this exposure and risk assessment, which includes air quality characterization, review of relevant 5 scientific evidence on health effects, modeling of exposure, modeling of risk, and risk 6 characterization. Each element identified in the diagram is described in a specific, identified 7 chapter of this exposure and risk assessment. 8 In this O₃ exposure and risk assessment, modeling of personal exposure and estimation of 9 risks which rely on personal exposure estimates, are implemented using the Air Pollution Exposure model (APEX)¹ (U.S. EPA, 2012 a, b). Modeling of population level risks for 10 11 endpoints based on application of results of epidemiological studies, is implemented using the environmental Benefits Mapping and Analysis Program (BenMAP),² a peer reviewed software 12 13 tool for estimating risks and impacts associated with changes in ambient air quality (U.S. EPA, 14 2013). The overall characterization of risk draws from the results of the exposure assessment and 15 both types of risk assessment. 16 The remainder of this chapter includes summary discussions of each of the main elements 17 of Figure 2-1, including policy-relevant exposure and risk questions (Section 2.1), characterization of ambient O₃, including important sources of O₃ precursors, and its relation to 18 19 population exposures, as well as simulation of just meeting existing and potential alternative O_3 20 standards (Section 2.2), review of health evidence identified in the literature describing 21 associations with ambient O_3 (Section 2.3), key components of exposure modeling (Section 2.4), 22 key components of risk modeling (Section 2.5), and risk characterization (Section 2.6). 23 Specific details related to the scope of the exposure and risk assessments and how each 24 element will be addressed in the quantitative exposure and risk analysis are provided in Chapter 25 3. 26

¹ APEX is available for download at http://www.epa.gov/ttn/fera/human_apex.html

² BenMAP is available for download at http://www.epa.gov/air/benmap/



2 Figure 2-1 Overview of Exposure and Risk Assessment Design

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4 2.1 POLICY-RELEVANT EXPOSURE AND RISK QUESTIONS

5 The first step in the design is to determine the set of policy-relevant exposure and risk 6 questions that will be informed by the assessment. Consistent with recommendations from the 7 recent National Academy of Sciences report "Science and Decisions: Advancing Risk 8 Assessment" (NAS, 2009), these exposure and risk assessments have been designed to address 9 the risk questions identified in the Integrated Review Plan for the Ozone National Ambient Air 10 Quality Standards (U.S. EPA, 2011). We have focused on designing the exposure and risk assessments to inform consideration of those risk-related policy-relevant questions in the 11 12 separately developed O₃ NAAQS Policy Assessment. The risk-related policy-relevant questions 13 identified in the Integrated Review Plan are related to two main activities, evaluation of the 14 adequacy of the existing standards and, if appropriate, evaluation of potential alternative 15 standards (U.S. EPA, 2011). With regard to evaluation of the adequacy of the existing standards, 16 the risk-related policy-relevant questions are:

"To what extent do risk and/or exposure analyses suggest that exposures of
concern for O₃-related health effects are likely to occur with existing ambient
levels of O₃ or with levels that just meet the O₃ standard? Are these
risks/exposures of sufficient magnitude such that the health effects might
reasonably be judged to be important from a public health perspective? What are
the important uncertainties associated with these risk/exposure estimates?"

7 With regards to evaluation of potential alternative standards, the risk-related policy-relevant8 questions are:

- 9 "To what extent do alternative standards, taking together levels, averaging times 10 and forms, reduce estimated exposures and risks of concern attributable to O3 11 and other photochemical oxidants, and what are the uncertainties associated with 12 the estimated exposure and risk reductions? What conclusions can be drawn 13 regarding the health protection afforded at-risk populations?"
- 14

15 This risk and exposure assessment is designed to inform consideration of these questions 16 through application of exposure and risk modeling for a set of urban case study areas. Exposure 17 and risk estimates will be generated for recent O3 concentrations, O3 concentrations after 18 simulating just meeting the existing standards, and O3 concentrations after simulating just 19 meeting potential alternative standards. Careful consideration will be given to addressing 20 variability and uncertainty in the estimates, and to the degree to which at-risk populations 21 experience exposures and risks. Exposure modeling is discussed in Chapter 5 (Urban-Scale 22 Assessment of Individual Exposure), while risk modeling is discussed in Chapter 6 23 (Characterization of Health Risks Based on Clinical Studies) and Chapter 7 (Characterization of 24 Health Risks Based on Epidemiological Studies). Chapter 8 (National-Scale Risk Assessment 25 and Representativeness Analysis) provides a national-scale assessment of risks under recent O3 26 concentrations to provide context for the urban-scale analyses and to help characterize the 27 representativeness of the urban-scale analyses.

28 In order to inform consideration of the risk-related policy-relevant questions, the first step 29 for all of the exposure and risk analyses is simulation of meeting the existing and alternative 30 standards. To do this, recent air quality measurements of O₃ are adjusted such that they mimic a 31 realistic and general atmospheric response to changes in precursor emissions for the specific 32 urban area and so that they just meet the existing and alternative standard levels. Conceptually, 33 there is an almost infinite set of combinations of precursor emissions reductions that will result 34 in just meeting the existing or alternative standards. The specific combinations of reductions that 35 might actually be implemented are not relevant for the exposure and risk analyses, as those will 36 result from the implementation processes which follow the establishment of a standard.

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1 However, it is appropriate to ask the question of how the patterns of ambient O₃ on multiple

- 2 temporal scales (hourly, daily, monthly, seasonally) and across each urban area, may respond to
- 3 precursor emissions reductions that result in meeting the existing and potential alternative

4 standards, and how these different patterns of O_3 could affect the exposure and risk results. The

- 5 answers to these questions are critical inputs to the exposure and risk analyses. Consideration of
- 6 the available methods for simulating just meeting existing and alternative standards is discussed
- 7 in Chapter 4 (Air Quality Characterization).

8 Analyses presented in this document to inform the policy-relevant risk questions 9 regarding potential alternative standards, are focused on alternative levels for an 8-hour standard. 10 Other elements of the standard (indicator, averaging time, and form),³ are addressed in the Policy 11 Assessment as part of the overall evaluation of the health protection afforded by the primary O₃ 12 standards.

13 With regard to potential alternative levels for an 8-hour O_3 standard, the quantitative risk 14 assessment evaluates the range of levels in 5 ppb increments from 60 to 70 ppb. These levels 15 were selected based on the evaluations of the evidence provided in the first draft PA, which 16 received support from the CASAC in their advisory letter on the first draft PA (Frey and Samet, 17 2012). For a subset of urban areas, we also evaluated a standard level of 55 ppb, consistent with 18 recommendations from CASAC to also give consideration to evaluating a level somewhat below 19 60 ppb. Thus, for most areas, we evaluate exposures and risks for potential alternative standard 20 levels of 70, 65, and 60 ppb. Some additional analyses were also included for evaluation of 21 exposures and risks for a potential alternative 8-hour standard level of 55 ppb.

22 2.2 AIR QUALITY CHARACTERIZATION

In order to address the policy-relevant questions discussed in Section 2.1, the first step is characterizing O_3 concentrations relevant to estimation of exposure and risk. This requires characterization of recent O_3 concentrations, O_3 concentrations after simulating just meeting the existing standards, and O_3 concentrations after simulating just meeting potential alternative standards. This section provides conceptual information on O_3 formation and responsiveness of O_3 to changes in precursor emissions, that inform the simulations of just meeting existing and

alternative standards.

 $^{^{3}}$ The "form" of a standard defines the air quality statistic that is compared to the level of the standard in determining whether an area attains the standard. The existing form of the 8-hour O₃ standard is the 4th highest daily maximum 8-hour average O₃, averaged over 3 years. The "indicator" of a standard defines the chemical species or mixture that is to be measured in determining whether an area attains the standard.

1

2.2.1 O₃ chemistry and response to changes in precursor emissions

O₃ occurs naturally in the stratosphere where it provides protection against harmful solar ultraviolet radiation, and it is formed closer to the surface in the troposphere from precursor emissions from both natural and anthropogenic sources. O₃ is created when its two primary precursors, volatile organic compounds (VOC) and oxides of nitrogen (NO_x), combine in the presence of sunlight. VOC and NO_x are, for the most part, emitted directly into the atmosphere. Carbon monoxide (CO) and methane (CH₄) can also be important for O₃ formation (U.S. EPA, 2013, section 3.2.2).

9 Rather than varying directly with emissions of its precursors, O_3 changes in a nonlinear 10 fashion with the concentrations of its precursors. NO_x emissions lead to both the formation and 11 destruction of O₃, depending on the local concentrations of NO_x, VOC, and radicals such as the 12 hydroxyl (OH) and hydroperoxy (HO₂) radicals. In areas dominated by fresh emissions of NO_x, 13 these radicals are removed via the production of nitric acid (HNO₃), which lowers the O₃ 14 formation rate. In addition, the depletion of O_3 by reaction with NO is called "titration" and is 15 often found in downtown metropolitan areas, especially near busy streets and roads, and in 16 power plant plumes. This "titration" results in O_3 concentrations that can be much lower than in 17 surrounding areas. Titration is usually confined to areas close to strong NO_x sources, and the 18 NO₂ formed can lead to O₃ formation later and further downwind. Consequently, O₃ response to 19 reductions in NO_x emissions is complex and may include O_3 decreases at some times and 20 locations and increases of O_3 in other times and locations. In areas with low NO_x concentrations, 21 such as those found in remote continental areas and rural and suburban areas downwind of urban 22 centers, the net production of O₃ typically varies directly with NO_x concentrations, and increases 23 with increasing NO_x emissions. 24 In general, the rate of O_3 production is limited by either the concentration of VOCs or

 NO_x , and O_3 formation, using these two precursors relies on the relative sources of OH and NO_x .

26 When OH radicals are abundant and are not depleted by reaction with NO_x and/or other species,

27 O₃ production is referred to as being "NO_x-limited" (U.S. EPA, 2013, section 3.2.4). In this

situation, O₃ concentrations are most effectively reduced by lowering NO_x emissions, rather than

- 29 lowering emissions of VOCs. When the abundance of OH and other radicals is limited either
- 30 through low production or reactions with NO_x and other species, O_3 production is sometimes
- 31 called "VOC-limited" or "radical limited" or "NO_x-saturated" (Jaegle et al., 2001), and O₃ is
- 32 most effectively reduced by lowering VOCs. However, even in NO_x-saturated conditions, very
- 33 large decreases in NO_x emissions can cause the O₃ formation regime to become NO_x-limited.
- 34 Consequently, reductions in NO_x emissions (when large), can make further emissions reductions
- 35 more effective at reducing O_3 . Between the NO_x -limited and NO_x -saturated extremes there is a
- 36 transitional region, where O_3 is less sensitive to marginal changes in either NO_x or VOCs. In

- rural areas and downwind of urban areas, O₃ production is generally NO_x-limited. However,
 across urban areas with high populations, conditions may vary. For contrast, while data from
- 3 monitors in Nashville, TN, suggest NO_x-limited conditions exist there, data from monitors in Los
- 4 Angeles suggest NO_x-saturated conditions (U.S. EPA, 2013, Figure 3-3).
- 5

2.2.2 Sources of O₃ and O₃ Precursors

 O_3 precursor emissions can be divided into anthropogenic and natural source categories, with natural sources further divided into biogenic emissions (from vegetation, microbes, and animals), and abiotic emissions (from biomass burning, lightning, and geogenic sources). The anthropogenic precursors of O_3 originate from a wide variety of stationary and mobile sources.

10 In urban areas, both biogenic and anthropogenic VOCs, as well as CO, are important for 11 O₃ formation. Hundreds of VOCs are emitted by evaporation and combustion processes from a large number of anthropogenic sources. Based on the 2005 national emissions inventory (NEI), 12 13 solvent use and highway vehicles are the two main anthropogenic sources of VOCs, with 14 roughly equal contributions to total emissions (U.S. EPA, 2013, Figure 3-2). The emissions 15 inventory categories of "miscellaneous" (which includes agriculture and forestry, wildfires, 16 prescribed burns, and structural fires), and off-highway mobile sources are the next two largest 17 contributing emissions categories with a combined total of over 5.5 million metric tons a year 18 (MT/year).

19 On the U.S. and global scales, emissions of VOCs from vegetation are much larger than those from anthropogenic sources. Emissions of VOCs from anthropogenic sources in the 2005 20 21 NEI were ~17 MT/year (wildfires constitute ~1/6 of that total), compared to emissions from 22 biogenic sources of 29 MT/year. Vegetation emits substantial quantities of VOCs, such as 23 isoprene and other terpenoid and sesqui-terpenoid compounds. Most biogenic emissions occur 24 during the summer because of their dependence on temperature and incident sunlight. Biogenic 25 emissions are also higher in southern and eastern states than in northern and western states for 26 these reasons and because of species variations.

27 Anthropogenic NO_x emissions are associated with combustion processes. Based on the 28 2005 NEI, the three largest sources of NO_x are on-road and off-road mobile sources (e.g., 29 construction and agricultural equipment), and electric power generation plants (EGUs) (U.S. 30 EPA, 2013, Figure 3-2). Emissions of NO_x therefore are highest in areas having a high density of 31 power plants and in urban areas having high traffic density. However, it is not possible to make 32 an overall statement about their relative impacts on O₃ in all local areas because EGUs are 33 sparser than mobile sources, particularly in the west and south and because of the nonlinear 34 nature of O_3 chemistry discussed in Section 2.2.1.

- Major natural sources of NO_x in the U.S. include lightning, soils, and wildfires. Biogenic
 NO_x emissions are generally highest during the summer and occur across the entire country,
 including areas where anthropogenic emissions are low. It should be noted that uncertainties in
 estimating natural NO_x emissions are much larger than for anthropogenic NO_x emissions.
 O₃ concentrations in a region are maintained by a balance between photochemical
- 6 production and transport of O_3 into the region; and loss of O_3 by chemical reactions, deposition
- 7 to the surface and transport out of the region. O_3 transport occurs on many spatial scales
- 8 including local transport between cities, regional transport over large regions of the U.S. and
- 9 international/long-range transport. In addition, O₃ is also transfered into the troposphere from the
- 10 stratosphere, which is rich in O_3 , through stratosphere-troposphere exchange (STE). STE occurs
- 11 in tropopause "foldings" that occur behind cold fronts, bringing stratospheric air with them (U.S.
- 12 EPA, 2013, section 3.4.1.1). Contributions to O₃ concentrations in an area from STE are defined
- 13 as being part of background O_3 (U.S. EPA, 2013, section 3.4).

14 **2.2.3** Simulation of Meeting Existing and Alternative Standards

15 Conceptually, simulation of meeting existing and alternative standards should reflect the physical and chemical processes of O₃ formation in the atmosphere and estimate how hourly 16 17 values of O_3 at each monitor in an urban area would change in response to reductions in 18 precursor emissions, allowing for nonlinearities in response to emissions reductions and allowing 19 for nonlinear interactions between reductions in NO_x and VOC emissions. For this assessment, 20 we have employed sophisticated air quality models to conduct simulations of hourly O₃ 21 responses to reductions in precursor emissions. This modeling incorporates all known emissions, 22 including emissions from both natural and anthropogenic sources within and outside of the U.S. 23 By using the model-adjustment methodology we are able to more realistically simulate the 24 temporal and spatial patterns of O₃ response to precursor emissions. We chose to simulate just 25 meeting the existing and alternative standards, by applying equal proportional decreases in U.S. 26 anthropogenic emissions of NOx and VOC, in order to avoid any suggestion that we are 27 approximating a specific emissions control strategy that a state or urban area might adopt to meet 28 a standard. These analyses allow us to apply an adjustment to ambient O₃ measurements in the 29 urban case study areas, to better represent how air quality concentrations at each monitor would 30 change to meet the existing and alternative standard levels. The details of the specific approach 31 used to simulating attainment for the existing and alternative standards, are discussed in greater 32 detail in Chapter 4 and in the Chapter 4 appendices. 33 It is fundamentally a policy decision, as to which sources of precursor emissions are most 34 appropriate to decrease to simulate just meeting existing and alternative O₃ standards. In

- 35 addressing the policy-relevant questions regarding the evaluation of alternative standards,
 - 2-7

1 consistent with previous reviews of the O₃ standards, this analysis is focused on simulating

- 2 reductions in risk associated with precursor emissions originating from anthropogenic sources
- 3 within the U.S. In doing so, we recognize that the CAA provides mechanisms primarily for
- 4 reducing emissions from U.S. emissions sources. As such, we estimate changes in exposure and
- 5 risks likely to result from just meeting alternative standards relative to just meeting the existing
- 6 standards, by simulating changes in atmospheric concentrations that represent atmospheric
- 7 response to reductions in U.S. anthropogenic emissions. However, we recognize that, in this
- 8 approach, we are simulating attainment of existing and alternative standard levels, based on
- 9 recent air quality concentrations and the chemical environment and emissions in those years. We
- have not mimicked the future-year atmospheric conditions and emissions inventory as would bedone for the implementation process.
- 12 In addition, while it is possible to decrease O₃ concentrations using decreases in either
- 13 NOx or VOC or both NO_x and VOC, the specific combination of the reductions in those
- 14 emissions is a policy decision, with recognition that atmospheric chemistry considerations will
- 15 make NO_x and VOC decreases more or less effective in specific urban areas, depending on the
- 16 degree to which O₃ formation is NO_x or VOC limited. As discussed above, in most locations,
- 17 decreases in NO_x are the most effective means to decrease ambient O_3 concentrations. However,
- 18 in some downtown urban areas, O_3 formation is VOC-limited, and therefore smaller decreases in
- 19 NO_x will not decrease O_3 .
- 20

2.2.4 Consideration of Health Evidence

21 A critical input for both the exposure and risk assessments is the health evidence 22 summarized in the Integrated Science Assessment (ISA) (U.S. EPA, 2013). This health evidence 23 provides the basis for evaluating the significance of exposures to O_3 , by informing health 24 benchmarks for estimating exposures of concern. The evidence also provides the basis for 25 selecting health endpoints that will be modeled in the risk assessment. This evidence includes 26 controlled human exposure studies and observational epidemiology studies. The health evidence 27 is also the source of the specific studies that are used to develop exposure-response (E-R) and 28 concentration-response (C-R) functions, used in the risk assessment. Finally, the health evidence 29 provides information on at-risk populations to guide the selections of study populations used in 30 the exposure and risk assessments. The following subsections summarize key conceptual aspects 31 regarding exposures of concern, health endpoints, E-R and C-R functions, and at-risk 32 populations.

1 2.2.5 Exposures of Concern

2 The O₃ ISA identifies health effects associated with exposures to varying concentrations 3 of O_3 . However, not all of the evidence is suitable for evaluation in a quantitative risk 4 assessment. Estimating exposures to ambient O_3 concentrations at and above benchmark levels 5 where health effects have been observed in studies provides a perspective on the public health 6 impacts of O₃-related health effects that have been demonstrated in human clinical and 7 toxicological studies but cannot currently be evaluated in quantitative risk assessments, such as 8 lung inflammation, increased airway responsiveness, and decreased resistance to infection. 9 To inform the selection of benchmark levels for O_3 exposure, it is appropriate to consider

10 the evidence from clinical studies which have evaluated individual controlled levels of O_3

11 exposure. There is substantial clinical evidence demonstrating a range of O₃-related effects

12 including lung inflammation and airway responsiveness in healthy individuals at an exposure

13 level of 0.080 ppm. There is additional evidence that asthmatics have larger and more serious

14 effects than healthy people at 0.070 ppm, as well as a substantial body of epidemiological

15 evidence of associations with O_3 levels that extend well below 0.080 ppm. There is a more

16 limited set of evidence based on clinical studies of healthy individuals exposed at 0.060 ppm in

17 which O_3 -related effects have been observed. This is the lowest level at which any O_3 -related

18 effects have been observed in clinical studies of healthy individuals (U.S. EPA, 2013, section

19 6.2.1).

Thus, benchmark levels of 0.060 ppm, 0.070 ppm, and 0.080 ppm are used in this
assessment to characterize exposures of concern for a range of potential health effects in healthy
and at-risk populations exposed to O₃.

23 2.2.6 Health Endpoints

The O₃ ISA identifies a wide range of health outcomes associated with short-term
exposure to ambient O₃, including an array of morbidity effects as well as
premature mortality. The ISA also identifies several morbidity effects and some
evidence for premature mortality associated with longer-term exposures to O₃. In

- 27 Evidence for premainre mortainty associated with tonger-term exposures to 03. If
- 28 *identifying health endpoints for risk assessment, we have focused on endpoints*
- 29 that pertain to at-risk populations, have public health significance, and for which
- 30 *information is sufficient to support a quantitative concentration-response*

1 2

relationship, in the case of epidemiological studies, or exposure-response relationship, in the case of controlled human exposure studies.⁴

3 In considering such endpoints for O3, we draw from two types of studies: controlled 4 human exposure and epidemiological studies. Each study type informs our characterization of 5 O3 risk and can do so in different ways. Estimates of risk based on results of controlled human 6 exposure studies are valuable because they provide clear evidence of the detrimental effects of 7 controlled (and measured) exposures to O₃ over multiple hours on lung function at moderate 8 levels of exertion. Results of these studies can be applied to modeled estimates of population 9 exposure to provide insights into population exposure characteristics, including types of activity 10 patterns and microenvironments, which are associated with high levels of risk. Controlled human 11 exposure studies, however, cannot directly provide relationships for endpoints such as premature 12 death or hospitalizations, focusing more on intermediate biological endpoints including 13 inflammatory, blood, neurological, cardiovascular, and respiratory biomarkers or symptoms. 14 Estimates of risk based on concentration-response functions from observational epidemiology 15 studies can provide insights on risk for more serious or chronic health endpoints. For example, epidemiological studies of O₃ described in the ISA have evaluated associations between O₃ and 16 17 various endpoints including respiratory symptoms, respiratory-related hospitalizations and emergency department (ED) visits, and premature mortality (U.S. EPA, 2013, sections 6.2.9 and 18 19 6.3.4). Epidemiological studies also generally focus on a population residing in specific area, 20 which may reflect a broad range of susceptibilities and sensitivities. Controlled human exposure 21 studies typically involve a smaller number of individuals over a more limited range of health 22 status, in some cases focused on at-risk populations, such as asthmatics and individuals with 23 COPD. Lastly, while controlled human exposure studies directly measure the exposures eliciting 24 the recorded effects, epidemiology studies have not traditionally been based on observations of 25 personal exposure to ambient O_3 , relying instead on surrogate measures of population exposure. 26 Such surrogates are often based on simple averages of ambient O₃ monitor observations. Thus, 27 with attention to their differing strengths and limitations, risk analyses based on each type of 28 study can inform the risk characterization.

29 30 The O_3 ISA makes overall causal determinations based on the full range of evidence including epidemiological, controlled human exposure, and

⁴ The distinction between concentration-response and exposure-response functions reflects the typical use of ambient concentrations as measured at monitor locations as surrogates for population exposures in observational epidemiology studies, as compared to the personal exposures to controlled concentrations of O₃ that are typically used in controlled human exposure studies. Both types of studies are intended to produce an exposure-responserelationship, however, the epidemiology studies are actually providing a concentration-response relationship, which captures the exposure-response relationship with errors in exposure measurement.

1	toxicological studies. Figure 2-1 shows the O_3 health effects which have been
2	categorized by strength of evidence for causality in the O_3 ISA (U.S. EPA, 2013,
3	chapter 2). The ISA determined there to be causal relationships between short-
4	term exposure to ambient O_3 and respiratory effects, including respiratory-related
5	morbidity and mortality and a likely causal relationship with all-cause total
6	mortality and with cardiovascular effects; the evidence was concluded to be
7	suggestive of a causal relationship between short-term exposure to ambient O_3
8	and central nervous system effects. The ISA determined to also be a likely causal
9	relationship between long-term O_3 exposures and respiratory effects (including
10	respiratory symptoms, new-onset asthma, and respiratory mortality), and
11	determined the evidence to be suggestive of causal relationships between long-
12	term O_3 exposures and total mortality as well as cardiovascular, reproductive and
13	developmental, and central nervous system effects.

Short-term O₃ exposures



1 The ISA identifies several specific respiratory responses to short-term O_3 exposure that 2 have been evaluated in controlled human exposure studies (U.S. EPA, 2013, section 6.2.1). 3 These include decreased inspiratory capacity, decreased forced vital capacity (FVC) and forced 4 expiratory volume in 1 second (FEV1); mild bronchoconstriction; rapid, shallow breathing 5 patterns during exercise; symptoms of cough and pain on deep inspiration (PDI); and pulmonary 6 inflammation. While such studies document quantitative relationships between short-term O_3 7 exposure and an array of respiratory-related effects, exposure-response data across a range of 8 concentrations sufficient for developing quantitative risk estimates are only available for O₃-9 related decrements in FEV1 (U.S. EPA, 2013, section 6.2.1).

10 Within the broad category of respiratory morbidity effects, the epidemiology literature 11 has provided effect estimates for a wide range of health endpoints associated with short-term O_3 12 exposures which we have considered for risk assessment. These health endpoints include lung 13 function, respiratory symptoms and medication use, respiratory-related hospital admissions, and 14 emergency department visits. In the case of respiratory symptoms, the evidence is most 15 consistently supportive of the relationship between short-term ambient O₃ metrics and 16 respiratory symptoms and asthma medication use in children with asthma, but not for a 17 relationship between O₃ and respiratory symptoms in children without asthma (U.S. EPA, 2013, 18 section 6.2.9). In the case of hospital admissions, there is evidence of associations between short-19 term ambient O_3 metrics and general respiratory-related hospital admissions as well as more 20 specific asthma-related hospital admissions (U.S. EPA, 2013, section 6.2.7.2).

21 With regard to mortality, studies have evaluated associations between short-term ambient 22 O₃ metrics and all-cause, non-accidental, and cause-specific (usually respiratory or 23 cardiovascular) mortality. The evidence from respiratory-related morbidity studies provides 24 strong support for respiratory-related mortality for which a causal determination has been made 25 (U.S. EPA, 2013, Table 2-3). There are also a number of large studies that have found 26 associations between O₃ and all-cause and all non-accidental mortality for which a likely causal 27 determination has been made (U.S. EPA, 2013, Table 2-3). Thus, it is appropriate to assess risks 28 for respiratory-related mortality as well as for all-cause total mortality associated with O₃ 29 exposure. The ISA also reports a likely causal determination for short-term O_3 and 30 cardiovascular effects, including cardiovascular mortality (U.S. EPA, 2013, Table 2-3). This 31 determination is supported by studies relating total and cardiovascular mortality, coupled with 32 evidence from animal toxicological studies and controlled human exposure studies which find 33 effects of O₃ on systemic inflammation and oxidative stress. Cardiovascular mortality effects are 34 covered through the estimation of risks associated with total mortality, which is dominated by

35 cardiovascular mortality. There are not sufficient epidemiological studies of cardiovascular

- 1 morbidity showing consistent associations to justify inclusion of any cardiovascular morbidity
- 2 endpoints in the quantitative risk assessment.

3 With regard to effects associated with long-term O_3 exposures, the ISA states that the 4 relationship between O_3 and respiratory-related effects, including respiratory symptoms, new-5 onset asthma, and respiratory mortality is likely causal (U.S. EPA, 2013, Table 2-3). This 6 suggests that for long-term exposures, when comparing the evidence for respiratory-related 7 mortality and total mortality, the evidence is strongest for respiratory-related mortality, which is 8 supported by the strong evidence for respiratory morbidity. As a result, it is appropriate to 9 include respiratory mortality rather than total mortality in the risk assessment and to give 10 consideration to inclusion of additional respiratory-related health endpoints.

11 2.2.7 Exposure and Concentration-response Functions for Health Endpoints

12 Estimation of risk requires characterization of the E-R and C-R functions along the full 13 range of potential exposures. For E-R functions, the evidence from individual controlled human 14 exposure studies provides responses for exposures at and above 60 ppb. McDonnell et al. (2012) 15 develop an integrated model of FEV1 response that is fit to the results from controlled human 16 exposure studies and find that a model with a threshold provides the best fit to the data. In 17 addition, the ISA notes that it is difficult to characterize the E-R relationship at and below 40 ppb 18 due to the dearth of data at these lower concentrations (U.S. EPA, 2013, section 2.5.4.4). Thus, 19 for the portion of the risk assessment based on application of results of controlled human 20 exposure studies, the threshold model is applied.

The evidence for a threshold in the C-R functions for mortality and morbidity outcomes derived from the epidemiological literature is limited. In general, the epidemiological evidence suggests a generally linear C-R function with no indication of a threshold. However, evaluation of evidence for a threshold in the C-R function is complicated by the high degree of heterogeneity between cities in the C-R functions and by the sparse data available at lower ambient O3 concentrations (U.S. EPA, 2013, sections 2.5.4.4 and 2.5.4.5).

The ISA also evaluated whether the magnitude of the relationship between short-term exposures to O₃ and mortality changes at lower concentrations (e.g., whether the C-R function is non-linear). The ISA concludes that epidemiologic studies that examined the shape of the C-R curve and the potential presence of a threshold have indicated a generally linear C-R function

- 31 with no indication of a threshold in analyses that have examined 8-h max and 24-h avg O3
- 32 concentrations, and that the evidence supports less certainty in the shape of the C-R function at
- 33 the lower end of the distribution of O_3 concentrations, e.g., 24-hour average O_3 below 20 ppb,
- due to the low density of data in this range (U.S. EPA, 2013, section 2.5.4.4). In the absence of
- 35 information in the scientific literature on alternative forms of C-R functions at low O_3

1 concentrations, the best estimate of the C-R function is a linear, no-threshold function. The

- 2 scientific literature does not provide sufficient information with which to quantitatively
- 3 characterize any potential additional uncertainty in the C-R functions at lower O₃ concentrations
- 4 for use in the quantitative risk assessment.
- 5 Multiple exposures to elevated O_3 levels over the course of an O_3 season may result in 6 adaptation within exposed population. Evidence suggests that repeated or chronic exposures to 7 elevated O₃ can result in morphologic and biochemical adaptation which reduces the impacts of 8 subsequent O₃ exposures (U.S. EPA, 2013, section 6.2.1.1). This has implications for exposure 9 modeling, in that the effects of modeled repeat exposures on risk may be attenuated relative to 10 the effects of the initial exposures. The ISA notes that "neither tolerance nor attenuation should 11 be presumed to imply complete protection from the biological effects of inhaled O3, because 12 continuing injury still occurs despite the desensitization to some responses (U.S. EPA, 2013, 13 section 6.2.1.1)." The ISA reports that there are limited epidemiological studies evaluating 14 adaptation to the mortality effects of O₃, although the limited evidence does suggest that mortality effects are decreased in later months during the O3 season relative to earlier months 15 16 (U.S. EPA, 2013, section 6.3.3). The impact of this phenomenon on risks based on application of 17 results from epidemiological studies is likely to be small, because the relative risk estimates from 18 those studies already incorporate any adaptive phenomenon.

19 2.2.8 At-risk Populations

20 The O₃ ISA refers to "at-risk" populations as an all-encompassing term used for groups 21 with specific factors that increase the risk of an air pollutant- (e.g., O_3) related health effect in a 22 population group (U.S. EPA, 2013, chapter 8). Populations or lifestages can experience elevated 23 risks from O₃ exposure for a number of reasons. These include high levels of exposure due to 24 activity patterns which include a high duration of time in high-O₃ locations, e.g., outdoor 25 recreation or work, high levels of activity which increase the dose of O₃, e.g., high levels of 26 exercise, genetic or other biological factors, e.g., life stage, which predispose an individual to 27 sensitivity to a given dose of O₃, pre-existing diseases, e.g., asthma or COPD, and 28 socioeconomic factors which may result in more severe health outcomes, e.g., low access to 29 primary care that can lead to increased emergency department visits or hospital admissions. To 30 consider risks to these populations, modeling of exposures to O₃ needs to incorporate 31 information on time spent by potentially at-risk populations in high O_3 locations. This requires 32 identification of populations with the identified exposure-related risk factors, e.g. children or 33 adults engaging in activities involving moderate to high levels of outdoor exertion, especially on 34 a repeated basis typical of student athletes or outdoor workers, as well as identifying populations 35 with high sensitivity to O_3 , e.g. asthmatic children. It also requires that information on O_3

1 concentrations be mapped to locations where at-risk populations are likely to be exposed, e.g.

near roadways where running may occur, or at schools or parks where children are likely to be
engaged in outdoor activities.

4 In addition to consideration of factors that lead to increased exposure to O_3 , modeling of 5 risk from O_3 exposures should incorporate additional information on factors that can lead to 6 increased dose of O_3 for a given exposure, e.g., increased breathing rates during periods of 7 exertion. These factors are especially important for risk estimates based on application of the 8 results of controlled human exposure studies. For risk modeling based on application of 9 observational epidemiology results, it is also important to understand characteristics of study 10 populations that can impact observed relationships between ambient O_3 and population health 11 responses.

12 The O_3 ISA identifies a number of factors which have been associated with modifications 13 of the effect of ambient O_3 on health outcomes. Building on the causal framework used

14 throughout the O_3 ISA, conclusions are made regarding the strength of evidence for each factor

15 that may contribute to increased risk of an O_3 -related health effect based on the evaluation and

16 synthesis of evidence across scientific disciplines. The O₃ ISA categorizes potential risk

17 modifying factors by the degree of available evidence. These categories include "adequate

18 evidence," "suggestive evidence," "inadequate evidence," and "evidence of no effect." See

19 Table 8-1 of the O₃ ISA for a discussion of these categories (U.S. EPA, 2013, chapter 8).

20 Factors categorized as having adequate evidence include asthma, lifestage (children less 21 than 18 years of age, adults older than 65 years of age), diets with nutritional deficiencies, and 22 working outdoors. For example, children are the group considered to be at greatest risk because 23 they breathe more air per unit of body weight, are more likely to be active outdoors when O_3 24 levels are high, are more likely than adults to have asthma, and are in a critical time period of 25 rapid lung growth and organ development. Factors categorized as having suggestive evidence 26 include genetic markers, sex (some studies have shown that females are at greater risk of 27 mortality from O_3 compared to males), low socioeconomic status, and obesity. Factors 28 characterized as having inadequate evidence include influenza and other respiratory infections, 29 COPD, cardiovascular disease, diabetes, hyperthyroidism, race, and smoking (U.S. EPA, 2013,

30 section 8.5, Table 8-6).

31

1 2.3 URBAN-SCALE MODELING OF INDIVIDUAL EXPOSURE

Estimates of human exposure to O₃ provide important information to inform consideration of policy-relevant questions identified in Section 2.2 regarding the occurrence of exposures of concern under air quality conditions that meet existing and potential alternative standards, and also to provide inputs to the portion of the risk assessment based on application of

2-15

1 results of controlled human exposure studies. Studies that measure human exposure to O₃ are

2 limited. More commonly, human exposure is estimated using sophisticated models which

3 combine information on ambient O₃ concentrations in various microenvironments, e.g. near

4 roads, in schools, etc., with information on activity patterns for individuals sampled from the

5 general population or specific subpopulations, e.g. children with asthma.

O₃ exposure is highly dependent on the ambient O₃ concentrations in an urban area.
Given that these concentrations are variable from year to year, it is important to model multiple

8 years representing the range of variability on O₃ concentrations to provide a better

9 characterization of potential exposures of concern. In addition, other important sources of

10 variability and uncertainty affecting the exposure estimates should be characterized, including

11 uncertainty and variability in the data on time-activity patterns, O₃ concentrations, and

12 population inputs. This can be accomplished in part by modeling exposure for multiple urban

13 areas selected to represent variability in these underlying sources of variability.

14 This section briefly describes the conceptual foundation for key components of exposure 15 modeling, characterization of microenvironmental O₃ concentrations, and characterization of 16 human activity patterns, including behaviors intended to avert exposures to O₃. In addition, a 17 brief discussion of key factors to consider in selecting urban case study areas for the exposure 18 analysis is provided. The specific exposure model used in this assessment, APEX, is described 19 more fully in Chapters 3 and 5. Characterization of ambient O₃ concentrations is discussed 20 earlier in this chapter and in greater detail in Chapter 4.

21

2.3.1 Microenvironmental O₃ Concentrations

22 Human exposure to O_3 involves the contact (via inhalation) between a person and the pollutant in the various locations (or microenvironments) in which people spend their time. O₃ 23 24 concentrations in some indoor microenvironments, such as within homes or offices, are 25 considerably lower than O_3 concentrations in similarly located outdoor microenvironments, 26 primarily due to deposition processes and the transformation of O_3 into other chemical 27 compounds within those indoor microenvironments. Concentrations of O₃ may also be quite 28 different in roadway environments, such as might occur while an individual is in a vehicle. 29 Thus, three important classes of microenvironments that should be considered when 30 evaluating population exposures to ambient O_3 are indoors, outdoors, and in-vehicle. Within 31 each of these broad classes of microenvironments, there are many subcategories, reflecting types 32 of buildings, types of vehicles, etc. The O₃ ISA evaluated the literature on indoor-outdoor O₃ 33 concentration relationships and found that studies consistently show that indoor concentrations 34 of O₃ are often substantially lower than outdoor concentrations unless indoor sources are present. 35 This relationship is greatly affected by the air exchange rate, which can be affected by open

2-16

- 1 windows, use of air conditioning, and other factors. Ratios of indoor to outdoor O₃
- 2 concentrations generally range from about 0.1 to 0.4 (U.S. EPA, 2013, section 4.3.2). In some
- 3 indoor locations, such as schools, there can be large temporal variability in the indoor-outdoor
- 4 ratios because of differences in air exchange rates over the day. For example, during the school
- 5 day, there is an increase in open doors and windows, so the indoor-outdoor ratio is higher during
- 6 the school day compared with an overall average across all hours and days. In-vehicle
- 7 concentrations are also likely to be lower than ambient concentrations, although the literature
- 8 providing quantitative estimates is smaller. Studies of personal exposure to O₃ have identified
- 9 that O₃ exposures are highest when individuals are in outdoor microenvironments, such as
- 10 walking outdoors midday, moderate when in vehicle microenvironments, and lowest in
- 11 residential indoor microenvironments (U.S. EPA, 2013, section 4.3.3). Thus the time spent
- 12 indoors, outdoors, and in vehicles is likely to be a critical component in estimating O_3 exposures.
- 13 Because of localized chemistry, O₃ concentrations on or near roadways can be much
- 14 lower than away from roadways. This is due to the high levels of NO_X emissions from motor
- 15 vehicles, which can lead to NOx titration of O_3 , reducing O_3 levels during times of peak traffic.
- 16 The ISA reports evidence that concentrations of NO, NO₂, and NOx are negatively correlated
- 17 with concentrations of O₃ near busy roadways. Because few monitors are located in direct
- 18 proximity to roadways, it is important to account for differences between near-road O₃
- 19 concentrations and ambient O₃ measurements in modeling exposure.
- 20

0 **2.3.2 Human Activity Patterns**

Human exposure can be measured using several metrics. Exposure to ambient concentrations is one such metric. It is also possible to model dose, which combines exposure information with physiological parameters related to activity levels. In order to model exposure to ambient concentrations, detailed information on the patterns of time spent in different microenvironments is critical. In order to model O₃ dose, additional information on the activities conducted while in those microenvironments is needed, along with data on physiological parameters associated with different activities.

Several large-scale databases of human time-activity-location patterns have been compiled. The most comprehensive of these databases in the Consolidated Human Activity Database (CHAD), which has been the basis of several previous exposure analyses for previous NAAQS reviews. These databases compile large numbers of diaries of time spent at different activities in different locations collected as part of smaller studies. The ISA notes the high degree of variability in activity patterns across the population, as well as the variability in time spent in different microenvironments. Time-activity-location patterns vary by age group, as well as by region of the U.S. Children generally spend more time in outdoor locations and also generally
 have higher activity levels in those environments.

3 The dose of O3 received for any given exposure in a microenvironment depends not only 4 on the activity levels and O_3 concentrations in the microenvironment, but also on ventilation 5 rates, which are related to age, body weight, and other physiological parameters. Children 6 generally have lower ventilation rates than adults when considering the volume of air breathed 7 per unit time; however, they tend to have a greater oral breathing contribution than adults, and 8 due to smaller lung volumes and generally greater breathing frequencies, children breathe at 9 higher body mass or surface area normalized minute ventilation rates, relative to their lung 10 volumes. Both of these factors tend to increase their applied or intake dose normalized to lung 11 surface area. For example, when comparing daily body mass normalized ventilation rates, 12 children can have up to a factor of two greater ventilation rates when compared to that of adults. 13 During periods of high activity, ventilation rates for children and young adults can be nearly 14 double those during moderate activity. Thus, it is important to model levels of activity and 15 associated ventilation rate as well as time spent in different microenvironments.

In addition to modeling daily exposures, it may also be important to understand the patterns of exposure over an O₃ season, including multiple repeated exposures for the same individuals. Some individuals or subpopulations may exhibit multiple high daily exposures due to persistent patterns of high activity in microenvironments with high O₃ concentrations. For example, children engaged in numerous outdoor sports over a summer O₃ season may have multiple exposures to elevated O₃ levels.

22 Another important issue in characterizing exposure involves consideration of the extent 23 to which people in relevant population groups modify their behavior for the purpose of 24 decreasing their personal exposure to O_3 based on information about predicted air quality levels 25 made public through the Air Quality Index (AQI). The AQI is the primary tool EPA has used to 26 communicate information on predicted occurrences of high levels of O₃ and other pollutants. The 27 AQI provides both the predicted level of air quality in an area along with a set of potential 28 actions that individuals and communities can take to reduce exposure to air pollution and thus 29 reduce the risk of health effects associated with breathing ambient air pollution. There are 30 several studies, discussed in the O₃ ISA, that have evaluated the degree to which populations are 31 aware of the AQI and what actions individuals and communities take in response to AQI values 32 in the unhealthy range. These studies suggest that at-risk populations, such as children, older 33 adults, and asthmatics, modify their behavior in response to days with bad air quality, most 34 commonly by reducing their time spent outdoors or limiting their outdoor activity exertion level. 35 A challenge remains in how to consider existing averting behaviors within the assessment tools 36 we use and how best to use improved knowledge of participation rates, the varying types of

1 actions performed particularly by potentially at-risk individuals, and the duration of these 2 averting behaviors to quantify the impact on estimated exposures and health risks.

3 2.3.3 Modeling of Exposures Associated with Simulating Just Meeting O₃ Standards

4 In order to address policy-relevant questions regarding changes in exposure associated 5 with potential alternative standards, the exposure assessment evaluates changes in the O_3 6 concentrations, and the resulting changes in exposure, associated with simulating just meeting 7 alternative standards relative to just meeting the existing standards. The new, model-adjustment 8 methodology being implemented in this risk and exposure assessment provides for more realistic 9 responses of hourly O₃ concentrations to changes in the precursor emissions that lead to O₃ 10 formation. Characterization of exposure and changes in exposure when simulating just meeting the alternative standards are discussed in greater detail in Chapter 5. 11

12

2.3.4 Considerations in Selecting Urban Case Study Areas for the Exposure Analysis

13 The goal of the urban area exposure analysis is to characterize the variability in exposures 14 for different locations, taking into account variability in essential factors that affect exposures. 15 Important factors identified earlier that may influence exposure include time activity patterns, 16 especially activities occurring in outdoor environments; demographics of the exposed 17 population, e.g., age and income level; and O_3 concentrations. In addition to these factors, the 18 selection of urban areas to include in the exposure analysis takes into consideration the location 19 of O_3 epidemiological studies (for comparability with the risk assessments), the availability of 20 ambient O₃ data and specific exposure information (e.g., air conditioning prevalence), and the 21 desire to represent a range of geographic areas. To make the exposure analysis most useful in 22 addressing the key policy-relevant questions, urban case study areas were also chosen such that 23 most of them exceeded the existing 8-hr O_3 standards and potential alternative standards during 24 the time period of interest.

25

2.4 **RISK ASSESSMENT**

26 Assessment of risk entails joint consideration of the exposure to a hazard, frequency of 27 adverse outcomes given exposure, and severity of resulting adverse outcomes. A risk assessment 28 for O₃ requires characterization of exposures to ambient O₃ for relevant populations,

- 29 identification of appropriate dose-response or concentration-response functions linking O_3 with
- 30 adverse health outcomes, and characterizing risks for individuals and populations.

31 As discussed above, there are two classes of studies that have provided information to 32 inform the risk modeling: controlled human exposure studies and observational epidemiology

33 studies. The conceptual approach to risk assessment varies based on which type of study result is

34 being applied. This section briefly describes the conceptual foundation for several aspects of risk modeling, including the concept of attributable risk, modeling of total risk and incremental risk
 reductions, development of risk estimates based on controlled human exposure studies, and
 development of risk estimates based on results of observational epidemiology studies.

- 4 This section briefly describes the conceptual foundation for key elements of risk 5 modeling, including a discussion of the concept of attributable risk, modeling of risk for total O_3 6 exposure and the distribution of risk over O_3 concentrations, modeling of risk reductions 7 associated with alternative standards, and key factors to consider in selecting urban case study 8 areas for the risk analysis. Characterization of ambient O₃ concentrations is discussed earlier in 9 this chapter and in greater detail in Chapter 4. The specific risk models used in the urban case study area risk analyses, APEX for analyses based on application of controlled human exposure 10 11 studies and BenMAP for analyses based on application of observational epidemiology studies, 12 are described more fully in Chapters 6 and 7, respectively. Chapter 8 provides an additional 13 national-scale assessment of mortality risk associated with recent O_3 concentrations, to provide 14 context for evaluating the magnitude of health risks in the urban case study areas and to evaluate 15 the representativeness of the urban case study areas in estimating O₃ risks.
- 16 2.4.1 Attributable Risk

17 This risk and exposure assessment relies on the concept of attributable risk in evaluating 18 both total risk and incremental changes in risk associated with just meeting existing and potential 19 alternative O_3 standards. Attributable risk is defined as the difference in incidence of an adverse 20 effect between an exposed and unexposed population for a specific stressor. Attributable risk is 21 an important concept when addressing risks that are associated with multiple causes, such as 22 mortality and respiratory hospital admissions.

Estimates of attributable risk require either an exposure-response (E-R) function (for
 analyses based on results of controlled human exposure studies) or a concentration-response (C R) function (for analyses based on results of epidemiology studies).

E-R functions require estimates of exposure, in this case supplied by the APEX modeling described above. In the case of the lung function endpoint evaluated in this risk analysis, the E-R function also requires information on age and exertion levels to predict the impact of O₃

- 29 exposure on decrements in lung function. E-R functions may provide estimates of the incidence
- 30 of an endpoint or the probability of exceeding benchmark decrement levels.

C-R functions derived from relative risk estimates reported in the epidemiological
 literature generally require estimates of ambient O₃ concentrations, baseline incidence rates, and
 estimates of exposed populations. Ambient O₃ concentrations should generally be constructed to

34 match the spatial and temporal averaging used in the underlying epidemiology study; e.g., a

study may have used a spatial average over a metropolitan statistical area of the 8-hour daily
 maximum.

3 As with exposure, attributable risk is highly dependent on the ambient O_3 concentrations 4 in an urban area. Given that these concentrations are variable from year to year, it is important to 5 model multiple years representing the range of variability of O_3 concentrations to provide a 6 better characterization of risk. In addition, other important sources of variability and uncertainty 7 affecting the risk estimates should be characterized, including uncertainty and variability in the 8 C-R and E-R functions, O_3 concentrations and O_3 exposure, and population inputs. This can be 9 accomplished in part by modeling risk for multiple urban areas selected to represent variability in 10 these underlying risk drivers.

11 2.4.2 Modeling of Risk for Total Exposure to O₃

12 As discussed earlier in this chapter, ambient O_3 is contributed to by emissions from a 13 variety of sources, including natural, U.S. anthropogenic, and non-U.S. anthropogenic sources. 14 Once in the atmosphere, O_3 molecules created from these different sources of emissions are not 15 distinguishable. Individuals and populations are exposed to total O₃ from all sources, and risks 16 associated with O_3 exposure are due to total O_3 exposure and do not vary for O_3 exposure 17 associated with any specific source. Given the absence of a detectable threshold in the available 18 C-R functions, total risk attributable to O_3 will thus be the risk associated with total exposure to 19 O₃, with no threshold or cutpoint applied. To address certain policy-related questions, it is 20 possible to approximately attribute risk to specific sources through the use of air quality 21 modeling techniques, and this is explored in the Policy Assessment. However, these techniques 22 are based on applying model results to total O_3 risk, rather than on directly modeling risk 23 attributable to specific sources. 24 As discussed earlier in this chapter, a critical policy-relevant risk question is the O_3

As discussed earlier in this chapter, a critical policy-relevant fisk question is the O_3 attributable risk remaining after just meeting the existing O_3 standards. This risk includes risks associated with O_3 from all sources after we have simulated just meeting the existing daily 8hour maximum standard level of 75 ppb. The estimates of total risk remaining after meeting the existing standard form the reference values for evaluating reductions in risk associated with just meeting alternative levels of the standard.

In addition to providing risk estimates for urban case study areas, it is also useful to evaluate O_3 risks across the entire U.S., both to better understand the total magnitude of the health burden associated with O_3 and to evaluate the representativeness of selected urban case study areas in characterizing the range and variability in risks across the U.S. The national-scale risk assessment presented in Chapter 8 is focused on estimating risk associated with recent O_3 concentrations, rather than on risk after just meeting existing or alternative standards. This is the

2-21

1 appropriate focus for the national analysis, because the techniques used to simulate just meeting

2 existing and alternative standards in urban case study areas are less certain in a national context

3 due to concerns about interdependence between air quality responses in different urban areas;

4 e.g., just meeting a standard in one urban area would likely have impacts on O₃ air quality in

5 surrounding urban areas. It is beyond the scope of this REA to attempt to simulate control

6 strategies that would result in national attainment of existing or alternative primary health

7 standards.

8 2.4.3 Distributions of Risk Across O₃ concentrations

9 Total O_3 risk for the O_3 season is calculated by summing daily risks across all days in the 10 O_3 season. Because of the high degree of variability in daily O_3 concentrations across an O_3 11 season, total O₃ risk will include risks calculated for some days with high O₃ concentrations as 12 well as for some days with very low O_3 concentrations. Therefore it is appropriate to provide the 13 distribution of total risk over the range of daily O₃ concentrations to allow for an understanding 14 of how O_3 concentrations on different days are contributing to the estimates of total risk. In 15 addition, as noted in the ISA and discussed above, because of the relatively lower density of data 16 on days with low concentrations of O₃, there is decreased confidence in the shape of the C-R 17 function at lower O₃ concentrations, and therefore lower confidence in risk estimates for days 18 with lower O_3 concentrations, especially in the range below 20 ppb. As a result, it is appropriate 19 to provide the distribution of total risk over the range of daily O₃ concentrations to allow for 20 better characterization of confidence in the estimates of total risk.

21 2.5 MODELING OF RISKS ASSOCIATED WITH SIMULATING JUST MEETING O₃ 22 STANDARDS

23 In order to address policy-relevant questions regarding changes in risk associated with 24 potential alternative standards, the risk assessment evaluates changes in the distribution of O_3 25 concentrations, and the resulting changes in risk, associated with simulating just meeting 26 alternative standards relative to just meeting the existing standards. The new, model-adjustment 27 methodology being implemented in this risk and exposure assessment provides for more realistic 28 responses of hourly O_3 concentrations to changes in the precursor emissions that lead to O_3 29 formation. As noted earlier there are multiple combinations of reductions in precursor emissions 30 that can result in just meeting alternative standards. As a result, there is variability in the 31 potential changes in the distribution of O_3 concentrations and risk that would result from just 32 meeting existing and alternative standards. Characterization of this variability, as well as 33 uncertainties in the simulation of just meeting the standards, will be included in Chapters 6 and 34 7.

12.6CONSIDERATIONS IN SELECTING URBAN CASE STUDY AREAS FOR THE2RISK ANALYSIS

3 The goal of the urban area risk analysis is to characterize the magnitude of risk and the 4 impact on risk of meeting existing and potential alternative standards. The selection of specific 5 urban case study areas is based on a set of factors reflecting both variability in factors that affect 6 risk and availability of high quality input data, to provide risk estimates that have higher overall 7 confidence. Important factors identified earlier that may influence risk include O₃ concentrations, 8 demographics, exposure factors, and magnitude of the effect estimate in the C-R function. In 9 addition to consideration of variability in these factors, urban areas are preferentially selected if 10 they have O₃ concentrations that are above the existing standards and potential alternative standards, if they have suitable epidemiological studies to provide C-R functions for mortality or 11 12 morbidity, if they have adequate monitoring data available to characterize population exposures, 13 and if they have appropriate baseline health incidence data available.

14 **2.7**

2.7 RISK CHARACTERIZATION

Risk characterization is the process of communicating the results of risk (and exposure) modeling in metrics that have meaning to decision makers. In the specific context of this review, this translates into providing metrics that are most useful in the Policy Assessment to assess the adequacy of the existing O₃ standards in protecting public health with an adequate margin of safety and to evaluate the additional protection provided by potential alternative standards.

20 Risk characterization requires careful translation of very complex outputs of exposure 21 and risk models into simpler metrics, for example, translating hourly O_3 exposures in various 22 microenvironments into estimates of population exposures above alternative exposure 23 benchmarks. Risk characterization also requires the condensation of a large number of analytical 24 steps and results to (a) summarize the results of the risk analysis, usually taking detailed results 25 and condensing them into a more aggregate interpretation while still providing information about 26 heterogeneity across space and time; (b) communicate the sensitivity of results to different 27 modeling assumptions; and (c) characterize the qualitative and quantitative uncertainty in results. 28 As described more fully in Chapter 5 and in the Policy Assessment, EPA has selected, 29 based on providing a reasonable measure of exposures of concern for at-risk populations and 30 lifestages, aggregate exposure metrics including the number and percent of certain highly 31 vulnerable populations exposed to levels of O_3 above exposure levels that have been identified in 32 the scientific literature as associated with adverse respiratory responses. As noted in section 33 2.3.1, these benchmark exposure levels are 0.060 ppm, 0.070 ppm, and 0.080 ppm. Highly 34 vulnerable populations include active children, older adults, and outdoor workers.

As described more fully in Chapters 6 and 7 and in the Policy Assessment, EPA has selected, based on providing characterization of risks to the public including at-risk populations and lifestages, aggregate risk metrics including the number and percent of vulnerable populations experiencing adverse respiratory responses based on application of results of controlled human exposure studies and the attributable incidence and percent of baseline incidence of mortality and morbidity endpoints based on application of results of epidemiology studies.

8 For all three types of metrics (exposure, risk based on controlled human exposure studies, 9 and risk based on epidemiology studies) and for the purpose of evaluating the adequacy of the 10 existing standards, the focus is on the exposure and risk remaining upon just meeting the existing 11 standards. For the purpose of evaluating alternative standards, the focus in on the changes in 12 exposure and risk after simulating just meeting the alternative standards, compared to exposures 13 and risk after simulating just meeting the existing standards.

14 As detailed in Chapter 3, quantitative sensitivity analyses are provided to evaluate the 15 impacts of critical inputs to the exposure and risk modeling. Limited quantitative uncertainty 16 analyses are also included, along with a comprehensive qualitative uncertainty assessment. The 17 overall treatment of uncertainty is guided by the WHO guidelines for uncertainty assessment 18 (World Health Organization, 2008). These guidelines recommend a tiered approach in which 19 progressively more sophisticated methods are used to evaluate and characterize sources of 20 uncertainty depending on the overall complexity of the risk assessment. 21 In order to inform considerations of overall confidence in the risk estimates derived from 22 application of C-R functions derived from the epidemiological literature, we provide the 23 distributions of total risk across the entire range of daily 8-hour maximum O₃ concentrations. In 24 addition, we provide an assessment of the representativeness of the urban areas selected for the 25 risk and exposure analysis in characterizing the overall distribution of risk across the U.S. This

26 assessment evaluates how well the selected urban areas capture important characteristics that are

27 associated with risk, including demographics, air quality levels, and factors affecting exposure

28 such as air conditioning prevalence.

1 **2.8 REFERENCES**

- Frey, C. and J. Samet. 2012. "CASAC Review of the EPA's Policy Assessment for the Review
 of the Ozone National Ambient Air Quality Standards (First External Review Draft –
 August 2012)." U.S. Environmental Protection Agency Science Advisory Board, EPACASAC-13-003.
- U.S. Environmental Protection Agency. 2012a. Total Risk Integrated Methodology (TRIM) Air
 Pollutants Exposure Model Documentation (TRIM.Expo / APEX, Version 4.4) Volume I:
 User's Guide. Research Triangle Park, NC: EPA Office of Air Quality Planning and
 Standards. (EPA document number EPA-452/B-12-001a).
- 10 <<u>http://www.epa.gov/ttn/fera/human_apex.html></u>.
- U.S. EPA. 2012 b. "Total Risk Integrated Methodology (TRIM) Air Pollutants Exposure Model
 Documentation (TRIM.Expo / APEX, Version 4.4) Volume II: Technical Support
 Document." Research Triangle Park, NC: Office of Air Quality Planning and Standards.
- 14 (EPA document number EPA-452/B-12-001b).
- 15 <<u>http://www.epa.gov/ttn/fera/human_apex.html</u>>.
- U.S. EPA. 2013. Integrated Science Assessment of Ozone and Related Photochemical Oxidants
 (Final Report). Washington, DC: EPA Office of Air and Radiation. (EPA document
 number EPA/600/R-10/076F).
- U.S. EPA. 2013. "Environmental Benefits Mapping Analysis Program Community Edition
 (BenMAP-CE v1.0)," posted on December 03, 2013,
- 21 <u><http://www.epa.gov/air/benmap/></u>.
- 22 World Health Organization. 2008. "Part 1: Guidance Document on Characterizing and
- 23 Communicating Uncertainty in Exposure Assessment, Harmonization Project Document
- 24 No. 6." Published under joint sponsorship of the World Health Organization, the
- 25 International Labor Organization and the United Nations Environment Programme. WHO
- 26 Press, World Health Organization, 20 Avenue Appia, 1211 Geneva 27, Switzerland (tel.:

27 +41 22 791 2476).

28

3 SCOPE

This chapter provides an overview of the scope and key design elements of this
quantitative exposure and health risk assessment. The design of this assessment began with a
review of the exposure and risk assessments completed during the last O₃ NAAQS review (U.S.
EPA, 2007a,b), with an emphasis on considering key limitations and sources of uncertainty
recognized in that analysis.

7 As an initial step in the current O₃ NAAQS review in October 2009, EPA invited outside 8 experts, representing a broad range of expertise (e.g., epidemiology, human and animal 9 toxicology, statistics, risk/exposure analysis, atmospheric science), to participate in a workshop 10 with EPA staff to help inform EPA's plan for the review. The participants discussed key policy-11 relevant issues that would frame the review and the most relevant new science that would be 12 available to inform our understanding of these issues. One workshop session focused on planning 13 for quantitative risk and exposure assessments, taking into consideration what new research 14 and/or improved methodologies would be available to inform the design of quantitative exposure 15 and health risk assessment. Based in part on the workshop discussions, EPA developed a draft 16 IRP (U.S. EPA, 2009) outlining the schedule, process, and key policy-relevant questions that 17 would frame this review. On November 13, 2009, EPA held a consultation with CASAC on the 18 draft IRP (74 FR 54562, October 22, 2009), which included opportunity for public comment. 19 The final IRP incorporated comments from CASAC (Samet, 2009) and the public on the draft 20 plan, as well as input from senior Agency managers. The final IRP included initial plans for 21 quantitative risk and exposure assessments for both human health and welfare (U.S. EPA, 2011a, 22 chapters 5 and 6).

As a next step in the design of these quantitative assessments, OAQPS staff developed more detailed planning documents, the O₃ *National Ambient Air Quality Standards: Scope and*

25 Methods Plan for Health Risk and Exposure Assessment (Health Scope and Methods Plan, U.S.

26 EPA, 2011b) and the O₃ National Ambient Air Quality Standards: Scope and Methods Plan for

27 Welfare Risk and Exposure Assessment (Welfare Scope and Methods Plan, U.S. EPA, 2011c).

28 These Scope and Methods Plans was the subject of a consultation with CASAC on May 19-20,

29 2011 (76 FR 23809, April 28, 2011). Based on consideration of CASAC (Samet, 2011) and

30 public comments on the Scope and Methods Plans, and information in the second draft ISA, we

31 modified the scope and design of the quantitative risk assessment and provided a memo with

32 updates to information presented in the Scope and Methods Plans (Wegman, 2012). The Scope

33 and Methods Plans together with the update memo provide the basis for the discussion of the

34 scope of this exposure and risk assessment provided in this chapter. This chapter also reflects

comments received from CASAC based on their review of the first draft Risk and Exposure
 Assessment on September 11-12, 2012 (Frey and Samet, 2012).

3 In presenting the scope and key design elements of the current risk assessment, this 4 chapter first provides a brief overview of the quantitative exposure and risk assessment 5 completed for the previous O₃ NAAOS review in section 3.1, including key limitations and 6 uncertainties associated with that analysis. The remaining sections describe the current exposure 7 and risk assessment, following the general conceptual framework described in Chapter 2. Section 8 3.2 provides a summary of the design of the urban-scale exposure assessment. Section 3.3 9 provides a summary of the design of the urban-scale risk assessment based on application of 10 results of human clinical studies. Section 3.4 provides a summary of the design of the urban-11 scale risk assessment based on application of results of epidemiology studies. Section 3.5 12 provides a summary of the design of the national-scale risk burden assessment and

13 representativeness analysis.

14 **3.1 OVERVIEW OF EXPOSURE AND RISK ASSESSMENTS FROM LAST REVIEW**

15 The exposure and health risk assessment conducted in the review, completed in March 16 2008, developed exposure and health risk estimates for 12 urban areas across the U.S. which 17 were chosen based on the location of O_3 epidemiological studies and availability of ambient O_3 18 data and to represent a range of geographic areas, population demographics, and O_3 climatology. 19 That analysis was in part based upon the exposure and health risk assessments included in the review completed in 1997.¹ The exposure and risk assessment incorporated air quality data (i.e., 20 21 2002 through 2004), and provided annual or O₃ season-specific exposure and risk estimates for 22 these recent years of air quality and for air quality scenarios simulating just meeting the existing 23 8-hour O₃ standard and several alternative 8-hour O₃ standards.

24 **3.1.1** Overview of exposure assessment from last review

Exposure estimates were used as an input to the risk assessment for lung function responses (a health endpoint for which exposure-response functions were available from controlled human exposure studies). Exposure estimates were developed for the general population and population groups including school-age children with asthma as well as all school-age children. The exposure estimates also provided information on population exposures

¹ In the 1994-1997 O₃ NAAQS review, EPA conducted exposure analyses for the general population, children who spent more time outdoors, and outdoor workers. Exposure estimates were generated for 9 urban areas for as is air quality and for just meeting the existing 1-hour standard and several alternative 8-hour standards. Several reports that describe these analyses can be found at: http://www.epa.gov/ttn/naaqs/standards/O₃/s_O₃-pr.html.

exceeding potential health effect benchmark levels that were identified based on the observed
 occurrence of health endpoints not explicitly modeled in the health risk assessment (e.g., lung
 inflammation, increased airway responsiveness, and decreased resistance to infection) associated
 with 6-8 hour exposures to O₃ in controlled human exposure studies.

5 The exposure analysis took into account several important factors including the 6 magnitude and duration of exposures, frequency of repeated high exposures, and breathing rate 7 of individuals at the time of exposure. Estimates were developed for several indicators of 8 exposure to various levels of O_3 air quality, including counts of people exposed one or more 9 times to a given O_3 concentration while at a specified breathing rate and counts of person-10 occurrences (which accumulate occurrences of specific exposure conditions over all people in 11 the population groups of interest over an O_3 season).

12 As discussed in the 2007 Staff Paper (U.S. EPA, 2007c) and in Section II a of the O₃ 13 Final Rule (73 FR 16440 to 16442, March 27, 2008), the most important uncertainties affecting 14 the exposure estimates were related to modeling human activity patterns over an O_3 season, 15 modeling of variations in ambient concentrations near roadways, and modeling of air exchange 16 rates that affect the amount of O₃ that penetrates indoors. Another important uncertainty, 17 discussed in more detail in the Staff Paper (U.S. EPA, 2007c, section 4.3.4.7), was the 18 uncertainty in energy expenditure values which directly affected the modeled breathing rates. 19 These were important since they were used to classify exposures occurring when children were 20 engaged in moderate or greater exertion. Health effects observed in the controlled human 21 exposure studies generally occurred under these exertion levels for 6 to 8-hour exposures to O_3 22 concentrations at or near 0.08 ppm. Reports that describe these analyses (U.S. EPA, 2007a, c; 23 Langstaff, 2007) can be found at: http://www.epa.gov/ttn/naaqs/standards/O₃/s O₃ index.html.

24 **3.1.2** Overview of risk assessment from last review

25 The human health risk assessment presented in the review completed in March 2008 was 26 designed to estimate population risks in a number of urban areas across the U.S., consistent with 27 the scope of the exposure analysis described above (U.S. EPA, 2007b, c). The risk assessment 28 included risk estimates based on both controlled human exposure studies and epidemiological 29 and field studies. O₃-related risk estimates for lung function decrements were generated using 30 probabilistic exposure-response relationships based on data from controlled human exposure 31 studies, together with probabilistic exposure estimates from the exposure analysis. For several 32 other health endpoints, O₃-related risk estimates were generated using concentration-response relationships reported in epidemiological or field studies, together with ambient air quality 33 34 concentrations, baseline health incidence rates, and population data for the various locations 35 included in the assessment. Health endpoints included in the assessment based on

3-3

epidemiological or field studies included hospital admissions for respiratory illness in four urban
areas, premature mortality in 12 urban areas, and respiratory symptoms in asthmatic children in 1
urban area.

4 In the health risk assessment conducted in the previous review, EPA recognized that there 5 were many sources of uncertainty and variability in the inputs to the assessment and that there 6 was significant uncertainty in the resulting risk estimates. The statistical uncertainty surrounding 7 the estimated O₃ coefficients in epidemiology-based concentration-response functions as well as 8 the shape of the exposure-response relationship chosen for the lung function risk assessment 9 were addressed quantitatively. Additional uncertainties were addressed through sensitivity 10 analyses and/or qualitatively. The risk assessment conducted for the previous O_3 NAAQS review 11 incorporated some of the variability in key inputs to the assessment by using location-specific 12 inputs (e.g., location-specific concentration-response functions, baseline incidence rates and 13 population data, and air quality data for epidemiological-based endpoints, location specific air 14 quality data and exposure estimates for the lung function risk assessment). In that review, several 15 urban areas were included in the health risk assessment to provide some sense of the variability 16 in the risk estimates across the U.S.

17 Key observations and insights from the O_3 risk assessment, in addition to important 18 caveats and limitations, were addressed in Section II.B of the Final Rule notice (73 FR 16440 to 19 14 16443, March 27, 2008). In general, estimated risk reductions associated with going from 20 then-current O₃ levels to just meeting the then-existing and alternative 8-hour standards showed 21 patterns of decreasing estimated risk associated with just meeting the lower alternative 8-hour 22 standards considered. Furthermore, the estimated percentage reductions in risk were strongly 23 influenced by the baseline air quality year used in the analysis, which was due to significant 24 year-to-year variability in O₃ concentrations. There was also noticeable city-to-city variability in 25 the estimated O_3 -related incidence of morbidity and mortality across the 12 urban areas. Uncertainties associated with estimated policy-relevant background (PRB) concentrations² were 26 27 also addressed and revealed differential impacts on the risk estimates depending on the health 28 effect considered as well as the location. EPA also acknowledged that at the time of the previous 29 review there were considerable uncertainties surrounding estimates of O₃ C-R coefficients and 30 the shape of concentration-response relationships and whether or not a population threshold or 31 non-linear relationship exists within the range of concentrations examined in the epidemiological 32 studies.

²Policy-relevant background (PRB) O₃ has been defined in previous reviews as the distribution of O₃ concentrations that would be observed in the U.S. in the absence of anthropogenic (man-made) emissions of O₃ precursor emissions (e.g., VOC, CO, NOx) in the U.S., Canada, and Mexico.

1 3.2 PLAN FOR THE CURRENT EXPOSURE AND RISK ASSESSMENTS

2 The Scope and Methods Plan, including updates (U.S. EPA, 2011b; Wegman, 2012), 3 outlined a planned approach for conducting the current quantitative O_3 exposure and risk 4 assessments, including broad design issues as well as more detailed aspects of the analyses. A 5 critical step in designing the quantitative risk and exposure assessments is to clearly identify the 6 goals for the analysis based on the policy-relevant questions identified in Chapter 2. We have 7 identified the following goals for the urban area exposure and risk assessments: (1) to provide 8 estimates of the percent of people in the general population and in sensitive populations with O_3 9 exposures above health-based benchmark levels; (2) to provide estimates of the percentage of the 10 general population and in sensitive populations with impaired lung function (defined based on 11 decrements in FEV_1) resulting from exposures to O_3 ; (3) to provide estimates of the potential 12 magnitude of premature mortality associated with both short-term and long-term O₃ exposures, 13 and selected morbidity health effects associated with short-term O₃ exposures; (4) to evaluate the 14 influence of various inputs and assumptions on risk estimates to the extent possible given 15 available methods and data; (5) to gain insights into the spatial and temporal distribution of risks 16 and patterns of risk reduction and uncertainties in those risk estimates. For the exposure and risk 17 analyses, we will estimate exposures and risks for recent ambient levels of O₃ and for O₃ 18 concentrations after simulating just meeting the existing O₃ standard and potential alternative 19 standards.

20 With regard to selecting alternative levels for the 8-hour O_3 standards for evaluation in 21 the quantitative risk assessment, we base the range of levels on the evaluations of the evidence 22 provided in the first draft PA, which received support from the CASAC in their advisory letter 23 on the first draft PA. The first draft PA recommended evaluation of 8-hour maximum 24 concentrations in the range of 60 to 70 ppb, with possible consideration of levels somewhat 25 below 60 ppb. The upper end of this range is supported by the clear evidence from both clinical 26 and epidemiological studies of effects at exposures of 70 ppb reported in the ISA and 27 summarized in the first draft PA. The lower end of this range is based on considerations of 28 evidence from clinical studies that have shown lung function decrements in healthy adult 29 populations at 60 ppb O_3 exposures, and that 10 percent of healthy adults exposed to 60 ppb O_3 30 experienced lung function decrements that could be adverse to asthmatics. The evidence showing 31 effects in healthy adults at exposures of 60 ppb supports the consideration of risks to sensitive 32 populations at exposure levels below 60 ppb, although specific exposure levels below 60 ppb at 33 which risks may be occurring are not supported by the evidence. An important distinction is that 34 the evidence from controlled human exposure studies is based on exposures, while the standard

- addresses ambient concentrations. Typically, exposures are lower than ambient concentrations
 because people spend a large fraction of their time indoors where O₃ concentrations are lower.³
- Because of the year-to-year variability in O₃ concentrations that results from temporal variability in meteorology and emissions that drive O₃ formation, the exposure and risk assessments evaluate scenarios for meeting the existing and alternative standards based on multiple years of O₃ data. O₃ concentrations from 2006-2010 are used in estimating exposure and risk. This range of years captures a high degree of variability in meteorological conditions, as well as reflecting years with higher and lower emissions of O₃ precursors.
- 9 In order to provide greater confidence in the exposure and risk estimates, this REA uses 10 an urban case study approach for assessing both exposure and risk. This approach provides 11 greater confidence in estimates by allowing us to make use of air quality data, population 12 information, health data, and epidemiology results that are well matched, and it does not require 13 extrapolation of results to locations without these data. In addition, the urban case study 14 approach allows us to simulate just meeting existing and alternative O_3 standards for each urban area, which is not currently feasible for health risk assessment at the national scale.⁴ Specific 15 16 selection criteria for case study urban areas included in the exposure and risk assessments are 17 described in the following sections. In order to gain an understanding of how well the urban case study areas represent risks at a national level and to provide context for the urban case study 18 19 results, we also include two national level analyses, 1) estimation of the national mortality 20 burden associated with recent ambient O_3 and 2) characterization of how well the risk estimates 21 for the set of urban areas modeled reflect the national distribution of mortality risk. 22 Throughout the exposure and risk analyses, we recognize that there are many sources of 23 variability and uncertainty. Each analysis considers carefully the potential sources and 24 significance of variability and uncertainties and, where data are available, provides quantitative
- assessment of variability and uncertainties, either through probabilistic analyses or through
- 26 sensitivity or scenario analyses. In general the analyses follow the WHO guidelines for
- 27 uncertainty assessment (World Health Organization, 2008), which recommend a tiered approach

³ While almost all people spend a large fraction of their time indoors, there is high variability in this fraction between children and adults, and between outdoor workers and indoor workers. The ratio of exposures to ambient concentrations will likely be higher for children than adults, and for outdoor workers compared to indoor workers.

 $^{^4}$ In order to simulate just meeting alternative standards everywhere nationwide using the model-based adjustment approach employed in this REA, some areas would see O₃ design values decreased below the targeted standard level due to O₃ transport between locations. We were not able to devise an approach that would just meet the standard in every location simultaneously. Using the urban case study approach, we can, acknowledging the counterfactual nature of the analysis, assume independence of attainment for each urban case study area, which allows us to simulate just meeting the standards in each urban case study area.

1 in which progressively more sophisticated methods can be used to evaluate and characterize

2 sources of uncertainty depending on the overall complexity, end use of the assessment, and

3 resources and data available to conduct particular uncertainty characterizations.

The planned approaches for conducting the exposure and risk analyses are briefly summarized below. We begin with a general discussion of how uncertainty and variability are addressed in the different elements of the exposure and risk assessment. This is followed by a discussion of the air quality data that will be used in both the exposure and risk assessments and then discussions of each component of the exposure and risk assessments.

9 10

3.3

CHARACTERIZATION OF UNCERTAINTY AND VARIABILITY IN THE CONTEXT OF THE O₃ EXPOSURE AND RISK ASSESSMENT

11 An important component of this population exposure and health risk assessment is the 12 characterization of both uncertainty and variability. Variability refers to the heterogeneity of a 13 variable of interest within a population or across different populations. For example, populations 14 in different regions of the country may have different behavior and activity patterns (e.g., air 15 conditioning use and time spent indoors) that affect their exposure to ambient O_3 and thus the 16 population health response. The composition of populations in different regions of the country 17 may vary in ways that can affect the population response to exposure to $O_3 - e.g.$, two 18 populations exposed to the same levels of O₃ might respond differently if one population is older 19 than the other. Variability is inherent and cannot be reduced through further research. 20 Refinements in the design of a population risk assessment are often focused on more completely 21 characterizing variability in key factors affecting population risk – e.g., factors affecting 22 population exposure or response – in order to produce risk estimates whose distribution 23 adequately characterizes the distribution in the underlying population(s).

24 Uncertainty refers to the lack of knowledge regarding the actual values of inputs to an 25 analysis. Models are typically used in analyses, and there is uncertainty about the true values of 26 the parameters of the model (parameter uncertainty) – e.g., the value of the coefficient for O_3 in a 27 C-R function. There is also uncertainty about the extent to which the model is an accurate 28 representation of the underlying physical systems or relationships being modeled (model 29 uncertainty) – e.g., the shapes of C-R functions. In addition, there may be some uncertainty 30 surrounding other inputs to an analysis due to possible measurement error—e.g., the values of 31 daily O₃ concentrations in a risk assessment location or the value of the baseline incidence rate for a health effect in a population.⁵ 32

⁵ It is also important to point out that failure to characterize variability in an input used in modeling can also introduce uncertainty into the analysis. This reflects the important link between uncertainty and variability

1 In any risk assessment, uncertainty is, ideally, reduced to the maximum extent possible 2 through improved measurement of key variables and ongoing model refinement. However, 3 significant uncertainty often remains, and emphasis is then placed on characterizing the nature of 4 that uncertainty and its impact on risk estimates. The characterization of uncertainty can be both 5 qualitative and, if a sufficient knowledge base is available, quantitative. 6 The characterization of uncertainty associated with risk assessment is ideally addressed in 7 the regulatory context using a tiered approach in which progressively more sophisticated 8 methods are used to evaluate and characterize sources of uncertainty depending on the overall 9 complexity and intended use of the risk assessment (WHO, 2008). Guidance documents 10 developed by EPA for assessing air toxics-related risk and Superfund Site risks as well as recent 11 guidance from the World Health Organization specify multitier approaches for addressing 12 uncertainty. 13 Following the approach used for previous NAAQS risk and exposure assessments (U.S. 14 EPA, 2008c, 2009b, 2010a, b), for the O_3 risk assessment, we are using a tiered framework 15 developed by WHO to guide the characterization of uncertainty. The WHO guidance presents a 16 four-tiered approach, where the decision to proceed to the next tier is based on the outcome of 17 the previous tier's assessment. The four tiers described in the WHO guidance include: 18 Tier 0: recommended for routine screening assessments, uses default uncertainty factors 19 (rather than developing site-specific uncertainty characterizations); 20 Tier 1: the lowest level of site-specific uncertainty characterization, involves qualitative 21 characterization of sources of uncertainty (e.g., a qualitative assessment of the general magnitude 22 and direction of the effect on risk results); 23 Tier 2: site-specific deterministic quantitative analysis involving sensitivity analysis, 24 interval-based assessment, and possibly probability bounded (high-and low-end) assessment; and 25 Tier 3: uses probabilistic methods to characterize the effects on risk estimates of sources 26 of uncertainty, individually and combined. With this four-tiered approach, the WHO framework provides a means for systematically 27 28 linking the characterization of uncertainty to the sophistication of the underlying risk assessment. 29 Ultimately, the decision as to which tier of uncertainty characterization to include in a risk

with the effort to accurately characterize variability in key model inputs actually reflecting an effort to reduce uncertainty.

assessment will depend both on the overall sophistication of the risk assessment and the
 availability of information for characterizing the various sources of uncertainty.

3 This risk and exposure assessment for the O₃ NAAQS review is relatively complex, 4 possibly warranting consideration of a full probabilistic (WHO Tier 3) uncertainty analysis. For 5 the exposure assessment, we include probabilistic representations of important sources of 6 variability; however, due to lack of information regarding reasonable alternative parameter settings for model input variable distributions, we are not able to include a complete probabilistic 7 8 analysis incorporating both variability and uncertainty. Instead, we provide sensitivity analyses 9 to explore the impact of specific model assumptions, and we include a comprehensive qualitative 10 discussion of uncertainty regarding the model inputs and outputs.

11 While a full probabilistic uncertainty analysis is not undertaken for the epidemiology-12 based risk assessment due to limits in available information on distributions of model inputs, we 13 provide a limited assessment using the confidence intervals associated with effects estimates 14 (obtained from epidemiological studies) to incorporate statistical uncertainty associated with 15 sample size considerations in the presentation of risk estimates. Technically, this type of 16 probabilistic simulation represents a Tier 3 uncertainty analysis, although as noted here, it will be 17 limited and only address uncertainty related to the fit of the C-R functions. Incorporation of 18 additional sources of uncertainty related to key elements of C-R functions (e.g., competing lag 19 structures, alternative functional forms, etc.) into a full probabilistic WHO Tier 3 analysis would 20 require that probabilities be assigned to each competing specification of a given model element 21 (with each probability reflecting a subjective assessment of the probability that the given 22 specification is the correct description of reality). However, for most model elements there is 23 insufficient information on which to base these probabilities. One approach that has been taken 24 in such cases is expert elicitation; however, this approach is resource- and time-intensive, and, 25 consequently, it is not feasible to use this technique in support of this O_3 risk assessment.

26 For most elements of the quantitative risk assessments, rather than conducting a full 27 probabilistic uncertainty analysis, we include a qualitative discussion of the potential impact of uncertainty on risk results (WHO Tier 2). For some critical elements of the epidemiology-based 28 29 risk assessment, e.g., the effect-estimate in the C-R function, we include sensitivity analyses to 30 explore the potential impact of our assumptions. This falls under the WHO Tier 2 classification, 31 although we are not able to assign probabilities to the sensitivity analyses. For these sensitivity 32 analyses, we will include only those alternative specifications for input parameters or modeling 33 approaches that are deemed to have scientific support in the literature (and so represent 34 alternative reasonable input parameter values or modeling options). This means that the array of 35 risk estimates presented in this assessment is expected to represent reasonable risk estimates that
1 can be used to provide some information regarding the potential impacts of uncertainty in the2 model elements.

3 3.4 AIR QUALITY CHARACTERIZATION

4 Figure 3-1 diagrams the basic information used in developing the air quality inputs for 5 the REA. Air quality inputs to the urban area exposure and risk assessments include (1) recent air quality data developed from O_3 ambient monitors in each selected urban study area and (2) 6 7 simulated air quality that reflects changes in the distribution of O_3 air quality estimated to occur 8 when the urban area just meets the existing or alternative O_3 standards under consideration. In 9 addition, O₃ air quality surfaces for recent years covering the entire continental U.S. were 10 generated for use in the national-scale assessment. Details of the air quality data used in the REA 11 are discussed in Chapter 4. 12



13

14 Figure 3-1 Conceptual Diagram for Air Quality Characterization in the Health REA

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16 The urban case study area exposure and risk analyses are based on five recent years of air 17 quality data, 2006-2010. We are including 5 years to reflect the considerable variability in 18 meteorological conditions and the variation in O_3 precursor emissions that have occurred in 19 recent years. The analyses focus on the O₃ season, which ranges from April to October in much 20 of the nation but is longer in some warmer areas such as Los Angeles and Houston. The required 21 O₃ monitoring seasons for the urban case study areas are described in more detail in Chapter 4. 22 In developing the O_3 air quality surfaces for the national-scale analysis, a combination of 23 monitoring data and modeled O_3 concentrations are used to provide greater coverage across the

U.S. The procedure for fusing O₃ monitor data with modeling results is described further in
 Chapter 4.

3 Several O_3 metrics are generated for use in the urban area exposure and risk analyses. 4 The exposure analyses use hourly O_3 concentrations, while the risk analyses use several different 5 averaging times. The specific metrics used in each analysis are discussed further in following 6 chapters. For the exposure analysis, hourly O_3 concentrations are interpolated to census tracts 7 using Voronoi neighbor averaging (VNA), a distance weighted interpolation method (Gold, 8 1997; Chen et al., 2004). For the epidemiology-based risk analysis, we developed a composite of 9 all monitors in the urban area for application with the epidemiology studies. We also evaluated 10 several different definitions of the spatial boundaries of the urban areas that determined the 11 monitors included in the spatial average. Some of the epidemiological studies specify a relatively 12 narrow set of counties within an urban area, while others use a broader definition, such as all 13 counties in a core based statistical area (CBSA) as defined by the Census Bureau. For those 14 epidemiological studies that used a relatively narrow set of counties, most were based on 15 counties in the center of the urban area. In most of these areas, the non-attaining O₃ monitors are 16 not located in the center of the urban area, but instead in the surrounding areas, reflecting the 17 transport and atmospheric chemistry governing O_3 formation. As a result, using a monitor set 18 that exactly reflects the specific counties used in the epidemiology studies can exclude counties 19 in an urban area that would realize the most risk reduction resulting from just meeting the O_3 20 standard. To better represent the changes in risk that could be experienced in the urban areas, the 21 core risk estimates for all endpoints will be based on the CBSA definition. Sensitivity analyses are included to evaluate the effect of using only the counties in each urban area that specifically 22 23 match the county set used in the epidemiology studies. 24 Simulation of just meeting the existing and alternative O_3 standards is accomplished by

25 adjusting hourly O₃ concentrations measured over the O₃ season using a model-based adjustment methodology that estimates O₃ sensitivities to precursor emissions changes.⁶ These sensitivities, 26 27 which estimate the response of O₃ concentrations to reductions in anthropogenic NOx and VOC 28 emissions, are developed using the Higher-order Decoupled Direct Method (HDDM) capabilities 29 in the Community Multi-scale Air Quality (CMAQ) model. This modeling approach incorporates 30 all known emissions, including sources of natural and anthropogenic emissions in and outside of 31 the U.S. By using the model-based adjustment methodology we are able to more realistically 32 simulate the temporal and spatial patterns of O₃ response to precursor emissions. We chose to

 $^{^{6}}$ In the first draft of this REA, we used a statistical quadratic rollback approach to simulate just meeting the existing O_{3} standards. In that draft, we proposed using the model based approach that is being used in this draft, and received support for the model based approach from CASAC.

1 simulate just meeting the existing and alternative standards in the urban cast study areas by

2 decreasing U.S. anthropogenic emissions of NOx and VOC throughout the U.S using equal

- 3 proportional decreases in emissions throughout the U.S., in order to avoid any suggestion that we
- 4 are approximating a specific emissions control strategy that a state or urban area might choose to
- 5 meet a standard. More details on the HDDM-adjustment approach are presented in Chapter 4 of
- 6 this REA and in Simon et al. (2013).

7 In the previous review, background O_3 (referred to in that review as policy relevant 8 background, or PRB) was incorporated into the REA by calculating risk only in excess of PRB. 9 CASAC members recommended that EPA move away from using PRB in calculating risks 10 (Henderson, 2007). In addition, comments received from CASAC, based on their review of the 11 first draft Risk and Exposure Assessment on September 11-12, 2012 (Frey and Samet, 2012), 12 agreed with the development of risk estimates with reference to zero O_3 concentration. Based on 13 these recommendations and comments, the second draft REA includes risks associated with O_3 14 from all sources after we have simulated just meeting the existing standard and estimates of total 15 risk remaining after meeting alternative levels of the standards. EPA believes that presenting 16 total risk is most relevant given that individuals and populations are exposed to total O_3 from all 17 sources, and risks associated with O_3 exposure are due to total O_3 exposure and do not vary for 18 O_3 exposure associated with any specific source. In addition, background O_3 is fully represented 19 in estimates of total risk given that the measured and adjusted air quality concentrations being 20 used in the risk and exposure analyses include O_3 produced from precursor emissions from both 21 anthropogenic and background sources. The evidence and information on background O_3 that is 22 assessed in the Integrated Science Assessment (ISA) is considered in the Policy Assessment 23 (PA) in conjunction with the total risk estimates provided in this second draft REA. With regard 24 to background O₃ concentrations, the PA will consider available information on ambient O₃ 25 concentrations resulting from natural sources, anthropogenic sources outside the U.S., and 26 anthropogenic sources outside of North America.

27 In providing a broader national characterization of O_3 air quality in the U.S., this REA 28 draws upon air quality data analyzed in the O₃ ISA as well as national O₃ databases and 29 modeling of O₃ using the Community Multiscale Air Quality (CMAQ) model. This information, 30 along with additional analyses, is used to develop a broad characterization of recent air quality 31 across the nation. This characterization includes O_3 levels in the urban case study areas for the 32 time periods relevant to the risk analysis and information on the spatial and temporal 33 characterization of O₃ across the national monitoring network. This information is then used to 34 place the relative comparative attributes of the selected study areas into a broader national 35 comparative context to help judge the overall representativeness of the selected study areas in 36 characterizing O_3 risk for the nation. In addition, to better characterize the spatial patterns of

1 responses of the distribution of O_3 to just meeting existing and alternative O_3 standards, we also 2 provide assessments of the historical patterns of responses of O_3 to emissions changes over time 3 and an assessment of national patterns of responses to emissions changes relative to the spatial 4 distribution of populations. These analyses are presented in more detail in Chapter 8 and Chapter 5 8 appendices.

6 3.5 EXPOSURE ASSESSMENT

17 18

7 Figure 3-2 diagrams the basic structure of the population exposure assessment. Basic

- 8 inputs to the exposure assessment include the following: (1) recent measurements of O_3
- 9 concentrations from monitors in each selected urban study area; (2) O_3 concentrations that reflect
- 10 changes in the distribution of O_3 air quality estimated to occur when an area just meets the

11 existing or alternative O₃ standards under consideration; (3) population and demographic

12 information, e.g., age, gender, etc.; (4) time-location activity pattern data; and (5) physiological

- 13 data, e.g., body mass index, ventilation rates, life-stage development, etc. Basic outputs include
- 14 numbers and percent of persons with O₃ exposures exceeding health-based benchmark levels and
- 15 time-series of O₃ exposures and ventilation rates for individuals (for use in the lung function risk
- 16 analysis). Details of the exposure modeling are discussed in Chapter 5.



19 Figure 3-2 Conceptual Diagram for Population Exposure Assessment

1 The scope of the exposure assessment includes 15 urban case study areas.⁷ These areas 2 3 were selected to be generally representative of U.S. populations, geographic areas, climates, and 4 different O₃ and co-pollutant levels, and they include all of the urban case study areas used in the 5 epidemiology-based risk analysis (see Chapter 7). Three additional cities are included in the 6 exposure modeling beyond those included in the epidemiology-based risk analysis. These cities 7 are included to provide additional information on heterogeneity in exposure but could not be 8 included in the epidemiology-based risk analysis because those analyses require additional 9 information not available in the three additional cities. In addition to providing population 10 exposures for estimation of lung function effects, the exposure modeling provides a 11 characterization of urban air pollution exposure environments and activities resulting in the 12 highest exposures. 13 Population exposure to ambient O_3 levels is evaluated using version 4.5 of the APEX 14 model. The model and updated documentation are available at 15 http://www.epa.gov/ttn/fera/apex_download.html. Exposures are estimated using recent ambient 16 O_3 concentrations, based on 2006-2010 air quality data, and for O_3 concentrations resulting from 17 simulations of just meeting the existing 8-hour O₃ standard and alternative O₃ standards, based 18 on adjusting 2006-2010 air quality data. Because the O_3 standard is based on the 3-year average of the 4th highest daily 8-hour maximum, we simulate just meeting the standard for two periods, 19 20 2006-2008 and 2008-2010. Exposures are estimated for school-age children (ages 5 to 18), 21 asthmatic school-age children, asthmatic adults (ages 19-95), and older persons (ages 65-95). 22 This choice of population groups includes a strong emphasis on children, asthmatics, and persons 23 \geq 65 years old and reflects the finding of the last O₃ NAAQS review (EPA, 2007a) and the ISA 24 (EPA, 2013, Chapter 8) that these are important at-risk groups. 25 In addition to estimating exposures exceeding health-based exposure benchmarks, the 26 exposure estimates are used as an input to the portion of the health risk assessment that is based 27 on exposure-response relationships derived from controlled human exposure studies. The 28 exposure analysis also provides a characterization of populations with high exposures in terms of 29 exposure environments and activities. In addition, the exposure analysis offers key observations 30 based on the results of the APEX modeling, viewed in the context of factors such as averting 31 behavior and key uncertainties and limitations of the model.

⁷ These cities are Atlanta, GA; Baltimore, MD; Boston, MA; Chicago, IL; Cleveland, OH; Dallas, TX; Denver, CO; Detroit, MI; Houston, TX; Los Angeles, CA; New York, NY; Philadelphia, PA; Sacramento, CA; St. Louis, MO; and Washington, D.C. We also considered included Seattle; however, the available monitoring data was not sufficient to accurately characterize O₃ exposures for most populations in the Seattle area.

3.6 URBAN-SCALE LUNG FUNCTION RISK ANALYSES BASED ON APPLICATION OF RESULTS FROM CONTROLLED HUMAN EXPOSURE STUDIES

4 The major components in the lung function risk assessment are shown in Figure 3-3.

5 Basic inputs to the analysis include 1) personal exposure to ambient O₃ derived from the

- 6 exposure modeling described in Section 3.2.3., 2) data from controlled human exposure studies,
- 7 used to construct exposure-response functions, 3) physiological data, including body mass index,
- 8 age, etc., and 4) exercise levels, which determine breathing rates and affect dose. Basic outputs
- 9 include the percentage of total population and sub-populations, e.g., children with asthma, with
- 10 predicted lung function decrements (measured as decrements in forced expiratory volume in one
- 11 second, or FEV₁), greater than or equal to 10, 15, and 20 percent, for recent O_3 levels and for O_3
- 12 levels after just meeting existing and alternative standards.
- 13



14

Figure 3-3 Conceptual Diagram of O₃ Lung Function Health Risk Assessment Based on Controlled Human Exposure Studies

17

18 Prior EPA risk assessments for O_3 have included risk estimates for lung function

19 decrements and respiratory symptoms based on analysis of individual data from controlled

20 human exposure studies. The current assessment applies probabilistic exposure-response

- 21 relationships which are based on analyses of individual data that describe the relationship
- 22 between a measure of personal exposure to O_3 and the measure(s) of lung function recorded in

1 the study. The current quantitative lung function risk assessment presents only a partial picture of

2 the risks to public health associated with short-term O₃ exposures, as there are additional

3 controlled human exposure studies that have evaluated cardiovascular and neurological outcomes

4 due to O_3 exposure. However, these studies do not provide sufficient information with which to

5 generate exposure-response functions and therefore are not suitable for quantitative risk

6 assessment.

7 Modeling of risks of lung function decrements is based on application of results from 8 controlled human exposure studies. These studies involve volunteer subjects who are exposed 9 while engaged in different exercise regimens to specified levels of O_3 under controlled 10 conditions for specified amounts of time. The responses measured in such studies have included 11 measures of lung function, such as forced expiratory volume in one second (FEV₁), respiratory 12 symptoms, airway hyper-responsiveness, and inflammation. The lung function risk assessment 13 includes lung function decrement risk estimates, using FEV_1 , for the adult population, school-age 14 children (ages 5-18), and asthmatic school-age children (ages 5-18).

15 In addition to estimating lung function decrements for healthy adults that were the study 16 groups in the controlled human exposure studies, this lung function risk assessment estimates 17 lung function decrements (≥ 10 , ≥ 15 , and $\geq 20\%$ changes in FEV₁) in children 5 to <18 years 18 old. The lung function estimates for children are based on applying data from young adult 19 subjects (18-35 years old) to children aged 5-18. This is based on findings from other chamber 20 studies and summer camp field studies documented in the 1996 O₃ Staff Paper (U.S. EPA, 21 1996a) and 1996 O₃ Criteria Document (U.S. EPA, 1996b), that lung function changes in healthy 22 children are similar to those observed in healthy young adults exposed to O₃ under controlled 23 chamber conditions.

Risk metrics estimated for lung function risk include the numbers of school-age children and other population groups experiencing one or more occurrences of a lung function decrement $\geq 10, \geq 15, \text{ and } \geq 20\%$ in an O₃ season and the total number of occurrences of these lung function decrements in school-age children and active school-age children.

28 The risk assessment includes two different modeling approaches. The first approach 29 employs a model that estimates FEV_1 responses for individuals associated with short-term 30 exposures to O₃ (McDonnell et al., 2012). This model is based on the data from controlled 31 human exposure studies included in the prior lung function risk assessment as well as additional 32 data sets for different averaging times and breathing rates. These data were from 23 controlled 33 human O₃ exposure studies that included exposure of 742 volunteers aged 18–35 years (see 34 McDonnell et al., 2007 and McDonnell et al., 2012, for a description of these data). Outputs from 35 this model include FEV_1 decrements for each simulated individual for each day, which can be

36 used to calculate the population distribution of FEV₁ decrements, and the percent of the

1 population with FEV₁ decrements $\geq 10, \geq 15$, and $\geq 20\%$ after just meeting existing and 2 alternative standards.

In addition, we are applying the approach used in the last review and in the first draft of
the REA, which employs a probabilistic population-level exposure-response function derived
from the results of a number of controlled human exposure studies.

6 This modeling approach uses a smaller set of controlled human exposure studies and the 7 population distribution of O_3 exposures to directly estimate the percent of the population with 8 moderate levels of exertion with lung function decrements $\geq 10, \geq 15$, and $\geq 20\%$.

9 Controlled human exposure studies, carried out in laboratory settings, are generally not 10 specific to any particular real-world location. A controlled human exposure studies-based risk 11 assessment can therefore appropriately be carried out for any locations for which there are 12 adequate air quality data on which to base the modeling of personal exposures. For this 13 assessment, we have selected 15 urban case study areas (matching the areas used in the exposure 14 analysis), representing a range of geographic areas, population demographics, and O_3 15 climatology. These 15 areas also include the 12 urban case study areas evaluated in the risk 16 analyses based on concentration-response relationships developed from epidemiological or field

17 studies.

In the controlled human exposure study based risk assessment, there are two broad sources of uncertainty to the risk estimates. One of the important sources of uncertainty is the estimation of the population distribution of individual time series of O₃ exposures and ventilation rates; these uncertainties are addressed as part of the exposure assessment. The second broad source of uncertainty in the risk calculation results from uncertainties in the lung function risk model. Sensitivity analyses are conducted to inform a qualitative discussion of these uncertainties.

25

3.7

URBAN CASE STUDY AREA EPIDEMIOLOGY-BASED RISK ASSESSMENT

26 The major components of the portion of the urban case study area health risk assessment 27 based on data from epidemiological studies are illustrated in Figure 3-4. Basic inputs to this 28 analysis include 1) measured O₃ concentrations for recent conditions and adjusted air quality 29 representing O₃ concentrations after just meeting existing and alternative standards, 2) C-R 30 functions derived from epidemiological studies evaluating associations between O₃ 31 concentrations and mortality and morbidity endpoints and 3) population counts and baseline 32 incidence rates for mortality and morbidity endpoints. Basic outputs for each urban area include 33 estimates of O₃-attributable incidence and percent O₃-attributable incidence for selected 34 mortality and morbidity endpoints and changes and percent changes in O₃-attributable incidence.



Figure 3-4 Conceptual Diagram of Urban Case Study Area Health Risk Assessment Based on Results of Epidemiology Studies

4 Epidemiological and field studies provide estimated concentration-response relationships 5 based on data collected in real--world settings. Ambient O₃ concentrations used in these studies 6 are typically spatial averages of monitor-specific measurements, using population-oriented 7 monitors. Population health responses for O₃ have included population counts of school 8 absences, emergency room visits, hospital admissions for respiratory and cardiac illness, 9 respiratory symptoms, and premature mortality. Risk assessment based on epidemiological 10 studies typically requires baseline incidence rates and population data for the risk assessment locations. To minimize uncertainties introduced by extrapolation, a risk assessment based on 11 12 epidemiological studies can be performed for the locations in which the studies were carried out, 13 rather than extrapolating results to urban areas where studies for a particular health endpoint 14 have not been conducted. 15 The set of urban case study areas included in this portion of the risk assessment was chosen in order to provide population coverage and to capture the observed heterogeneity in O₃-16 17 related risk across selected urban study areas. In addition, locations had to have at least one 18 epidemiological study conducted in order for the location to be included for a specific endpoint. 19 This assessment also evaluates the mortality risk results for the selected urban areas within a 20 broader national context to better characterize the nature, magnitude, extent, variability, and 21 uncertainty of the public health impacts associated with O₃ exposures. This national-scale

assessment is discussed in the next section.

We selected 2007 and 2009 as analysis years for the urban case study area risk analysis. 1 2 These two years are the midpoint years in the two three-year periods 2006-2008 and 2008-2010. 3 2007 represents a year with generally higher O₃ concentrations, and 2009 represents a year with 4 generally lower O3 concentrations. Analyses for these two years will provide a good 5 representation of the effects of baseline O₃ concentrations on the risk estimates. 6 This risk assessment is focused on health effect endpoints for which the weight of the 7 evidence as assessed in the O₃ ISA supports the causal determination that a likely causal or 8 causal relationship exits between a specific health effect category to be due to exposure to O_3 . 9 The analysis includes estimates of mortality risk associated with short-term 8-hour maximum or 10 8-hour mean O₃ concentrations in all 12 urban case study areas, as well as risk of hospitalization for chronic obstructive pulmonary disease and pneumonia. In addition, the analysis includes 11 12 analysis of hospitalizations for additional respiratory diseases in Los Angeles, New York City, 13 and Detroit, due to limited availability of epidemiological studies covering these endpoints 14 across the 12 urban areas. The analysis also evaluates risks of respiratory related emergency 15 department visits in Atlanta and New York City and risks of respiratory symptoms in Boston, 16 again based on availability of epidemiological studies in these locations. Table 3-1 summarizes 17 the endpoints evaluated for each of the 12 urban case study areas.

Urban Case Study	Mortality	COPD and	Other	Respiratory	Respiratory
Area	1.101000109	Pneumonia	respiratory	Related ED	Symptoms
		hospitalizations	hospitalizations	visits	
Atlanta, GA	Х	X		Х	
Baltimore, MD	X	X			
Boston, MA	Х	X			X
Cleveland, OH	X	X			
Denver, CO	Х	X			
Detroit, MI	Х	X	Х		
Houston, TX	X	X			
Los Angeles, CA	Х	X	X		
New York, NY	X	Х	Х	Х	
Philadelphia, PA	Х	X			
Sacramento, CA	X	X			
St. Louis, MO	X	X			

Table 3-1 Short-term O₃ Exposure Health Endpoints Evaluated in Urban Case Study Areas

3

4 This analysis will also estimate the respiratory mortality risks associated with longer-term exposures to O₃. This is supported by the O₃ ISA, which concluded that the evidence for long-5 6 term exposures to O₃ as likely to be causally related to respiratory effects, including respiratory 7 mortality and morbidity, indicates causal relationship with. There is one national study of long-8 term exposures and respiratory mortality which provides a C-R function for use in the risk 9 assessment. Several other studies have examined long-term exposures and cardiopulmonary 10 mortality, but consistent with the ISA, we focused on respiratory mortality because of the 11 additional supporting evidence related to long-term exposure and morbidity. Because the long-12 term exposure C-R function is based on comparing O_3 and mortality across urban areas, the same 13 C-R function is applied in each of the 12 urban case study areas. The available epidemiological 14 studies evaluating long term O₃ exposures and morbidity endpoints do not provide information 15 that can be used to develop suitable C-R functions. As a result, we are not including quantitative 16 risk estimates for morbidity associated with long-term exposures. 17 We have identified multiple options for specifying the concentration-response functions

18 for particular health endpoints. This risk assessment provides an array of reasonable estimates for

3-20

each endpoint based on the available epidemiological evidence. This array of results provides a
 limited degree of information on the variability and uncertainty in risk due to differences in study
 designs, model specification, and analysis years, amongst other differences.

4 As part of the risk assessment, we address both uncertainty and variability. We provide a 5 limited probabilistic characterization of uncertainty in the national-scale mortality risk estimates 6 using the confidence intervals associated with effects estimates (obtained from epidemiological 7 studies). However, this addresses only one source of uncertainty. For other sources of 8 uncertainty, we include a number of sensitivity analyses to evaluate the impact of alternative 9 approaches to simulating just meeting existing and alternative standards, alternative C-R 10 functions, definitions of O₃ seasons to which C-R functions are applied, and definitions of urban areas to which the C-R functions are applied. In addition, we evaluate the impact in a subset of 11 12 locations of using co-pollutant C-R functions. In the case of variability, we identify key sources 13 of variability associated with O₃ risk (for both short-term and long-term exposure-related 14 endpoints included in the risk assessment) and discuss the degree to which these sources of 15 variability are reflected in the design of the risk assessment. Finally, we also include a 16 comprehensive qualitative assessment of uncertainty and variability.

17 We also provide a representativeness analysis (see Chapter 8) designed to support the 18 interpretation of risk estimates generated for the set of urban study areas included in the risk 19 assessment. The representativeness analysis focuses on comparing the urban study areas to 20 national-scale distributions for key O₃-risk related attributes (e.g., demographics including 21 socioeconomic status, air-conditioning use, baseline incidence rates and ambient O_3 levels). The 22 goal of these comparisons is to assess the degree to which the urban study areas provide 23 coverage for different regions of the country as well as for areas likely to experience elevated O₃-24 related risk due to their specific mix of O₃-risk related attributes.

25 The risk assessment based on application of results of epidemiological studies is 26 implemented using the environmental Benefits Mapping and Analysis Program Community 27 Edition (BenMAP-CE) (U.S. EPA, 2013), EPA's GIS-based computer program for the 28 estimation of health impacts associated with air pollution. BenMAP-CE draws upon a database 29 of population, baseline incidence and effect estimates (regression coefficients) to automate the 30 calculation of health impacts. EPA has traditionally relied upon the BenMAP program to 31 estimate the health impacts avoided and economic benefits associated with adopting new air 32 quality rules. It is also suitable for estimating risks associated with ambient concentrations of O_3 33 and changes in risk resulting from just meeting existing and alternative O₃ standards.

1 3.8 NATIONAL-SCALE MORTALITY RISK ASSESSMENT

The major components of the national-scale mortality risk assessment are shown in
Figure 3-5. Basic inputs to this analysis are similar to those for the urban case study area
epidemiology--based assessment and include 1) gridded O₃ concentrations over the continental
U.S. for recent conditions, 2) C-R functions relating long-term and short-term exposures to O₃ to
mortality, and 3) population and baseline mortality rates. Basic outputs include county and
national estimates of incidence and percent of mortality attributable to O₃.
The national-scale mortality risk assessment serves two primary purposes. First, it serves

9 as part of the representativeness analysis discussed above, providing an assessment of the degree

- 10 to which the urban study areas included in the risk assessment provide coverage for areas of the
- 11 country expected to experience elevated mortality rates due to O₃-exposure. Second, it provides a
- 12 broader perspective on the distribution of risks associated with recent O_3 concentrations
- 13 throughout the U.S., and provides a more complete understanding of the overall public health
- 14 burden associated with O_3 .⁸ We note that a national-scale assessment such as this was completed
- 15 for the risk assessment supporting the latest PM NAAQS review (US EPA, 2010) with the results
- 16 of the analysis being used to support an assessment of the representativeness of the urban study
- 17 areas assessed in the PM NAAQS risk assessment, as described here for O₃.

⁸ In the previous O₃ NAAQS review, CASAC commented that "There is an underestimation of the affected population when one considers only twelve urban "Metropolitan Statistical Areas" (MSAs). The CASAC acknowledges that EPA may have intended to illustrate a range of impacts rather than be comprehensive in their analyses. However, it must be recognized that O₃ is a regional pollutant that will affect people living outside these 12 MSAs, as well as inside and outside other urban areas." Inclusion of the national-scale mortality risk assessment partially addresses this concern by providing a broader characterization of risk for an important O₃ health endpoint.



Figure 3-5 Conceptual Diagram of National O₃ Mortality Risk Assessment Based on Results of Epidemiology Studies

5

1 2

6 The national-scale risk assessment focuses on mortality only, due to the availability of 7 large multi-city epidemiology studies for short-term mortality and the availability of a long-term 8 mortality study which provides information to develop a suitable C-R function. As noted in the 9 discussion of the urban case study area analyses, the available epidemiological studies evaluating 10 long-term O_3 exposures and morbidity endpoints do not provide information that can be used to 11 develop suitable C-R functions. In the case of short-term morbidity endpoints, the available 12 epidemiological studies are generally located in only a few urban areas and, even in the case of 13 the multi-city hospitalization studies, cover only a small fraction of the urban areas in the U.S. In 14 addition, baseline mortality rates are available for every county in the U.S., while baseline hospitalization rates are available in only a small subset of counties. For these reasons, the 15 16 national-scale risk assessment includes only mortality associated with short- and long-term O₃ 17 exposures.

18 We provide a limited probabilistic characterization of uncertainty in the national-scale 19 mortality risk estimates using the confidence intervals associated with effects estimates (obtained 20 from epidemiological studies). However, this addresses only one source of uncertainty. To 21 address some other key potential sources of uncertainty in the national assessment, we conduct 22 sensitivity analyses. Risk estimates are provided for two alternative C-R functions for short-term 23 exposure, reflecting two multi-city epidemiological studies. For short-term exposure-related 24 mortality, the assessment provides several estimates of national mortality risk, including a full 25 national-scale estimate including all counties in the continental U.S., and an analysis restricted to 26 the set of urban areas included in the time-series studies that provide the effect estimates. We

3-23

1 have greater confidence in the analysis based on the large urban areas included in the

- 2 epidemiological studies, but the information from the full analysis of all counties is useful to gain
- 3 understanding of the potential magnitude of risk in less urbanized areas. In addition, the national-
- 4 scale mortality risk assessment evaluates the sensitivity of the nationwide estimates to
- 5 assumptions about the transferability of effect estimates from the cities included in the
- 6 underlying epidemiological studies to other cities in the U.S. Finally, the assessment includes a
- 7 sensitivity analysis evaluating the use of regional priors city--rather than using a national prior in
- 8 developing the city specific Bayesian adjusted effect estimates.⁹ These sensitivity analyses are
- 9 described in detail in Chapter 8.
- 10 The national-scale risk assessment is conducted only for recent O_3 conditions. We do not 11 attempt to simulate nationwide O₃ concentrations that would result from just meeting the existing 12 or alternative O_3 standards everywhere in the U.S. Such a simulation would require detailed 13 modeling of attainment strategies in all potential non-attainment areas and would need to take 14 into account the interdependence of O_3 concentrations across urban areas. This type of analysis is 15 beyond the scope of this risk assessment. Analyses of nationwide attainment are included as part 16 of the Regulatory Impact Analyses that accompany proposed and final rulemaking packages and 17 will likely be included in the rulemaking portion of this review.

18 3.9 PRESENTATION OF EXPOSURE AND RISK ESTIMATES TO INFORM THE O₃ 19 NAAQS POLICY ASSESSMENT

20 We present exposure estimates in three ways: person-occurrences, number, and percent 21 of persons in different populations (e.g., adults, all school-age children, asthmatic school-age 22 children, outdoor workers) with at least one 8-hour average exposure at or above benchmark 23 levels of 60 ppb, 70 ppb, and 80 ppb. In addition, the same types of results are shown for persons 24 with multiple exposures at or above the benchmark levels. The results are presented in summary 25 tables and graphics, while detailed tables of results are provided in an appendix. The focus in the 26 presentation of results is on exposures occurring after simulating just meeting the existing 27 standard and on the change in number and percent of exposures between meeting the existing 28 standard and meeting alternative standards. Results are presented for the five modeled years, for 29 all 15 urban case study areas. 30 Quantitative risk estimates from the analyses based on application of controlled human

31 exposure studies are presented for the two different risk models. For each model, we provide

⁹ In multi-city Bayesian analyses, it is necessary to specify initial values or "priors" which are then "updated" using information from the individual city specific estimates. These priors are generally a mean value across all of the cities, in this case, cities in regions or cities across the nation.

1 estimates of the percent of different populations (adults, all children, children with asthma) with

2 lung function decrements greater than or equal to 10, 15, and 20 percent. As with exposure, the

3 focus in the presentation of results is on risk occurring after simulating just meeting the existing

- 4 standards and on the change in risk occurring between meeting the existing standard and meeting
- 5 alternative standards.

6 Results from the epidemiology-based risk assessment are presented in two ways: (1) total 7 (absolute) health effects incidence for recent air quality and simulations of air quality just 8 meeting the existing and alternative standards under consideration and (2) risk reduction 9 estimates, reflecting the change in the distribution of O_3 between scenarios of just meeting the 10 existing standard and just meeting alternative standards. In addition, risks are presented as the 11 percent of baseline incidence, and risks per 100,000 population, to allow for comparisons 12 between urban areas with very different population sizes. We include risk modeled across the 13 full distribution of O_3 concentrations, as well as core risk estimates for O_3 concentrations down 14 to 0 ppb.

We present an array of risk estimates in order to provide additional context for understanding the potential impact of uncertainty on the risk estimates. For core estimates and sensitivity analyses, we provide the statistical confidence intervals, demonstrating the relative precision of estimates. The graphical presentation of sensitivity analyses focuses on the differences from the core estimates in terms of risk per 100,000 population.

The results of the representativeness analysis are presented using cumulative probability plots (for the national-level distribution of O_3 risk-related parameters) with the locations where the individual urban study areas fall within those distributions noted in the plots using vertical lines. Similar types of plots are used to present the distribution of national-scale mortality estimates based on the national-scale risk assessment, showing the location of the urban case study areas within the overall national distribution.

26 Chapter 9 of this risk and exposure assessment provides a synthesis of the results from 27 the four assessments (urban case study area exposure, urban case study area lung function risk, 28 urban case study area epidemiology-based risk, and national mortality risk). Chapter 9 focuses 29 on comparing patterns of results across locations, years, and alternative standards. Chapter 9 also 30 provides perspective on the overall degree of confidence of the analyses and the 31 representativeness of the set of results in characterizing patterns of exposure and risk and 32 patterns of changes in exposure and risk from just meeting alternative standards relative to just 33 meeting the existing standards.

1 **3.10 REFERENCES**

- Chen, J., R. Zhao and Z. Li. 2004. Voronoi-based k-order neighbor relations for spatial analysis.
 ISPRS J Photogrammetry Remote Sensing, 59(1-2), 60-72.
- Folinsbee, L.J.; W.F. McDonnell and D.H. Horstman. 1988. Pulmonary function and symptom
 responses after 6.6-hour exposure to 0.12 ppm O₃ with moderate exercise. *JAPCA*. 38:
 28-35.
- Gold, C. 1997. Voronoi methods in GIS. In: Algorithmic Foundation of Geographic Information
 Systems (va Kereveld M.; J. Nievergelt; T. Roos; P. Widmayer, eds). Lecture Notes in
 Computer Science, Vol 1340. Berlin: Springer-Verlag, 21-35.
- Henderson, R. 2007. Clean Air Scientific Advisory Committee's (CASAC) Review of the
 Agency's Final O₃ Staff Paper. EPA-CASAC-07-002. March 26.
- Horstman, D.H.; L.J. Folinsbee; P.J. Ives; S. Abdul-Salaam and W.F. McDonnell. 1990. O₃
 concentration and pulmonary response relationships for 6.6-hr exposures with five hrs of
 moderate exercise to 0.08, 0.10, and 0.12 ppm. *Am Rev Respir Dis.* 142:1158-1163.
- Kim, Chong S.; N.E. Alexis; A.G. Rappold; H. Kehrl; J. Milan; J.C. Hazucha; J.C. Lay; M.T.
 Schmitt; C. Martin; R.B. Devlin; D.B. Peden and D. Diaz-Sanchez. 2011. "Lung
 Function and Inflammatory Responses in Healthy Young Adults Exposed to 0.06 ppm O₃
 for 6.6 Hours." American Journal of Respiratory and Critical Care Medicine 183, no. 9
 (2011): 1215-1221.
- Langstaff, J.E. 2007. OAQPS Staff Memorandum to O₃ NAAQS Review Docket (EPA HQ OAR-2005-0172). Subject: Analysis of Uncertainty in O₃ Population Exposure
 Modeling. [January 31, 2007].
- $23 \qquad \qquad < \underline{http://www.epa.gov/ttn/naaqs/standards/O_3/s_O_3_cr_td.html} >.$
- McDonnell, W.F., et al. 1991. "Respiratory response of humans exposed to low levels of O₃ for
 6.6 hours." *American Review of Respiratory Disease*, 147:804-810.
- McDonnell, W.F.; P.W. Stewart and M.V. Smith. 2007. "The temporal dynamics of O₃ –induced
 FEV₁ changes in humans: an exposure-response model." *Inhalation Toxicology*, 19:483–
 494.
- McDonnell W.F.; P.W. Stewart and M.V. Smith. 2010. "Prediction of O₃ -induced lung function
 responses in humans." *Inhalation Toxicology*, 22(2):160-8.
- McDonnell, W.F.; P.W. Stewart; M.V. Smith; C.S. Kim and E.S. Schelegle. 2012. Prediction of
 lung function response for populations exposed to a wide range of ozone conditions.
 Inhalation Toxicology 24: 619-633.
- Samet, J. 2009. Consultation on EPA's Draft Integrated Review Plan for the National Ambient
 Air Quality Standards for Particulate Matter. (EPA document number EPA-CASAC-10 004. December 3).
- Samet, J. 2011. Consultation on EPA's O₃ National Ambient Air Quality Standards: Scope and
 Methods Plan for Health Risk and Exposure Assessment (April 2011) and O₃ National
 Ambient Air Quality Standards: Scope and Methods Plan for Welfare Risk and Exposure
 Assessment (April 2011). EPA-CASAC-11-008. June 21.

1 2 3	 Schelegle, E.S.; C.A. Morales; W.F. Walby; S. Marion and R.P. Allen. 2009. "6.6-Hour Inhalation of O₃ Concentrations from 60 to 87 Parts per Billion in Healthy Humans." American Journal of Respiratory and Critical Care Medicine 180 (2009): 265-272.
4 5 6 7	 Simon H.; K.R. Baker; F. Akhtar; S.L. Napelenok; N. Possiel; B. Wells and B. Timin. 2013. A Direct Sensitivity Approach to Predict Hourly Ozone Resulting from Compliance with the National Ambient Air Quality Standard. Environmental Science and Technology. 2013 Feb. 12. [Epub ahead of print]
8 9 10 11 12	U.S. Environmental Protection Agency. 1996a. "Review of National Ambient Air Quality Standards for O ₃ : Assessment of Scientific and Technical Information - OAQPS Staff Paper." (EPA document number EPA/452/R-96-007). Research Triangle Park, NC: Office of Air Quality Planning and Standards. Available from: NTIS, Springfield, VA; PB96–203435. < <u>http://www.epa.gov/ttn/naaqs/standards/O₃ /s_O₃ pr.html</u> >.
13 14 15	U.S. EPA. 1996b. "Air Quality Criteria for O ₃ and Related Photochemical Oxidants." (EPA document number EPA/600/P-93/004aF-cF). Research Triangle Park, NC: Office of Research and Development, National Center for Environmental Assessment.
16	< http://cfpub.epa.gov/ ncea/cfm/recordisplay.cfm?deid=2831>.
17 18 19	U.S. EPA. 2007a. O ₃ Population Exposure Analysis for Selected Urban Areas. Research Triangle Park, NC: Office of Air Quality Planning and Standards. (EPA document number EPA-452/R-07-010, July).
20 21 22	U.S. EPA. 2007b. O ₃ <i>Health Risk Assessment for Selected Urban Areas</i> . Research Triangle Park, NC: Office of Air Quality Planning and Standards. (EPA document number EPA 452/R-07-009, July).
23 24 25 26	U.S. EPA. 2007c. "Review of the National Ambient Air Quality Standards for O ₃ : Policy Assessment of Scientific and Technical Information. OAQPS Staff Paper." Research Triangle Park, NC: Office of Air Quality Planning and Standards. (EPA report number EPA-452/R-07-007a. < <u>http://www.epa.gov/ttn/naaqs/standards/O₃/s_O₃_cr_sp.html</u> >.
27 28 29	U.S. EPA. 2008a." National Air Quality: Status and Trends Through 2007." Research Triangle Park, NC: Office of Air Quality Planning and Standards. (EPA document EPA-454/R-08-006, November). < <u>http://www.epa.gov/airtrends/2008/index.html</u> >.
30 31 32	U.S. EPA. 2008b. "EPA's Report on the Environment." Washington, DC. (EPA document number EPA/600/R-07/045F, May 2008). < <u>http://cfpub.epa.gov/ncea/cfm/recordisplay.cfm?deid=190806</u> >.
33 34 35 36	U.S. EPA. 2008c. "Risk and Exposure Assessment to Support the Review of the NO2 Primary National Ambient Air Quality Standard." (EPA document number EPA-452/R-08-008a, November 2008). < <u>http://www.epa.gov/ttn/naaqs/standards/nox/data/20081121_NO2_REA_final.pdf>.</u>
37 38 39 40 41	U.S. EPA. 2009a. <i>Integrated Review Plan for the</i> O ₃ <i>National Ambient Air Quality Standards Review: External Review Draft.</i> Research Triangle Park, NC: Environmental Media Assessment Group, National Center for Environmental Assessment and Health and Environmental Impacts Division, Office of Air Quality Planning and Standards. (EPA document number EPA 452/D-09-001, September).

1 2 3 4	U.S. EPA. 2009b. "Risk and Exposure Assessment to Support the Review of the SO2 Primary National Ambient Air Quality Standard." (EPA document number EPA-452/R-09-007, August 2009). < <u>http://www.epa.gov/ttn/naaqs/standards/so2/data/200908SO2REAFinalReport.pdf</u> .>
5 6 7 8	U.S. EPA. 2010a. "Quantitative Risk and Exposure Assessment for Carbon Monoxide – Amended." EPA Office of Air Quality Planning and Standards. (EPA document number EPA-452/R-10-009, July). < <u>http://www.epa.gov/ttn/naaqs/standards/co/data/CO-REA-</u> <u>Amended-July2010.pdf</u> >.
9 10 11 12	U.S. EPA. 2010b. "Quantitative Health Risk Assessment for Particulate Matter." Research Triangle Park, NC: Office of Air Quality Planning and Standards. (EPA document number EPA-452/R-10-005). < <u>http://www.epa.gov/ttn/naaqs/standards/pm/data/PM_RA_FINAL_June_2010.pdf</u> >.
13	U.S. EPA. 2011a. Integrated Review Plan for the O ₃ National Ambient Air Quality Standards.
14 15 16	Research Triangle Park, NC: National Center for Environmental Assessment, Office of Research and Development and Office of Air Quality Planning and Standards, Office of Air and Radiation. (EPA document number EPA 452/R-11-006, April).
17 18 19	U.S. EPA. 2011b. O ₃ National Ambient Air Quality Standards: Scope and Methods Plan for Health Risk and Exposure Assessment. Research Triangle Park, NC: Office of Air Quality Planning and Standards. (EPA document number EPA-452/P-11-001, April).
20 21 22	U.S. EPA. 2011c. O ₃ National Ambient Air Quality Standards: Scope and Methods Plan for Welfare Risk and Exposure Assessment. Research Triangle Park, NC: Office of Air Quality Planning and Standards. (EPA document number EPA-452/P-11-002, April).
23 24 25 26	U.S. EPA. 2013. "Environmental Benefits Mapping Analysis Program Community Edition (BenMAP-CE v1.0)." Research Triangle Park, NC: Health and Environmental Impacts Division, Office of Air Quality Planning and Standards. < <u>http://www.epa.gov/air/benmap/</u> >.
27 28 29 30 31 32	 Wegman, L. 2012. Updates to information presented in the Scope and Methods Plans for the O₃ NAAQS Health and Welfare Risk and Exposure Assessments. Memorandum from Lydia Wegman, Division Director, Health and Environmental Impacts Division, Office of Air Quality Planning and Standards, Office of Air and Radiation, US EPA to Holly Stallworth, Designated Federal Officer, Clean Air Scientific Advisory Committee, US EPA Science Advisory Board Staff Office. May 2, 2012.
33 34	World Health Organization. 2008. Harmonization Project Document No. 6. Part 1: Guidance Document on Characterizing and Communicating Uncertainty in Exposure Assessment.
35	< <u>http://www.who.int/ipcs/methods/harmonization/areas/exposure/en/.</u> >.
36	
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4 AIR QUALITY CONSIDERATIONS

2 4.1 INTRODUCTION

3 Air quality information is used in Chapters 5-8 to assess risk and exposure resulting from 4 recent O_3 concentrations, as well as to estimate the relative change in risk and exposure that 5 could result from just meeting the existing O₃ standard of 75 ppb and the potential alternative standard levels of 70 ppb, 65 ppb, and 60 ppb¹. The same air quality data are used to examine 6 fifteen² urban case study areas in the population exposure analyses discussed in Chapter 5 and 7 8 the lung function risk assessment based on application of results from clinical studies discussed 9 in Chapter 6: Atlanta, GA; Baltimore, MD; Boston, MA; Chicago, IL; Cleveland, OH; Dallas, 10 TX; Denver, CO; Detroit, MI; Houston, TX; Los Angeles, CA; New York, NY; Philadelphia, 11 PA; Sacramento, CA; St. Louis, MO; and Washington, DC. The epidemiology-based risk assessment discussed in Chapter 7 examines twelve³ of the fifteen urban case study areas 12 evaluated in the population exposure analyses. Finally, Chapter 8 includes an assessment of the 13 14 national-scale O₃ mortality risk burden associated with recent O₃ concentrations, and 15 characterizes the representativeness of the 15 urban case study areas compared to the rest of the 16 U.S. This chapter describes the air quality information developed for these analyses, providing 17 an overview of monitoring data and air quality (section 4.2) and an overview of air quality inputs 18 to the risk and exposure assessments (section 4.3).

19

1

20 4.2 OVERVIEW OF O₃ MONITORING AND AIR QUALITY DATA

To determine whether or not the NAAQS have been met at an ambient O₃ monitoring site, a statistic commonly referred to as a "design value" must be calculated based on 3 consecutive years of data collected from that site. The form of the existing O₃ NAAQS design value statistic is the 3-year average of the annual 4th highest daily maximum 8-hour O₃ concentration in parts per billion (ppb), with decimal digits truncated. The existing primary and secondary O₃ NAAQS are met at an ambient monitoring site when the design value is less than

¹ For a subset of urban areas and analyses, the REA evaluates a standard level of 55 ppb, consistent with recommendations from CASAC to also give consideration to evaluating a level somewhat below 60 ppb.

² In the first draft REA, we proposed to include 16 urban areas in the second draft REA. However, further analysis of the air quality information available for Seattle, WA has prompted us to not include that city. This decision and supporting analysis are discussed in more detail in Appendix 4-E.

³ These cities are Atlanta, GA; Baltimore, MD; Boston, MA; Cleveland, OH; Denver, CO; Detroit, MI; Houston, TX; Los Angeles, CA; New York, NY; Philadelphia, PA; Sacramento, CA; and St. Louis, MO.

or equal to 75 ppb.⁴ In counties or other geographic areas with multiple monitors, the area-wide
design value is defined as the design value at the highest individual monitoring site, and the area
is said to have met the NAAQS if all monitors in the area are meeting the NAAQS.

- 4 Air quality monitoring data from 1,468 U.S. ambient O_3 monitoring sites were retrieved
- 5 by EPA staff for use in the risk and exposure assessments. The initial dataset consisted of hourly 6 O_3 concentrations in ppb collected between 1/1/2006 and 12/31/2010 from these monitors. Data
- 7 for nearly 1,400 of these monitors were extracted from EPA's Air Quality System (AQS)
- 8 database⁵, while the remaining data came from EPA's Clean Air Status and Trends Network
- 9 CASTNET) database which consists of primarily rural monitoring sites. While CASTNET
- 10 monitors did not begin reporting regulatory data to AQS until 2011, it is generally agreed that
- 11 data collected from these monitors prior to 2011 is of comparable quality to the data reported to12 AQS.

13 These data were split into two design value periods, 2006-2008 and 2008-2010, and all

14 subsequent analyses based on these data were conducted independently for these two periods.

15 Observations flagged in AQS as having been affected by exceptional events were included the

16 initial dataset, but were not used in design value calculations in accordance with EPA's

17 exceptional events policy. Missing data intervals of 1 or 2 hours in the initial dataset were filled

18 in using linear interpolation. These short gaps often occur at regular intervals in the ambient data

- 19 due to an EPA requirement for monitoring agencies to perform routine quality control checks on
- 20 their O₃ monitors. Quality control checks are typically performed between midnight and 6:00

21 AM when O_3 concentrations are low. Missing data intervals of 3 hours or more were not

22 replaced. Interpolated data values were not used in design value calculations.

- Figures 4-1 and 4-2 show the design values for the existing O₃ NAAQS for all regulatory monitoring sites in the U.S. for the 2006-2008 and 2008-2010 periods, respectively. In general,
- 25 O₃ design values were lower in 2008-2010 than in 2006-2008, especially in the Eastern U.S.

26 There were 518 O₃ monitors in the U.S. with design values above the existing standard in 2006-

27 2008, compared to only 179 in 2008-2010.

⁴ For more details on the data handling procedures used to calculate design values for the current O₃ NAAQS, see 40 CFR Part 50, Appendix P.

⁵ EPA's Air Quality System (AQS) database is a national repository for many types of air quality and related monitoring data. AQS contains monitoring data for the six criteria pollutants dating back to the 1970's, as well as more recent additions such as PM2.5 speciation, air toxics, and meteorology data. At present, AQS receives hourly O₃ monitoring data collected from nearly 1,400 monitors operated by over 100 state, local, and tribal air quality monitoring agencies.



1 2

Figure 4-1 Map of Monitored 8-hour O₃ Design Values for the 2006-2008 Period



Figure 4-2 Map of Monitored 8-hour O₃ Design Values for the 2008-2010 Period

4 4.3 OVERVIEW OF URBAN-SCALE AIR QUALITY INPUTS TO RISK AND 5 EXPOSURE ASSESSMENTS

6 The air quality information input into the urban-scale risk and exposure assessments 7 includes both recent air quality data from the years 2006-2010, as well as air quality data 8 adjusted to reflect just meeting the existing and potential alternative standard levels. In this 9 section, we summarize these air quality inputs and discuss the methodology used to adjust air 10 quality to meet the existing and potential alterative standards.

Figure 4-3 presents a flowchart of air quality data processing steps for the urban-scale
analyses. The rest of section 4.3.1 will provide more details on each step depicted in the flow
diagram. Additional information is provided in Appendices 4-A, 4-B and 4-D.



Figure 4-3 Flowchart of Air Quality Data Processing for Different Parts of the Urban scale Risk and Exposure Assessments

7

1

6 4.3.1 Urban Case Study Areas

4.3.1.1 Exposure Modeling and Controlled Human Study Based Lung Function Risk Assessment

The 15 urban case study areas in the exposure modeling and lung function risk

8 assessments covered a large spatial extent, with boundaries generally similar to those covered by

9 the respective Combined Statistical Areas (CSA) defined by the U.S. Census Bureau. Table 4-1

- 10 gives some basic information about the 15 urban case study areas in the exposure assessment,
- 11 including the number of ambient monitoring sites, the required O₃ monitoring season, and the
- 12 2006-2008 and 2008-2010 design values for each study area. All 15 of the urban case study areas
- 13 had 8-hour O₃ design values above the existing standard in 2006-2008, while 13 urban areas had

⁶ Composite monitors do not always include the highest design value monitor in every urban area.

⁷ 4800 VNA surfaces were created for each urban area/alternative standard level pair: 24 hrs \times 365 days \times 5 years.

- 1 8-hour O₃ design values above the existing standard in 2008-2010. Chicago (74 ppb) and Detroit
- 2 (75 ppb) had design values meeting the existing standard during the 2008-2010 period. The
- 3 design values in the 15 urban areas decreased by an average of 6 ppb between 2006-2008 and
- 4 2008-2010, ranging from no change in Sacramento to a decrease of 15 ppb in Atlanta.
- 5

6Table 4-1 Monitor and Area Information for the 15 Urban Case Study Areas in the Exposure
 7 Modeling and Clinical Study Based Risk Assessment

	# of	$\# \text{ of } O_3$	Population	Required O ₃	2006-2008	2008-2010
Area Name	Counties	Monitors	(2010)	Monitoring Season	DV (ppb)	DV (ppb)
Atlanta	33	13	5,618,431	March - October	95	80
Baltimore	7	7	2,710,489	April - October	91	89
Boston	10	14	5,723,468	April - September	83	77
Chicago	16	26	9,686,021	April - October	78	74
Cleveland	8	13	2,881,937	April - October	82	77
Dallas	11	20	6,366,542	January - December	89	86
Denver	13	26	3,390,504	March - September	86	77
Detroit	9	12	5,218,852	April - September	81	75
Houston	10	22	5,946,800	January - December	91	85
Los Angeles	5	54	17,877,006	January - December	119	112
New York	27	31	21,056,173	April - October	90	84
Philadelphia	15	19	7,070,622	April - October	92	83
Sacramento	7	26	2,755,972	January - December	102	102
St. Louis	17	17	2,837,592	April - October	85	77
Washington	26	22	5,838,518	April - October	87	81

9

10 Since O₃ design values are based on the annual 4th highest 8-hour daily maximum O₃ 11 concentrations from 3-consecutive years, it is useful to look at inter-annual variability. In 12 general, the annual 4th highest 8-hour O₃ concentrations decreased in 11 of the 15 urban areas 13 from 2006 to 2010, while remaining relatively constant in the other 4 areas (Figure 4-4). The 14 average decrease in the annual 4th highest daily maximum concentration from 2006 to 2010 was 15 8 ppb. However, there was significant year-to-year variability, and some areas showed increases 16 in some years relative to 2006, even though the 2010 values were generally lower.



Figure 4-4Trends in Annual 4th Highest 8-hour Daily Maximum O3
Concentrations in ppb for the 15 Urban Case Study Areas for 2006-
2010. Urban areas are grouped into 3 regions: Eastern (top), Central
(middle), and Western (bottom).



4.3.1.2 Epidemiology Based Risk Assessment

1 Table 4-2 gives some basic information on the 12 urban case study areas in the 2 epidemiology-based risk assessment for each set of area boundaries. The spatial extent of each urban case study area was based on the respective Core Based Statistical Area (CBSA)⁸. The 3 4 CBSAs were generally smaller than the study areas used in the exposure modeling and clinical 5 study based risk assessments, except for Baltimore and Houston, where the two study areas were 6 identical. The rationales for the definitions of the spatial areas used in each type of analysis are provided in the corresponding chapters. The final two columns in Table 4-2 show the annual 4th 7 8 highest daily maximum 8-hour O₃ concentration in ppb for the monitors within each urban case 9 study area in 2007 and 2009.

It should be noted that the CBSA boundaries used for the urban case study areas in this 10 assessment are different than those used in the 1st draft of the REA, where the study areas were 11 12 derived from the Zanobetti and Schwartz (2008) study. The change to the CBSA boundaries was 13 intended to capture a larger portion of the urban area populations by including some surrounding 14 suburban counties, rather than focusing strictly on the urban population centers. Two sensitivity 15 analyses were conducted to determine the effect of changing the spatial extent of the urban case study areas on the epidemiology-based risk estimates. These sensitivity analyses are presented in 16 17 Chapter 7, and a summary of the two alternative sets of boundaries for the 12 urban case study 18 areas are provided in Appendix 4-A. 19 Since O₃ is not directly emitted but is formed through photochemical reactions, precursor

20 emissions may continue to react and form O₃ downwind of emissions sources, thus the highest 21 O₃ concentrations are often found downwind of the highest concentrations of precursor 22 emissions near the urban population center. There were some instances where the highest 23 monitor occurred outside of the CBSA, but within the exposure area, which was designed to 24 always include the monitor associated with the area-wide design value. For example, in Los 25 Angeles, the CBSA includes Los Angeles and Orange counties, but the highest O₃ concentrations 26 are typically measured further downwind in Riverside and San Bernardino counties. Thus, the 27 values reported in Table 4-2 may not match the values shown in Figure 4-4. 28

- 20
- 29
- 30
- 31

⁸ Core Based Statistical Areas (CBSAs) are used by the Office of Management and Budget (OMB) to group U.S. counties into urbanized areas. These groupings are updated by OMB every 5 years. The CBSAs used in the epidemiology based risk assessment are based on the OMB deliniations from 2008. For more information see: http://www.whitehouse.gov/sites/default/files/omb/assets/bulletins/b10-02.pdf

					2009
		# of O ₃	Population	2007 4 th high	4 th high
Area Name	# of Counties	Monitors	(2010)	(ppb)	(ppb)
Atlanta	28	13	5,268,860	102	77
Baltimore	7	7	2,710,489	92	83
Boston	7	11	4,552,402	89	75
Cleveland	5	10	2,077,240	83	72
Denver	10	16	2,543,482	97	79
Detroit	6	8	4,296,250	93	73
Houston	10	22	5,946,800	90	91
Los Angeles	2	21	12,828,837	105	108
New York	23	22	18,897,109	94	81
Philadelphia	11	15	5,965,343	102	74
Sacramento	4	17	2,149,127	93	96
St. Louis	16	17	2,812,896	94	74

1Table 4-2Monitor and Area Information for the 12 Urban Case Study Areas in the2Epidemiology Based Risk Assessment

3

4 4.3.2 Recent Air Quality

5 6

The sections below summarize the recent air quality data input into the epidemiological study-based risk assessment, and the exposure and controlled human exposure study-based risk assessment. Additional details on these inputs are provided in Appendix 4-A.

8 9

7

4.3.2.1 Exposure Modeling and Controlled Human Exposure Study Based Risk Assessment

10 As discussed in more detail in Chapter 5, the REA uses the Air Pollutants Exposure

11 (APEX) model (U.S. EPA, 2012a, b) to simulate exposure and to estimate lung function

12 decrements based on application of results of controlled human exposure studies to populations

13 in the 15 urban case study areas. The APEX model uses spatial fields of hourly O_3

14 concentrations at each census tract within an urban area to simulate exposure. In the first draft

15 REA, these hourly spatial fields were generated for four urban areas using the concentrations

16 from the nearest neighboring O₃ monitor. In this draft, we use Voronoi Neighbor Averaging

17 (VNA) (Gold, 1997; Chen et al, 2004) to estimate hourly O_3 concentrations at each census tract

18 in all 15 urban case study areas, for recent measured air quality, air quality meeting the existing

19 standard of 75 ppb, and air quality meeting potential alternative standards. The VNA fields were

estimated using ambient hourly O₃ concentrations from monitors in each urban area, as well as
monitors within a 50 km buffer region around the boundaries of each area. Additional details on
the procedure used to generate the VNA fields, and a technical justification for the change from
nearest neighbor fields to VNA fields are included in Appendix 4-A.
Figure 4-5 shows county-level maps of the 15 urban case study areas. Counties colored

pink indicate the study area boundaries used in the Zanobetti & Schwartz (2008) and/or Smith et
 al (2009b) studies⁹, where applicable. Counties colored gray indicate additional counties within

9 the CBSA boundaries, and counties colored peach indicate any additional counties included in

- 10 the exposure and lung function risk assessments. The X's indicate locations of the O₃ monitors
- 11 used in the risk and exposure assessments, including those within the 50 km buffer region used
- 12 to create the VNA fields.

⁹ The Zanobetti and Schwartz (2008) and Smith et al (2009) study area boundaries were identical for 6 of the 12 urban case study areas, and had at least one county in common for all 12 urban case study areas. The 'Epidemiology Study Area' labels in figures 4-5 refer to counties included in either of these two studies.



Figure 4-5a Maps of the 5 Eastern U.S. Urban Case Study Areas Including O₃ Monitor
 Locations

4-11



Figure 4-5b Maps of the 5 Central U.S. Urban Case Study Areas Including O₃ Monitor
 Locations

4-12



Figure 4-5c Maps of the 5 Western U.S. Urban Case Study Areas Including O₃ Monitor
 Locations

4.3.2.2 Epidemiology Based Risk Assessment

1	We input O_3 air quality concentration data for the epidemiology-based risk analyses into
2	the environmental Benefits Mapping and Analysis Program Community Edition (BenMAP-CE)
3	(U.S. EPA, 2013) for assessment. We used BenMAP to analyze four different daily O_3 metrics in
4	12 of the 15 urban case study areas, which were the basis for concentration-response
5	relationships derived in various epidemiology studies:
6	1. Daily maximum 1-hour concentration
7	2. Daily maximum 8-hour concentration
8	3. Daytime 8-hour average concentration (10:00AM to 6:00PM)
9	4. Daily 24-hour average concentration
10	The air quality monitoring data used in BenMAP were daily time-series of "composite
11	monitor" values for each of the 12 urban areas for years 2007 and 2009, which were chosen to
12	represent years with high and low O ₃ concentrations, respectively. The composite monitor values
13	were calculated by first averaging the hourly O3 concentrations for all monitors within the area-
14	of-interest (resulting in a single hourly time-series for each urban area), then calculating the four
15	daily metrics listed above. More details on the composite monitor value calculations and a
16	presentation of the resulting concentrations can be found in Appendices 4-A and 4-D,
17	respectively.
10	

18

4.3.3 Air Quality Adjustments for "Just Meeting" Existing and Potential Alternative O₃ Standards

The focus of the risk and exposure assessments is the evaluation of risks and exposures after just meeting existing and alternative standards, and the change in risk between just meeting existing standards and just meeting alternative standards. These evaluations require estimation of the change in hourly O₃ concentrations that may occur in each urban area when "just meeting" the existing and potential alternative O₃ standards.

26 The first draft REA and the previous O₃ NAAQS review used the "quadratic rollback" 27 method to adjust ambient O_3 concentrations to simulate just meeting existing and alternative 28 standards (U.S. EPA, 2007; Wells et al., 2012). Although the quadratic rollback method replicates 29 historical patterns of air quality changes better than some alternative methods (e.g. simply 30 shaving peak concentrations off at the NAAQS level and the proportional rollback technique), its 31 implementation relies on a statistical relationship instead of on a mechanistic characterization of 32 physical and chemical processes in the atmosphere. Because of its construct as a statistical fit to 33 measured O₃ values, the quadratic rollback technique cannot capture spatial and temporal 34 heterogeneity in O_3 response and also cannot account for nonlinear atmospheric chemistry that

causes increases in O₃ during some hours and in some locations as a result of emissions
 reductions under some circumstances.

3 Photochemical grid models are better able to simulate these phenomena and therefore the 4 first draft REA proposed to replace quadratic rollback with a model-based O₃ adjustment 5 methodology and presented a test case for Atlanta and Detroit using modeling for July/August 6 2005 (Simon et al., 2012). The section below summarizes the methodology applied in this 7 second draft REA to adjust air quality for attainment of existing and alternative standards. This 8 new methodology applies Higher-Order Decoupled Direct Method (HDDM) capabilities in the 9 Community Multi-scale Air Quality (CMAQ) model to simulate the response of O_3 10 concentrations to reductions in US anthropogenic NO_x and VOC emissions. The model 11 incorporates anthropogenic U.S., Canadian, Mexican and other international emissions, as well 12 as emissions from non-anthropogenic sources. Since sources of background O_3 are incorporated 13 explicitly in the modeling, specifying U.S. background concentrations is unnecessary. 14 Application of this approach also addresses the recommendation by the National Research 15 Council of the National Academies (NRC, 2008) to explore how emissions reductions might 16 effect temporal and spatial variations in O₃ concentrations, and to include information on how 17 NO_x versus VOC control strategies might affect risk and exposure. 18

4.3.3.1 Methods

19 The EPA has developed an HDDM-adjustment methodology to estimate hourly O_3 20 concentrations that could occur at each monitor location if urban case study areas were to meet 21 the existing and various alternative levels of the O_3 standard. An early version of this 22 methodology was proposed in the first draft REA (Simon et al., 2012). The methodology was 23 later improved and published in a peer-reviewed journal (Simon et al., 2013). The methodology 24 and its application to hourly O_3 concentrations in the urban case study areas is summarized 25 below and described in more detail in Appendix 4-D.

The HDDM-adjustment methodology uses the CMAQ photochemical model to determine monitoring site-specific response of hourly O_3 concentrations to reductions in US anthropogenic NOx and VOC emissions. These responses are then applied to ambient data to create a 5-year time-series of hourly O_3 concentrations at each monitor location which is consistent with meeting various potential levels of the O_3 NAAQS for the two three-year attainment periods 2006-2008 and 2008-2010. The steps are outlined in Figure 4-6 and summarized below:

Step 1: Run CMAQ simulation with HDDM to determine hourly O₃ sensitivities to NO_x
 emissions and VOC emissions for the grid cells containing monitoring sites in an urban area.

1	• Inputs: Model-ready emissions and meteorology data
2	• Outputs: O ₃ concentrations and sensitivities at locations of monitoring sites for
3	each hour in January and April-October, 2007
4	• Step 2: For each monitoring site, season, and hour of the day use linear regression to
5	relate first order sensitivities of NO _x and VOC (S_{NOx} and S_{VOC}) to modeled O ₃ and
6	second order sensitivities to NOx and VOC (S^2_{NOx} and S^2_{VOC}) to the first order
7	sensitivities.
8	• Inputs: Step 1 outputs
9	• Outputs: Functions to calculate typical sensitivities based on monitor location,
10	O_3 concentration, season, and hour of the day
11	• Step 3: For each measured hourly O ₃ value between 2006 and 2010, calculate the first
12	and second order sensitivities based on monitoring site-, season-, and hour-specific
13	functions derived in Step 2.
14	• Inputs: Step 2 outputs and hourly ambient data for 2006-2010.
15	• Outputs: Hourly O ₃ observations paired with modeled sensitivities for all hours
16	in 2006-2010 at all monitor locations
17	• Step 4: Adjust measured hourly O ₃ concentrations for incrementally increasing levels of
18	emissions reductions using assigned sensitivities and then recalculate design values until
19	an emissions reduction level is reached at which all monitors in an urban area are below
20	the existing and potential alternative levels of the standard.
21	• Inputs: Step 3 outputs
22	• Outputs: Adjusted hourly O ₃ values for 2006-2010 at monitor locations to show
23	compliance with the existing and potential alternative standard levels based on the
24	three year average of the 4^{th} highest 8-hour daily max O_3 value. For each
25	standard, two sets of data are created: 2006-2008 and 2008-2010. Because the
26	emissions reductions used to attain standards in the two time periods might be
27	different, adjusted 2008 O_3 values are different for the two sets of data.
28	



Figure 4-6 Flowchart of HDDM adjustment methodology to inform risk and exposure assessments.

1

5 We chose to adjust air quality for just meeting the existing and alternative standards by 6 decreasing U.S. anthropogenic emissions of NOx and VOC throughout the U.S. For the purpose 7 of this analysis we used the Community Multiscale Air Quality (CMAQ) model version 4.7.1 8 equipped with HDDM to simulate 8 months in 2007 (January and April-October). This time 9 period was chosen to cover the full O₃ season and also includes at least one month from each 10 season of the year. A full description of the model inputs, model set-up, and operational model 11 evaluation against ambient data is available in Appendix 4-B. Sensitivities derived from the 12 2007 model simulation were applied to the two 3-year periods of ambient data (2006-2008 and 13 2008-2010) described in section 4.3.1.1. By applying equal proportional decreases in emissions 14 throughout the U.S., we were able to estimate how O₃ would respond to changes in ambient NOx 15 and VOC concentrations without simulating a specific control strategy. The model was set up to 16 track response in hourly O_3 concentrations to these across-the-board changes in US 17 anthropogenic NOx and VOC emissions. In choosing to apply across the board reductions 18 throughout the modeling domain, we recognize that not all emissions across the domain 19 contribute equally to nonattainment in each urban area. However, by decreasing emissions
1 across the domain, we allow for the possibility of contribution from both regional and local 2 emissions sources to nonattainment and to the overall distribution of O_3 concentrations in urban 3 areas. The modeling included sources which contribute to background O₃ such as biogenic 4 emissions, wildfire emissions, and transport of O₃ and its precursors from international source 5 regions. In addition, the HDDM tool was set-up to specifically calculate the changes in O₃ that 6 would occur from changes in US anthropogenic emissions alone, yet to account for the effects of 7 background sources on this response. Consequently, it is not necessary to set a "floor" 8 background O_3 concentration as was done for quadratic rollback because background sources are 9 explicitly accounted for in the model estimates of O₃ response to US anthropogenic emissions.

10 As described in more detail in Appendix 4-D, the HDDM adjustment methodology 11 estimates hourly O₃ concentrations that would be associated with attaining a targeted level of the 12 standard either though reductions in US anthropogenic NOx emissions alone or through 13 reductions of both US anthropogenic NOx and VOC emissions in equal percentages. Because 14 the combined NOx/VOC cuts are constrained to equal percentage cuts of both precursors, this is not an optimized NOx/VOC control scenario but rather a sensitivity analysis to characterize the 15 16 range of results that could be obtained with alternate assumptions. In most of the urban areas, 17 although the NOx/VOC scenario affected O_3 response on some days, it did not affect O_3 18 response at the highest design value (or controlling) monitor in such a way to reduce the total 19 required emissions cuts. However, for the two cities of Chicago and Denver, the NOx/VOC 20 scenarios allowed for lower percentage emissions cuts (applied to both NOx and VOC) to reach 21 targeted standard levels than the NOx only scenario. Because of this, the core analyses presented 22 in Chapters 5, 6, and 7 were based on the NOx only assumption for all cities except for Chicago 23 and Denver which used the NOx/VOC equal percentage reduction assumption. Sensitivity 24 analyses were performed to compare the NOx only and the NOx/VOC cases in 9 cities: Denver, 25 Detroit, Houston, Los Angeles, New York, Philadelphia, and Sacramento. The effects of these 26 sensitivity analyses on air quality and on the epidemiology-based risk assessment are discussed 27 in more detail in Appendix 4-D and Chapter 7, respectively.

28 For New York and Los Angeles it should also be noted that a somewhat different 29 approach was used for the HDDM-adjustment application. The HDDM adjustment methodology 30 produces estimates of hourly O₃ concentrations with standard error bounds for every potential 31 emission reduction scenario. Uncertainties in the application of the methodology to very large emissions perturbations along with the fact that the mean estimate does not capture the 32 33 variability in modeled responses on similar days resulted in the inability of this methodology to 34 estimate O₃ distributions in these two cities which would meet lower alternative standard levels 35 (65 ppb for New York, 60 ppb for Los Angeles). This does not indicate that these two areas 36 would not be able to meet these lower standard levels in reality, but simply reveals the

limitations of this adjustment methodology. Consequently for these two cities, we used the 95th percent confidence interval lower bound estimate of hourly O₃ concentrations to capture a scenario in which these cities could meet lower standard levels based on the range of responses in O₃ concentrations to emissions reduction predicted by the model for each city (See Appendix 4-D for more details). Estimates of risk for these two cities for these alternative standards will be significantly more uncertain, reflecting the use of the lower bound O₃ predictions.

4.3.3.2 Resulting Air Quality

8 The HDDM adjustment technique tended to have several effects on the distribution of air 9 quality values. First, adjusted hourly O_3 concentrations at night and during the morning rush-10 hour tended to be higher than the recent observed concentrations (additional details are provided 11 in Appendix 4-D). The CMAQ model predicts that, in general, these times have NOx titration 12 conditions meaning that a reduction in NOx causes an increase in O₃ concentrations. The NOx 13 titration effect was most pronounced in urban core areas which have higher volume of mobile 14 source NOx emissions from vehicles than do the surrounding areas. Response of daytime 15 concentrations was more varied. In general, O_3 tended to increase on low days and decrease on high days. However, specific monitors that were either always heavily VOC limited or always 16 17 heavily NOx limited showed consistent increases and decreases respectively regardless of 18 whether O_3 concentrations were high or low on a particular day. It should be noted that locations 19 which were heavily VOC limited tended to have much lower observed O₃ concentrations than 20 downwind areas. The tendency of the model to predict O₃ increases on lower concentration days 21 and decreases on higher concentration days also leads to more compressed O_3 distributions in the 22 HDDM adjustment cases. The variability in predicted daily O_3 concentrations decreased when 23 meeting lower standard levels. The following paragraphs summarize a comparison of O_3 24 distributions from application of the quadratic rollback and HDDM adjustment approach for a 25 case where the existing standard is estimated to be met, characterize the distribution of 26 composite monitor O₃ values at different standard levels, and provide a discussion of the spatial 27 distribution of O₃ changes in several cities. More details and figures for other case-study areas 28 are provided in Appendix 4-D. 29 Figures 4-7 and 4-8 show a comparison of April-October composite monitor O₃ 30 distributions for recent conditions (2006-2008) and for meeting the existing standard using the 31 quadratic rollback technique versus the HDDM adjustment methodology. The composite monitor

- 32 values in these plots are based on the monitors included in the composite monitor from the
- 33 Zanobetti and Schwartz (2008) study which was used in the 1st draft REA and do not include all
- 34 monitors in the CBSA as used in the main Chapter 7 analysis. In general, the O₃ distribution in

- 1 the HDDM adjustment case is shifted upward compared to the quadratic rollback case. The
- 2 upward shift is more pronounced in the lower parts of the O₃ distribution. In all cities displayed
- 3 in Figure 4-7, the 25^{th} percentile, median, and mean of the 8-hour daily maximum O_3
- 4 concentrations are higher in the HDDM adjustment case than the quadratic rollback. In some
- 5 cities (Sacramento and St. Louis) the 75th percentile values appear approximately equivalent in
- 6 the two cases while in other cities the 75th percentile values are slightly higher in the HDDM
- 7 adjustment case. In Houston, the very highest portion of the O_3 distribution is lower in the
- 8 HDDM adjustment case than in the quadratic rollback case but in many cities the upper parts of
- 9 the distributions for these two cases are roughly equivalent. Similar results are seen in the 2008-
- 10 2010 time period; however there are more cases during this time period where HDDM
- 11 adjustment and quadratic rollback have similar values in the upper half of the O₃ distribution. A
- 12 comparison of Figure 4-7 and 4-8 shows that there is some seasonality to this effect. The two
- 13 techniques appear to give very similar 8-hour daily maximum O₃ composite monitor
- 14 distributions during the summer months (June-August) and most of the situations with higher O₃
- 15 levels with the HDDM adjustment come from cooler, lower O₃ time periods (April, May,
- 16 September, and October). Although here we discuss composite monitor distributions based on
- 17 April-October, the risk analyses in Chapter 7 are based on the required O₃ monitoring season,
- 18 which is longer than April October for some cities. We expect that the O_3 increases shown for
- 19 spring and fall months here are also representative of the type of response in other "cool season"
- 20 months. The exceptions to this occur in Denver, Houston, New York and Los Angeles which
- 21 have higher composite monitor O₃ values from the HDDM adjustment compared to quadratic
- 22 rollback even in the summer time period.

Boston: Z & S, April-October, 2006-2008



Detroit: Z & S, April-October, 2006-2008

quadratic rollbac ddm adjustment



Baltimore: Z & S, April-October, 2006-2008



base

Atlanta: Z & S, April-October, 2006-2008





Houston: Z & S, April-October, 2006-2008



Philadelphia: Z & S, April-October, 2006-2008



base



NewYork: Z & S, April-October, 2006-2008





Sacramento: Z & S, April-October, 2006-2008

SaintLouis: Z & S, April-October, 2006-2008





Figure 4-7 Distributions of composite monitor 8-hour daily maximum O3 concentrations from ambient measurements (black), quadratic rollback (blue), and the HDDM adjustment methodology (red) for meeting the existing standard. Values are based on the Zanobetti & Schwartz study areas for April-October of 2006-2008.

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4-21







LosAngeles: Z & S, April-October, 2006-2008

8

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(ppb) 60

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Baltimore: Z & S, June-August, 2006-2008





Denver: Z & S, June-August, 2006-2008

Detroit: Z & S, June-August, 2006-2008





LosAngeles: Z & S, June-August, 2006-2008



Cleveland: Z & S, June-August, 2006-2008



Houston: Z & S, June-August, 2006-2008







Philadelphia: Z & S, June-August, 2006-2008



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SaintLouis: Z & S, June-August, 2006-2008



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Figure 4-8 3 4 5 6 2006-2008.

Distributions of composite monitor 8-hour daily maximum O₃ concentrations from ambient measurements (black), quadratic rollback (blue), and the HDDM adjustment methodology (red) for meeting the existing standard. Values are based on the Zanobetti & Schwartz study areas for June-August of

Boston: Z & S, June-August, 2006-2008

2 Figures 4-9 and 4-10 show "box-and-whisker" plots of the April-October composite 3 monitor daily maximum 8-hour O₃ concentration distributions for the 12 urban case study areas 4 evaluated in the epidemiology-based risk assessment; for recent air quality, and air quality 5 adjusted to meet the existing and potential alternative standards. Figure 4-9 shows values from 6 2007, while figure 4-10 shows values from 2009. Appendix 4-D contains additional plots 7 comparing the changes in the distribution of composite monitor values in each urban area due to 8 the air quality adjustments across varying spatial extents, season lengths, and years. In general, 9 the range of the composite monitor distributions decreased (i.e. the minimum value increased, 10 while the maximum value decreased) in all 12 urban case study areas as the air quality data were 11 adjusted to meet lower standard levels. However, the changes within the inter-quartile range of 12 these distributions (represented by the "boxes") varied in response to the model-based air quality 13 adjustments across the 12 urban areas. Three different types of responses are highlighted in the 14 boxplots for Atlanta, New York, and Houston.

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15 The Atlanta boxplots provide an example of an urban area in which all but the lowest 16 composite monitor values decreased as the air quality data was adjusted to simulate compliance 17 with progressively lower levels of the standard. The upper tail of the distribution (represented by 18 the top whisker in each boxplot) decreased more quickly than the remainder of the distribution, 19 resulting in less total variability in the composite monitor values with each progressively lower 20 standard level. This type of response was also seen Sacramento and St. Louis, and to a lesser 21 extent in Baltimore, Denver, and Philadelphia.

In New York, the boxplots showed an initial increase in the 25th percentile and median 22 23 composite monitor values when the observed O_3 concentrations were adjusted to meet the 24 existing standard. However, the median composite monitor value decreased relative to the existing standard as O₃ concentrations were adjusted to meet the 70 ppb standard, and both the 25 median and 25th percentile values decreased when air quality were further adjusted to meet the 26 27 65 ppb standard. When the air quality were adjusted to meet 65 ppb, the median and mean 28 composite monitor values were lower than under observed conditions. This type of response was 29 also observed in Cleveland, Detroit, and Los Angeles.

In Houston, the median composite monitor value also increased between observed air quality and air quality adjusted to meet the existing standards. However, the pattern in Houston differed from New York and other cities as air quality was further adjusted to reflect meeting the potential alternative standards. The median value remained relatively constant relative to the existing standard, while the 25th percentile values continued to increase. Thus, in Houston, the air quality adjustments always resulted in a median composite monitor value higher than what

- 1 was seen in the observed data. The composite monitor distributions in Boston also exhibited this
- 2 type of behavior.
- 3







Figure 4-9 Distributions of composite monitor 8-hour daily maximum values for the 12 urban case study areas in the epidemiology-based risk assessment. Plots depict values based on ambient measurements (base), and values obtained with the HDDM adjustment methodology showing attainment of 75, 70, 65 and 60 ppb standards. Values shown are based on CBSAs for April-October of 2007. Note that the HDDM adjustment technique was not able to adjust air quality to show attainment of a 60 ppb standard in New York, so no boxplot is shown for that case.





Figure 4-10 Distributions of composite monitor 8-hour daily maximum values for the 12 urban case study areas in the epidemiology-based risk assessment. Plots depict values based on ambient measurements (base), and values obtained with the HDDM adjustment methodology showing attainment of 75, 70, 65 and 60 ppb standards. Values shown are based on CBSAs for April-October of 2009. Note that Detroit air quality was meeting 75 ppb in 2008-2010, and the HDDM adjustment technique was not able to adjust air quality to show attainment of a 60 ppb standard in New York, so no boxplots are shown for those cases.

2 The exposure modeling and the clinical-based risk assessments used spatially varying 3 surfaces of hourly O₃ concentrations estimated at the centroid of each census tract within the 15 4 urban case study areas. The maps in Figures 4-11, 4-12, and 4-13 depict the spatial distributions of the 2006-2008 average 4th highest (top) and May – September mean (bottom) daily maximum 5 6 8-hour (MDA8) O₃ concentrations for 3 of the 15 urban case study areas; for observed air quality 7 (left), air quality adjusted to meet the existing standard (center), and air quality adjusted to meet 8 the 65 ppb alternative standard (right). Appendix 4-A contains additional maps of the observed 9 4th highest MDA8 and May – September mean MDA8 concentrations in all 15 urban case study 10 areas for 2006-2008 and 2008-2010. Appendix 4-D contains maps and related figures showing 11 the changes in air quality that resulted from the HDDM adjustments for just meeting the existing 12 standard, and just meeting the potential alternative standard of 65 ppb.

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These maps portray the general pattern seen in all 15 urban case study areas for the 4th 13 14 highest concentrations, which decreased when observed air quality were adjusted to meet the 15 existing standard, and continued to decrease as the air quality were further adjusted to meet the 16 various alternative standards. The May-September average values also generally decreased in 17 suburban and rural areas surrounding the urban population center in all 15 areas. However, three 18 different types of general behavior which were seen in the seasonal average values near the 19 urban population centers, which are exemplified in Figures 4-11 (Atlanta), 4-12 (New York), and 20 4-13 (Houston).

In Atlanta, the observed May - September average were nearly constant across the entire study area. The observed values decreased nearly uniformly across the entire study area when observed air quality was adjusted to meet the existing standard, and continued to do so when air quality was further adjusted to meet the alternative standard of 65 ppb. The magnitudes of these decreases were slightly larger in suburban and rural areas than near the urban population center. This type of behavior was also seen in Sacramento and Washington, D.C.

27 In New York, the observed May – September average values were lower near the urban 28 population center than in the surrounding suburban areas. When the observed air quality was 29 adjusted to meet the existing standard, the seasonal average values increased near the urban 30 population center and decreased in the suburban areas, so that the spatial pattern was reversed. 31 When air quality was further adjusted to meet the 65 ppb alternative standard, large area-wide 32 decreases in the seasonal average values were seen relative to the existing standard. While New 33 York represents one of the most extreme examples, similar behavior was observed in 7 other 34 urban areas: Baltimore, Cleveland, Dallas, Detroit, Los Angeles, Philadelphia, and St. Louis. 35 Houston started out in a similar fashion as New York. The observed May – September 36 average concentrations were lower near the urban population center than in the surrounding

- 1 areas, and a similar pattern of increasing and decreasing seasonal average values occurred when
- 2 observed air quality was adjusted to meet the existing standard. However, unlike New York, the
- 3 seasonal average values near the Houston city center remained nearly constant relative to the
- 4 existing standard when air quality were further adjusted to meet the 65 ppb standard. Boston,
- 5 Chicago, and Denver exhibited this same type of behavior.
- 6



Atlanta, 2006 - 2008

- 7 8
- Figure 4-11 Maps showing the 4th highest (top) and May-September average (bottom) daily maximum 8-hour O₃ concentrations in Atlanta based on 2006-2008 ambient measurements (left), HDDM adjustment to meet the existing standard (center), and HDDM adjustment to meet the alternative standard of 65 ppb (right). Squares represent measured values at monitor locations; circles represent VNA estimates at census tract centroids.
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Figure 4-12 Maps showing the 4th highest (top) and May-September average (bottom) daily maximum 8-hour O₃ concentrations in New York based on 2006-2008 ambient measurements (left), HDDM adjustment to meet the existing standard (center), and HDDM adjustment to meet the alternative standard of 65 ppb (right). Squares represent measured values at monitor locations; circles represent VNA estimates at census tract centroids.



Figure 4-13Maps of 4th highest (top) and May-September average (bottom) daily
maximum 8-hour O3 concentrations in Houston for 2006-2008 ambient
measurements (left), HDDM adjustment to meet the existing standard
(center), and HDDM adjustment to meet the alternative standard of 65 ppb
(right). Squares represent measured values at monitor locations; circles
represent VNA estimates at census tract centroids.

10 4.4 OVERVIEW OF NATIONAL-SCALE AIR QUALITY INPUTS

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12 The national-scale analysis, presented in Chapter 8, is focused only on evaluating the 13 total national burden of mortality risk associated with recent O₃ conditions. As such it uses a 14 different approach to characterize air quality conditions throughout the U.S. The national-scale 15 analysis employs a data fusion approach that takes advantage of the accuracy of monitor 16 observations and the comprehensive spatial information of the CMAQ modeling system to create 17 national-scale "fused" spatial surfaces of seasonal average O₃ concentrations. Measured O₃ 18 concentrations from 2006-2008 were fused with modeled concentrations from a 2007 CMAQ

1 model simulation, run for a 12 km domain covering the contiguous U.S. In the first draft of the 2 REA, the spatial surfaces were created using the enhanced Voronoi Neighbor Averaging (eVNA) 3 technique (Timin et al, 2010), using the EPA's Model Attainment Test Software (MATS; Abt 4 Associates, 2010b). In this draft, the spatial surfaces are created using EPA's Downscaler 5 software (Berrocal et al, 2012). More details on the ambient measurements, the 2007 CMAQ 6 model simulation, the Downscaler fusion technique, and a technical justification for changing 7 from eVNA to Downscaler can be found in Appendix 4-C. 8 Three national "fused" spatial surfaces were created for: 9 1) the May-September average of the 8-hour daily maximum O_3 concentrations 10 (consistent with the metric used by Smith et al. 2009); 11 2) the June-August average of the daily 10am-6pm mean O₃ concentrations (consistent 12 with the metric used by Zanobetti and Schwartz 2008); and 13 3) the April-September average of the 1-hour daily maximum O₃ concentrations 14 (consistent with the metric used by Jerrett et al 2009). 15 Figures 4-14 to 4-16 show the geographic distributions of these spatial surfaces. The 16 spatial distributions of these three surfaces are very similar, with the highest levels occurring in

17 Southern California for all three surfaces.



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Figure 4-17 shows the frequency and cumulative distributions of these three seasonal
average O₃ surfaces based on all grid cells in the 12 km CMAQ modeling domain. The
minimum, median, mean, 95th percentile, and maximum values for all three surfaces are shown
in Table 4-3, and correlation coefficients between the three metrics are given in Table 4-4.

10 The May-September average 8-hour daily maximum concentrations were most frequently 11 in the 30-60 ppb range, while the June-August average daily 10am–6pm mean concentrations 12 were more evenly distributed across a range of 20-60 ppb. The April-September average 1-hour 13 daily maximum concentrations were about 5 ppb higher on average than the May-September 14 average 8-hour daily maximum concentrations, and about 8 ppb higher on average than the June-15 August average daily 10am-6pm mean concentrations. The correlation coefficients between 16 these three metrics were all very high (R > 0.97).



1Table 4-3Summary Statistics Based on the Three Fused Seasonal Average O3 Surfaces2Based on all CMAQ 12 km Grid Cells

	May-September average	June-August average daily	April-September average
	8-hour daily maximum	10am–6pm mean	1-hour daily maximum
Statistic	concentration (ppb)	concentration (ppb)	concentration (ppb)
Minimum	21.8	14.9	26.2
Median	43.6	41.7	48.8
Mean	43.2	40.9	48.2
95 th Percentile	54.3	54.8	59.0
Maximum	76.1	80.1	84.2

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Table 4-4 Correlation Coefficients Between the Three Fused Seasonal Average O₃ Surfaces Based on all CMAQ 12 km Grid Cells

Seasonal metrics compared	Correlation coefficient	
May-September average 8-hour daily maximum vs.	0.974	
June-August average daily 10am-6pm mean	0.274	
May-September average 8-hour daily maximum vs.	0.995	
April-September average 1-hour daily maximum	0.335	
June-August average daily 10am-6pm mean vs.	0.072	
April-September average 1-hour daily maximum	0.772	

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8 These seasonal average metrics are not equivalent to the form of the existing standard, which is based on the 4th highest value rather than on the seasonal mean. Thus, the values shown 9 10 in the three fused surfaces should not be directly compared to the existing standard. Figure 4-18 11 shows comparisons between these three metrics and the 2006-2008 O₃ design values based on 12 CMAQ 12 km grid cells containing O₃ monitors, and Table 4-5 presents correlation coefficients 13 and summary statistics based on the ratios between the design values and these three metrics. 14 The design values were, on average, approximately 50% higher than the seasonal average values, 15 with substantial spatial heterogeneity, and some variation across the seasonal average metrics. 16 The April-September average 1-hour daily maximum was the most strongly correlated with the design values (R = 0.75), followed by the May-September average 8-hour daily maximum (R =17 18 0.71), and then the June-August average daily 10am-6pm mean (R = 0.69). 19



1Table 4-5Correlation Coefficients and Ratios of the 2006-2008 O3 Design Values to the22006-2008 Fused Seasonal Average O3 Levels for the CMAQ 12km Grid Cells3Containing O3 Monitors

	May-September average	June-August average	April-September average
Statistic	8-hour daily maximum	daily 10am-6pm mean	1-hour daily maximum
Correlation	0.71	0.69	0.75
Ratios			
Minimum	1.1	1.1	1.0
2.5 th Percentile	1.3	1.3	1.2
Median	1.5	1.5	1.4
Mean	1.6	1.6	1.4
97.5 Percentile	2.0	2.2	1.6
Maximum	2.4	3.0	1.9

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6 4.5 UNCERTAINITIES IN MODELING OF RESPONSES TO EMISSION 7 REDUCTIONS TO JUST MEET EXISTING AND POTENTIAL ALTERNATIVE 8 STANDARDS

9 We recognize that there are sources of uncertainty in air quality measurements and the air 10 quality estimates for each air quality scenario. These sources of uncertainty are described below 11 and in Table 4-6 which discusses qualitatively the magnitude of uncertainty and potential for 12 directional bias.

There is inherent uncertainty in all deterministic air quality models, such as CMAQ, the photochemical grid model which was used to develop the model-based O₃ adjustment methodology. Evaluations of air quality models against observed pollutant concentrations build confidence that the model performs with reasonable accuracy despite both structural and parametric uncertainties. A comprehensive model performance evaluation provided in Appendix 4-B shows generally acceptable model performance which is equivalent to or better than typical state-of-the science regional modeling simulations as summarized in Simon et al (2012). The use

20 of the Higher Order Decoupled Direct Method (HDDM) within CMAQ to estimate O₃ response

- 21 to emissions perturbations adds uncertainty to that inherent in the model itself. HDDM allows for
- 22 the approximation of O₃ concentrations under alternate emission scenarios without re-running
- 23 the model simulation with different inputs. This approximation becomes less accurate for larger
- 24 emissions perturbations. To accommodate increasing uncertainty at larger emissions
- 25 perturbations, the HDDM modeling was performed at three distinct emissions levels to allow for
- 26 a better characterization of O_3 response over the entire range of emissions levels. The accuracy

1 of the HDDM estimates can be quantified at distinct emissions levels by re-running the model

- 2 with modified emissions inputs and comparing the results. This method was applied to quantify
- 3 the accuracy of 3-step HDDM O₃ estimates for 50% and 90% NOx cut conditions for each urban
- 4 case study areas (as shown in Appendix 4-D). At 50% NOx cut conditions, HDDM using
- 5 information from these multiple simulations predicted hourly O₃ concentrations with a mean bias
- 6 and a mean error less than +/- 1 ppb in all case study areas compared to brute force model
- 7 simulations. At 90% NOx cut conditions, HDDM using information from these multiple
- 8 simulations predicted hourly O₃ concentrations with a mean bias less than +/- 3ppb and a mean
 9 error less than +/- 4 ppb in all case study areas. These small bias and error estimates show that
- 10 uncertainty due to the HDDM approximation method is small up to 90% emissions cuts.
- 11 In order to apply modeled O_3 response to ambient measurements, regressions were 12 developed which relate O₃ response to emissions perturbations with ambient O₃ concentrations 13 for every season, hour-of-the-day, and monitor location. Applying O_3 responses based on this 14 relationship adds uncertainty. Preliminary work showed that the relationships developed with 15 these regressions were generally statistically significant for most season, hour-of-the-day, and monitor location combinations for 2005 modeling in Detroit and Charlotte (Simon et al, 2012). 16 17 Statistical significance was not evaluated for each regression in this analysis since there were 18 over 460,000 regressions created (322 monitors \times 5 sensitivity coefficients \times 3 emissions levels 19 \times 4 seasons \times 24 hours = 463,680 regressions). Statistics can quantify the goodness of fit for the 20 modeled relationships and can quantify the uncertainty in response at any given O_3 concentration 21 based on variability in model results at that portion of the distribution for each regression. The 22 regression model provided both a central tendency and a standard error value for O₃ response at 23 each measured hourly O₃ concentration. The base analysis in all case study areas except New 24 York and Los Angeles used the central tendency which will inherently dampen some of the 25 variability in O_3 response. The standard error of each sensitivity coefficient was propagated 26 through the calculation of predicted O₃ concentrations at various standard levels. These standard errors reflect the amount of variability that is lost due to the use of a central tendency. Since 27 28 emissions reductions increased for lower standard levels the standard errors were larger for adjustments to lower standards. Mean (95th percentile) standard errors for the 75 ppb adjustment 29 case ranged from 0.13 (0.26) to 1.18 (2.87) ppb in the 15 case study areas. Mean (95th percentile) 30 31 standard errors for the 65 ppb adjustment case ranged from 0.54 (1.07) to 1.39 (2.98) ppb. The 32 largest standard errors occurred in Los Angeles and New York due to the large emissions 33 reductions applied in these cases. In cases where the use of the central tendency of response 34 reduced the total estimated emissions reductions required to achieve a given standard level, in 35 general we expect that the benefits of reducing high O_3 concentrations and the disbenefits of 36 increasing low O_3 would both be underestimated. For the exposure assessment which estimates

1 health outcomes that occur at O₃ concentrations above 60, this would lead to an underestimation 2 of risks. For the epidemiology-based risk assessment which is effected by the entire range of O_3 3 concentrations, the impact is undetermined since changes at both ends of the O₃ distribution in 4 opposite directions would affect the results. The opposite would be true in cases where the use of 5 the central tendency of response increased the total estimated emissions reductions required to 6 achieve a given standard. However, given the small standard error values even in the case study 7 areas with the greatest uncertainty (i.e. less than 1.5 ppb mean standard error), this source of 8 uncertainty is not expected to substantially impact results.

9 Relationships between O₃ response and hourly O₃ concentration were developed based on 10 8 months of modeling: January and April-October 2007. These relationships were applied to 11 ambient data from 2006-2010. Some locations monitor for months not included in this modeling 12 (i.e., February, March, November, and December) while others do not. Seasonal relationships 13 were developed between O_3 response to emissions reductions and O_3 concentration. Summer was 14 the only season for which modeling data was created for all months (June, July, August). The 15 winter relationships were developed based on January modeling, the spring relationships were developed based on April/May modeling, and the autumn relationships were developed based on 16 17 September/October modeling. The reduction in data points (31 or 61 instead of ~90) increases 18 uncertainty in the statistical fit for these seasons. In addition, the modeling generally showed 19 more O_3 disbenefits to NOx decreases in cooler months. So applying April/May relationships to 20 March and September/October relationships to November could potentially underestimate O_3 21 increases that would happen in those two months in the five case study areas which measure O_3 22 during March and/or November: Dallas, Denver, Houston, Los Angeles, and Sacramento. The 23 eight months that were modeled capture a variety of meteorological conditions. In cases where 24 other years have more frequent occurances of certain types of meteorological conditions, the 25 regressions should be able to account for this. For instance, if a monitor only had 2-3 high O_3 26 days associated with sunny, high pressure conditions in the 2007 modeling but had 30-40 of 27 those days in another year, the regression may be more uncertain at those high O_3 values but 28 should still be able to capture the central tendency which can be applied to the more frequent 29 occurances in other years. If, on the other hand, the meteorology/ O_3 conditions in another year 30 were completely outside the range of conditions captured in the model, then the regression based 31 on modeled conditions might not be able to capture those conditions. Finally, if emissions 32 change drastically between the modeled period and the time of the ambient data measurements 33 this could also change the relationship between O₃ response and O₃ concentrations. The 34 regressions derived from the 2007 modeling period are only applied to measurements made 35 within 3 years of the modeled time period. Although some emissions changes did occur over this

time period, we believe it is still reasonable to apply 2007 modeling to this relatively small
 window of measurements which occurs before and after the modeling.

3 O₃ response is modeled for across-the-board reductions in U.S. anthropogenic NOx (and 4 VOC). These across-the-board cuts do not reflect actual emissions control strategies. The form, 5 locations, and timing of emissions reductions that would be undertaken to meet various levels of 6 the O_3 standard are unknown. The across-the-board emissions reductions bring levels down 7 uniformly across time and space to show how O₃ would respond to changes in ambient levels of 8 precursor species but do not reflect spatial and temporal heterogeneity that may occur in local 9 and regional emissions reductions. In cases where VOC reductions were modeled, equal 10 percentage NOx and VOC reductions were applied in the adjustment methodology. Regional 11 NOx reductions are likely to be the primary means used to reduce high O₃ concentrations at DV 12 monitors. In limited cases, VOC emissions reductions may also help lower high O₃ 13 concentrations at these locations. In actual control strategies, NOx and VOC reductions may be 14 applied in combination but are unlikely to be applied in equal percentages. The available 15 modeling constrained the NOx/VOC case to this type of control scenario. The across-the-board 16 cuts and the equal percentage NOx and VOC reductions scenario does not optimize the lowest 17 cost or least total emissions combinations as state and local agencies will likely attempt to 18 achieve.

		Potential influ	ience of		
		uncertainty on risk		Knowledge-	
		estimates		Base	Comments (KB: knowledge base, INF: influence
Source	Description	Direction	Magnitude	uncertainty*	of uncertainty on risk estimates)
A. O ₃ measurements	O_3 concentrations measured by ambient monitoring instruments have inherent uncertainties associated with them. Additional uncertainties due to other factors may include: - monitoring network locations - O_3 monitoring seasons - monitor malfunctions - wildfire and smoke impacts - interpolation of missing data	Both	Low	Low	KB: O ₃ measurements are assumed to be accurate to within $\frac{1}{2}$ of the instrument's Method Detection Limit (MDL), which is 2.5 ppb for most instruments. EPA requires that routine quality assurance checks are performed on all instruments, and that all data reported to AQS are certified by both the monitoring agency and the corresponding EPA regional office. The CASTNET monitoring data were subject to their own set of QA requirements, and these data are generally believed to be of comparable quality to the data stored in AQS. KB: Monitor malfunctions sometimes occur causing periods of missing data or poor data quality. Monitoring data affected by malfunctions are usually flagged by the monitoring agency and removed from AQS. In addition, the AQS database managers run several routines to identify suspicious data for potential removal. KB: There is a known tendency for smoke produced from wildfires to cause interference in O ₃ instruments. Measurements collected by O ₃ analyzers were reported to be biased high by 5.1– 6.6 ppb per 100 µg/m ³ of PM2.5 from wildfire smoke ,EPA, 2007). However, smoke concentrations high enough to cause significant interferences are infrequent and the overall impact is believed to be minimal.

1 Table 4-6 Summary of Qualitative Uncertainty Analysis of Key Air Quality Elements in the O₃ NAAQS Risk Assessment

		Potential influence of			
		uncertainty on risk		Knowledge-	
		estimates		Base	Comments (KB: knowledge base, INF: influence
Source	Description	Direction	Magnitude	uncertainty*	of uncertainty on risk estimates)
Source	Description	Direction	Magnitude	uncertainty*	of uncertainty on risk estimates) KB: Missing intervals of 1 or 2 hours in the measurement data were interpolated, which may cause some additional uncertainty. However, due to the short length of the interpolation periods, and the tendency for these periods to occur at night when O ₃ concentrations are low, the overall impact is believed to be minimal. INF: EPA's current O ₃ monitoring network requirements have an urban focus. Rural areas where O ₃ concentrations are lower tend to be under-represented by the current monitoring network. The network requirements also state that at least one monitor within each urban area must be sited to capture the highest O ₃ concentrations in that area, which may cause some bias toward higher measured concentrations. INF: Each state has a required O ₃ monitoring season which varies in length from May – September to year-round. Some states turn their O ₃ monitors off during months outside of the required season, while others leave them on. This can cause discrepancies in the amount of data available, especially in months outside of the
					required monitoring season. The risk estimates attempt to minimize these impacts by focusing only on months where O_3 monitoring is required.

		Potential influence of			
		uncertainty on risk		Knowledge-	
		estimates		Base	Comments (KB: knowledge base, INF: influence
Source	Description	Direction	Magnitude	uncertainty*	of uncertainty on risk estimates)
B. Veronoi Neighbor Averaging (VNA) spatial fields	VNA is a spatial interpolation technique used to estimate O_3 concentrations in unmonitored areas, which has inherent uncertainty	Both	Low- Medium	Low- Medium	KB: VNA interpolates monitored hourly O ₃ concentrations to provide estimates of O ₃ exposure at each census tract in the 15 urban areas. The VNA estimates are weighted based on distance from neighboring monitoring sites, thus the amount of uncertainty tends to increase with distance from the monitoring sites. KB: The 15 urban areas each had fairly dense monitoring networks which were generally sufficient to capture spatial gradients in O ₃ concentrations. The use of hourly data to create the VNA fields instead of daily or other aggregates also served to reduce uncertainty by better capturing relationships in the diurnal patterns between O ₃ monitors.
C.CMAQ modeling	Model predictions from CMAQ, like all deterministic photochemical models, have both parametric and structural uncertainty associated with them	Both	Low- Medium	Low- Medium	KB: Structural uncertainties are uncertainties in the representation of physical and chemical processes in the model. These include: choice of chemical mechanism used to characterize reactions in the atmosphere, choice of land surface model and choice of planetary boundary layer model. KB: Parametric uncertainties include uncertainties in model inputs (hourly meteorological fields, hourly 3-D gridded emissions, initial conditions, and boundary conditions) KB: Uncertainties due to initial conditions are minimized by using a 10 day ramp-up period

		Potential influ	ience of		
		uncertainty or	n risk	Knowledge-	
		estimates		Base	Comments (KB: knowledge base, INF: influence
Source	Description	Direction	Magnitude	uncertainty*	of uncertainty on risk estimates)
					from which model results are not used.
					KB: Evaluations of models against
					observed pollutant concentrations build
					confidence that the model performs with
					reasonable accuracy despite the uncertainties listed
					above. A comprehensive model evaluation
					provided in Appendix 4-B shows generally
					acceptable model performance which is equivalent
					or better than typical state-of-the science regional
					modeling simulations as summarized in Simon et
					al (2012). However, both under-estimations and
					over-estimations do occur at some times and
					locations. Generally the largest mean biases occur
					on low O_3 days during the summer season. In
					addition, the model did not fully capture rare
					wintertime high O ₃ events occurring in the
					Western U.S.
	HDDM allows for the				KB: To accommodate increasing
	approximation of O_3				uncertainty at larger emissions perturbations, the
	concentrations under				HDDM modeling was performed at three distinct
	alternate emissions				emissions levels to allow for a better
D Higher Order	scenarios without re-				characterization of O ₃ response over the entire
Decoupled Direct	running the model	Both	Low-	Low-	range of emissions levels. The replication of brute
Method (HDDM)	simulation multiple times	Dom	Medium	Medium	force hourly O ₃ concentration model results by the
	using different emissions				HDDM approximation was quantified for 50%
	inputs. This				and 90% NOx cut conditions for each urban case
	approximation becomes				study areas (as shown in Appendix 4-D). At 50%
	less accurate for larger				NOx cut conditions, HDDM using information
	emissions perturbations				from these multiple simulations predicted hourly

		Potential influ	ience of		
		uncertainty or	n risk	Knowledge-	
		estimates		Base	Comments (KB: knowledge base, INF: influence
Source	Description	Direction	Magnitude	uncertainty*	of uncertainty on risk estimates)
	especially under nonlinear chemistry conditions.				O_3 concentrations with a mean bias and a mean error less than +/- 1 ppb in all urban case study areas compared to brute force model simulations. At 90% NOx cut conditions, HDDM using information from these multiple simulations predicted hourly O_2 concentrations with a mean
					bias less than $+/-$ 3ppb and a mean error less than $+/-4$ ppb in all urban case study areas.
E. Application of HDDM sensitivities to ambient data	In order to apply modeled sensitivities to ambient measurements, regressions were developed which relate O ₃ response to emissions perturbations with ambient O ₃ concentrations for every season, hour-of- the-day and monitor location. Applying O ₃ responses based on this relationship adds uncertainty.	Both	Medium	Medium	KB: Preliminary work showed that the relationships developed with these regressions were generally statistically significant for most season, hour-of-the-day, and monitor location combinations for 2005 modeling in Detroit and Charlotte. Statistical significance was not evaluated for each regression in this analysis since there were over 460,000 regressions created (322 monitors \times 5 sensitivity coefficients \times 3 emissions levels \times 4 seasons \times 24 hours = 463,680 regressions). Statistics can quantify the goodness of fit for the modeled relationships and can quantify the uncertainty in response at any given O ₃ concentration based on variability in model results at that portion of the distribution for each regression. However it is not possible to quantify the applicability of this modeled relationship to the actual atmosphere. KB: The regression model provided both a central tendency and a standard error value for O ₃ response at each measured hourly O ₃

		Potential influence of			
		uncertainty or	n risk	Knowledge-	
		estimates		Base	Comments (KB: knowledge base, INF: influence
Source	Description	Direction	Magnitude	uncertainty*	of uncertainty on risk estimates)
					concentration. The base analysis used the central
					tendency which will inherently dampen some of
					the variability in O_3 response. The standard error
					of each sensitivity coefficient was propagated
					through the calculation of predicted O ₃
					concentrations at various standard levels. These
					standard errors reflect the amount of variability
					that is lost due to the use of a central tendency.
					Since emissions reductions increased for lower
					standard levels the standard errors were larger for
					adjustments to lower standards. Mean (95 th
					percentile) standard errors for the 75 ppb
					adjustment case ranged from 0.13 (0.26) to 1.18
					(2.87) ppb in the 15 case study areas. Mean (95^{th})
					percentile) standard errors for the 65 ppb
					adjustment case ranged from 0.54 (1.07) to 1.39
					(2.98) ppb. The largest standard errors occurred in
					Los Angeles and New York.
					INF: The NOx emissions reductions
					resulted in both increases and decreases in O_3
					depending on the time and location. In cases
					where the use of the central tendency of response
					reduced the total estimated emissions reductions
					required to achieve a given standard level, in
					general we expect that the benefits of reducing
					high O_3 concentrations and the disbenefits of
					increasing low O_3 would be underestimated. For
					the exposure assessment which estimates health
					outcomes that occur at O_3 concentrations above

		Potential influ	ience of		
		uncertainty or	n risk	Knowledge-	
		estimates		Base	Comments (KB: knowledge base, INF: influence
Source	Description	Direction	Magnitude	uncertaintv*	of uncertainty on risk estimates)
					60, this would lead to an underestimation of risks. For the epidemiology-based risk assessment which is effected by the entire range of O_3 concentrations, the impact is undetermined since changes at both ends of the O_3 distribution in opposite directions would affect the results. The opposite would be true in cases where the use of the central tendency of response increased the total estimated emissions reductions required to achieve a given standard
F. Applying modeled sensitivities to un- modeled time periods	Relationships between O ₃ response and hourly O ₃ concentration were developed based on 8 months of modeling: January and April-October 2007. These relationships were applied to ambient data from 2006-2010. Some locations monitor for months not included in this modeling (February, March, November, and December) while others do not.	Both	Low- Medium	Low- Medium	KB: The eight months that were modeled capture a variety of meteorological conditions. In cases where other years have more frequent occurances of certain types of conditions, the regressions should be able to account for this. For instance, if a monitor only had 2-3 high O ₃ days associated with sunny, high pressure conditions in the 2007 modeling but had 30-40 of those days in another year, the regression may be more uncertain at those high O ₃ values but should still be able to capture the central tendency which can be applied to the more frequent occurances in other years. If, on the other hand, the meteorology/O ₃ conditions in another year were completely outside the range of conditions captured in the model, then the regression based on modeled conditions might not be able to capture those conditions. KB: If emissions change drastically

		Potential influence of			
		uncertainty or	n risk	Knowledge-	
		estimates		Base	Comments (KB: knowledge base, INF: influence
Source	Description	Direction	Magnitude	uncertainty*	of uncertainty on risk estimates)
					between the modeled period and the time of the
					ambient data measurements this could also change
					the relationship between O ₃ response and O ₃
					concentrations. The regressions derived from the
					2007 modeling period are only applied to
					measurements made within 3 years of the modeled
					time period. Although some emissions changes did
					occur over this time period, we believe it is still
					reasonable to apply 2007 modeling to this
					relatively small window of measurements which
					occurs before and after the modeling.
					INF: Seasonal relationships were
					developed between O ₃ response to emissions
					reductions and O ₃ concentration. Summer was the
					only season for which modeling data was created
					for all months (June, July, August). The winter
					relationships were developed based on January
					modeling, the spring relationships were developed
					based on April/May modeling, and the autumn
					relationships were developed based on
					September/October modeling. The reduction in
					data points (31 or 61 instead of ~90) increases
					uncertainty in the statistical fit for these months. In
					addition, the modeling generally showed more O_3
					disbenefits to NOx decreases in cooler months. So
					applying April/May relationships to March and
					September/October relationships to November
					could potentially underestimate O ₃ increases that
					would happen in those two months in the five

		Potential influence of uncertainty on risk			
				Knowledge-	
		estimates		Base	Comments (KB: knowledge base, INF: influence
Source	Description	Direction	Magnitude	uncertainty*	of uncertainty on risk estimates)
					urban case study areas which measure O ₃ during
					March and/or November: Dallas, Denver,
					Houston, Los Angeles, and Sacramento.
	O ₃ response is modeled				KB: The form, locations, and timing of emissions
	for across-the-board				reductions that would be undertaken to meet
G Assumptions	reductions in U.S.				various levels of the O_3 standard are unknown.
of across-the-	anthropogenic NOx (and		Low-	Low-	The across-the-board emissions reductions bring
board emissions	VOC). These across-the-	Both	Medium	Medium	levels down uniformly across time and space to
reductions	board cuts do not reflect		Wiedrum	Wearum	show how O ₃ would respond to changes in
reductions	actual emissions control strategies.				ambient levels of precursor species but do not
					reflect spatial and temporal heterogeneity that may
					occur in local and regional emissions reductions.
					KB: NOx reductions are likely to be the primary
	In cases where VOC reductions were modeled, equal percentage NOx and VOC reductions were applied in the adjustment				means used to reduce high O_3 concentrations at
					DV monitors. In limited cases, VOC emissions
					reductions may also help lower high O ₃
H Assumption of					concentrations at these locations. NOx and VOC
equal percentage			Low		reductions may be applied in combination but are
NOx and VOC		Both	Medium	Medium	unlikely to be applied in equal percentages. The
reductions			Wiedrum		available modeling constrained the NOx/VOC
reductions	methodology				case to this unrealistic scenario. The equal
	memodology.				percentage NOx and VOC reductions scenario
					does not optimize the lowest cost or least total
					emissions combinations as state and local agencies
					will likely attempt to achieve.
	Downscaler combines				KB: Downscaler combines modeled and
I Downscaler	monitored and modeled	Both	Low-	Low-	monitored concentrations to provide estimates of
	concentrations to produce	Dom	Medium	Medium	O ₃ concentrations in unmonitored areas while
	a "fused" air quality				correcting model biases near monitors. The cross-

		Potential influence of			
		uncertainty on risk		Knowledge-	
		estimates		Base	Comments (KB: knowledge base, INF: influence
Source	Description	Direction	Magnitude	uncertainty*	of uncertainty on risk estimates)
	surface. Uncertainties may				validation analysis in Appendix 4-A shows that
	occur in sparsely				Downscaler generally gives more accurate
	monitored regions, or in				estimates of air quality in monitored locations than
	urban areas with dense				either the monitored or modeled values alone.
	monitoring networks and				However, it is not possible to quantify the
	large spatial gradients.				uncertainty associated with the estimates in
					unmonitored locations.
					KB: The air quality surfaces modeled by
					Downscaler for the national-scale risk assessment
					were seasonal average concentrations, which tend
					to have smaller spatial gradients than other metrics
					such as peak concentrations, and thus less
					uncertainty.
					INF: The cross-validation analysis in Appendix 4-
					A also shows that Downscaler tends to over-
					estimate low concentrations and under-estimate
					high concentrations. The mean bias in the
					estimates in monitored locations is nearly zero, but
					monitor locations are often chosen to capture the
					highest concentrations, thus there might be some
					bias towards higher concentrations in umonitored
					areas.

* Refers to the degree of uncertainty associated with our understanding of the phenomenon, in the context of assessing and characterizing its uncertainty. Sources classified as having a "low" impact would not be expected to impact the interpretation of risk estimates in the context of the O₃ NAAQS review; sources classified as having a "medium" impact have the potential to change the interpretation; and sources classified as "high" are likely to influence the interpretation of risk in the context of the O₃ NAAQS review.

1 4.6 REFERENCES

2 Abt Associates, Inc. 2010a. "Environmental Benefits and Mapping Program (Version 4.0)." 3 Bethesda, MD. Prepared for U.S. Environmental Protection Agency Office of Air Quality Planning and Standards. Research Triangle Park, NC. Available on the Internet at 4 5 <http://www.epa.gov/air/benmap>. 6 Abt Associates, Inc. 2010b. "Model Attainment Test Software (Version 2)." Bethesda, MD. 7 Prepared for the U.S. Environmental Protection Agency Office of Air Quality Planning 8 and Standards. Research Triangle Park, NC. Available on the Internet at: 9 http://www.epa.gov/scram001/modelingapps.mats.htm. 10 Bell, M.L.; A. McDermott; S.L. Zeger; J.M. Samet and F. Dominici. 2004. "Ozone and Shortterm Mortality in 95 U.S. Urban Communities," 1987-2000. JAMA, 292:2372-2378. 11 12 Berrocal, V.J.; A. E. Gelfand and D.M. Holland. 2012. "Space-Time Data Fusion Under Error in 13 Computer Model Output: An Application to Modeling Air Quality." *Biometrics*, 68(3), 14 837-848. 15 Chen, J. R.; Zhao; Z. Li. 2004. "Voronoi-based k-order Neighbor Relations for Spatial Analysis." 16 ISPRS J Photogrammetry Remote Sensing, 59(1-2), 60-72. 17 Duff, M.; R. L. Horst; T.R. Johnson, 1998. "Quadratic Rollback: A Technique to Model 18 Ambient Concentrations Due to Undefined Emission Controls." San Diego, CA: 19 Presented at the Air and Waste Management Annual Meeting, June 14-18, 1998. 20 Fann, N.; A.D. Lamson; S.C. Anenberg; K. Wesson; D. Risley; B.J. Hubbell. 2012. "Estimating 21 the National Public Health Burden Associated with Exposure to Ambient PM_{2.5} and 22 Ozone." Risk Analysis, 32:81-95. Gold, C. 1997. "Voronoi Methods in GIS," Vol. 1340. In Algorithmic Foundation of Geographic 23 24 Information Systems (Kereveld M., J. Nievergelt, T. Roos, P. Widmayer eds). Lecture 25 notes in Computer Science, Berlin: Springer-Verlag, 21-35. 26 Hall, E.; A. Eyth; S. Phillips. 2012. "Hierarchical Bayesian Model (HBM)-Derived Estimates of 27 Air Quality for 2007: Annual Report." (EPA document number EPA/600/R-12/538). 28 < http://www.epa.gov/heasd/sources/projects/CDC/AnnualReports/2007_HBM.pdf>.

1	Jerrett, M.; R.T. Burnett; C.A. Pope, III; K. Ito; G. Thurston; D. Krewski; Y. Shi, E. Calle and
2	M. Thun. 2009. "Long-term O ₃ Exposure and Mortality." N. Eng. J. Med., 360:1085-
3	1095.
4	Johnson, T. 2002. "A Guide to Selected Algorithms, Distributions, and Databases Used in
5	Exposure Models Developed by the Office of Air Quality Planning and Standards,"
6	prepared by TRJ Environmental, Inc. for the U.S. Environmental Protection Agency.
7	Research Triangle Park, NC: Office of Research and Development.
8	National Research Council of the National Academies. 2008. "Estimating Mortality Risk
9	Reduction and Economic Benefits from Controlling Ozone Air Pollution." Washington,
10	DC: The National Academies Press.
11	Rizzo, M. 2005. "A Comparison of Different Rollback Methodologies Applied to Ozone
12	Concentrations," posted on November 7, 2005,
13	http://www.epa.gov/ttn/naaqs/standards/ozone/s_O3_ cr_td.html.
14	Rizzo, M. 2006. "A Distributional Comparison between Different Rollback Methodologies
15	Applied to Ambient Ozone Concentrations," posted on May 31, 2006,
16	http://www.epa.gov/ttn/naaqs/standards/ozone/s_O3_ cr_td.html.
17	Simon, H.; K. Baker; N. Possiel; F. Akhtar; S. Napelenok; B. Timin; B. Wells. 2012. "Model-
18	based Rollback Using the Higher Order Direct Decoupled Method (HDDM)," posted at
19	< <u>http://www.epa.gov/ttn/naaqs/standards/ozone/s_O₃_td.html</u> >.
20	Simon, H.; K. R. Baker; F. Akhtar; S.L. Napelenok; N. Possiel; B. Wells and B. Timin. 2013. "A
21	Direct Sensitivity Approach to Predict Hourly Ozone Resulting from Compliance with
22	the National Ambient Air Quality Standard" Environmental Science and Technology,
23	Vol. 47, 2304-2313.
24	Smith, R.L., B. Xu, P. Switzer. 2009b. "Reassessing the Relationship Between Ozone and Short-
25	term Mortality in U.S. Urban Communities," Inhale Toxicol, Vol. 21: 37-61.
26	Timin, B.; K. Wesson and J. Thurman. 2010. "Application of Model and Ambient Data Fusion
27	Techniques to Predict Current and Future Year PM _{2.5} Concentrations in Unmonitored
28	Areas, " in D.G. Steyn and St Rao (eds), Air Pollution Modeling and Its Application XX,
29	Netherlands: Springer, pp. 175-179.
1	U.S. Environmental Protection Agency. 2007. Review of the National Ambient Air Quality
----	---
2	Standards for Ozone: Policy Assessment of Scientific and Technical Information OAQPS
3	Staff Paper. Washington, DC: EPA Office of Air and Radiation. (EPA document number
4	U. S. EPA. 2012a. Integrated Science Assessment for Ozone and Related Photochemical
5	Oxidants: Third External Review Draft. Research Triangle Park, NC: EPA Office of Air
6	Quality Planning and Standards. (EPA document number EPA-452/R-07-007;
7	EPA/600/R-10/076C).
8	U.S. EPA. 2012b. Total Risk Integrated Methodology (TRIM) - Air Pollutants Exposure Model
9	Documentation (TRIM.Expo / APEX, Version 4.4) Volume I: User's Guide. Research
10	Triangle Park, NC: Office of Air Quality Planning and Standards. (EPA document
11	number EPA-452/B-12-001a). < <u>http://www.epa.gov/ttn/fera/human_apex.html</u> >.
12	U.S. EPA. 2012c. Total Risk Integrated Methodology (TRIM) - Air Pollutants Exposure Model
13	Documentation (TRIM.Expo / APEX, Version 4.4) Volume II: Technical Support
14	Document. Research Triangle Park, NC: Office of Air Quality Planning and Standards.
15	(EPA document number EPA-452/B-12-001b).
16	< <u>http://www.epa.gov/ttn/fera/human_apex.html</u> >.
17	Wells, B.; K. Wesson and S. Jenkins. 2012. "Analysis of Recent U.S. Ozone Air Quality Data to
18	Support the O_3 NAAQS Review and Quadratic Rollback Simulations to Support the First
19	Draft of the Risk and Exposure Assessment."
20	< <u>http://www.epa.gov/ttn/naaqs/standards/ozone/s_0₃_td.html</u> >.
21	Zhang, L.; D.J. Jacob; N.V. Smith-Downey; D.A. Wood; D. Blewitt; C.C. Carouge; A. van
22	Donkelaar; D.B. A. Jones; L.T. Murray and Y. Wang. 2011. "Improved Estimate of the
23	Policy-relevant Background Ozone in the United States Using the GEOS-Chem Global
24	Model with 1/2°x2/3° Horizontal Resolution Over North America." Atmos Environ, Vol.
25	45, pp. 6769-6776.
26	Zanobetti, A. and J. Schwartz. 2008. "Mortality Displacement in the Association of Ozone with
27	Mortality: An Analysis of 48 Cities in the United States." American Journal of
28	Respiratory and Critical Care Medicine, 177:184-189.
29	

4-54

5 CHARACTERIZATION OF HUMAN EXPOSURE TO OZONE

3 5.0 OVERVIEW

4 As part of the previous 2007 O₃ NAAQS review, EPA staff conducted exposure analyses for the general population, all school-age children (ages 5-18), all active school-age children,¹ 5 and asthmatic school-age children (U.S. EPA, 2007a,b). Exposure estimates were generated for 6 7 12 urban study areas² for recent years of air quality and for just meeting the existing 8-hr standard and several alternative 8-hr standards. EPA also conducted a health risk assessment that 8 9 produced risk estimates for the number and percent of all school-age children experiencing 10 impaired lung function and other respiratory symptoms associated with the exposures estimated for these same 12 study areas. 11

12 The exposure analysis conducted for this current NAAQS review builds upon the 13 methodology and lessons learned from the exposure analyses conducted in previous O₃ reviews (U.S. EPA, 1996a, 2007a,b) and information provided in the final ISA (U.S. EPA, 2013). Here, 14 we estimate exposures for people residing in 15 urban study areas in the U.S.³ The population 15 exposures to ambient O₃ concentrations were modeled using EPA's Air Pollutants Exposure 16 17 (APEX) (US EPA, 2012a,b). Exposures were calculated considering O₃ concentrations in recent 18 years, using 2006 to 2010 spatially interpolated ambient monitoring data. Exposures were also 19 estimated considering alternative air quality scenarios, that is, where O₃ concentrations just meet 20 the existing 8-hr O₃ NAAQS and at several other standard levels considering the same indicator, 21 form, and averaging time, based on adjusting data as described in Chapter 4. Exposures were 22 modeled for 1) all school-age children (ages 5-18), 2) asthmatic school-age children (ages 5-18), 3) asthmatic adults (ages 19-95), and 4) all older adults (ages 65-95), each while at moderate or 23 greater exertion level at the time of exposure.⁴ The strong emphasis on children, asthmatics, and 24 older adults reflects the finding of the last O₃ NAAQS review (U.S. EPA, 2007a) and the ISA 25 26 (U.S. EPA, 2013, Chapter 8) that these are important at-risk groups. Exposure model output of 27 interest for this chapter are the percent (and number) of persons exposed at or above 8-hr average

¹ In the previous 2007 exposure assessment, a study group of active school-age children was identified as children having their median daily physical activity index (PAI) over the exposure period ≥ 1.75, an activity level characterized by exercise physiologists as being "moderately active" or "active" (McCurdy, 2000).

² The twelve study areas evaluated in the 2007 exposure assessment were Atlanta, Boston, Chicago, Cleveland, Detroit, Houston, Los Angeles, New York, Philadelphia, Sacramento, St. Louis, Washington DC (an area which at that time was modeled to include Baltimore as part of the Baltimore-Northern Virginia MSA).

³ In addition to the twelve study areas identified in the 2007 exposure assessment, staff has added Dallas and Denver, while also separately modeling Baltimore (from Washington DC) in this current assessment. Inclusion of Seattle, WA was considered but not included due to a lack of appropriate monitoring data.

⁴ The "all school-age children" study group includes both asthmatic and non-asthmatic children ages 5 to 18. The "all older adults" includes both asthmatic and non-asthmatic older adults ages 65 to 95. Note also that the 8-hr average exposure of interest in both this and the previous assessment was concomitant with moderate or greater exertion for all study groups.

1 O₃ concentrations of concern, all while at moderate or greater exertion levels, based on adverse

2 effects observed in human clinical exposure studies. Further, the complete time series of

- 3 individual exposures estimated by APEX serves as input to a module that estimates human health
- 4 risk (Chapter 6).

5 This chapter first provides a brief overview of human exposure and exposure modeling 6 using APEX (section 5.1), the scope of this O_3 exposure assessment and key inputs used to 7 model exposure in the 15 U.S. study areas selected (section 5.2), and followed by the main body 8 exposure results (section 5.3). Then, section 5.4 presents an assemblage of targeted analyses 9 designed to provide additional insight to the main body of exposure results by focusing on 10 important data inputs, additional at-risk populations, lifestages, or scenarios, influential attributes 11 in estimating exposures, and performance evaluations. The results of these and other exposure 12 model targeted analyses are integrated in an uncertainty characterization section (section 5.5) 13 along with a final section summarizing the key observations for this chapter (section 5.6).

14 5.1 SYNOPSIS OF O₃ EXPOSURE AND EXPOSURE MODELING

15 **5.1.1 Human Exposure**

Human exposure to a contaminant is defined as "contact at a boundary between a human
and the environment at a specific contaminant concentration for a specific interval of time," and
has units of concentration times duration (National Research Council, 1991). For air pollutants
the contact boundary is nasal and oral openings in the body, and *personal exposure* of any
individual to a chemical in the air for a discrete time period is fundamentally quantified as (Lioy,
1990; National Research Council, 1991):

(5-1)

- 22
- 23

23 24 $E_{[t_1, t_2]} = \int_{t_1}^{t_2} C(t) dt$

where $E_{[t_1,t_2]}$ is the personal exposure or *exposure concentration* during the time period from t_1 to t_2 , and C(t) is the concentration at time t in the breathing zone. The breathing rate at the time of exposure will influence the dose received by the individual. While we do not directly estimate dose in this assessment, *intake* is the total O₃ inhaled (i.e., exposure concentration, duration, and ventilation combined).⁵

30

31

⁵ In chapter 6, the estimation of risk combines the time series of both the personal exposure concentrations and ventilation rate, among other variables in essentially calculating a dose, though not explicitly output from the model.

1 5.1.2 Estimating O₃ Exposure

2 Exposure to O_3 can be directly estimated by monitoring the concentration of O_3 in a 3 person's breathing zone (close to the nose/mouth) using a personal exposure monitor. Studies 4 employing this measurement approach have been reviewed in the ISA and EPA O₃ Air Quality 5 Criteria Documents (U.S. EPA, 1986, 1996b, 2006, 2013). Personal exposure measurements 6 from these studies are useful in describing a general range of exposure concentrations (among 7 other reported measurement data) and in identifying factors that may influence varying exposure 8 levels. However, these measurement studies are largely limited by the disparity between sample 9 measurement duration and exposure concentration averaging-times of interest and in 10 appropriately capturing variability in population exposure occurring over large geographic areas, 11 particularly when considering both concentration (e.g., spatial variability) and population (e.g., 12 age, sex) attributes that influence exposure.

13 O₃ exposure for individuals, small groups of individuals or large populations can be 14 calculated indirectly (or *modeled*) using Equation 5-1. When employing such an approach in a 15 population exposure assessment, two basic types of input data are needed; a time-series of O_3 16 concentrations that appropriately represents spatial heterogeneity in O_3 concentrations and a 17 corresponding time-series of locations visited by the persons exposed. When considering air 18 pollutant concentrations, population exposure models are commonly driven by ambient 19 concentrations. These ambient concentrations may be provided by monitoring data, by air quality 20 model estimates, or perhaps by a combination of these two data sources. Then, an understanding 21 of the relationships between ambient pollutants and the locations people occupy is needed. This 22 is because human exposure, regardless of the pollutant or whether one is interested in individual 23 or population exposure, depends on where an individual is located, how long they occupy that 24 location, and what the pollutant concentration at the point of contact is. Furthermore, if interested 25 in air pollutant intake rate or dose, one needs to know what activity the person is performing 26 while exposed.

27 Thus, the types of measurement and modeling studies that provide information for more 28 realistically estimating exposure to O_3 can be augmented from the above list to include studies 29 of: 1) O₃ formation, deposition, and decay, 2) people's locations visited and activities performed, 30 3) human physiology, and 4) local scale meteorological measurements and/or modeling. Useful 31 data derived from these varied studies are O_3 concentrations (i.e., fixed site, personal exposure, 32 indoor and outdoor locations), built environment physical factors (i.e., air exchange rates 33 (AERs), infiltration rates, decay and deposition rates), human time-location-activity patterns 34 (minute-by-minute, hourly, daily, and longer-term), time-averaged or activity-specific breathing 35 rates among varying sexes and/or lifestages, and hourly ambient temperatures.

1 When integrating these varied data (among others such as population demographics and 2 disease prevalence) and understanding factors affecting exposure, exposure models can extend 3 beyond the limited information given by measurement data alone. For example, an exposure 4 model can reasonably estimate exposures for any perceivable at-risk population (e.g., asthmatics 5 living in a large urban area) and considering any number of hypothetical air quality conditions 6 (e.g., just meeting a daily maximum 8-hr average concentration of 70 ppb). Exposure models that 7 account for variability in human physiology can also realistically estimate pollutant intake by 8 using activity-specific ventilation rates. These types of measurements cannot realistically be 9 performed for a study group or population of interest, particularly when considering time, cost, 10 and other constraints. The following section provides an overview of how such exposure 11 modeling can be done using APEX, the model developed by EPA to perform such calculations

12 and used to estimate O_3 exposures in this REA.

13

5.1.3 Modeling O₃ Exposure Using APEX

EPA has developed the APEX model for estimating human population exposure to criteria and air toxic pollutants, used most recently in estimating exposures for the O₃ (U.S. EPA, 2007b), nitrogen dioxide (U.S. EPA, 2008), sulfur dioxide (U.S. EPA, 2009a), and carbon monoxide (U.S. EPA, 2010) NAAQS reviews. APEX is a probabilistic model designed to account for the numerous sources of variability that affect people's exposures. An overview of the approaches used by APEX to estimate exposure concentrations is found in Appendix 5A with details provided in U.S. EPA (2012a,b).

21 Briefly, APEX simulates the movement of individuals through time and space and 22 estimates their exposure to a given pollutant while occupying indoor, outdoor, and in-vehicle 23 locations. The model stochastically generates simulated individuals in selected study areas using 24 census-derived probability distributions for demographic characteristics. Population demographics are drawn from the 2000 Census data⁶ at a tract level, and a national commuting 25 database based on 2000 Census data provides home-to-work commuting flows between tracts.⁷ 26 27 Any number of individuals can be simulated, and collectively they approximate a random 28 sampling of people residing in a particular study area. 29 Daily activity patterns for individuals in a study area, an input to APEX, are obtained 30 from detailed daily time-location-activity pattern survey data that are compiled in the 31 Consolidated Human Activity Database (CHAD) (McCurdy et al., 2000; U.S. EPA, 2002). These

32 daily diaries are used to construct a sequence of locations visited and activities performed for

33 APEX simulated individuals consistent with their demographic characteristics, day-type (e.g.,

⁶ Due to resource limitations and data availability, the 2010 Census data have not yet been processed to include in this 2nd draft REA.

⁷ There are approximately 65,400 census tracts in the $\sim 3,200$ counties in the U.S.

1 weekend or weekday), and season of the year, as defined by ambient temperature regimes

- 2 (Graham and McCurdy, 2004). The time-location-activity data input to APEX are linked with
- 3 personal attributes of the surveyed individuals' such as age, sex, employment status, day-of-
- 4 week surveyed, and daily maximum and daily mean temperature. These specific personal
- 5 attribute data are then used by APEX to best match the daily diary with the simulated persons of
- 6 interest, using the same variables as first-order diary selection characteristics. The approach is
- 7 designed to capture the important attributes contributing to an individuals' time-location-activity
- 8 pattern, and of particular relevance here, time spent outdoors (Graham and McCurdy, 2004). In
- 9 using a diverse collection of time-location-activity diaries that capture the duration and
- 10 frequency of occurrence of visitations/activities performed, APEX can simulate expected
- 11 variability in human behavior, both within and between individuals. This, combined with
- 12 exposure concentrations, allows for the reasonable estimation of the magnitude, frequency,
- 13 pattern, and duration of exposures an individual experiences.
- 14 A key concept in modeling exposure using APEX is the *microenvironment*, a term that 15 refers to the immediate surroundings of an individual at a particular time. APEX has a flexible 16 approach for modeling micro-environmental concentrations whereas the model user defines the 17 type, number and characteristics of the microenvironments to be modeled. Typical 18 microenvironments include indoors at home, indoors at school, near roadways, inside cars, and 19 outside home. In this exposure assessment, all microenvironmental O_3 concentrations are 20 derived from ambient O₃ concentrations input to APEX and are estimated using either a mass-21 balance or transfer factors approach, selected by the user. The mass balance approach assumes 22 that the air in an enclosed microenvironment is well-mixed and that the air concentration is 23 spatially uniform at a given time within the microenvironment. The approach employs indoor-to-24 outdoor AERs (i.e., number of complete air exchanges per hour) and considers removal 25 mechanisms such as deposition to building surfaces and chemical decay rates. The transfer 26 factors model is simpler than the mass balance model, and employs two variables, a *proximity* 27 *factor*, used to account for proximity of the microenvironment to sources or sinks of pollution, or 28 other systematic differences between concentrations just outside the microenvironment and the 29 ambient concentrations, and a *penetration factor*, which quantifies the degree to which the 30 outdoor air penetrates into the microenvironment.
- Activity-specific simulated breathing rates of individuals are used in APEX to characterize intake received from an exposure. This is done because controlled human exposure studies have shown adverse health outcomes are associated with both elevated concentrations and study participant exertion levels. The breathing rates calculated by APEX are derived from the energy expenditure associated with each simulated persons' activity performed, adjusted for age- and sex-specific physiological parameters (Graham and McCurdy, 2005). The energy
 - 5-5

1 expenditure estimates themselves are derived from distributions of METS⁸ (or metabolic

2 equivalents of work) associated with every activity performed (McCurdy et al., 2000, using

3 Ainsworth et al., 1993).

4 An important feature of APEX is the ability to account for variability in exposure by 5 representing input variables as statistical distributions along with dependent conditional 6 variables, where appropriate. For example, the distribution of AERs in a home, office, or motor 7 vehicle can depend on the type of heating and air conditioning present, which are also stochastic 8 inputs to the model, as well as the ambient temperature on a given day. The user can choose to 9 keep the value of a stochastic parameter constant for the entire simulation (appropriate for the 10 volume of a house), or can specify that a new value shall be drawn hourly, daily, or seasonally 11 from specified distributions.

12 Finally, APEX calculates a unique time-series of exposure concentrations on the order of 13 minutes or smallest diary event duration that each simulated person may experience during the 14 modeled time period, based in that individual's estimated microenvironmental concentrations 15 and the time spent in each of sequence of microenvironments visited according to the time-16 location-activity diary of each individual. Then, hourly average exposures of each simulated 17 individual are estimated using time-weighted averages of the within-hour exposures. From 18 hourly exposures, APEX calculates any other time averaged exposure of interest (e.g., 8-hr or 19 daily average) that a simulated individual experiences during the modeled period.

20 5.2 SCOPE OF THE EXPOSURE ASSESSMENT

21 This section broadly presents the scope of the exposure assessment including descriptions 22 of the modeling domains, ambient concentrations used, time periods and populations modeled, as 23 well as identifying key approaches, inputs and outputs used by APEX in estimating population 24 O₃ exposures. Detailed descriptions regarding APEX modeling, model inputs and other 25 supporting information are provided in Appendix 5A-5E and the APEX user's guide and 26 technical support documents (U.S. EPA 2012a,b). Figure 5-1 illustrates the general conceptual 27 framework for generating our population exposure concentrations, including the time series of 28 exposure and ventilation rate output generated as input to population risk calculations in Chapter 29 6.

30 5.2.1 Urban Areas Selected

The selection of urban areas to include in the exposure assessment considered the
 location of O₃ epidemiological studies, the availability of ambient O₃ monitoring data, and the

⁸ METS are a dimensionless ratio of the activity-specific energy expenditure rate to the basal or resting energy expenditure rate. The metric is used by exercise physiologists and clinical nutritionists to estimate work undertaken by individuals as they go through their daily activities (Montoye et al., 1996).

- 1 desire to represent a range of geographic areas, encompassing variability in climate and
- 2 population demographics. Specifically, the criteria included the following:
 - The overall set of urban locations should represent a range of geographic areas, urban population demographics, and climate, beginning with study areas selected in the 2007 O₃ NAAQS review.
 - The locations should be focused on areas that do not meet or are close to not meeting the existing 8-hr O₃ NAAQS and should include areas having O₃ non-attainment designations.
 - There must be sufficient O_3 ambient air quality data for the recent 2006-2010 period.
 - The study areas should include the 12 cities modeled in the epidemiologic-based risk assessment (Chapter 7).

3 Based on these criteria, we chose the 15 study areas listed in Table 5-1 to develop our 4 population exposure estimates. We then defined an *air quality domain* for each study area, 5 broadly bounding the ambient concentration field where exposures were to be estimated. To do 6 this, we evaluated 1) counties modeled in the previous 2007 O₃ NAAOS review common to 7 current study areas, 2) political/statistical county aggregations (e.g., whether in a metropolitan 8 statistical areas or MSAs), and 3) if the study area was designated as a non-attainment area 9 (NAA), the counties that were part of the NAA list. We identified a final list of 215 counties⁹ to 10 comprise the air quality domain for the 15 study areas, the names of which are provided in 11 Appendix 5B.

12 **5.2.2 Time Periods Simulated**

- 13 The exposure periods modeled are the O_3 seasons for which routine hourly O_3 monitoring
- 14 data were available for years 2006 to 2010 (Table 5-1), and defined by 40 CFR part 58,
- 15 Appendix D, Table D-3. These periods are designed to reasonably capture year-to-year
- 16 variability in ambient concentrations and meteorology and include most of the high
- 17 concentration events occurring in each area. Having this wide range of air quality data across
- 18 multiple years allows us to more realistically estimate a range of exposures, rather than using a
- 19 single year of air quality data. While the number of available O₃ monitors may vary slightly from
- 20 year to year, we assumed constant representation by the available monitors and associated
- 21 statistically interpolated data for each year over the simulation period (see section 5.2.3).

⁹ Of the 215 counties defining the air quality domain, 207 remained in the exposure model domain.



Figure 5-1 Conceptual Framework Used for Estimating Study Area Population O₃ Exposure Concentrations

Table 5-1 General Characteristics of the Population Exposure Modeling Domain Comprising Each Study Area

		Study Area Number of:					
						Persons	
Study Area (Abbreviation)	$\mathrm{O}_3Season^1$	Counties	Ambient Monitors	APEX Air Districts	US Census Tracts	All School Age Children (age 5-18)	All Ages (age 5-95)
Atlanta (ATL)	Mar 1-Oct 31	32	14	664	678	860,649	3,850,951
Baltimore (BAL)	Apr 1-Oct 31	7	12	603	618	505,140	2,209,226
Boston (BOS)	Apr 1-Sep 30	7	13	1,005	1,028	905,208	4,449,291
Chicago (CHI)	Apr 1-Oct 31	16	28	1,882	2,055	1,899,073	8,345,373
Cleveland (CLE)	Apr 1-Oct 31	8	16	802	879	578,733	2,692,846
Dallas (DAL)	Mar 1-Oct 31	11	21	1,012	1,036	1,097,004	4,698,392
Denver (DEN)	Mar 1-Sep 30	12	25	655	675	560,137	2,626,239
Detroit (DET)	Apr 1-Sep 30	9	13	1,419	1,454	1,016,896	4,572,479
Houston (HOU)	Jan 1-Dec 31	10	19	779	802	970,528	3,925,054
Los Angeles (LA)	Jan 1-Dec 31	5	50	2,000	3,352	3,620,972	14,950,340
New York (NY)	Apr 1-Oct 31	27	32	1,900	4,889	3,843,450	18,520,868
Philadelphia (PHI)	Apr 1-Oct 31	15	19	1,452	1,555	1,231,052	5,506,954
Sacramento (SAC)	Jan 1-Dec 31	7	18	447	461	466,169	1,926,598
St. Louis (STL)	Apr 1-Oct 31	15	16	494	518	527,755	2,340,325
Wash., DC (WAS)	Apr 1-Oct 31	26	28	1,013	1,037	966,791	4,498,374
All Study Areas	-	207	324	16,127	21,037	19,049,557	85,113,310

4 ¹ Each study area's O₃ monitoring season is defined by 40 CFR part 58, Appendix D, Table D-3.

5 5.2.3 Ambient Concentrations Used

We used the available hourly ambient monitor concentration data within and around each study area along with a statistical interpolation technique (Chapter 4) to estimate hourly census tract concentrations within the counties comprising each study area. These concentrations served as the 'base' air quality input for each study area year. Ambient concentrations were also

- 10 adjusted to just meet the existing standard (75 ppb, 4th highest 8-hr average, averaged over a 3-
- 11 year period) and alternative standard levels (70, 65, 60, and 55 ppb) using an air quality model
- 12 and the statistical interpolation technique (Chapter 4).

13 These estimated hourly census tract O_3 concentrations served as the APEX *air districts*,

14 the basic ambient concentrations from which each simulated persons microenvironmental

15 concentrations are estimated. Having these temporally and spatially resolved air districts in each

study area allows for better utilization of APEX spatial and temporal capabilities in estimating
 exposure. Because APEX simulates where individuals are located at specific times of the day,
 more realistic exposure estimates are obtained in simulating the contact of individuals with these
 spatially and temporally diverse concentrations.

5 Even though we estimated O_3 ambient concentrations at all census tracts in each county-6 level study area, the study area *exposure modeling domain* was defined as a subset of these 7 census tracts by using the ambient monitoring sites within the urban core of each study area's air 8 quality domain and a 30 km radius of influence. This zone of influence is consistent with what 9 was done in the 1st draft O₃ REA, though in that exposure assessment, only the ambient 10 monitoring data sites themselves were used to represent the APEX air districts, hence 11 concentrations measured at a particular monitoring site would be directly extrapolated outwards 12 to all census tracts within 30 km of that site. In contrast, by incorporating the VNA estimated 13 concentrations and retaining the same 30 km radius of influence, we are stressing the 14 significance of the monitor information in defining the urban core air quality while also 15 reasonably estimating concentration gradients (where such gradients exist) with increasing 16 distance from monitoring locations.

Thus, all air districts¹⁰ and census tracts that fall within the 30 km radius of each ambient monitor were used to estimate the exposures, defining the final exposure modeling domain in each study area (Table 5-1). The monitor IDs used to select the census tracts to be modeled are provided in Appendix 5B, while the complete list of census tract IDs where exposures are modeled are within the APEX control files for each study area (and are the same for each simulation year).

23 **5.2.4 Meteorological Data Used**

24 APEX uses study area temperature data to select representative diaries for a particular 25 day and in selecting an appropriate air exchange rate used to calculate indoor residential 26 microenvironmental concentrations. APEX uses the data from the closest weather station to each 27 Census tract. To ensure reasonable coverage for each study area, a few to several meteorological 28 stations recording hourly surface temperature measurements were identified using data obtained from the National Weather Service ISH data files.¹¹ Details regarding the meteorological stations 29 30 selected and data processing are given in Appendix 5B. Briefly, APEX requires the temperature 31 input data to be 100% complete. In general, any missing values were filled using a linear

¹⁰ The original number of air quality districts for New York and Los Angeles needed to be reduced by about half due to exceeding personal computer memory capacity when APEX used > 2,000 air districts. See Appendix 5B for details.

¹¹ <u>http://www.ncdc.noaa.gov/oa/climate/surfaceinventories.html</u>

interpolation or regression approach that employs information from proximal meteorological
 stations.

3 5.2.5 Populations Simulated

4 Exposure was estimated for four at-risk study groups residing in each study area: all 5 school-age children (ages 5-18), asthmatic school-age children, asthmatic adults (ages 19-95), 6 and all older adults (ages 65-95). Due to the increased amount of time spent outdoors engaged in 7 relatively high levels of physical activity (which increases intake), school-age children as a group 8 are particularly at risk for experiencing O_3 -related health effects (U.S. EPA, 2013, Chapter 8). 9 We report results for all school-age children down to age five, recognizing an increasing trend 10 for younger children to attend school. Some U.S. states allow 4-year-olds to attend kindergarten, 11 and most states have preschool programs for children younger than five. In 2000, six percent of 12 U.S. children ages 3 to 19 who attend school were younger than five years old (2000 Census 13 Summary File 3, Table QT-P19: School Enrollment). Currently we do not estimate exposure for 14 these younger children due to a lack of information that would let us confidently characterize 15 these younger aged children. While EPA guidance recommends, for certain instances, an upper 16 age group of children ages 16 through 21 (U.S. EPA, 2005), we restricted our upper age 17 classification of children through age 18. In considering the expected variability in activity 18 patterns over the span of ages 16 through 21 (e.g., time spent outdoors, time in school, each in 19 contrast to time spent working) and the relatively small difference in respiratory physiology over that same age span compared with that of adults (e.g., Figure 5-17), factors critical for high O₃ 20 exposure and dose, we assumed simulated persons age 19 to 21 would be best included in our 21 22 adult study group. The number of persons represented in each of the 15 study areas is given in 23 Table 5-1 and, considering all study areas together, captures approximately 32.8 % of all 24 children ages 5 to 18 and 32.0 % of the total U.S. population ages 5 to 95. 25 The number of asthmatic school-age children and asthmatic adults in each study area was 26 estimated using asthma prevalence from the Center for Disease Control (CDC) and Prevention's

27 National Health Interview Survey (NHIS).¹² Briefly, years 2006-2010 NHIS survey data were

combined to calculate asthma prevalence, defined as the probability of a "*Yes*" response to the

29 question: "*do you still have asthma?*" among those that responded "*Yes*" to the question "*has a*

30 *doctor ever diagnosed you with asthma?*". The asthma prevalence was first stratified by NHIS

defined regions (Midwest, Northeast, South, and West), sex, age (single years for ages 0-17) or

32 age groups (ages \geq 18), and by a family income/poverty ratio.¹³ These new asthma prevalence

33 estimates were then linked to U.S. census tract level poverty ratio probabilities (U.S. Census

¹³ The income/poverty ratio threshold used was 1.5, that is the surveyed person's family income was considered either \leq or > than a factor of 1.5 of the U.S. Census estimate of poverty level for the given year.

¹² See <u>http://www.cdc.gov/nchs/nhis.htm</u> (accessed October 4, 2011).

- 1 Bureau, 2007), also stratified by age and age groups, to generate a final database consisting of
- 2 census tract level asthma prevalence for the entire U.S. A detailed description of how the data
- 3 base was developed is presented in Appendix 5C, while the estimated asthma prevalence used
- 4 for each census tract is provided in the APEX asthma prevalence input file. A summary of the
- 5 asthma prevalence calculated for each study area simulation is provided here in Table 5-2.

	Asthma Prevalence (%)					
Study Area	Children (5-18)	Adults (18-95)	All Persons (5-95)			
Atlanta	9.6	6.5	7.2			
Baltimore	9.7	6.6	7.3			
Boston	11.4	7.9	8.6			
Chicago	10.7	7.8	8.4			
Cleveland	10.9	7.7	8.4			
Dallas	9.9	6.5	7.3			
Denver	8.9	7.7	7.9			
Detroit	11.1	7.7	8.5			
Houston	10.1	6.5	7.4			
Los Angeles	9.0	7.7	8.0			
New York	12.2	8.1	9.0			
Philadelphia	11.3	7.9	8.7			
Sacramento	9.0	7.8	8.1			
St. Louis	11	7.6	8.4			
Washington DC	9.5	6.4	7.1			
All Areas	10.5	7.6	8.2			

Table 5-2 Asthma Prevalence for Children and Adults Estimated by APEX in Each Simulated Study Area

8

9 All simulated persons (either asthmatic or non-asthmatic) used time-location-activity data 10 from CHAD, the most complete, high quality source of human activity data for use in exposure 11 modeling. The current CHAD database contains over 53,000 individual daily diaries including 12 time-location-activity patterns for individuals of both sexes across a wide range of ages (<1 to 13 94). The database is geographically diverse, containing diaries from individuals residing in 14 several major cities, suburban, and rural areas across the U.S. Time spent performing activities 15 within particular locations can be on a minute-by minute basis, thus avoiding the smoothing of 16 potential peak exposures longer event durations would yield.

- 1 Table 5-3 summarizes the studies and number of diaries in CHAD used in this 2 assessment, noting that the total CHAD diaries used by APEX is restricted to just over 41,000 given our simulation age range (5-95) and additionally selected usability requirements.¹⁴ 3 4 Additional context regarding the representativeness of the CHAD data in estimating exposure is 5 provided in section 5.3.1 and Appendix 5G. 6 APEX creates a sequence of daily diaries across the entire O_3 season for each simulated 7 individual using a method designed to capture the tendency of individuals to repeat activities, 8 based on reproducing realistic variation in a key diary variable (Glen et al., 2008). For this O₃ 9 analysis, the key variable selected is the amount of time an individual spends outdoors each day, 10 one of the most important determinants of exposure to high levels of O_3 (see section 5.3.2). The
- 11 longitudinal method targets two statistics, a population diversity statistic (D) and a within-person
- 12 autocorrelation statistic (*A*). Values of 0.2 for *D* and 0.2 for *A* were initially developed based on
- 13 analyses by Geyh et al. (2000) and Xue et al. (2004), with both studies evaluating groups of
- 14 children ages 7 to 12 in a single study area. We adjusted values for D upwards to 0.5 to reflect a
- 15 broader range of ages and to better estimate repeated activities.¹⁵ Further details regarding the
- 16 development of the longitudinal methodology can be found in U.S. EPA (2012a, b).

18

¹⁴ In this assessment, the CHAD diaries must be from persons having a known age, sex, day-of-week, and daily temperature. In addition, diaries must have no more than 3 hours total of missing location and/or activity data.

¹⁵ A small *D* means that the overall variability between people in the key diary statistic is smaller than the variability observed over days within the same person. A *D* closer to 1 means that each person shows little variation over time relative to the variability between persons.

Table 5-3 Consolidated Human Activity Database (CHAD) Study Information and Diary-days Used by APEX

Activity Pattern Study (Abbrev.)	General Study Area	Study Years	Subject Ages	Diary-days (ages 4-94)	Diary-days (ages 4-18)
Baltimore Retirement Home (BAL)	Baltimore, MD	1997-98	72 - 93	304	0
California Youth (CAY)	California	1987-88	12 - 17	182	182
California Adults (CAA)	California	1987-88	18 - 94	1,555	36
California Children (CAC)	California	1989-90	<1 - 11	771	771
Cincinnati (CIN)	Cincinnati, OH	1985	<1 - 86	2,259	727
Detroit Exposure and Aerosol Research (DEA) ^{1,2}	Detroit, MI	2005-06	18 - 74	331	5
Denver CO Personal Exposure (DEN)	Denver, CO	1982-83	18 - 70	714	7
EPA Longitudinal (EPA) ^{1,2}	RTP, NC	1999-2000, 2002, 06-08	<1 - 60	1,386	0
LA O_3 Exposure: Elementary School (LAE)	Los Angeles, CA	1989	10 - 12	50	50
LA O_3 Exposure: High School (LAH)	Los Angeles, CA	1990	13 - 17	42	42
National Human Activity Pattern Study: Air (NHA)	National	1992-94	<1 - 93	4,129	693
National Human Activity Pattern Study: Water (NHW)	National	1992-94	<1 - 93	4,099	745
National-Scale Activity Survey (NSA)	7 US metro. areas	2009	35 - 92	6,825	0
Population Study of Income Dynamics I (ISR) ¹	National	1997	<1 - 13	3,507	3,507
Population Study of Income Dynamics II (ISR) ¹	National	2002-03	5 - 19	4,800	4,793
Population Study of Income Dynamics III (ISR) ^{1,2}	National	2007-08	10 - 19	2,619	2614
RTI O_3 Averting Behavior (OAB) ¹	35 US metro. areas	2002-03	2 - 12	2,187	2,187
RTP Panel (RTP) ¹	RTP, NC	2000-01	55 - 85	871	0
Seattle (SEA) ¹	Seattle, WA	1999-2002	6 - 91	1,624	317
Study of Use of Products and Exposure Related Behavior (SUP) ^{1,2}	Sac/San Fran, CA Counties	2006-10	1 - 88	2,533	994
Washington, D. C. (WAS)	Wash., DC	1982-83	18 - 71	686	10
Totals		1982 - 2010	<1 - 94	41,474	17,680

3 ¹ Study data added after 2007 O_3 NAAQS review.

 $4 \qquad ^{2} \mbox{ Study data added after 2012 1}^{st} \mbox{ Draft } O_{3} \mbox{ REA}.$

2

5.2.6 Key Physiological Processes And Personal Attributes Modeled

3 The modeling of physiological processes relevant to the O_3 exposure and intake is 4 complex, particularly when representing inter- and intra-personal variability in energy 5 expenditure (EE) and ventilation rates (VE). APEX has a module capable of estimating several 6 variables associated with every activity performed by simulated individuals. Briefly, the module 7 links the diary indicated activities to specific energy expended, the rate of oxygen consumed 8 (VO_2) and the associated ventilation rate, all considering the unique sequence of events 9 individuals go through each simulated day. The activity-specific time-series of VE estimates 10 ultimately serve as an important variable used in estimating O₃ intake as well as in identifying 11 when simulated individuals performing activities at moderate or greater exertion. In addition, 12 age, sex, and body mass related physiological differences are specifically taken into account by 13 the ventilation algorithm, derived using ventilation data obtained from several human studies 14 (see Graham and McCurdy, 2005): $\ln(V_E/BM) = b_0 + b_1 \ln(V_{O2}/BM) + b_2 \ln(1 + age) + b_3 sex + e_b + e_w$ 15 (5-2) 16 where, 17 18 ln = natural logarithm of variable \dot{V}_E / BM 19 = activity specific ventilation rate, body mass normalized (liter air/kg) = see Table 5-4 20 b_i • V o 2 / BM 21 = activity specific oxygen consumption rate, body mass normalized 22 $(liter/O_2/kg)$ 23 age = age of the individual (years) 24 = sex (-1 for males, +1 for females) sex 25 = randomly sampled error term for between persons $N\{0, se\}$, (liter air/kg) e_{h} 26 = randomly sampled error term for within persons $N\{0, se\}$, (liter air/kg) e_w 27 28 As indicated by Equation 5-2, the random error (ε) is allocated to two variance 29 components used to estimate the between-person (inter-individual variability) residuals 30 distribution (e_b) and within-person (intra-individual variability) residuals distribution (e_w) . The 31 regression parameters b_0 , b_1 , b_2 , and b_3 are assumed constant over time for all simulated persons, 32 e_b is sampled once per person by APEX, while whereas e_w varies from event to event. Point 33 estimates of the regression coefficients and standard errors of the residuals distributions are

34 given in Table 5-4. See Appendix 5A, Isaacs et al. (2008), and Chapter 7 of the APEX TSD (US

- 1 EPA, 2012b) for further discussion of this module. See also section 5.4.4 for a limited
- 2 performance evaluation of this module in estimating ventilation rates.
- 3

7

Age		Random Error ¹				
group	b ₀	b ₁	b ₂	b ₃	e _b	e _w
<20	4.3675	1.0751	-0.2714	0.0479	0.0955	0.1117
20-<34	3.7603	1.2491	0.1416	0.0533	0.1217	0.1296
34-<61	3.2440	1.1464	0.1856	0.0380	0.1260	0.1152
61+	2.5826	1.0840	0.2766	-0.0208	0.1064	0.0676

4 Table 5-4 Ventilation equation coefficient estimates (b_i) and residuals distributions (e_i)

¹ These are values of the coefficients and residuals distributions described by Equation (5-2) and described in Graham and McCurdy (2005).

8 Two key personal attributes determined for each simulated individual in this assessment 9 are body mass (BM) and body surface area (BSA). Each simulated individual's body mass is 10 randomly sampled from age- and sex-specific body mass distributions generated from National 11 Health and Nutrition Examination Survey (NHANES) data for the years 1999-2004.¹⁶ Details in 12 their development and the parameter values are provided by Isaacs and Smith (2005). Then age-13 and sex-specific body surface area can be estimated for each simulated individual based on 14 logarithmic relationships developed by Burmaster (1998) using body mass as an independent

15 variable as follows:

16

$$BSA = e^{-2.2781} BM^{0.6821}$$
(5-3)

17 5.2.7 Microenvironments Modeled

18 APEX is designed to estimate human exposure by using algorithms that attempt to 19 capture the full range of O_3 concentrations expected within several microenvironments. Broadly 20 aggregated, these can be either indoor, inside a motor vehicle, near road, or outdoor locations. 21 The two methods available in APEX for calculating pollutant concentrations within 22 microenvironments are a mass balance model and a transfer factor approach. Table 5-5 lists the 23 28 microenvironments selected for this analysis and the exposure calculation method used for 24 each. 25 The importance of modeling indoor microenvironments (e.g., homes, offices, schools) is

²⁵ The importance of modeling indoor microenvironments (e.g., homes, offices, schools) is 26 underscored by research indicating that personal exposure measurements of O_3 may not be well-

¹⁶ Demographic (Demo) and Body Measurement (BMX) datasets for each of the NHANES studies were obtained from http://www.cdc.gov/nchs/nhanes/nhanes_questionnaires.htm.

- 1 correlated with ambient measurements and indoor concentrations are usually much lower than
- 2 ambient concentrations (U.S. EPA, 2013, Section 4.3.3). We used mass balance modeling to
- 3 estimate O₃ concentrations in all indoor microenvironments, considering probabilistic
- 4 distributions of temperature-dependent (where data were available) building air exchange and
- 5 chemical decay rates. Parameter settings for each of these variables are provided in Appendix
- 6 5B, while additional discussion regarding updates made to air exchange rates using more recent
- 7 study data is given in Appendix 5E.

The remaining microenvironments were modeled using a transfer factors approach.

- 9 Outdoor microenvironmental concentrations were assumed equivalent to ambient concentrations,
- 10 near-road concentrations were adjusted considering whether or not O₃ concentrations were
- 11 reduced by atmospheric reactions (e.g., scavenging by NO_X) or other processes, and vehicular
- 12 microenvironments considered both the outdoor concentration attenuation and
- 13 infiltration/removal in the concentration estimation. Specific parameter settings for each of these
- 14 variables are provided in Appendix 5B.

Calculation Method Microenvironment Variables Indoor: Residence, Community Center or Auditorium, Restaurant, Mass Building air Hotel/Motel, Office building/Bank/Post Office, Bar/Night Club/Café, School, balance exchange & Shopping Mall/Non-Grocery Store, Grocery Store/Convenience Store, chemical Metro-Subway-Train Station, Hospital/Medical/Care Facility, Industrial decay rates Factory/Warehouse, Other Indoor Outdoor: Residential, Park/Golf Course, Restaurant/Café, School Grounds, Factors None Boat, Other Outdoor Non-Residential Near-road: Metro-Subway-Train Stop, Within 10 Yards of street, Parking Factors Proximity Garage (covered or below ground), Parking lot (open)/Street parking, factors Service Station Vehicle: Cars/Light Duty Trucks, Heavy Duty Trucks, Bus, Train/Subway Factors Proximity & penetration factors

15 Table 5-5 Microenvironments Modeled, Calculation Method Used, and Variables Included

16 **5.2.8 Model Output**

- 17 APEX estimates the complete time series of exposure concentrations for every simulated
- 18 individual and can summarize data using standardized time metrics (e.g., hourly or daily average,
- 19 daily maximum 8-hr average) or can output the minute-by-minute exposure concentrations (as is
- 20 needed for the risk estimation in Chapter 6). As an indicator of exposure to O_3 air pollution, we
- 21 selected the daily maximum 8-hr average O₃ exposure¹⁷ for every simulated individual and

¹⁷ It is important to stress here that only the maximum 8-hr exposure concentration is retained for each day simulated, per person. While every day could contain twenty-four unique 8-hr averages and that it is entirely

1 stratified these exposures by exertion level at the time of exposure. This indicator was selected

- 2 based on controlled human exposure studies where reported adverse health responses were
- 3 associated with exposure to O_3 and while the study subject was exercising.¹⁸ Factors important in
- 4 calculating this indicator includes the magnitude, duration, frequency of exposures, and the
- 5 breathing rate of individuals at the time of exposure. As a reminder, the calculated daily
- 6 maximum 8-hr average exposure concentrations are distinct from that of daily maximum 8-hr
- 7 average ambient concentrations by accounting for simulated individual's time-location-activity
- 8 patterns and O₃ concentration decay/variation occurring within the occupied microenvironments.

9 Benchmark levels used in this assessment include 8-hr average O₃ exposure
 10 concentrations of 60, 70 and 80 ppb; the same benchmark levels used for the 2007 O₃ exp

- 10 concentrations of 60, 70 and 80 ppb; the same benchmark levels used for the 2007 O_3 exposure
- 11 assessment (U.S. EPA, 2007b). Estimating exposures to ambient O_3 concentrations at and above
- 12 these benchmark levels is intended to provide perspective on the public health impacts of O_3 -
- related health effects observed in human clinical and toxicological studies, but that cannot
- 14 currently be evaluated in quantitative risk assessments (e.g., lung inflammation, increased airway
- 15 responsiveness, and decreased resistance to infection). The 80 ppb-8hr benchmark level
- 16 represents an exposure level where there is substantial clinical evidence demonstrating a range of
- 17 O₃-related effects including lung inflammation and airway responsiveness in healthy individuals.
- 18 The 70 ppb-8hr benchmark level reflects evidence that asthmatics have larger and more serious
- 19 effects than healthy people as well as a substantial epidemiological evidence of adverse effects
- 20 associated with O₃ levels that extend below 80 ppb-8hr. The 60 ppb-8hr benchmark level
- 21 represents the lowest exposure level at which O₃-related effects have been observed in clinical
- studies of healthy individuals. See ISA section 6.2.1 for further discussions regarding the body of
 evidence supporting the selection of these benchmark levels.
- The level of exertion of individuals engaged in particular activities is approximated by an equivalent ventilation rate (EVR), that is, ventilation normalized by body surface area (BSA, in
- 25 equivalent ventuation rate (EVR), that is, ventuation normalized by body surface area (DSA, in
- $26 m^2$) and is calculated as VE/BSA, where VE is the ventilation rate in liters/minute. For
- 27 identifying moderate or greater exertion occurring during any 8-hr average exposure period in
- this assessment, we used the lower bound EVR value of 13 (liters/min-m²) based on a range of
- 29 EVRs used by Whitfield et al. (1996) to categorize persons engaged in moderate exertion
- 30 activities for an 8-hr period. Whitfield et al. (1996) developed this range from EVR data reported
- 31 in a 6.6-hr controlled human exposure study conducted by McDonnell et al. (1991).

possible multiple benchmark exceedances could occur for an individual on certain high O_3 concentration days, staff judge this is not a practical output for the purposes of this assessment.

¹⁸ It is worth noting that the adverse health responses in the human clinical studies are generally based on 6.6 hour exposure to O_3 . Therefore, it is possible that the number of benchmark exceedances is underestimated because of the lesser likelihood of an 8-hr exposure above the same threshold due to the longer averaging time.

1 APEX then calculates two general types of exposure estimates for the population of 2 interest: the estimated number of people exposed to a specified O_3 concentration level and, the 3 number of days per O_3 season that they are so exposed; the latter metric is expressed in terms of 4 *person-days.* The former highlights the number of individuals exposed *one or more* times per O_3 5 season at or above a selected benchmark level. The person-days measure estimates the number of 6 times per season the simulated individuals are exposed at or above a selected benchmark level 7 and summed across individuals comprising the population. We note that a person-days metric 8 conflates people and occurrences: one occurrence for each of 10 people would be counted the 9 same as 10 occurrences for one person (i.e., 10 person-days at or above benchmark level). In this 10 assessment we are more interested in reporting *multiday* exposures rather than total person-days, 11 that is, the number of times an individual experiences multiple exposures at or above a 12 benchmark level during an O₃ season. Given the complexities of the exposure modeling, the four 13 study groups considered, the 15 study areas, the 5 years of ambient air quality, the multiple air 14 quality scenarios simulated, and ultimately the output data generated, including both single and 15 multiday exposures for simulated individuals, the consolidation of the results and the related 16 graphic depictions used in this assessment requires additional discussion.

17 To begin, a simple example of summary results is the estimated percent of asthmatic 18 school-age children experiencing exposures at or above a single 8-hr benchmark level when 19 considering base air quality stratified by year (e.g., Figure 5-2, left panel). This presentation 20 largely depicts the variability in O_3 exposure across the 15 study areas within years, along with 21 an illustration of broad year-to-year temporal variability. A general finding regarding temporal 22 variability extracted from this graph would be that fewer asthmatic school-age children exceed 23 daily maximum 8-hr average exposures of 60 ppb considering 2009 base air quality when 24 compared with other simulation years. An observation regarding the spatial variability could 25 include the range of exposures within years (i.e., the study area variability) spans between 15 to 26 35 percentage points, dependent on the particular simulation year (Figure 5-2, left panel). One 27 could also stratify the same exposure results by study area (e.g., Figure 5-2, right panel), thus 28 depicting variability in estimated exposures across years within each study area, along with 29 having broad study area comparisons. A general finding regarding temporal variability in this 30 type of presentation would be that the range of exposures within study areas spans about 20 31 percentage points, though some study areas have a generally small range (<5 percentage points) 32 for most simulated years. An observation regarding spatial variability could be that Chicago 33 largely has the fewest asthmatic school-age children at or above benchmark levels, having a 34 mean about 15%, while Los Angeles consistently has the most asthmatic school-age children, 35 having a mean about 35%, at or above benchmark levels while at moderate or greater exertion. 36





5 While these boxplots are an efficient tool that summarize potentially complex data sets 6 by illustrating important statistical aspects of data analysis results (e.g., means, ranges, 7 occasional upper percentile data values), at times important features of the data may be masked 8 (e.g., trends or patterns within consolidated variables) and the presentation of other aspects of the 9 exposure results would require the generation of additional graphs (e.g., results for additional 10 benchmark levels). A tabular format could be one way to present all possible data, though given 11 the number of APEX simulations performed (i.e., > 1,000) and aforementioned dimensions of 12 the assessment, linking the trends and patterns across all study areas, years, and benchmark 13 levels from the numerous output tables would be visually challenging. 14 This discussion regarding properly representing temporal and spatial variability in the 15 exposure results can be further extended to include the added dimension of the five air quality 16 scenarios (base, existing standard, and three alternative standard levels). Mindful of these

17 complexities, we elected to use a multi-panel graphing approach to succinctly summarize the

18 exposure output data, while also retaining as much information as possible in a single page

19 format to allow for visual analysis of trends and patterns. As an example, Figure 5-3 (top panels)

1 illustrates boxplots for Atlanta similar to those presented above, though the exposure results are 2 for the three exposure benchmark levels of interest, with each stratified by the particular adjusted 3 air quality scenario. As expected with increasing stringency of the 8-hr standard level, fewer 4 asthmatic school-age children are exposed at or above a given benchmark level. Also expected is 5 the fewer percent of asthmatic school-age children exposed to higher benchmark levels when compared with lower benchmark levels. While these three graphs can provide a clear depiction 6 7 of the exposure results for a single study area, the six years encompassing the two averaging 8 periods 2006-2008 and 2008-2010 are combined in the graphic and difficulty would remain in 9 simultaneously exhibiting all 15 areas. 10

To overcome these limitations, Figure 5-3 (lower panel) exhibits all of the dimensions of 11 the exposure results mentioned above (i.e., year, benchmark level, study area) along with 12 distinguishing between the two standard averaging periods for each the existing (75 ppb-8hr) and 13 alternative standard levels (60, 65, 70 ppb-8hr). The nomenclature above each subgraph 14 indicating the particular air quality scenario requires defining. For example, a panel heading of 15 "75" contains the exposures estimated when air quality was adjusted to just meet the existing standard level of 75 ppb-8hr (4th highest daily maximum 8-hr average O₃ concentration averaged 16 17 over a three year period) either using years 2006, 2007, and 2008 ambient air quality data or for a 18 second averaging period that extended from 2008 through 2010 (with results for each given by 19 two separate lines on the same plot). Exposure results are readily observed for any air quality 20 scenario, year, or benchmark level of interest. For example, when considering the 75ppb 21 standard 2006-2008 averaging time scenario, 20% of asthmatic school-age children in Atlanta 22 experience at least one daily maximum 8-hr average exposure of 60 ppb occurs when 23 considering year 2006 air quality, while only about 5% experience exposures at or above the 24 same benchmark level considering 2008 air quality (though when considering the 2008-2010 25 averaging period, approximately 20% of asthmatic children are estimated experience at least one 26 exposure at or 60 ppb-8hr). Fewer than 5% of asthmatic school-age children in Atlanta 27 experience at least one benchmark exposure of 70 ppb-8hr considering any year and any air 28 quality scenario, including just meeting the existing O₃ standard. 29

Because APEX simulates the complete time series of exposure for every simulated individual, also output is the number of times an individual experiences a benchmark exceedance over the duration of the simulation (i.e., the entire O₃ season simulated in each study area). These data can also be summarized in a similar multi-panel format, though differ slightly in composition from that of Figure 5-3. Instead of displaying the percent of persons with at least one exceedance of each of the three benchmarks, presented are the percent of persons with multiple exposures at or above a single benchmark within an O₃ season. For example, Figure 5-4 illustrates the percent of asthmatic school-age children in Atlanta having multiple days where

1 exposures ($\geq 2, \geq 4$, and ≥ 6 per O₃ season) were at or above 60 ppb-8hr considering the 2006-2 2010 air quality adjusted to just meet the existing and alternative standards levels. When 3 considering 2006 air quality adjusted to just meet the existing standard, approximately 10% of 4 asthmatic school-age children experienced at least two days where their daily maximum 8-hr 5 average exposure was at or above 60 ppb, though fewer than 5% experienced such exposures in 6 2009. When collectively considering all simulated air quality scenarios and years, fewer than 3% 7 of asthmatic school-age children experienced at least four exposures at or above 60 ppb and 8 virtually no asthmatic school-age children experienced six or more such exposures over the O_3 9 season.

10



11

Figure 5-3 Percent of asthmatic school-age children in Atlanta with at least one O₃ exposure at or above 60 ppb-8hr (left top panel), 70 ppb-8hr (middle top panel), and 80 ppb-8hr (right top panel while at moderate or greater exertion, years 2006-2010 air quality adjusted to just meet the existing and alternative O₃ standard levels. The multi-panel display (bottom) illustrates the same exposure results expanded to reflect individual data points by year, standard averaging period, and benchmark level.

18

19 Also worth discussing is the appearance of a similar pattern between the benchmark level 20 results (Figure 5-3) and the number of exceedances of a single benchmark (Figure 5-4). Because 21 the ambient concentration is an important determinant in exposure concentrations, it is not 22 surprising to see that the trend over years for persons having at least one exposure at or above a 23 particular benchmark level (e.g., 60 ppb-8hr) is similar to those experiencing at least two 24 exposures above 60 ppb-8hr (though a smaller percentage of persons). This is because years 25 having the highest peak concentrations will yield the greatest percent of persons above 26 benchmark levels, and when one year has a day with the highest concentration, it is likely that

- 1 year also has a second day with a similarly and relatively high concentration, and so on. Using
- 2 the same logic, one might also conclude that there could be a pattern between the percent of
- 3 persons experiencing a single exceedance of 70 ppb-8hr and multiple exceedances (e.g., four) of
- 4 60 ppb-8hr also driven by the overall ambient concentration distribution. However, given that
- 5 very few persons experience these types of benchmark exceedances, determining the relationship
- 6 between the two (if present) may not be of practical significance. For brevity, the complete
- 7 multiday exposure results for all APEX simulations are presented in Appendix 5F, with results
- 8 presented for one study group (e.g., all school-age children) in the main body of the REA.
- 9



Figure 5-4 Percent of asthmatic school-age children in Atlanta with multiple O₃ exposures at or above 60 ppb-8hr while at moderate or greater exertion, years 2006-2010 air quality adjusted to just meet the existing and alternative O₃ standard levels.

14

15 **5.3 EXPOSURE ASSESSMENT RESULTS**

16 **5.3.1 Overview**

17 The results of the exposure analysis are presented as a series of figures focusing on the 18 defined range of benchmark levels (i.e., persons experiencing daily maximum 8-hr average O_3 19 exposure concentrations at or above 60 ppb, 70 ppb, and 80 ppb), noted as being of particular 20 health concern (Section 5.2.8). A range of concentrations in the air quality data over the five year 21 period (2006-2010) were used in the exposure model, providing a range of estimated exposures 22 output by APEX. The adjusted air quality was developed using two distinct 3-year period design values (2006-2008 and 2008-2010), as described in Chapter 4.¹⁹ Exposures were estimated for 23 24 four study groups of interest (i.e., all school-age children (5-18), asthmatic school-age children, 25 asthmatic adults (19-95), and older adults (65-95)) in each of the 15 study areas. 26 In this exposure assessment, we are primarily interested in O_3 exposures associated with

27 the ambient air quality adjusted to just meet the existing and potential alternative O₃ standards.

¹⁹ Thus, the year 2008 will have two sets of estimated exposures, one from each of the two sets of design values. In Figure 5-2, the greater temporal variability observed for 2008 is driven in part by differences in some study areas resulting from the air quality adjustment period. Exposure results for both 2008 averaging periods are provided when presenting data by year. Where mean results are presented in subsequent results sections, the two values given for year 2008 were first averaged to give a single exposure value for 2008 before averaging across all years.

1 Thus, most of the exposure results presented and discussed are for where ambient air quality was

2 adjusted to just meet these particular scenarios. While understanding exposures and health risks

3 associated with historical and existing air quality is important, the primary goal of this and any

4 REA is to evaluate to what extent the existing NAAQS, and its associated air quality, protect

5 health and to what extent alternative NAAQS protect health. Exposure results associated with

6 recent (base) air quality are briefly discussed here first, though largely reported in Appendix 5F.

7

5.3.2 Exposure Modeling Results for Base Air Quality

8 The exposure results for the base air quality are distinguished from the other air quality 9 scenario results primarily due to the wide ranging variability in estimated exposures across the 10 study areas and years. The variability in exposures are the result of the wide ranging variability 11 in ambient concentration levels, with perhaps some years in some study areas exhibiting air 12 quality at or near that just meeting the current 8-hr standard, while other study areas and years 13 exhibiting air quality levels much higher than the existing 8-hr standard. These exposures are 14 informative in describing the existing or recent health risks associated with a unique air quality 15 scenario, but because they variably diverge from a set concentration level of interest (such as the 16 existing 8-hr standard), they are of limited relevance in evaluating the adequacy of either the 17 existing NAAQS as well as potential alternative air quality standards. That said, detailed tabular 18 and graphic presentations of exposure results associated with the base air quality (years 2006-19 2010) are provided in Appendix 5F, with only key findings summarized in the following 20 discussion.

21 Consistent with the previously discussed observations regarding year-to-year variability 22 in ambient concentrations (Chapter 4), most study areas have the greatest percent of all school-23 age children experiencing concentrations at or above the three benchmark levels during 2006 or 24 2007 along with having the lowest percent of all school-age children exposed during 2009. In 25 general, between 20-40% of all school-age children experience at least one O_3 exposure at or 26 above 60 ppb-8hr, 10-20% experience at least one O₃ exposure at or above 70 ppb-8hr, and 0-27 10% experience at least one O_3 exposure at or above 80 ppb-8hr, all while at moderate or greater exertion (i.e., an 8-hr EVR \geq 13 L/min-m²) and considering the base air quality (2006-2010). 28 29 Year-to-year variability observed for asthmatic school-age children and the percent of asthmatic 30 school-age children were similar to exposure results for all school-age children, largely a 31 function of having both simulated study groups using an identical time-location-activity diary 32 pool to construct each simulated individual's time series of activities performed and locations 33 visited. 34 The overall year-to-year pattern of exposure for asthmatic adults is similar to that

35 observed for all school-age children, though the percent of the asthmatic adult study group

1 exposed is lower by a factor of about three or more. Having a lower percent of asthmatic adults 2 exposed is expected given that outdoor time expenditure is an important determinant of O_3 3 exposure (section 5.4.2) and that adults spend less time outdoors than children (section 5.4.1), as 4 well as adults having a lower outdoor participation rate. The percent of all older adults 5 experiencing exposures at or above the selected benchmark levels is lower by a fewer percentage 6 points when compared with the results for asthmatic adults. Again, older adults, on average, 7 would tend to spend less time outdoors and do so with less frequency when compared with both 8 adults and children (section 5.4.1), in addition to fewer older adults performing activities at 9 moderate or greater exertion for extended periods of time, thus leading to fewer persons exposed 10 to O_3 concentrations of concern.

11 The year-to-year patterns of the single and multiple exposure occurrences considering 12 base air quality (2006-2010) were similar among the four exposure study groups, therefore only 13 results for all school-age children will be summarized here. Depending on the year and study 14 area, about 10-25% of all school-age children could experience at least two exposures above the 15 60 ppb-8hr benchmark during the O₃ season, while about 5-10% school-age children could 16 experience at least four. Most study areas and years are estimated to have fewer than 5% of all 17 school-age children experience six or more exposures above 60 ppb-8hr considering the base air 18 quality. When considering the multi-day exposures for all school-age children at or above the 70 19 ppb-8hr benchmark, about 2-10% of all school-age children could experience at least two 20 exposures during the O_3 season, while four or more exposures were generally limited to fewer 21 than 4% of all school-age children. Almost half of the study area-year combinations had no 22 school-age children experiencing two or more exposures at or above the 80 ppb-8hr benchmark, 23 with the other half estimated to have about 1% of all school-age children experiencing two or 24 more exposures at or above the 80 ppb-8hr benchmark. School-age children having four or more 25 80 ppb-8hr benchmark exceedances were limited to only a few study area years and, where a 26 non-zero value was estimated, were limited to $\leq 0.5\%$ of the study group.

5.3.3 Exposure Modeling Results for Simulations of Just Meeting Existing and Alternative O₃ Standards

29 In this section, we present the exposures estimated when considering the air quality 30 adjusted to just meeting the existing O₃ NAAQS standard, as well as when considering potential 31 alternative standard levels (55, 60, 65, 70 ppb 8-hr) of the existing standard. Comprehensive 32 multi-panel displays of exposure results are presented for each of the study groups of interest, 33 i.e., all school-age children (5-18), asthmatic school-age children, asthmatic adults (19-95), and 34 all older adults (ages 65-95; Figure 5-5 to Figure 5-8, respectively). Included in each display are 35 the three benchmark levels (60, 70, and 80 ppb-8hr), the five years of air quality (2006-2010), for 36 the 15 study areas. A single multi-panel display is used to present the results for each of the four

study groups, beginning with the estimated percent of persons exposed at least one time at or above the selected benchmark levels. Modeled exposures in the 15 study areas and considering each benchmark level are presented on the same scale to allow for direct comparisons across the multi-panel display. The most notable patterns in the exposure results are described here using one study group (i.e., all school-age children), as there is a general consistency in the year-to-

6 year variability within each study area across all four study groups. Any deviation from the

7 observed pattern will be discussed for the subsequent study group.

8 We note that after adjusting to just meet a potential 8-hr ambient standard level of 55 9 ppb, there were nearly no persons exposed at or above any of the selected benchmark levels, thus 10 these data, while modeled, are not presented in detail here. In addition, in one study area 11 (Chicago), O₃ ambient monitor design values were below that of the existing standard during the 12 2008-2010, therefore APEX simulations could not be performed for meeting the existing 13 standard for that 3-year period. And finally, we were not able to simulate just meeting a standard 14 level of 60 ppb-8hr or below in the New York study area (see Chapter 4 for details), thus APEX 15 simulations for these air quality scenarios could not be performed in New York.

16 Figure 5-5 illustrates the exposures estimated for all school-age children in each study 17 area with general observations as follows. After adjusting air quality to just meet the existing and 18 alternative standards, there are virtually no school-age children exposed at or above 80 ppb-8hr, 19 with very few school-age children exposed at or above the 70 ppb-8hr benchmark. For example, 20 out of 87 possible study area and year combinations considering air quality adjusted to just meet 21 the existing standard (the least stringent standard level considered here), only 29 resulted in >22 0.1% estimated percent of all school-age children exposed at least once at or above the 80 ppb-23 8hr benchmark with the maximum percent of all school-age children exposed estimated for St. 24 Louis (1.1%). Ninety-four percent of study area and year combinations had fewer than 5% of all 25 school-age children experiencing at least one daily maximum 8-hr average exposure ≥ 70 ppb 26 considering ambient air quality adjusted to just meeting the existing standard, again with a 27 maximum of 8.1% occurring in St. Louis. When considering air quality adjusted to just meet an 28 8-hr ambient standard level of 70 ppb, $\leq 0.2\%$ of all school-age children experience at least one 29 80 ppb-8hr exposure benchmark exceedance for all study area and year combinations, while for 30 76 or 90 study area and year combinations, $\leq 1\%$ of all school-age children experience a 70 ppb-31 8-hr exposure benchmark exceedance. This pattern of having very few school-age children 32 experiencing exposures at or above 70 and 80 ppb-8hr is as expected given the nature of the air 33 quality adjustment procedure that limits 8-hr ambient concentrations at or above the selected 34 potential alternative standard level. 35 In contrast, approximately 10-20% percent of all school-age children are estimated to be

an contrast, approximately 10-20% percent of all school-age children are estimated to be
 exposed to at least one 60 ppb-8hr concentration when considering air quality just meeting the

associated with the base air quality, a general year-to-year exposure pattern emerges with respect
to study area and year. For the Northeastern (Boston, New York), Mid-Atlantic (Philadelphia,
Washington DC, Cleveland) and Mid-Western (Chicago, Detroit, and St. Louis) study areas, the
maximum percent of all school-age children exposed generally occurs during year 2007. For the
Southern (Atlanta, Dallas, Houston) and Western (Denver, Los Angeles, Sacramento) study

existing standard (Figure 5-5). And similar to that mentioned above regarding exposures

7 areas, the maximum exposure occurs during year 2006. Deviations from this temporal exposure

8 pattern appear mostly as a result of the standard averaging period, with the 2008-2010 period

9 producing equal or greater maximum exposures during either 2008, 2010, or both years and most

10 prevalent in the Northeastern and Mid-Atlantic study areas (Baltimore, Boston, New York,

11 Philadelphia, Washington DC; note also a trend in Atlanta, Denver, St. Louis).

1

12 These 60 ppb-8hr exposure patterns remain consistent when considering air quality 13 adjusted to just meet a 70 ppb-8hr ambient standard, though the percent of all school-age 14 children exposed is less than that observed when considering the air quality adjusted to just meet 15 existing standard. Further, 75 of 90 study area and year combinations are estimated to have \leq 16 10% of all school-age children experience a 60 ppb-8hr or greater exposure, though between 10-20% of all school-age children were estimated to be exposed for a few study area and year

18 combinations (e.g., Atlanta-2006, Chicago-2007 and -2010, and Houston-2009). When

considering air quality adjusted to just meet a 65 ppb standard level, the percent of all school-age
children experiencing an exposure at or above 60 ppb-8hr diminishes to 5% or less for most
study areas and years (i.e., 81 of 90 study area year combinations).

All of what has been described regarding the estimated exposures to school-age children (i.e., the year-to-year and benchmark level patterns, and the percent of the study group exposed) also applies to the exposures estimated for asthmatic school-age children (Figure 5-6). Different however would be the relative number of asthmatic school-age children exposed in each study area if compared with all school-age children, as the asthma prevalence rates vary by study area (Table 5-2), though on average are about 10% of the population of children.

28 The percent of asthmatic adults (Figure 5-7) experiencing daily maximum 8-hr average 29 exposures above the selected benchmark levels is sharply lower than that estimated for all 30 school-age children. For example, only three of a possible 84 study area and year combinations 31 (Chicago-2007, Houston-2009, and St. Louis-2007) were estimated have > 0.1% of asthmatic 32 adults experience a daily maximum 8-hr average exposure ≥ 80 ppb, and only six of a possible 33 84 study area and year combinations were estimated have >1% of asthmatic adults experience an 34 daily maximum 8-hr average exposure \geq 70 ppb, all occurring when considering air quality just 35 meeting the existing standard. No study area or year combination has more than 10% of 36 asthmatic adults estimated to experience an exposure at or above 60 ppb-8hr when considering

air quality just meeting the existing standard, with 67 of 84 study area and year combinations
 estimated to have 5% or less asthmatic adults experiencing such exposures.

3 When considering air quality adjusted to just meeting a standard level of 70 ppb-8hr, no 4 asthmatic adults experience an exposure at or above 80 ppb-8hr and $\leq 0.6\%$ experience a daily 5 maximum 8-hr average exposure ≥ 70 ppb for any study area or year combination. Less than 5% 6 of asthmatic adults could experience an exposure at or above 60 ppb-8hr when considering air 7 quality adjusted to just meet a standard level of 70 ppb-8hr for 88 or 90 possible study area year 8 combinations, with the maximum percent of adult asthmatics exposed outside this range 9 occurring in Denver (6.8%-2008) and St. Louis (5.5%-2007).

10 Older adults are estimated to have the fewest exposures above the two highest benchmark 11 levels when considering the adjusted air quality. For example, only two of a possible 84 study 12 area and year combinations (St. Louis-2007 and Washington DC-2008) were estimated have > 13 0.1% of asthmatic adults experience a daily maximum 8-hr average exposure \geq 80 ppb, and only 14 six of a possible 84 study area and year combinations were estimated have > 1% of asthmatic adults experience a daily maximum 8-hr average exposure \geq 70 ppb, all occurring when 15 16 considering air quality just meeting the existing standard (Figure 5-8). Also, exceeding the 60 17 ppb-8hr exposure benchmark appears to be limited to fewer than 5% of all older adults when 18 considering air quality adjusted to just meet the existing standard and a standard level of 70 ppb-19 8hr, and occurs in < 2% of all older adults when considering a standard level of 65 ppb-8hr.

20 An example of multi-day exposure results associated with adjusted air quality is provided 21 in Figure 5-9. The percent of all school-age children estimated to experience multi-day exposures 22 above benchmark levels during each study area's O₃ season is largely limited to two air quality 23 scenarios: the existing standard and air quality adjusted to just meeting a standard level of 70 24 ppb-8hr. This is because of the small percent of school-age children experiencing even a single 25 exposure above the lowest benchmark level when considering standard levels at or below 65 26 ppb-8hr. In addition, when experiencing multiple exposures, most school-age children appear to 27 have at most two days above benchmark levels per O₃ season, even when considering the lowest 28 benchmark level of 60 ppb-8hr. For example, 81 of 87 possible study area and year combinations 29 have < 10% of all school-age children experiencing two or more exposures \geq 60 ppb-8hr when 30 considering an ambient standard level of 75 ppb-8hr, while 83 of 90 possible study area and year 31 combinations have < 5% of all school-age children experiencing two or more exposures ≥ 60 32 ppb-8hr when considering an ambient standard level of 70 ppb-8hr. With increasing stringency 33 in the standard level to 65 ppb-8hr, 81 of 90 possible study area and year combinations have < 34 1% of all school-age children experiencing two or more exposures ≥ 60 ppb-8hr. 35 Multi-day exposure to the higher exposure benchmarks (either the 70 or 80 ppb-8hr) is a 36 rare occurrence, even when considering the air quality adjusted to the existing O_3 standard. For

5-28

- 1 example, there were no school-age children experiencing two or more exposures above 80 ppb-
- 2 8hr in all but one study area year combination and, and when considering that one study year
- 3 having a non-zero value (St. Louis-2007), the estimated percent of all school-age children at or
- 4 above the exposure benchmark was only 0.1%. Further, 83 of 87 possible study area and year
- 5 combinations have < 1% of all school-age children experiencing two or more exposures ≥ 70
- 6 ppb-8hr, also when considering an ambient standard level of 75 ppb-8hr.
- 7
- 8



Figure 5-5 Percent of all school-age children with at least one daily maximum 8-hr average O₃ exposure at or above 60, 70, and 80 ppb while at moderate or greater exertion, years 2006-2010, air quality adjusted to just meet the existing and potential alternative standards.



Figure 5-6 Percent of asthmatic school-age children with at least one daily maximum 8-hr average O₃ exposure at or above 60, 70, and 80 ppb while at moderate or greater exertion, years 2006-2010, air quality adjusted to just meet the existing and potential alternative standards.



Figure 5-7 Percent of all asthmatic adults with at least one daily maximum 8-hr average O₃ exposure at or above 60, 70, and 80 ppb-8hr while at moderate or greater exertion, years 2006-2010, air quality adjusted to just meet the existing and potential alternative standards.



Figure 5-8 Percent of all older adults with at least one daily maximum 8-hr average O₃ exposure at or above 60, 70, and 80 ppb-8hr while at moderate or greater exertion, years 2006-2010, air quality adjusted to just meet the existing and potential alternative standards.



Figure 5-9 Percent of all school-age children with multiple daily maximum 8-hr average O₃ exposures at or above 60 ppb while at moderate or greater exertion, years 2006-2010, air quality adjusted to just meet the existing and potential alternative standards.

15.4TARGETED EVALUATION OF EXPOSURE MODEL INPUT AND OUTPUT2DATA

3 This section summarizes the results of several targeted evaluations intended to provide 4 additional insights to APEX input data or approaches used to estimate exposures (CHAD data 5 attributes and activity pattern evaluations, comparison of CHAD outdoor time data with ATUS, 6 comparisons of asthmatic outdoor time expenditure and exertion levels to that of non-7 asthmatics), exposure results for additional exposure populations of interest (outdoor workers, 8 school-age children during summers, impact of averting), and model performance evaluations 9 (personal exposure measurements and independent ventilation rate estimates compared with 10 APEX estimates). Detailed analysis results are provided in Appendix 5G.

11 5.4.1 ANALYSIS OF TIME-LOCATON-ACTIVITY DATA

12 While CHAD is the most comprehensive and relevant source of time-location-activity 13 data available for use in our exposure modeling, there are a few limitations to the survey data 14 contained therein, many of which are founded in the individual studies from which activity 15 patterns were derived (Graham and McCurdy, 2004). CHAD is a collection of related survey 16 data, though individual study attributes can range widely (e.g., survey participant ages, region or 17 city of residence, time-of-year data collected). We note that many of the assumptions about use 18 of these activity patterns in exposure modeling are strengthened by the manner in which they are 19 used by APEX. This is done by focusing on selecting the most important individual attributes 20 that contribute to variability in human behavior (e.g., age, sex, day-of-week, ambient 21 temperature) and linking these attributes of simulated individuals to the population demographics 22 of each census tract (see section 5.2.5) and the study area temperatures (section 5.2.4). Further, 23 one key lifestyle attribute is also accounted for in generating longitudinal diary profiles by 24 simulating both the intra- and interpersonal variability in time spent outdoors (section 5.2.5; Glen 25 et al., 2008).

A few questions may arise as to the representativeness of the CHAD diaries to the simulated population. For example, the year of a particular survey study may differ from our simulated exposure population by as much as 30 years (i.e., some activity pattern data were generated in the 1980s). In addition, there are other personal attributes (e.g., ethnicity, income level, lifestyle factors²⁰), health conditions (e.g., asthma, cardiovascular disease), and situational factors (e.g., availability of parks and recreation areas) that are not used in creating the simulated persons that could be influential in estimating exposures. Considering this, a number of

²⁰ Examples of such factors for adults could include married/unmarried, having infants or young children/no children. Lifestyle factors for children could include whether the child is active/non-active or whether or not there is time spent outdoors.
evaluations were performed to answer questions regarding important personal attributes used in
 generating simulated individuals and the general representativeness of the CHAD time-location activity data. First though, we summarize the newly acquired activity pattern data now included
 in CHAD compared with data available and used in the 1st draft O₃ REA.

5 5.4.1.1 General Evaluation of CHAD Study Data: Historical and Recently Acquired Data

6 The number of diary days having complete information and used by APEX in the 2^{nd} 7 draft O₃ REA is 41,474 (Table 5-3). This is an increase of about 8,700 diaries currently used by 8 APEX compared with what was used by APEX in the 1st Draft O₃ REA. Further, there have been 9 eight new study data sets incorporated into CHAD and used in our current exposure assessment 10 since the previous O₃ NAAQS review conducted in 2007, most of which were from recently 11 conducted activity pattern studies (see Appendix 5B, Section 5B-4 for more information 12 regarding these studies). The diary data included from these new studies have more than doubled 13 the total activity pattern data used for 2007 O₃ exposure modeling and has increased the number 14 of children's diaries by about a factor of five. Currently, the majority of diaries (54%) from 15 CHAD are taken from surveys conducted in the past decade, while the pre-1990s diaries 16 represent less than 15% of the total diaries available by APEX.

17 5.4.1.2 Exposure-Relevant Personal Attributes Included in CHAD and APEX Simulated 18 Individuals

19 The survey participants whose diary data are within CHAD were asked a number of 20 questions regarding their personal attributes. The number and type of attributes present for 21 diaries in CHAD is driven largely by the original intent of the individual study. In our exposure 22 assessment, we have strict requirements to simulate individuals using several personal attributes, 23 namely age, sex, temperature (as a surrogate for seasonal variation in activity patterns), and day-24 of-week. These attributes are considered as important drivers influencing daily activity patterns 25 (Graham and McCurdy, 2004) and when diaries do not have these particular attributes for a 26 particular day, the diary day will not be used by APEX. We compared the representation of these 27 and other attributes in the current CHAD used by APEX with that in the 1st draft O₃ REA and 28 found strong similarities in the attribute distributions between both databases, suggesting little 29 change in the overall composition of the database regarding these influential attributes. 30 While there may be other personal or situational attributes that affect daily time 31 expenditure (e.g., socioeconomic status, occupation of an employed person), these attributes are 32 typically not included in our assessment to generate simulated individuals simply because the

33 response to the attribute is missing for most of the study participants/CHAD diary days. For

34 example, income level is missing for about two-thirds of the CHAD diaries because either the

35 original study did not have an income/occupation related survey question or perhaps the

36 participant refused to answer the question if it were posed. If one were to select this personal

1 attribute in developing a simulated individual's activity pattern (among using any other attribute

- 2 having missing responses), the pool of diaries available to simulate individuals may be extremely
- 3 limited, likely leading to repetition of diaries used for individuals or groups of similar individuals
- 4 and artificially reducing both intra- and inter-personal variability in time expenditure, or perhaps
- 5 resulting in model simulation failure altogether. This is why personal attributes are carefully
- 6 selected and prioritized according to both their prevalence in CHAD and whether the attribute
- 7 has a known significant influence on activity patterns.

8 5.4.1.3 Evaluation of Afternoon Time Spent Outdoors for CHAD and Survey Participants

9 There have been questions raised regarding the representativeness of the diaries from 10 studies conducted in the 1980s and whether there are any recognizable patterns in time 11 expenditure in the CHAD diaries across the time period when data were collected. Because time 12 spent outdoors is a significant factor influencing daily maximum 8-hr average O_3 exposures, we 13 evaluated the current collection of CHAD diaries used by APEX for two metrics and considering 14 two dimensions: outdoor participation rate (i.e., the percent of people who spent some time 15 outdoors during their survey day) and the mean time spent outdoors for where the persons spent 16 at least one minute outdoors or at least 2 hours outdoors. Because time spent outdoors is an 17 important determinant for highly exposed individuals, we summarize the results here for the 18 diaries having at least 2 hours of outdoor time here, while all other results are provided in 19 Appendix 5G. CHAD diaries were stratified by five age groups (4-18, 19-34, 35-50, 51-64, 65+) 20 and three decades (1980s, 1990s, and 2000s) using the year the particular activity pattern study 21 was conducted. We note that CHAD is composed of primarily cross-sectional data (single diary 22 days per person), thus the trend evaluated over the three decades is changes (if any) in 23 participation rate and the time spent outdoors by the composite study population, not within 24 individuals.

25 Regardless of decade and duration of time spent outdoors, children tended to have the 26 highest outdoor participation rate when compared with the other age groups, while the oldest 27 adults (aged 65 or greater) tend to have the lowest participation rate. The CHAD diaries from the 28 1980's studies for children ages 4-18 have the highest outdoor participation rate (50%) compared 29 to other decades (35-40%) and all other age groups and decade of collection. When considering 30 the pool of diaries available for this age group, these 1980's studies contribute to approximately 31 19% of diaries having two or more hours of time spent outdoors during the afternoon. This 32 translates to a small effect on the overall outdoor participation rate for diary pools that would 33 include these earlier studies (39% participation rate) compared to the participation rate excluding 34 these studies (36% participation rate). In general, these outdoor participation rates are similar to 35 the finding reported recently by Marino et al. (2012) of 37.5%, though estimated for pre-school 36 age children. Thus, when considering participation in outdoor activities and the

representativeness of the CHAD study data from the 1980s, it is unlikely that use of these oldest
 diaries would strongly influence exposure model estimates.

3 There is variability in the amount of outdoor time evaluated over the three decades, with 4 diaries from the 2000's studies exhibiting perhaps the lowest range of mean outdoor time (190-5 220 min/day) compared with the 1980's (210-240 min/day) and 1990's (212-258 min/day) 6 studies, a trend perhaps most notable when considering the children's diaries (a decrease in time 7 spent outdoors of about 30 minutes over the three decades). However, the coefficient of variation 8 (COV) for each of the age groups and across all decades for the cross-sectional data was 9 consistently about 40%, supporting a general conclusion of no appreciable differences in the 10 mean time spent outdoors over the three decades of data collection. Thus, when considering all 11 diaries having at least 2 hours of afternoon outdoors time and the representativeness of the 12 CHAD study data from the 1980s, inclusion of these earlier diaries is also unlikely to have a 13 strong adverse influence on exposure modeling outcomes. Though combined with the higher 14 participation rate for these earlier diaries, exposures estimated using these diaries may be higher 15 than when estimated when excluding these diaries from CHAD.

16 5.4.1.4 Evaluation of Afternoon Time Spent Outdoors for ATUS Survey Participants

We evaluated recent year (2002-2011) time expenditure data from the American Time Use Survey (ATUS) (US BLS, 2012). As was done with the CHAD data set, the purpose was to evaluate trends (if any) in outdoor time over the period of time data were collected. A few strengths of the ATUS data are (1) its recent and ongoing data collection efforts, (2) large sample size (totaling over 120,000 diary days), (3) national representativeness, and (4) that varying diary approaches would not be an influential or confounding factor in evaluating trends over time.

23 ATUS does however have a few noteworthy limitations when compared with the CHAD 24 data: (1) there are no survey participants under 15 years of age, (2) time spent at home locations 25 is neither distinguished as indoors or outdoors, and (3) missing or unknown location data can 26 comprise a significant portion of a persons' day (on average, about 40% (George and McCurdy, 27 2009)). To overcome the limitation afforded by the ambiguous home location, we identified 28 particular activity codes most likely to occur outdoors (e.g., participation in a sport) to better 29 approximate each ATUS individual's outdoor time expenditure. Missing time was circumvented 30 by our focused analysis: about 85% of missing time information occurs outside of the hours of 31 interest here (i.e., before 12:00 PM and after 8:00 PM). Data were stratified by the same five age 32 groups as was done for the CHAD data, though here the time trends were assessed over 33 individual survey years.

When considering person-days having at least 2 hours of time spent outdoors, there were no clear trends over the nine year ATUS study period regarding either the participation rate or the mean time spent outdoors for any of the age groups. Consistent with CHAD, the participation

1 rate of children was greater than that of the other age groups. The range in ATUS diary outdoor 2 participation rate (10-20%) for all age groups is lower than that observed for the CHAD data 3 (generally between 20-40%), while the range in mean time spent outdoors (190-240 minutes per 4 day) was similar to that of the CHAD data. The lower participation rate for ATUS participants is 5 not surprising given the lack of distinction regarding time indoors and outdoors while at home 6 for ATUS participants and possibly influenced in part by not having any activity patterns for 7 children under 15 years old. Overall, results of the ATUS data analysis generally support the 8 representativeness of the CHAD data, and while participation in outdoor activities calculated 9 using ATUS diaries was less than CHAD diaries, ATUS survey methods obfuscate the strength

10 of this finding.

5.4.1.5 Evaluation of Outdoor Time and Exertion Level for Asthmatics and Non-Asthmatics in CHAD

13 Due to limited number of CHAD diaries with survey requested health information, all 14 CHAD diaries are assumed appropriate for any APEX simulated individual (i.e., whether 15 asthmatic, non-asthmatic, or no compromising health condition was indicated), provided they 16 concur with age, sex, temperature, and day-of-week selection criteria. In general, the assumption 17 of modeling asthmatics similarly to healthy individuals (i.e., using the same time-location-18 activity profiles) is supported by the activity analyses reported by van Gent et al. (2007) and 19 Santuz et al. (1997), though other researchers, for example, Ford et al., (2003), have shown 20 significantly lower leisure time activity levels in asthmatics when compared with persons who 21 have never had asthma. To provide additional support to the assumption that any CHAD diary 22 day can be used to represent the asthmatic population regardless of the study participants' 23 characterization of having asthma or not, we first compared participation in afternoon outdoor 24 activities at elevated exertion levels among asthmatic, non-asthmatic, and unknown health status 25 using the CHAD diaries. We then compared compatible CHAD diary days with literature 26 reported outdoor time participation at varying activity levels.

27 In the first comparison, participation in afternoon outdoor activities for non-asthmatic 28 children and adults in CHAD were found similar when compared with their respective asthmatic 29 cohorts (both about 40-50%). Outdoor participation rate for persons having unknown asthma 30 status, a smaller fraction of the total diaries, varied $\pm 10\%$ from that having known asthma status 31 (children were higher, adults were lower). The amount of time spent outdoors by the persons that 32 did so varied little across the two populations and three asthma categories. On average, CHAD 33 diaries from children indicate approximately 2¹/₄ hours of afternoon time is spent outdoors, 80% 34 of which is at a moderate or greater exertion level, again regardless of their asthma status, known 35 or unknown. Slightly less afternoon time is spent outdoors by adults when compared with 36 children, and while their participation in moderate or greater exertion level activities is much less 1 (about 63%), there was little difference between asthmatic adults and non-asthmatic adults2 considering outdoor time or percent at moderate or greater exertion.

3 For the second comparison, the percentage of waking hours outdoors at varying activity 4 levels for asthmatics reported in three independent asthma activity pattern studies (Shamoo et al., 5 1994; EPRI, 1988; EPRI 1992) were compared to CHAD diary days having similar personal 6 attributes and stratified by asthma status. The range in the percent of waking hours outside at 7 moderate activity level for CHAD diaries was similar to that estimated using the three 8 independent literature sources (2-10%), however the range in percent of outdoor time associated 9 with strenuous activities using the CHAD asthmatic diaries extends beyond that of asthmatic 10 persons from the three independent studies by about a factor of two higher. At this time, the 11 reason for this difference is unknown. Overall, given the above mentioned similarities in outdoor 12 time, participation, and activity levels, use of a CHAD diary regardless of a persons' asthma 13 condition is reasonably justified based on the available data analyzed.

14 **5.4.2** Characterization of Factors Influencing High Exposures

15 We investigated the factors that influence persons experiencing the highest daily 16 maximum 8-hr average exposures. These exposure results in six selected study areas, Atlanta, 17 Boston, Denver, Houston, Philadelphia, and Sacramento, considering base air quality and air 18 quality just meeting the existing standard were combined with each simulated individual's 19 microenvironmental time expenditure during the afternoon hours (12:00 PM through 8:00 PM), 20 times of day commonly when daily peak high O_3 concentrations occur. We first evaluated the relative contribution seven variables²¹ had on the total explained variability in daily maximum 8-21 22 hr average exposures. We then evaluated the distribution of identified influential variables for 23 simulated individuals with the highest exposures. And finally, we identified the 24 microenvironmental locations highly exposed persons occupied and the activities performed 25 within them, given that within an 8-hr time frame most persons would likely visit multiple 26 locations and perform different activities. 27 When considering only person days having the highest daily maximum 8-hr average O_3 28 exposures at any of the six study areas and either air quality scenario and age groupings,

- 29 collectively the main effects of ambient concentrations and outdoor time combined with their
- 30 interaction similarly contribute to approximately 80% of the total explained variance results,
- 31 suggesting that for highly exposed persons, the most important influential factors are time spent
- 32 outdoors corresponding with high daily maximum 8-hr average ambient O₃ concentrations.

²¹ The seven variables include the main effects of (1) daily maximum 8-hr ambient O₃, (2-4) afternoon time spent outdoors, near-roads, and inside vehicles, and (5) physical activity index (PAI), while also including interaction effects from (6) afternoon time outdoors by daily maximum 8-hr ambient concentration and (7) PAI by afternoon time outdoors.

1 The distributions of afternoon outdoor time and ambient concentration for highly exposed 2 individuals were evaluated considering base air quality and air quality adjusted to just meeting 3 the existing standard. As an example, exposure results in Boston indicated that for about half of 4 the days, simulated school-age children experiencing high exposures spend about 240 minutes 5 outdoors during the afternoon hours along with experiencing daily maximum 8-hr average 6 ambient O_3 concentrations ≥ 75 ppb. In contrast when adjusting ambient concentrations to just 7 meeting the existing standard, for about half of the days, simulated school-age children 8 experiencing similar high exposures need to spend about 280 minutes outdoors during the 9 afternoon hours along with experiencing daily maximum 8-hr average ambient O₃ concentrations 10 \geq 60 ppb. Simply put, under conditions of lower ambient concentrations, persons need to spend a 11 significantly greater amount of time outdoors to experience similar exposures observed at higher 12 ambient concentration conditions. 13 When considering these highly exposed children, on average about half of children's total

afternoon time is spent outdoors on high exposed enhanced, on average about han of enhanced s total afternoon time is spent outdoors on high exposure days, 40% is spent indoors, while only 10% of time is spent near-roads or inside motor vehicles. In general, greater than half of the time highly exposed children spent outdoors specifically involves performing a moderate or greater exertion level activity, such as a sporting activity. While apportionment of afternoon microenvironmental time was similar for highly exposed adults in other age groups considered (e.g., 19-35),

19 important high exertion activities performed outdoors also included those associated with paid

20 work and performing chores.

5.4.3 Exposure Results for Additional At-Risk Populations and Lifestages, Exposure Scenarios, and Air Quality Input Data Used

23 5.4.3.1 Exposures Estimated for All School-age Children During Summer Months, Neither
 24 Attending School or Performing Paid Work

25 As mentioned earlier in describing the longitudinal approach used in the main body of the 26 exposure assessment, the sequence of activity diaries for all simulated individuals is determined 27 by a user-selected profile variable of interest. In this assessment our longitudinal diary approach 28 uses time spent outdoors to link together CHAD diary days, an attempt to appropriately balance 29 intra- and inter-personal variability in that variable. For the primary exposure results, all 30 available diaries were used in developing any one sample pool without restriction outside of the particular characteristics on interest in developing the pool (i.e., age, sex, day-of-week, 31 32 temperature, time spent outdoors). In this targeted simulation in Detroit during three summer 33 months of 2007 (June, July, and August), we restricted the diary pool of all school-age children 34 to include only those diary days that did not have any time spent inside a school nor had time

- 35 spent performing paid work during any day of the week. The results of this targeted simulation
- 36 were compared to an identical simulation, only differing in that all CHAD diary days were used

- i.e., including any diary day for persons having school time or paid work, and as was done forthe main body of this exposure assessment.
- 3 Figure 5-10 indicates that when restricting the CHAD diary pool to include only those 4 diaries having no time spent at school or performing paid work activities, there is about 1/3 or 5 33% increase in the number of all school-age children at or above the 60 ppb-8hr benchmark, a 6 relationship also consistent across the alternative standards and when considering multiple 7 exposures. A similar relationship was found for the other benchmarks (not shown, see Appendix 8 5-G). Clearly, based on the analysis results reported in section 5.4.2 regarding factors influencing 9 those highly exposed, using only activity pattern data that do not include school or work-related 10 events (which would likely occur more so indoors than outdoors) and sampling from a pool of 11 diaries consistent with summer temperatures would increase the likelihood simulated individuals 12 spend time outdoors and be exposed to concentrations at or above the selected benchmarks.



13

standard level (ppb) 60 65 70 75

- 14 Figure 5-10 Comparison of the percent of all school-age children having daily maximum 8-
- 15 hr average O₃ concentration at or above 60 ppb during June, July, and August in Detroit
- 16 2007: using any available CHAD diary ("All CHAD Diaries") or using CHAD diaries
- 17 having no time spent in school or performing paid work ("No School/Work Diaries").

1 5.4.3.2 Exposures Estimated for Outdoor Workers During Summer Months

2 A targeted APEX simulation was performed for the Atlanta study area to simulate 3 summertime exposures for two hypothetical outdoor worker study groups, persons between the 4 age 19-35 and 36-55, using 2006 air quality just meeting the existing standard. To do this, both 5 the daily and longitudinal activity patterns used by APEX were adjusted to best reflect patterns 6 expected for outdoor workers (e.g., a standardized work schedule during weekdays) while also 7 maintaining variability in those patterns across various occupation types. Briefly, the distribution 8 of all employed persons' occupations was estimated using data provided by the U.S. Bureau of Labor and Statistics (US BLS, 2012b)²² and linked with 144 occupation titles from the 9 Occupational Information Network (O*NET)²³ identified as having one or more days per week 10 11 where paid work was performed outdoors. These data were then aggregated to twelve broadly 12 defined BLS occupation groups, generating a data set containing the number of days per week 13 work time would be performed outdoors by that occupation group and properly weighted to 14 reflect the population distribution of persons employed in each outdoor work group. Then, 15 existing CHAD diary days reflecting outdoor paid work were identified, isolated and replicated 16 to reflect this BLS/O*NET outdoor participation rate and occupation group frequencies. A 17 10,000 person simulation was performed by APEX using this adjusted CHAD activity pattern 18 database designed to simulate outdoor workers and compared with exposure results generated 19 from an identical APEX simulation of all employed persons, though differing by using the 20 standard CHAD database and population-based modeling approach used in the main body REA. 21 Details regarding the development of CHAD activity patterns used as input to simulate outdoor 22 workers, as well as other settings and conditions for APEX is described in Appendix 5G. 23 Estimated exposures are presented in Figure 5-11 for one of two age study groups 24 investigated (results for both age groups were similar) and considering either a longitudinal 25 approach designed specifically to reflect an outdoor worker weekday schedule (left panel) or 26 when using our general population-based modeling approach (right panel). The results indicate 27 that when accounting for a structured schedule that includes repeated occurrences of time spent outdoors for a specified study group, all while simulated individuals are likely to be more 28 29 consistently performing work tasks that may be at or above moderate or greater exertion levels, 30 there are a greater percent of the study group experiences exposures at or above the selected 31 health effect benchmark levels than that estimated using our general population-based modeling 32 approach. Keep in mind outdoor workers are expected to experience more exposures at or above 33 benchmark levels, though represent a fraction of the total employed population. It is possible

 ²² U.S. employment data by SOC codes were obtained from: http://www.bls.gov/emp/#tables: Table 1.2
 Employment by occupation, 2010 and projected 2020.

²³ Additional information is available at <u>http://www.onetonline.org</u>.

- 1 that, in using the general population-based approach along with the longitudinal algorithm that
- 2 accounts for within and between variability in outdoor time, a number of outdoor workers are
- 3 incidentally simulated and represent a significant portion of those who experienced exposures at
- 4 or above benchmark levels.²⁴ However, the differences between exposures estimated for the two
- 5 longitudinal approaches become much greater when considering the percent of persons
- 6 experiencing multiple exposure days at or above benchmark levels, primarily when considering
- 7 the 60 ppb-8hr benchmark level. For example, $\leq 2\%$ of the general population-based exposure
- 8 group was estimated to have two or more exposures at or above 60 ppb-8hr, while >17% of
- 9 specifically simulated outdoor workers were estimated to experience exposures at or above that
- 10 same level.

²⁴ In this outdoor worker exposure scenario, approximately 30% of our outdoor worker study group ages 19-55 were estimated to experience at least one exposure at or above 60 ppb-8hr while at moderate or greater exertion. Assuming outdoor workers constitute approximately 12% of the workforce (Appendix G, Table 5G-8), outdoor workers experiencing at least one exposure at or above 60 ppb-8hr could contribute 3.6% to a total exposed population (i.e., outdoor and non-outdoor workers). For the same air quality scenario and using the general population-based approach, we estimated 5-8% of a total employed study group (incidentally comprised of outdoor and non-outdoor workers) would experience exposures at or above the same benchmark, suggesting between 48-75% of persons experiencing exposures above the 60 ppb benchmark have similar activity pattern characteristics as outdoor workers.



Figure 5-11 Percent of workers between ages 19-35 experiencing exposures at or above selected benchmark levels while at moderate or greater exertion using an outdoor worker approach (left panel) and a general population-based approach (right panel) considering air quality adjusted to just meet the existing standard in Atlanta, GA, Jun-Aug, 2006.

5

6 5.4.3.3 Exposures Estimated for All School-age Children When Accounting for Averting 7 Behavior

8 A growing area of air pollution research involves evaluating the actions persons might 9 perform in response to high O_3 concentration days (ISA, section 4.1.1). Most commonly termed 10 *averting behaviors*, they can be broadly characterized as personal activities that either reduce 11 pollutant emissions or limit personal exposure levels. The latter topic is of particular interest in 12 this REA due to the potential negative impact it could have on O₃ concentration-response (C-R) 13 functions used to estimate health risk and on time expenditure and activity exertion levels 14 recorded in the CHAD diaries used by APEX to estimate O₃ exposures. To this end, we have 15 performed an additional review of the available literature here beyond that summarized in the 16 ISA to include several recent technical reports that collected and/or evaluated averting behavior 17 data (Graham, 2012). The purpose was to generate a few reasonable quantitative approximations 18 that allow us to better understand how averting behavior might affect time-location-activity 19 patterns, and then simulate how such personal adjustments might affect our population exposure 20 estimates. 21 Based on the elements evaluated in our literature review (i.e., air pollution awareness, 22 prevalence and duration of averting response), we conclude that most people are aware of alert 23 notification systems (in particular those persons having compromised health and reside in an 24 urban area). We approximate that 30% of all asthmatics (or 15% of the general population) may 25 reduce their outdoor activity level on alert days (e.g., KS DOH, 2006; McDermott et al., 2006; 26 Wen et al., 2009; Zivin and Neidell, 2009) and that outdoor time/exertion during afternoon hours 27 may be reduced by about 20-40 minutes in response to an air quality alert notification

1 (Bresnahan et al., 1997; Mansfield et.al, 2006, Neidell, 2010; Sexton, 2011). We used these

2 literature derived estimates to generate an adjusted activity diary pool used by APEX to simulate

- 3 a 2-day exposure period (August 1-August 2, 2007) in Detroit to approximate the effect averting
- 4 may have on exceedances of exposure benchmarks.
- 5 When considering base air quality and our designed target to represent averting
- 6 performed by the general population 15.3 % of all simulated school-age children spent on
- 7 average 44 minutes less time outdoors resulting in approximately one percentage point or
- 8 fewer children experienced exposures at or above any of the selected benchmark levels (Figure
- 9 5-12, left panel). When considering base air quality and our designed target to represent an
- 10 averting response by the population of asthmatics 30.3% of simulated asthmatic school-age
- 11 children spent on average 44 minutes less time outdoors resulting in approximately two
- 12 percentage points or fewer experienced exposures at or above any of the selected benchmark
- 13 levels (Figure 5-12, right panel).



1 Figure 5-12 Percent of all school-age children (left panel) and asthmatic school-age

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5.4.3.4 Comparison of APEX Estimated Exposures Using Three Different Base Case Air Quality Data Sets: AQS, VNA, and EVNA

8 For this exposure assessment, we elected to use a modeling approach to estimate the 9 ambient input concentration field and better account for spatial gradients that may exist (Chapter 10 4). To support the selection of VNA, we compared exposure results separately generated using 11 ambient monitor (AOS) aVNA, and VNA as input to APEX for three study areas: Atlanta

- ambient monitor (AQS), eVNA, and VNA as input to APEX for three study areas: Atlanta,
- 12 Detroit, and Philadelphia. All APEX settings were generally consistent with the simulations
- 13 discussed previously, though the air quality data differed in that the year selected was 2005
- 14 (based on the available CMAQ data) and that a 4 Km grid was used to define the spatial area for
- 15 this evaluation rather than census tracts. Daily maximum 8-hr average exposures were estimated

<sup>children (right panel) having daily maximum 8-hr average O₃ concentration at or above
benchmark levels during a 2-day simulation in Detroit, base air quality, August 1-2, 2007.</sup>

 ⁴ Red bars indicate exposure results when considering effect of averting.

for asthmatic school-age children residing in the same census tracts comprising each air quality
 domain and summarized in Figure 5-13.

- 3 Exposure results for all three air quality input data sets were very comparable, with a few 4 notable differences. Using AQS monitor concentration data tended to result in a 1-3% greater 5 percent of asthmatic school-age children at or above each of the selected benchmark levels when compared with exposures estimated using VNA concentrations. While the VNA concentrations 6 7 are based on the AQS monitor data, the approach generates a concentration gradient with 8 distance from areas of known concentration that are typically less than the observed values, thus 9 yielding fewer persons exposed to the highest concentrations. Using the eVNA approach to 10 generate ambient concentrations tended to result in 2-5% greater percent of asthmatic school-age 11 children at or above each of the selected benchmark levels when compared with exposures 12 estimated using either the AQS or VNA approaches. This is because at times, the eVNA
- 13 approach estimated high concentrations in areas where no observations were present, based on
- 14 modeling which captures gradients in O_3 that may result from nearby sources (see Chapter 4).



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Figure 5-13 Comparison of APEX exposure results generated for three study areas (Atlanta, Detroit, and Philadelphia) using three different 2005 air quality input data sets: AQS, VNA, and eVNA.

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5.4.3.5 Comparison of APEX Estimated Exposures Using Two Different Adjusted Air Quality Data Sets: Quadratic Rollback and HDDM

3 We elected to use an air quality modeling based approach rather than the previously used 4 statistical approach to adjust air quality to just meet the current and alternative standard levels 5 (Chapter 4). To support the selection of the HDDM approach, we compared exposure results for 6 the scenario of just meeting the existing standard, separately generated using air quality inputs 7 obtained using the quadratic rollback and HDDM method to adjust air quality for the Atlanta 8 study area. All APEX settings were generally consistent with the simulations discussed 9 previously, though both the air quality data sets used in this comparison differed from that done 10 in the main exposure results above in that only the ambient monitor locations were used to define 11 the air districts and assumed a 30 km radius of influence, as was done for the first draft REA. 12 Daily maximum 8-hr average exposures were estimated for asthmatic school-age children in

- 13 census tracts within 30 km of each air district and summarized in Figure 5-14.
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Figure 5-14 Comparison of exposure results generated by APEX using two different air quality adjustment approaches to just meet the existing standard in Atlanta: quadratic rollback (left panel) and HDDM (right panel).

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The quadratic adjusted air quality resulted in slightly fewer percent of asthmatic schoolage children exposed at or above the highest benchmark (80 ppb-8hr) when compared with

22 exposures estimated using the HDDM model simulation approach, though a significantly greater

23 percent of asthmatic school-age children were exposed to the lowest benchmark (60 ppb-8hr)

24 using the quadratic approach. This is because the quadratic approach generally targets the highest

concentrations for adjustment, while the HDDM approach accounts for changes across the full
 concentration distribution to meet the adjusted concentration level of interest.

3 5.4.4 Limited Performance Evaluations

4 5.4.4.1 Personal Exposure Comparisons

5 A new evaluation of APEX was performed using a subset of personal O_3 exposure 6 measurements obtained from the Detroit Exposure and Aerosol Research Study (DEARS) (Meng 7 et. al, 2012). For five consecutive days, personal O_3 outdoor concentrations along with daily 8 time-location activity diaries were collected from 36 adult study participants in Wayne County 9 Michigan during July and August 2006. An APEX simulation was performed considering these 10 same geographic and temporal features, followed with the sub-setting of APEX output data 11 according to important personal attributes of the DEARS study participants (5-day collection 12 study periods, age/sex distributions, outdoor time, ambient concentrations, and air exchange 13 rate). A comparison sample was generated randomly from the complete simulation, selecting for 14 50 APEX simulated individuals. 15 For both data sets and considering the two output variables separately (outdoor time and 16 daily exposure), the median daily values for each study participant were ranked, then plotted 17 along with each individual's corresponding minimum and maximum value using each 18 individual's 5 person-days of data (Figure 5-15). In spite of the distinct matching of influential personal attributes, over 50% of APEX simulated individuals had median daily O3 exposure 19 20 concentrations above 10 ppb, while only 3% of DEARS participants' median values exceeded 10 21 ppb. The reason(s) for this difference is being investigated.



Figure 5-15 Distribution of daily average O₃ exposures (top panels) and daily afternoon
 outdoor time (bottom panels) and for DEARS study participants (left panels) and APEX
 simulated individuals (right panels) in Wayne County, MI, July-August 2006.

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5 APEX modeled exposures have previously been compared with personal exposure 6 measurements for O₃ (US EPA, 2007b). Briefly, APEX O₃ simulation results were compared 7 with 6-day personal O_3 concentration measurements for children ages 7-12 (Xue et al., 2004; 8 Geyh et al., 2000). Two separate areas of San Bernardino County were surveyed: urban Upland 9 CA, and the combined small mountain towns of Lake Arrowhead, Crestline, and Running 10 Springs, CA. Available ambient monitoring data for these locations during the same study years (1995-1996) were used as the air quality input to APEX. APEX predicted personal exposures, 11 12 averaged similarly across a 6-day period, matched reasonably well for much of the concentration 13 distribution considering both locations, but tended to underestimate exposures at the upper 14 percentiles of the distribution. The average difference between the 6-day means was less than 1 15 ppb, with a range of -11 ppb to +8 ppb, though predicted upper bounds for a few averaged 16 exposures having higher exposure concentrations were under-predicted by up to 24 ppb (e.g., 17 Figure 5-16). In addition, modeled exposure concentration variability was less than that observed 18 in the personal exposure measurements. At the time of analysis, these differences were proposed

1 to be largely driven by under-estimation of the spatial variability of the outdoor concentrations

2 used by APEX (US EPA, 2007b).

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Figure 5-16 Means (and range) of 6-day average personal O₃ exposures, measured and modeled (APEX), Upland Ca. Obtained from Figure 8-22 of US EPA (2007b).

7 5.4.4.2 Ventilation Rate Comparisons

8 The algorithm used by APEX to estimate minute-by-minute ventilation rate serves as the 9 basis for recent updates to the ventilation rate distributions provided in EPAs Exposure Factors 10 Handbook (U.S. EPA, 2009b; US EPA, 2011). During the development of the ventilation 11 distributions for EPA at that time, two peer-reviewed studies were identified as providing 12 somewhat relevant measurement data to evaluate the APEX energy expenditure and ventilation 13 algorithm (see Graham, 2009 for additional comparison details). The results of this evaluation 14 are summarized below.

15 Briefly, Brochu et al. (2006a,b) presents data for ventilation rates derived from tracking 16 doubly-labeled water (DLW) consumption/elimination to estimate energy expenditure in healthy 17 normal-weight males and females, ages from 1 month to 96 years (n=1,252). Estimates of energy 18 expended were combined with a fixed oxygen uptake factor (H=0.21) and using a fixed ventilatory equivalent (VQ)²⁵ of 27. The DLW measurement period ranged from 7-21 days, 19 20 resulting in time-averaged metrics that may in some instances provide reasonable estimates for a 21 mean daily ventilation rate, but not useful for estimating variability in an individual's ventilation 22 rate over shorter time periods (as is needed by APEX). Further, while DLW is considered by 23 some as a 'gold standard' for measuring energy expenditure, this characterization would not necessarily be directly transferable to approximations that use this measured value (i.e., 24 25 ventilation rate in Brochu et al. (2006a,b) is a calculated value, not measured). Reported

²⁵ The ventilatory equivalent (VQ) is the ventilation rate (VE) divided by the oxygen consumption rate (VO₂)

1 ventilation rates are daily averages for several age groupings (e.g. ages 1 to < 2, 2 to < 5, 5 to <2 7, etc.) along with derived percentiles, each assuming the existence of normally distributed data. 3 A 14-day APEX simulation was performed (i.e., the median of 7-21 days for the DLW 4 measurement study) to estimate daily ventilation rates for comparison with the time-averaged 5 Brochu et al (2006a) data. Twenty-five thousand persons were simulated by APEX to generate a reasonable number of persons within each year of age and other potential categorical variables 6 7 (e.g., 100-200, although a few older age groups resulted in having fewer persons). It is important 8 when comparing the two types of data for them to be similar as possible, particularly since age 9 and body mass are important influential variables in both estimation methods. A total of 9,613 10 normal-weight individuals were simulated by APEX and used for the following analysis. Multiday ventilation rates were averaged across the 14-day simulation period, yielding a mean daily 11 12 ventilation rate for each person to best represent the DLW time averaging done by Brochu et al. 13 (2006a). 14 Figure 5-17 compares the APEX simulated individuals body mass normalized mean daily 15 ventilation rates with those reported by Brochu et al. (2006a; Table 2, page 684) for several age 16 groupings of normal-weight individuals. The two largest differences appear for children of both

- 17 sexes less than age 10 (i.e., Brochu et. al (2006a) estimates are systematically lower than APEX
- 18 estimates) and for ages 16-33 (i.e., APEX estimates are lower than Brochu et al (2006a). Body
- 19 mass normalized ventilation rates also appear to be slightly higher using APEX when
- 20 considering persons above age 64 and for both sexes.



Figure 5-17 Comparison of body mass normalized mean daily ventilation rates estimated by APEX (closed symbols) and by Brochu et al., 2006 (open symbols).

1 2 One principal issue identified by us as potentially responsible for some of the above 3 differences in ventilation estimates is in the VQ used by Brochu et al. (2006a). A single value of 4 27 was used in estimating ventilation rates for both children and adults, however it is widely 5 recognized that while a VQ of 27 may be a reasonable approximation for estimating mean 6 ventilation rates of adults, it is not appropriate for use in estimating mean ventilation rates in 7 children. With this in mind, the Brochu et al. (2006a) ventilation estimates were modified here 8 using the VQ estimates offered by Arcus-Arth and Blaisdell (2007). Figure 5-18 illustrates the 9 comparison of APEX body mass normalized mean daily ventilation rates with that of Brochu et 10 al. (2006a) corrected ventilation estimates. The body mass normalized ventilation estimates for 11 school-age children are more similar to those generated by APEX when correcting the Brochu et 12 al (2006a) VQ parameter. Thus, mean ventilation rates generated by APEX are reasonably 13 correlated with independent measures from the Brochu et al. (2006a, b) estimates, particularly 14 when correcting the Brochu et al (2006a) ventilation estimates for children using a more 15 appropriate estimate of VQ for children.



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Figure 5-18 Comparison of body mass normalized mean daily ventilation rates in male and female school-age children (5-18) when correcting Brochu et al. (2006a) results with child appropriate VQ estimates.

1 In a second study identified for comparison with APEX estimates, Arcus-Arth and 2 Blaisdell (2007) provide ventilation estimates for children <19 years of age using energy intake 3 (EI, or calories consumed) and body mass data provided by the USDA's Continuing Survey of 4 Food Intake for Individuals (CSFII; USDA, 2000). Two-day daily average EIs were combined 5 with a values of H (i.e., 0.22 for infants, 0.21 for non infants) and VQ (i.e., 33.5 for children 0-8, 6 30.6 for boys 9-18, 31.5 for girls 9-18 years old). Again, time-averaging of the data may provide 7 reasonable estimates of a daily mean, but offer no variability in ventilation estimates for shorter 8 durations. Furthermore, data for both sexes are combined and reported by age, with stratified 9 results by sex reported only for aggregated age groups (males and females, 9-18 years old).

10 A 2-day model simulation was performed by APEX to generate ventilation estimates for children to compare with results of Arcus-Arth and Blaisdell (2007).²⁶ APEX ventilation 11 estimates were time-averaged to generate mean daily values, and since the data reported in 12 13 Arcus-Arth and Blaisdell (2007) were not separated by sex (outside of broad age categories), the 14 APEX estimates were also combined by sex to provide a comparable mean estimate for each 15 year of age (5-18). Body mass was also not used as a categorical variable in Arcus-Arth and 16 Blaisdell (2007), therefore all APEX simulated individuals were used, regardless of whether they 17 could be classified as overweight or of normal weight. In addition, daily ventilation rates for a 18 few age groups of children were obtained from Tables 3 and 4 of Brochu et al. (2006a), though 19 considering both estimates for normal and overweight individuals (there were no combined data 20 available). The Brochu et al. (2006a) results have been corrected for VQ as noted above using 21 VQ estimates of Arcus-Arth and Blaisdell (2007) and added for comparison. 22 Figure 5-19 illustrates ventilation rate estimates from the APEX simulation, along with 23 associated data for school-age children (ages 5-18) obtained from the two publications. Daily 24 mean ventilation estimates are quite similar at each year of age, with slightly higher estimates by 25

Arcus-Arth and Blaisdell (2007) at ages 9 and above, particularly when compared with APEX
 ventilation estimates. Ventilation estimates are remarkably similar for school-age children for all

three sources of data, particularly when considering the differences in the type of input data used

and the varied approaches of APEX, Brochu et al. (2006a), and Arcus-Arth and Blaisdell (2007).

29 This overall agreement suggests reasonable confidence can be conferred to the algorithm used by

30 APEX to estimate at a minimum, daily mean ventilation rates.

²⁶ Table III, page 103 of Arcus-Arth and Blaisdell (2007) provided body mass normalized ventilation rates.



Figure 5-19 Comparison of body mass normalized daily mean ventilation rates in school-age children (5-18) estimated using APEX and literature reported values.

5 5.4.4.3 Evaluation of Longitudinal Profile Methodology

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6 We evaluated the APEX approach used for linking together cross-sectional activity 7 pattern diaries to generate longitudinal profiles for our simulated individuals (Appendix 5G, 8 Section 5G-3). Of particular interest were how well variability in outdoor participation rate and 9 the amount of time expended were represented in our population-based exposure simulations. 10 Our goal in developing the most reasonable longitudinal profiles is to capture expected, 11 important features of population activity patterns, i.e., there is correlation within an individual's 12 day-to-day activity patterns (though neither exactly repeated nor entirely random for individuals) 13 and variability across the modeled study group in day-to-day activity patterns (i.e., not every 14 simulated individual in the study group does the same activity on the same day). 15 The simulated longitudinal profiles indicate the method for linking together cross-16 sectional diaries generates a diverse mixture of persons having variable, though expected, 17 activity patterns: A small fraction of the simulated population spend a limited amount of 18 afternoon time outdoors and occurring at a low frequency across an O₃ season, a small fraction 19 consistently spends a greater amount (> 2 hours) of time outdoors and occurring at greater 20 frequency (e.g., 4/5 days per week), while the remaining simulated individuals fall somewhere in 21 between regarding participation and total time. While we are not aware of a population database

1 available to compare with these simulated results, we are comfortable with the method

2 performance in representing the intended variability in longitudinal activity patterns (see section

3 5G-3 for details).

4 5.5 VARIABILITY AND UNCERTAINTY

5 An important issue associated with any population exposure or risk assessment is the 6 characterization of variability and uncertainty. Variability refers to the inherent heterogeneity in 7 a population or variable of interest (e.g., residential air exchange rates). The degree of variability 8 cannot be reduced through further research, only better characterized with additional 9 measurement. Uncertainty refers to the lack of knowledge regarding the values of model input 10 variables (i.e., *parameter uncertainty*), the physical systems or relationships used (i.e., use of 11 input variables to estimate exposure or risk or *model uncertainty*), and in specifying the scenario 12 that is consistent with purpose of the assessment (i.e., scenario uncertainty). Uncertainty is, 13 ideally, reduced to the maximum extent possible through improved measurement of key 14 parameters and iterative model refinement. The approaches used to assess variability and to 15 characterize uncertainty in this REA are discussed in the following two sections. Each section 16 also contains a concise summary of the identified components contributing to uncertainty and 17 how each source may affect the estimated exposures.

18 5.5.1 TREATMENT OF VARIABILITY

19 The purpose for addressing variability in this REA is to ensure that the estimates of 20 exposure and risk reflect the variability of ambient O₃ concentrations, population and lifestage 21 characteristics, associated O₃ exposure and dose, and potential health risk across the study area 22 and for the simulated at-risk study groups. In this REA, there are several algorithms that account 23 for variability of input data when generating the number of estimated benchmark exceedances or 24 health risk outputs. For example, variability may arise from differences in the population 25 residing within census tracts (e.g., age distribution) and the activities that may affect population 26 and lifestage exposure to O_3 (e.g., time spent inside vehicles, time performing moderate or 27 greater exertion level activities outdoors). A complete range of potential exposure levels and 28 associated risk estimates can be generated when appropriately addressing variability in exposure 29 and risk assessments; note however that the range of values obtained would be within the 30 constraints of the input parameters, algorithms, or modeling system used, not necessarily the 31 complete range of the true exposure or risk values. 32 Where possible, we identified and incorporated the observed variability in input data sets

33 to estimate model parameters within the exposure assessment rather than employing standard

34 default assumptions and/or using point estimates to describe model inputs. The details regarding

1 variability distributions used in data inputs are described in Appendix 5B. To the extent possible

2 given the data available for the assessment, we accounted for variability within the exposure

3 modeling. APEX has been designed to account for variability in some of the input data,

4 including the physiological variables that are important inputs to determining ventilation rates.

5 As a result, APEX addresses much of the variability in factors that affect human exposure.

6 Important sources of the variability accounted for in this analysis are summarized in Appendix

7 5D.

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CHARACTERIZATION OF UNCERTAINTY 5.5.2

9 While it may be possible to capture a range of exposure or risk values by accounting for 10 variability inherent to influential factors, the true exposure or risk for any given individual within 11 a study area is unknown, though can be estimated. To characterize health risks, exposure and risk 12 assessors commonly use an iterative process of gathering data, developing models, and 13 estimating exposures and risks, given the goals of the assessment, scale of the assessment 14 performed, and limitations of the input data available. However, significant uncertainty often 15 remains and emphasis is then placed on characterizing the nature of that uncertainty and its 16 impact on exposure and risk estimates.

17 The REA's for the previous O₃, NO₂, SO₂, and CO NAAQS reviews each presented a 18 characterization of uncertainty of exposure modeling (Langstaff, 2007; US EPA 2008, 2009a, 19 2010). The qualitative approach used in this and other REAs is described by WHO (2008). 20 Briefly, we identified the key aspects of the assessment approach that may contribute to 21 uncertainty in the exposure and risk estimates and provided the rationale for their inclusion. 22 Then, we characterized the *magnitude* and *direction* of the influence on the assessment results 23 for each of these identified sources of uncertainty. Consistent with the WHO (2008) guidance, 24 staff scaled the overall impact of the uncertainty by considering the degree of uncertainty as 25 implied by the relationship between the source of uncertainty and the exposure concentrations. A 26 qualitative characterization of low, moderate, and high was assigned to the magnitude of 27 influence and knowledge base uncertainty descriptors, using quantitative observations relating to 28 understanding the uncertainty, where possible. A summary of the key findings of those prior 29 characterizations that are most relevant to the current O_3 exposure assessment are provided in 30 Table 5-6.

			Is rating			
Sources of Uncertainty		Influence of Uncertainty on Exposure/Intake Dose Estimates		Knowledge-		current APEX O ₃
Category	Element	Direction	Magnitude	base Uncertainty	Comments	assessment?
	Database Quality	Both	Low	Low All ambient pollutant measurements available from AQS are both comprehensive and subject to guality control.		Yes. No further characterization needed.
	Instrument Measurement Error	Over	Low	Low	Mean bias estimated as 1.2% (CV of 4.4%). See Table 2 and Figure 6 of Langstaff (2007).	Yes. No further characterization needed.
Ambient Monitoring Concentrations	Missing Data Substitution Method	Both	Low	Low Overall completeness of data yield negligible mean bias (~0) along with an estimated standard deviation of 4 ppb when replacing missing values. See Table 3 of Langstaff (2007).		Yes. No further characterization needed.
	Temporal Representation	Both	Low	Low	Appropriately uses 1-hr time-series of O_3 concentrations for 5 years. No missing data for any hour input to APEX.	Yes. No further characterization needed.
	Spatial Representation: Large Scale	Both	Low	Low	Tens of monitors used in each study area.	Yes. No further characterization needed.
	Spatial Representation: Neighborhood Scale (1)	Both	Low	Low	Spatial interpolation using jackknife method (removal of a single monitor) yielded generally unbiased observed/predicted ratios (mean 1.06), having an estimated standard deviation of 0.2. Langstaff (2007).	Yes. For the uncertainties characterized, the historical rating is appropriate if and
	Spatial Representation: Neighborhood Scale (2)	Over	Low	Low	When reducing the APEX radius setting from an unlimited value (actual value used) to 10 km (i.e., the tendency would be to more accurately represent exposure), a smaller fraction (1-3 percentage points) of population exceeds benchmark levels. See Figures $7 - 9$ of Langstaff (2007).	when using ambient monitor data alone to represent air quality surface. However in this 2 nd draft REA, local- scale air quality was estimated using VNA (see below).

Table 5-6 Characterization of Key Uncertainties in Historical and Current APEX Exposure Assessments

			Is rating			
Sour	ces of Uncertainty	Influence of Uncertainty on Exposure/Intake Dose Estimates		Knowledge-		current APEX O ₃
Category	Element	Direction	Magnitude	base Uncertainty	Comments	assessment?
	Spatial Representation: Local Scale VNA estimates	Both	Low	Low - Moderate	Scenario-based evaluation in three study areas indicated small differences in exposure results when comparing ambient monitor data or statistically interpolated concentrations to 4 Km grid as an input to APEX (Figure 5-13). General dependencies of the approaches used could lead to observed lack of distinction in exposure results.	Yes. Newly evaluated.
	Spatial Representation: Vertical Profile	Both	Moderate	Moderate Differences between ground-level (0-3 meters) and building rooftop sited (25 meters monitor concentrations can be significant. Most importantly, use of higher elevation monitors would tend to overestimate ground- level exposures (i.e., persons outdoors).		Yes. Given judged impact to exposure, additional characterization is possibly warranted.
	Quadratic Approach	Both	Low - Moderate	Moderate	Variable differences (e.g., none to a factor of two or three) in the estimated number of persons exposed across study areas when using differing 3-year roll-back periods for a single year of air quality (Langstaff, 2007).	Yes. Uncertainty in the approach has resulted in use of HDDM approach.
Adjustment of Air Quality to Simulate Just Meeting the Current Standard	HDDM Simulation Approach	Both	Low - Moderate	Low - Moderate	Expected patterns in both air quality and exposure result from HDDM/emissions reduction approach (full distribution affected rather than only upper percentiles, Figure 5- 14). Variable differences remain (e.g., none to a factor of two or three) in the estimated percent of persons exposed across study areas when using differing 3-year roll-back periods for 2008 air quality (Figures 5-5 to 5- 9). New York study area could not be simulated to just meet 60 and 55 ppb alternative standards.	Yes. Newly evaluated.
APEX: General Input Databases	Population Demographics and Commuting (US Census)	Under	Low	Low	Comprehensive and subject to quality control. Differences in 2000 data versus modeled years (2006-2010) are likely small when estimating percent of population exposed.	Yes. No further characterization needed.

			Is rating			
Sources of Uncertainty		Influence of Uncertainty on Exposure/Intake Dose Estimates		Knowledge-		current APEX O ₃
Category	Element	Direction	Magnitude	base Uncertainty	Comments	assessment?
	Activity Patterns (CHAD)	Both	Low - Moderate	Low- Moderate	Comprehensive and subject to quality control. Significantly increased number of diaries used to estimate exposure from prior review and 1 st draft REA for this review (Table 5-3). Thoroughly evaluated trends and patterns in historical data – no major issues noted with use of historical data to represent current patterns (Figures 5G-1 and 5G-2). Compared outdoor participation and time with ATUS data base – CHAD participation is higher than ATUS, likely due to ATUS survey methods. Activity data for asthmatics generally similar to non-asthmatics (Tables 5G2-to 5G-5). Remaining uncertainty with other influential factors that cannot be accounted for (e.g., SES, region/local outdoor participation rates)	Yes. Newly evaluated.
	Meteorological (NWS)	Both	Low	Low	Comprehensive and subject to quality control, few missing values. Limited application in selecting CHAD diaries and AERs.	Yes. No further characterization needed.
	Poverty Status (US Census) Weighted Asthma Prevalence (CDC)	Both	Low	Low	Data used are from a peer-reviewed quality controlled source. Application accounts for variability in most important influential variables (age, sex, region, poverty) though possible that variability in microscale prevalence not entirely represented.	New. Could possibly use further characterization, though typically available local prevalence rates are not well stratified by influential variables.

			Is rating			
Sources of Uncertainty		Influence of Uncertainty on Exposure/Intake		Knowledge-		current APEX O ₃
Category	Element	Direction	Magnitude	base Uncertainty	Comments	exposure assessment?
APEX: Microenvironmental Concentrations	Outdoor Near-Road and Vehicular: Proximity Factors	Both	Low	Low- Moderate	Uncertainty in mean value used approximated as 15 percentage points. See Figure 10 and Table 7 of Langstaff (2007). May be of greater importance in certain study areas or under varying conditions, though even with this mean difference, in-vehicle penetration/decay decreases exposures and hence importance of in-vehicle microenvironments.	Yes. No further characterization needed.
	Indoor: Near-Road	Over	Low	Low	Expected reduction in O_3 for persons residing near roads not modeled here, but when included, there is a small reduction (~3%) in the number of persons experiencing exposure above benchmark levels (Langstaff, 2007).	Yes. No further characterization needed.
	Indoor: Air Exchange Rates	Both	Low	Moderate	Uncertainty due to random sampling variation via bootstrap distribution analysis indicated the AER GM and GSD uncertainty for a given study area tends range to at most from fitted ± 1.0 GM and ± 0.5 GSD hr ⁻¹ . Non-representativeness remains an important issue as city-to-city variability can be wide ranging (GM/GSD pairs can vary by factors of 2-3) and data available for city-specific evaluation are limited (Langstaff, 2007). Also, indoor exposures are estimated as not important to daily maximum 8-hr average O_3 exposure.	Yes. No further characterization needed.

Sources of Uncertainty		Influence of Uncertainty on Exposure/Intake Dose Estimates		Historical Un	certainty Characterization	Is rating appropriate for current APEX O ₃
Category	Element	Direction	Magnitude	base Uncertainty	Comments	assessment?
	Indoor: A/C Prevalence (AHS)	Both	Low	Low	Comprehensive and subject to quality control, estimated 95 th percentile confidence bounds range from a few to just over ten percentage points, though some cities use older year data (Table 9 of Langstaff, 2007). Note, variable indicates presence/absence not actual use. Also, indoor exposures are estimated here as limited in importance to daily maximum 8-hr average exposures and sensitivity analyses in NO ₂ REA (in-vehicle was most influential exposure ME) concluded indoor prevalence variable was of limited importance.	Yes. No further characterization needed.
	Indoor: Removal Rate	Both	Low	Low	Greatest uncertainty in the input distribution regarded representativeness, though estimated as unbiased but correct to within 10% (Langstaff, 2007).	Yes. No further characterization needed.
	Vehicular: Penetration Factors	Both	Low	Moderate	Input distribution is from an older measurement study though consistent with recent, albeit limited data.	Yes. No further characterization needed.

			Is rating			
Sources of Uncertainty		Influence of Uncertainty on Exposure/Intake Dose Estimates		Knowledge- base		current APEX O ₃ exposure
APEX: Simulated Activity Profiles	Longitudinal Profiles	Under	Low - Moderate	Moderate	Depending on the longitudinal profile method selected, the number of persons experiencing multiple exposure events at or above a selected level could differ by about 15 to 50% (see Appendix B, Attachment 4 of NO ₂ REA). Long-term diary profiles (i.e., monthly, annual) do not exist for a population, limiting the evaluation. The general population-based modeling approach used for main body REA results does not assign rigid schedules, for example explicitly representing a 5-day work week for employed persons. However, when considering such scheduling (e.g., outdoor workers or all children spending entire summer season not in-school), estimated exposures are greater than when not considering rigid weekly/seasonal schedules. For our hypothetical outdoor worker scenario, the number of multiday exposures at or above benchmark levels was primarily affected (though mainly the 60 ppb level, Figure 5-11), while both percent of children experiencing single and multiday exposures were increased by about 30% when simulating a rigid schedule (Figure 5-10).	Yes. Newly evaluated.
	Commuting	Both	Low	Moderate	New method used in this assessment is designed to link Census commute distances with CHAD vehicle drive times. Considered an improvement over the former approach that did not match distance and time. While vehicle time accounted for through diary selection, not rigidly scheduled. However, In- vehicle exposures are not important drivers for persons exceeding benchmark levels (section 5.3.2).	Yes. Newly evaluated.

			Is rating			
Sour	ces of Uncertainty	Influence of Uncertainty on Exposure/Intake		Knowledge-		current APEX O ₃
Category	Element	Direction	Magnitude	base Uncertainty	Comments	exposure assessment?
	At-Risk Population and Lifestages	Both	Low	Low – Moderate	An updated evaluation shows activity patterns of asthmatics are similar to that of non-asthmatics (section 5.3.1, Tables 5G-2 to 5G-5).	Yes. Newly evaluated.
APEX: Physiological Processes	Body Mass (NHANES)	Unknown	Low	Low	Comprehensive and subject to quality control, though older (1999-2004) than current simulated population, possible small regional variation is not represented by national data.	Yes. No further characterization needed.
	NVO _{2max}	Unknown	Low	Low	Upper bound control for unrealistic activity levels rarely used by model, thus likely not very influential.	Yes. No further characterization needed.
	RMR	Unknown	Low	Low	Approach from older literature (Schofield, 1985), used in ventilation equation. Note ventilation rate estimates are reasonable.	Newly identified. May need additional characterization.
	METS distributions	Over	Low - Moderate	Low - Moderate	APEX estimated daily mean METs range from about 0.1 to 0.2 units (between about 5- 10%) higher than independent literature reported values (Table 15 of Langstaff, 2007). Shorter-term values are of greater importance in this assessment.	Yes. Given judged impact to exposure, additional characterization is needed.
	Ventilation rates	Over	Low - Moderate	Low - Moderate	APEX estimated daily ventilation rates can be greater (2-3 m ³ /day) than literature reported measurement values (Table 25 of Langstaff, 2007), though if accounting for measurement bias this minimizes the discrepancy (Graham and McCurdy, 2005; see Figures 5-18 and 5- 19). Also, a shorter-term comparison (hours rather than daily), while more informative, cannot be performed due to lack of data.	Yes. Additional characterization would be warranted if minute or hourly ventilation rate data were available.
Exposure Benchmark Level	EVR characterization of moderate or greater exertion	Over	Moderate	Low - Moderate	Given that the EVR serves as a cut point for selecting persons performing at moderate or greater exertion and is a lower bound value ($\sim 5^{\text{th}}$ percentile), the simulated number of persons achieving this level of exercise is possibly overestimated.	Newly identified. May need additional characterization.

1 5.6 KEY OBSERVATIONS

2 Two additional tables are provided to additionally summarize the exposure results across 3 all study areas and years of air quality data: Table 5-7 contains the percent of all school-age 4 children experiencing at least one exposure at or above the three exposure benchmark levels, 5 while Table 5-8 contains the percent of all school-age children experiencing at least two 6 exposures at or above the three exposure benchmark levels, with both tables considering results associated with each of the adjusted air quality scenarios.²⁷ Two descriptive statistics are 7 8 provided from the exposure results for each study area: the mean percent of persons exposed in 9 each study area averaged across the 5 years simulated and the maximum percent of persons 10 exposed in each study area, representing the worst year of air quality simulated. Figure 5-20 11 illustrates the estimated mean and maximum percent of all school-age children exposed for each 12 study area when considering the 60 ppb-8hr benchmark and adjusted air quality scenarios, and 13 using the data provided in Table 5-7 and Table 5-8. 14 Presented below are key observations resulting from the O₃ exposure analysis: 15 General: The estimated percent of any study group exposed at least once at or above the 16 selected benchmark levels were highest considering the base air quality though percent 17 exposed varied by study area, year, and benchmark level (Appendix 5F). Very few 18 persons within any study group (all are estimated to be < 0.3%) experienced any 19 benchmark exceedances when considering an alternative standard level of 55 ppb-8hr 20 (data not shown). 21 Study Group: The percent of all school-age children exposed at or above the selected • 22 benchmark levels across all study areas, years, and air quality scenarios were similar to 23 exposures for asthmatic school-age children (e.g., Figure 5-5 and Figure 5-6, 24 respectively) with both of these study groups having consistently higher percent of 25 persons exposed than that estimated for asthmatic adults and all older adults (Figure 5-7 26 and Figure 5-8, respectively), generally by about a factor of three or more. The percent of 27 all older adults at or above any benchmark level tended to be only a few percentage 28 points or less when compared with corresponding benchmark exceedances for asthmatic 29 adults. 30 • **80 ppb-8hr Exposure Benchmark:** In general, less than 1% of any study group,

including all school-age children and any study area, was exposed at least once at or
 above the highest exposure benchmark, 80 ppb-8hr, when considering the existing

²⁷ The maximum sample size is 6 years based on years simulated, and for a few instances varied based on available air quality (e.g., Chicago does not have 3 years simulated for just meeting the current standard during 2008-2010 period because air quality was below the current standard, thus the total sample size for this study area is 3.

standard air quality scenario (Table 5-7). When considering a standard level of 70 ppb-8hr, $\leq 0.2\%$ of any study group and any study area was exposed at least once at or above that same benchmark.

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- 4 70 ppb-8hr Exposure Benchmark: Less than 10% of any study group, including all • 5 school-age children and any study area, was exposed at least once at or above an 6 exposure benchmark of 70 ppb-8hr, when considering the existing standard air quality 7 scenario (Table 5-7). When considering a standard level of 70 ppb-8hr, $\leq 3.5\%$ of any 8 study group and in any study area was exposed at least once at or above that same 9 benchmark. A standard level of 65 ppb-8hr is estimated to reduce the percent of persons 10 at or above an exposure benchmark of 70 ppb-8hr to $\leq 0.5\%$ of any study group and in 11 any study area.
- 60 ppb-8hr Exposure Benchmark: In general, no more than 26% of any study group in any study area was exposed at least once at or above the lowest exposure benchmark, 60 ppb-8hr, when considering the existing standard air quality scenario (Table 5-7, Figure 5-20). When considering a standard level of 70 ppb-8hr, < 20% of any study group in any study area was exposed at least once at or above that same benchmark. A standard level of 65 ppb-8hr is estimated to reduce the percent of persons at or above an exposure benchmark of 60 ppb-8hr to ≤ 10% of any study group and study area.
- Multi-day Benchmark Exceedances: When considering air quality adjusted to just meet 19 20 the existing standard, multi-day exposure benchmark exceedances are largely limited to 21 two or more exceedances at the 60 ppb-8hr benchmark, all occurring for < 15% of any 22 study group in any study area (e.g., Table 5-8, Figure 5-9). There were no persons 23 estimated to experience any multi-day exposures at or above 80 ppb-8hr for any study 24 group in any study area, while $\leq 2.2\%$ of persons were estimated to experience two or 25 more exposures at or above 70 ppb-8hr, each considering any adjusted air quality 26 scenario.
- 27 • **Targeted Data Evaluations:** Afternoon time spent outdoors, along with ambient O_3 28 concentrations are the most influential factors when considering those persons highest 29 exposed. There is no apparent temporal trend in the amount of outdoor time or 30 participation rate when comparing historical CHAD diaries (1980s studies) to recently 31 collected diary data (2000s studies); regardless, majority of CHAD data are from studies 32 conducted since 2000. Use of activity pattern data from non-asthmatics to represent 33 asthmatics appears reasonably justified based on an evaluation indicating their having 34 similar outdoor time expenditure and attaining similar activity levels. APEX estimated 35 daily exposures are somewhat comparable to personal exposure measurements; however, 36 both over- and under-estimations occurred to varying degrees (Figure 5-15; Figure 5-16).

- APEX estimated ventilation rates were comparable to literature provided estimates,
 particularly those of school-age children (Figure 5-19).
- 3 Targeted Exposure Scenarios: When considering a modeling approach that more • 4 rigidly schedules longitudinal time location activity patterns compared with the standard 5 longitudinal approach used by APEX, a greater percent of persons experience at least one 6 or more exposures at or above benchmark levels. For example, an APEX model 7 simulation using only summer time (no school) CHAD diary days for non-working 8 school-age children generated approximately 30% more persons at or above exposure 9 benchmark levels compared with exposures estimated using our population-based 10 modeling approach (Figure 5-10). When accounting for a fraction of the population to 11 avert in response to a bad air quality day, approximately 1-2 percentage point fewer 12 persons experienced exposures at or above benchmark levels compared with exposures 13 estimated using our population based modeling approach (Figure 5-12). 14

Table 5-7 Mean and Maximum Percent of all School-age Children Estimated to Experience at Least One Daily Maximum 8-hr Average Exposure to O₃ at or Above Selected Health Benchmark Levels

	Adjusted Air	P One	g At Least evel ¹				
	Quality	6	0 ppb-8hr	7	′0 ppb-8hr	8	0 ppb-8hr
Study Area	Scenario	mean	max	mean	max	mean	max
	75	14.8	19.3	2.8	4.4	0.3	0.7
Atlanta	70	7.5	10.8	0.7	1.4	0.1	0.2
	65	2.9	4.8	0.2	0.5	0	0
	75	12.2	19.0	2.0	4.0	0.2	0.4
Baltimore	70	7.1	11.8	0.7	1.2	0.1	0.1
	65	3.0	5.4	0.2	0.3	0	0
	75	13.8	21.9	2.8	6.6	0.3	1.0
Boston	70	9.0	15.7	1.2	3.2	0.1	0.2
	65	3.4	6.7	0.2	0.5	0	0
	75	13.7	24.7	3.2	7.5	0.2	0.7
Chicago	70	9.2	16.0	1.0	2.7	0	0.1
	65	4.2	8.1	0.2	0.4	0	0
	75	10.2	18	1.4	3.7	0.1	0.2
Cleveland	70	4.2	9.3	0.3	0.9	0	0
	65	1.1	3.0	0.1	0.2	0	0
	75	12.9	22.9	1.9	4.5	0.1	0.3
Dallas	70	7.5	16.0	0.6	1.5	0	0.1
	65	3.0	7.6	0.1	0.3	0	0
	75	17.0	25.6	1.7	4.1	0.1	0.5
Denver	70	10.2	18.9	0.5	1.7	0	0.1
	65	3.8	9.5	0.1	0.4	0	0
	75	14.1	19.1	2.4	4.2	0.1	0.2
Detroit	70	7.3	10.3	0.5	0.9	0	0
	65	2.9	4.6	0.1	0.2	0	0
	75	11.4	17.8	2.3	5.5	0.3	0.7
Houston	70	6.6	11.9	0.8	2.1	0	0.1
	65	2.7	5.7	0.1	0.4	0	0
	75	9.5	10.2	0.6	1.0	0	0.1
Los Angeles	70	4.4	5.0	0.1	0.2	0	0
	65	1.1	1.5	0	0	0	0
	75	10.9	19.0	1.6	3.7	0.1	0.3
New York	70	3.3	6.6	0.2	0.5	0	0
	65	0	0.1	0	0	0	0
	75	13.8	20.5	2.1	4.2	0.2	0.4
Philadelphia	70	7.1	11.8	0.6	1.5	0	0.1
	65	2.4	4.6	0.1	0.3	0	0
	75	10.3	16.5	1.6	2.7	0.1	0.2
Sacramento	70	5.8	10.0	0.4	0.9	0	0
	65	2.7	4.7	0.1	0.2	0	0
	75	16.3	25.8	3.3	8.1	0.3	1.1
St. Louis	70	10.2	16.9	1.0	2.7	0.1	0.2
	65	3.9	7.3	0.1	0.4	0	0
	75	13.2	23.4	2.4	6.0	0.2	0.8
vvashington	70	6.6	12.5	0.6	1.4	0	0.1
	65	2.3	5.0	0.1	0.2	0	0

¹ The mean is the arithmetic average of the estimated percent of all school-age children exposed across 2006-2010 year air quality; max is the highest estimated percent of all school-age children exposed in a year.



Figure 5-20 Incremental increases in percent of all school-age children exposed to O_3 at or above 60 ppb-8hr for each study area, year 2006-2010 air quality. Average percent (left panels), maximum percent (right panels), at least one exposure (top panels), at least two exposures (bottom panels) per year.

Table 5-8 Mean and Maximum Percent of All School-age Children Estimated to Experience at Least Two Daily Maximum 8-hr Average Exposures to O₃ At or Above Selected Health Benchmark Levels

	Adjusted Air	P Two	ercent of Al Exposures	Experiencing At Least Benchmark Level ¹			
	Quality	6	0 ppb-8hr	7	0 ppb-8hr	8) ppb-8hr
Study Area	Scenario	mean	max	mean	max	mean	max
	75	6.0	8.9	0.4	0.7	0	0
Atlanta	70	2.1	3.3	0	0.1	0	0
	65	0.4	0.8	0	0	0	0
	75	4.6	8.4	0.2	0.5	0	0
Baltimore	70	1.8	3.7	0	0.1	0	0
	65	0.4	0.9	0	0	0	0
	75	4.5	9.7	0.3	1.1	0	0
Boston	70	2.2	5.5	0.1	0.4	0	0
	65	0.4	1.1	0	0	0	0
	75	5.3	11.6	0.5	1.3	0	0
Chicago	70	2.5	5.7	0.1	0.2	0	0
-	65	0.8	1.8	0	0	0	0
	75	3.1	7.5	0.1	0.5	0	0
Cleveland	70	0.9	2.6	0	0	0	0
	65	0.1	0.5	0	0	0	0
	75	4.8	12.2	0.2	0.8	0	0
Dallas	70	2.2	7.1	0	0.1	0	0
	65	0.5	2.0	0	0	0	0
	75	7.6	14.4	0.2	0.4	0	0
Denver	70	3.5	9.2	0	0.1	0	0
	65	0.7	2.8	0	0	0	0
	75	5.0	8.6	0.3	0.8	0	0
Detroit	70	1.9	3.6	0	0.1	0	0
	65	0.4	1.1	0	0	0	0
	75	3.8	6.3	0.2	0.6	0	0
Houston	70	1.5	2.9	0	0.1	0	0
	65	0.3	0.7	0	0	0	0
	75	4.1	4.5	0.1	0.1	0	0
Los Angeles	70	1.6	1.8	0	0	0	0
	65	0.3	0.3	0	0	0	0
	75	3.4	8.0	0.1	0.4	0	0
New York	70	0.5	1.4	0	0	0	0
	65	0	0	0	0	0	0
	75	5.0	8.7	0.2	0.5	0	0
Philadelphia	70	1.7	3.3	0	0.1	0	0
	65	0.3	0.6	0	0	0	0
	75	3.7	7.4	0.2	0.5	0	0
Sacramento	70	1.5	3.4	0	0.1	0	0
	65	0.4	0.9	0	0	0	0
	75	7.0	13.8	0.6	2.2	0	0.1
St. Louis	70	3.2	7.0	0.1	0.3	0	0
	65	0.7	2.0	0	0	0	0
	75	5.5	12.5	0.4	1.4	0	0
Washington	70	2.0	5.0	0	0.1	0	0
	65	0.4	1.2	0	0	0	0

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¹ The mean is the arithmetic average of the estimated percent of all school-age children exposed across 2006-2010

year air quality; max is the highest estimated percent of all school-age children exposed in a year.

1 5.7 REFERENCES

- Ainsworth, B. E.; W.L. Haskell; A. S. Leon; D. R. Jacobs Jr. and H. J. Montoye; J. F. Sallis and
 R. S. Paffenbarger Jr. 1993. "Compendium of Physical Activities: Classification of Energy
 Costs of Human Physical Activities." *Medicine & Science in Sports Exercise*, 25:71-80.
- Arcus-Arth, A. and J. Blaisdell. 2007. "Statistical Distributions of Daily Breathing Rates for
 Narrow Age Groups of Infants and Children." *Risk Analysis*, 27(1):97-110.
- Bresnahan, B. W.; M. Dickie and S. Gerking. 1997. "Averting Behavior and Air Pollution." *Land Economics*, 73:340-357.
- Brochu, P.; J.F. Ducre-Robitaille and J. Brodeur. 2006a. "Physiological Daily Inhalation Rates
 for Free-living Individuals Aged 1 Month to 96 Years, Using Data from Doubly Labeled
 Water Measurements: A Proposal for Air Quality Criteria, Standard Calculations and
- 12 Health Risk Assessment." *Human and Ecological Risk Assessment*, 12:675-701.
- Brochu, P.; J. F. Ducre-Robitaille and J. Brodeur. 2006b. "Supplemental Material for
 Physiological Daily Inhalation Rates for Free-living Individuals Aged 1 Month to 96
 Years, Using Data from Doubly Labeled Water Measurements: A Proposal for Air Quality
 Criteria, Standard Calculations and Health Risk Assessment." *Human and Ecological Risk Assessment*, 12:1-12.
- Burmaster, D.E. 1998. "LogNormal Distributions for Skin Area as a Function of Body Weight."
 Risk Analysis, 18(1):27-32.
- EPRI. 1988. A Study of Activity Patterns Among a Group of Los Angeles Asthmatics. Research
 Project 940-5. Electric Power Research Institute. Prepared by Roth Associates, November
 1988.
- EPRI. 1992. A Survey of Daily Asthmatic Activity Patterns in Cincinnati. TR-101396. Research
 Project 940-05. Electric Power Research Institute. Prepared by Roth Associates. November
 1992.
- Ford, E. S.; G. W. Heath; D. M. Mannino and S. C. Redd. 2003. "Leisure-time Physical Activity
 Patterns Among U.S. Adults with Asthma." *Chest*, 124:432-437.
- George, B. J. and T. McCurdy. 2009. "Investigating the American Time Use Survey from an
 Exposure Modeling Perspective." *Journal of Exposure Science and Environmental Epidemiology*, 21:92-105.
- Geyh, A. S.; J. Xue, H. Özkaynak and J. D. Spengler. 2000. "The Harvard Southern California
 Chronic Ozone Exposure Study: Assessing Ozone Exposure of Grade-school-age Children
 in Two Southern California Communities." *Environmental Health Perspectives*, 108:265 270.
- Glen, G.; L. Smith; K. Isaacs; T. McCurdy and J. Langstaff. 2008. "A New Method of
 Longitudinal Diary Assembly for Human Exposure Modeling." *Journal of Exposure Science and Environmental Epidemiology*, 18:299-311.
- Graham, S. E. and T. McCurdy. 2004. "Developing Meaningful Cohorts for Human Exposure
 Models." *Journal of Exposure Analysis and Environmental Epidemiology*, 14:23-43.
- 3 Graham, S. E. and T. McCurdy. 2005. *Revised Ventilation Rate (VE) Equations for Use in*
- 4 *Inhalation-oriented Exposure Models*. Washington, DC: EPA Office of Air and Radiation.
- 5 (EPA document number EPA/600/X-05/008, Appendix A). Available at:
- 6 <<u>http://cfpub.epa.gov/ncea/cfm/recordisplay.cfm?deid=202543</u>>.
- 7 Graham, S. 2009. *Response to Peer-review Comments on Appendix A*, prepared by S. Graham
- 8 (U.S. EPA). Response is Appendix D of US EPA (2009b). Research Triangle Park, NC:
- 9 EPA Office of Air and Radiation. Available at:
- 10 <<u>http://cfpub.epa.gov/ncea/cfm/recordisplay.cfm?deid=202543</u>>.
- 11 Graham, S. E. 2012. Comprehensive Review of Published Averting Behavior Studies and
- Available Technical Documents. Memo to Bryan Hubbell, Group Leader, Risk and
 Benefits Group, Office of Air Quality Planning and Standards. (EPA Docket # EPA-HQ-
- 14 OAR-2008-0699-0085).
- Isaacs, K. and L. Smith. 2005. New Values for Physiological Parameters for the Exposure Model
 Input File Physiology, text memorandum submitted to the U.S. Environmental Protection
 Agency under EPA Contract EP-D-05-065. NERL WA 10. Alion Science and Technology.
 Found in US EPA . (2009). Risk and Exposure Assessment to Support the Review of the
 SO₂ Primary National Ambient Air Quality Standard. (EPA document number EPA-
- 20 452/R-09-007, August 2009). Available at:
- 21 <<u>http://www.epa.gov/ttn/naaqs/standards/so2/data/200908SO2REAFinalReport.pdf</u>>.
- Isaacs, K.; G. Glen; T. McCurdy and L. Smith. 2008. "Modeling Energy Expenditure and
 Oxygen Consumption in Human Exposure Models: Accounting for Fatigue and EPOC."
 Journal of Exposure Science and Environmental Epidemiology, 18:289–298.
- Kansas Department of Health and Environment. 2006. Environmental Factors, Outdoor Air
 Quality, and Activity Level: Results from 2005 Kansas Behavioral Risk Factor Surveillance System. Office of Health Promotion, KDOH, available at:
- 28 <<u>http://www.kdheks.gov/brfss/PDF/cste_report_final.pdf</u>>.
- Langstaff, J. E. 2007. *Analysis of Uncertainty in Ozone Population Exposure Modeling*,
 OAQPS Staff Memorandum to Ozone NAAQS Review, January 31. Washington, DC:
- 31 Office of Air Radiation. (EPA docket number OAR-2005-0172). Available at:
- 32 <<u>http://www.epa.gov/ttn/naaqs/standards/ozone/s_O₃ cr_td.html</u>>.
- Lioy, P.J. 1990. "The Analysis of Total Human Exposure for Exposure Assessment: A Multi discipline Science for Examining Human Contact with Contaminants." *Environmental Science and Technology*, 24: 938-945.
- Mansfield, C.; F. R. Johnson and G. Van Houtven. 2006. "The Missing Piece: Averting Behavior
 for Children's Ozone Exposures." *Resource Energy Economics*, 28:215-228.

Marino, A. J.; E. N. Fletcher; R. C. Whitaker and S. E. Anderson. 2012. "Amount and Environmental Predictors of Outdoor Playtime at Home and School: Across-sectional

- Analysis of a National Sample of Preschool-aged Children Attending Head Start." *Health* & *Place*, 18: 1224-1230.
- McCurdy, T. 2000. "Conceptual Basis for Multi-route Intake Dose Modeling Using an Energy
 Expenditure Approach." *Journal of Exposure Analysis and Environmental Epidemiology*, 10:1-12.
- McCurdy, T.; G. Glen; L. Smith and Y. Lakkadi. 2000. "The National Exposure Research
 Laboratory's Consolidated Human Activity Database." *Journal of Exposure Analysis and Environmental Epidemiology*, 10:566-578.
- McDermott, M.; R. Srivastava and S. Croskell. 2006. "Awareness of and Compliance with Air
 Pollution Advisories: A Comparison of Parents of Asthmatics with Other Parents." *Journal of Asthma*, 43:235-239.
- McDonnell, W. F.; H. R. Kehrl; S. Abdul-Salaam; P. J. Ives; L. J. Folinsbee; R. B. Devlin; J. J.
 O'Neil and D. H. Horstman. 1991. "Respiratory Response of Humans Exposed to Low
 Levels of Ozone for 6.6 Hours." *Archives of Environmental Health*, 46(3):145-150.
- Meng, Q.; R. Williams and J. P. Pinto. (2012). "Determinants of the Associations Between
 Ambient Concentrations and Personal Exposures to PM_{2.5}, NO₂, and O₃ During DEARS."
 Atmospheric Environment, 63:109-116.
- Montoye, H. J.; H. C. G. Kemper; W. H. N. Saris and R. A. Washburn. 1996. *Measuring Physical Activity and Energy Expenditure*. Human Kinetics: Champaign, IL.
- National Research Council. 1991. *Human Exposure Assessment for Airborne Pollutants: Advances and Opportunities*. Washington, DC: National Academy of Sciences.
- Neidell, M. 2010. "Air Quality Warnings and Outdoor Activities: Evidence from Southern
 California Using a Regression Discontinuity Approach Design." *Journal of Epidemiology Community Health*, 64:921-926.
- Santuz, P.; E. Baraldi; M. Filippone and F. Zacchello. 1997. "Exercise Performance in Children
 with Asthma: Is it Different from that of Healthy Controls?" *European Respiratory Journal*, 10:1254-1260.
- Schofield, W. N. 1985. "Predicting Basal Metabolic Rate, New Standards, and Review of
 Previous Work." *Human Nutrition Clinical Nutrition*, 39C(S1):5-41.
- Sexton, A. L. (2011). "Responses to Air Quality Alerts: Do Americans Spend Less Time
 Outdoors?" Available at:
 <<u>http://www.apec.umn.edu/prod/groups/cfans/@pub/@cfans/@apec/documents/asset/cfans</u>
 asset 365645.pdf>.
- Shamoo, D. A.; W. S. Linn; R. C. Peng; J. C. Solomon; T. L. Webb; J. D. Hackney and H. Hong.
 1994. "Time-activity Patterns and Diurnal Variation of Respiratory Status in a Panel of
 Asthmatics: Implications for Short-term Air Pollution Effects." *Journal of Exposure*
- Asthmatics: Implications for Short-term Air Pollution Effects." *Journal of Exp* Analysis and Environmental Epidemiology, 4(2):133-148.

- U.S. Bureau of Labor Statistics. 2012. American Time Use Survey User's Guide. "Understanding
 ATUS 2003 to 2011," August 2012. Data and documentation available at:
 <<u>http://www.bls.gov/tus/</u>>.
- U.S. Census Bureau. 2007. "2000 Census of Population and Housing. Summary File 3 (SF3)
 Technical Documentation," available at:
- 6 http://www.census.gov/prod/cen2000/doc/sf3.pdf. Individual SF3 files '30' (for
- 7 income/poverty variables pct50) for each state were downloaded from:
- 8 <<u>http://www2.census.gov/census_2000/datasets/Summary_File_3/</u>>.
- 9 U.S. Department of Agriculture. 2000. "Continuing Survey of Food Intake by Individuals
 10 (CSFII) 1994-96," 1998. Agricultural Research Service, CD-ROM, available at:
 11 http://www.ars.usda.gov/Services/docs.htm?docid=14531>.
- U.S. Environmental Protection Agency. 1986. Air Quality Criteria for Ozone and Other
 Photochemical Oxidants. Research Triangle Park, NC: Office of Health and Environmental
 Assessment, Environmental Criteria and Assessment Office. (EPA document number EPA-600/8-84-020aF-eF). Available from the National Technical Information Service,
- 16 Springfield, VA. (NTIS publication number PB87-142949).
- U.S. EPA. 1996a. *Review of National Ambient Air Quality Standards for Ozone: Assessment of Scientific and Technical Information OAQPS Staff Paper*. Research Triangle Park, NC:
 EPA Office of Air Quality Planning and Standards. (EPA document number EPA/452/R96-007). Available at: http://www.epa.gov/ttn/naags/standards/ozone/s_O3_pr_sp.html>.
- U.S. EPA. 1996b. *Air Quality Criteria for Ozone and Related Photochemical Oxidants*. Research
 Triangle Park, NC: EPA Office of Research and Development, National Center for
 Environmental Assessment. (EPA document number EPA/600/P-93/004aF-cF). Available
 at: <<u>http://cfpub.epa.gov/ncea/cfm/recordisplay.cfm?deid=2831></u>.
- U.S. EPA. 2002. Consolidated Human Activities Database (CHAD) Users Guide. Database and
 documentation available at: <<u>http://www.epa.gov/chadnet1/</u>>.
- U.S. EPA. 2005. Guidance on Selecting Age Groups for Monitoring and Assessing Childhood
 Exposures to Environmental Contaminants. (EPA document number EPA/630/P-03/003F).
 Available at: <<u>http://www.epa.gov/raf/publications/pdfs/AGEGROUPS.PDF</u>>.
- U.S. EPA. 2006. Air Quality Criteria for Ozone and Related Photochemical Oxidants (Final).
 Research Triangle Park, NC: EPA National Center for Environmental Assessment. (EPA document number EPA/600/R-05/004aF-cF). Available at:
 http://cfpub.apa.gov/ncea/cfm/recordisplay.cfm2daid=149923
- 33 <<u>http://cfpub.epa.gov/ncea/cfm/recordisplay.cfm?deid=149923</u>>.
- 34 U.S. EPA. 2007a. *Review of National Ambient Air Quality Standards for Ozone: Policy*
- 35 Assessment of Scientific and Technical Information OAQPS Staff Paper. Research
- Triangle Park, NC: EPA Office of Air Quality Planning and Standards. (EPA document
- 37 number EPA-452/R-07-007). Available at:
- 38 <<u>http://www.epa.gov/ttn/naaqs/standards/ozone/data/2007_07_ozone_staff_paper.pdf</u>>.

- 1 U.S. EPA. 2007b. Ozone Population Exposure Analysis for Selected Urban Areas. Research 2 Triangle Park, NC: EPA Office of Air Quality Planning and Standards. Available at: 3 <http://www.epa.gov/ttn/naaqs/standards/ozone/s O₃ cr td.html>. 4 U.S. EPA. 2008. Risk and Exposure Assessment to Support the Review of the NO₂ Primary 5 National Ambient Air Quality Standard. Washington, DC: EPA Office of Air and 6 Radiation. (EPA document number EPA-452/R-08-008a0, November). Available at: 7 <http://www.epa.gov/ttn/naaqs/standards/nox/data/20081121 NO2 REA final.pdf>. 8 U.S. EPA. 2009a. Risk and Exposure Assessment to Support the Review of the SO₂ Primary 9 National Ambient Air Quality Standard. (EPA document number EPA-452/R-09-007, 10 August). Available at: <http://www.epa.gov/ttn/naaqs/standards/so2/data/200908SO2REAFinalReport.pdf>. 11 12 U.S. EPA. 2009b. Metabolically Derived Human Ventilation Rates: A Revised Approach Based 13 Upon Oxygen Consumption Rates. Washington, DC: EPA National Center for 14 Environmental Assessment. (EPA document number EPA/600/R-06/129F). Available at: 15 <http://cfpub.epa.gov/ncea/cfm/recordisplay.cfm?deid=202543>. 16 U.S. EPA. 2010. Quantitative Risk and Exposure Assessment for Carbon Monoxide – Amended. 17 Research Triangle Park, NC: EPA Office of Air Quality Planning and Standards. (EPA 18 document number EPA-452/R-10-009, July). Available at: 19 <http://www.epa.gov/ttn/naaqs/standards/co/data/CO-REA-Amended-July2010.pdf>. 20 U.S. EPA. 2011. Exposure Factors Handbook, 2011 edition. Washington, DC: EPA National 21 Center for Environmental Assessment. (EPA document number EPA/600/R-09/052F). 22 Available at: <<u>http://www.epa.gov/ncea/efh/pdfs/efh-complete.pdf</u>>. 23 U.S. EPA. 2012a. Total Risk Integrated Methodology (TRIM) - Air Pollutants Exposure Model 24 Documentation (TRIM.Expo / APEX, Version 4.4) Volume I: User's Guide. Research 25 Triangle Park, NC: EPA Office of Air Quality Planning and Standards. (EPA document 26 number EPA-452/B-12-001a). Available at: 27 <http://www.epa.gov/ttn/fera/human apex.html>. 28 U.S. EPA. (2012b. Total Risk Integrated Methodology (TRIM) - Air Pollutants Exposure Model 29 Documentation (TRIM.Expo / APEX, Version 4.4) Volume II: Technical Support 30 Document. Research Triangle Park, NC: EPA Office of Air Quality Planning and 31 Standards. (EPA document number EPA-452/B-12-001b). Available at: 32 <http://www.epa.gov/ttn/fera/human_apex.html>. 33 U.S. EPA. 2013. Integrated Science Assessment of Ozone and Related Photochemical Oxidants. 34 Research Triangle Park, NC: EPA National Center for Environmental Assessment. (EPA document number EPA 600/R-10/076F). Available at: 35 36 <http://cfpub.epa.gov/ncea/isa/recordisplay.cfm?deid=247492#Download>. 37 van Gent, R.; K. van der Ent; L. E. M. van Essen-Zandvliet; M. M. Rovers; J. L. L. Kimpen; G. 38 de Meer and P. H. C. Klijn. 2007. "No Difference in Physical Activity in (Un)diagnosed
- 39 Asthma and Healthy Controls." *Pediatric Pulmonology*, 42:1018-1023.

- 1 Wen, X. J.; L. Balluz and A. Mokdad. 2009. "Association Between Media Alerts of Air Quality 2 Index and Change of Outdoor Activity Among Adult Asthma in Six States," BRFSS, 2005. 3 Journal of Community Health, 34:40-46.
- 4 Whitfield, R.; W. Biller; M. Jusko and J. Keisler. 1996. A Probabilistic Assessment of Health 5 Risks Associated with Short- and Long-Term Exposure to Tropospheric Ozone. Argonne,
- 6 IL: Argonne National Laboratory.
- 7 World Health Organization. 2008. Uncertainty and Data Quality in Exposure Assessment. "Part 8 1: Guidance Document on Characterizing and Communicating Uncertainty in Exposure 9 Assessment." Available at:
- 10 <http://www.who.int/ipcs/publications/methods/harmonization/exposure assessment.pdf.
- 11 Xue, J.; T. McCurdy; J. Spengler and H. Özkaynak. 2004. "Understanding Variability in Time Spent in Selected Locations for 7-12 Year Old Children." Journal of Exposure Analysis 12 13 and Environmental Epidemiology, 14(3):222-33.
- 14 Zivin, J. G. and M. Neidell. 2009. "Days of Haze: Environmental Information Disclosure and
- 15 Intertemporal Avoidance Behavior." Journal of Environmental Economic Management,
- 58(2):119-128. 16

6 CHARACTERIZATION OF HEALTH RISKS BASED ON CONTROLLED HUMAN EXPOSURE STUDIES

3 6.1 INTRODUCTION

1

2

4 This chapter presents information regarding the methods and results for a controlled 5 human exposure-based O_3 (O_3) health risk assessment that builds upon the methodology used in 6 the assessment conducted as part of the O_3 NAAQS review completed in 2008 and also 7 introduces a new method for estimating risk. In the previous review, EPA conducted a health risk 8 assessment that produced risk estimates for the number and percent of school-aged children, 9 asthmatic school-aged children, and the general population experiencing lung function 10 decrements associated with O_3 exposures for 12 urban areas, where lung function is measured as 11 forced expiratory volume in one second (FEV₁). That portion of the risk assessment was based 12 on exposure-response relationships developed from analysis of data from several controlled 13 human exposure studies which were combined with population-level exposure distributions 14 developed for children and adults. Risk estimates for lung function decrements were developed 15 for recent air quality levels and for just meeting the existing 8-hour standard and several 16 alternative 8-hour standards. The methodological approach followed in the last risk assessment 17 and risk estimates resulting from that assessment are described in the 2007 Staff Paper (U.S. 18 EPA, 2007a).

19 The goals of the current O_3 risk assessment are to provide estimates of the number and 20 percentage of persons that would experience adverse respiratory effects associated with recent O_3 21 levels and with meeting the existing and potential alternative O_3 standards in specific urban 22 areas; and to develop a better understanding of the influence of various inputs and assumptions 23 on the risk estimates. The current assessment includes estimates of risks of lung function 24 decrements in school-aged children (ages 5 to 18), asthmatic school-aged children, and the adult 25 population (19 and above). We recognize that there are many sources of uncertainty in the inputs 26 and approach used in this portion of the health risk assessment which make the specific estimates 27 uncertain, however, we have sufficient confidence in the magnitude and direction of the 28 estimates provided by the assessment for it to serve as a useful input to decisions on the 29 adequacy of the O_3 standard and risk reductions associated with alternative standards. 30 We are estimating lung function risk using two methodologies in this review. The

31 primary results are based on a new model that estimates FEV_1 responses for individuals 32 associated with short-term exposures to O₃ (McDonnell, Stewart, and Smith, 2012). We refer to

33 this model as the McDonnell-Stewart-Smith (MSS) model. We also provide estimates following

34 the methodology used in previous reviews which provides population level estimates of the

1 percent and number of people at risk. We refer to this model as the Exposure-Response (E-R)

2 model used in previous reviews. Both of these models are implemented in the air pollution

3 exposure model APEX (EPA, 2012b,c). Following this introductory section, this chapter

4 discusses the scope of the controlled human exposure study based risk assessment, describes the

5 risk models, and provides key results from the assessment. The results of sensitivity analyses are

6 reported and key uncertainties are identified and summarized. More detailed descriptions of

7 several parts of the analyses are included in appendices that accompany the REA.

8 6.1.1 Development of Approach for Current Risk Assessment

9 The lung function risk assessment described in this chapter builds upon the methodology 10 and lessons learned from the risk assessment work conducted for previous reviews (EPA, 1996, 11 2007a). The current risk assessment also is based on the information evaluated in the ISA (EPA, 12 2013a). The general approach used in the current risk assessment was described in the Scope and 13 Methods Plan for Health Risk and Exposure (EPA, 2011), that was released to the CASAC and 14 general public in April 2011 for review and comment and which was the subject of a 15 consultation with the CASAC O₃ Panel in May 2011. The first draft REA was reviewed by 16 CASAC in September 2012. The approach used in the current risk assessment reflects 17 consideration of the comments offered by CASAC members and the public on the Scope and

18 Methods Plan and the first draft REA.

19 Controlled human exposure studies involve volunteers who are exposed while engaged in 20 different exercise regimens to specified levels of O_3 under controlled conditions for specified 21 amounts of time. For the current health risk assessment, we are using probabilistic exposure-22 response relationships based on analysis of individual data that describe the relationship between 23 measures of personal exposure to O_3 and measures of lung function recorded in the studies. 24 Therefore, a risk assessment based on exposure-response relationships derived from controlled 25 human exposure study data requires estimates of personal exposure to ambient O_3 . Because data 26 on personal hourly exposures to O_3 of ambient origin are not available, estimates of personal 27 exposures to varying ambient concentrations are derived through exposure modeling, as 28 described in Chapter 5. While the quantitative risk assessment based on controlled human 29 exposure studies addresses only lung function responses, it is important to note that other 30 respiratory responses have been found to be related to O_3 exposures in these types of studies, 31 including increased lung inflammation, increased respiratory symptoms, increased airway 32 responsiveness, and impaired host defenses. Sufficient information is not available to 33 quantitatively model these other endpoints. Section 6.2 of the ISA provides a discussion of these 34 additional health endpoints which are an important part of the overall characterization of risks

35 associated with ambient O_3 exposures.

6.1.2 Comparison of Controlled Human Exposure- and Epidemiologic-based Risk Assessments

In contrast to the **exposure-response** relationships derived from controlled human exposure studies, epidemiological studies provide estimated **concentration-response** relationships based on data collected in real world community settings. The assessment of health risk based on epidemiological studies is the subject of Chapter 7. The characteristics that are relevant to carrying out a risk assessment based on controlled human exposure studies versus one based on epidemiology studies can be summarized as follows:

- The relevant controlled human exposure studies in the ISA provide data that can be used to estimate exposure-response functions, and therefore a risk assessment based on these studies requires as input (modeled) personal exposures to ambient O₃. The relevant epidemiological studies in the ISA provide concentration-response functions, and, therefore, a risk assessment based on these studies requires as input (actual monitored or adjusted based on monitored) ambient O₃ concentrations, and personal exposures are not required as inputs to the assessment.
- 16 • Epidemiological studies are carried out in specific real world locations (e.g., specific 17 urban areas). To minimize extrapolation uncertainty, a risk assessment based on 18 epidemiological studies is best performed in locations where the studies took place. 19 Controlled human exposure studies, carried out in laboratory settings, are generally not 20 specific to any particular real world location. A risk assessment based on controlled 21 human exposure studies can therefore appropriately be carried out for any location for 22 which there are adequate air quality and other data on which to base the modeling of 23 personal exposures.
- To derive estimates of risk from concentration-response relationships estimated in
 epidemiological studies, it is usually necessary to have estimates of the baseline
 incidences of the health effects involved. Such baseline incidence estimates are not
 needed in a controlled human exposure studies-based risk assessment.
- 28

6.2 SCOPE OF LUNG FUNCTION HEALTH RISK ASSESSMENT

29 The current controlled human exposure-based O_3 health risk assessment is one approach 30 used to estimate risks associated with exposure to ambient O₃ in a number of urban areas 31 selected to illustrate the public health impacts of this pollutant. The short-term exposure related 32 health endpoints selected for this portion of the O₃ health risk assessment include those for which the ISA concludes that the evidence as a whole supports the general conclusion that O₃, acting 33 34 alone and/or in combination with other components in the ambient air pollution mix is causal or 35 likely to be causally related to the endpoint. 36 In the 2007 O₃ NAAQS review, the controlled human exposure-based health risk

- assessment involved developing risk estimates for lung function decrements ($\geq 10, \geq 15$, and
- 20% changes in FEV₁) in school-aged children (ages 5 to 18 years old). The strong emphasis

1 on children reflects the finding of previous O₃ NAAQS reviews that children are an important at-

2 risk group. Due to the increased amount of time spent outdoors engaged in relatively high levels

3 of physical activity (which increases intake), school-aged children as a group are particularly at

4 risk for experiencing O₃-related health effects.

5 Outdoor workers and other adults who engage in moderate exertion for prolonged 6 periods or heavy exertion for shorter periods during the day also are clearly at risk for 7 experiencing similar lung function responses when exposed to elevated ambient O₃ 8 concentrations. In this second draft REA, we focus the quantitative risk assessment for lung 9 function decrements on all and asthmatic school-aged children (ages 5-18), and the adult 10 population (ages 19 and above).

For the second draft assessment, lung function risks are estimated for 15 cities, Atlanta,
 Baltimore, Boston, Chicago, Cleveland, Dallas, Denver, Detroit, Houston, Los Angeles, New

13 York, Philadelphia, Sacramento, St. Louis, and Washington, DC.

14

6.2.1 Selection of Health Endpoints

15 The ISA identifies several responses to short-term O_3 exposure that have been evaluated 16 in controlled human exposure studies (US EPA, 2013, sections 6.2.1.1, 6.2.2.1, 6.2.3.1, and 17 6.3.1). These include decreased inspiratory capacity; decreased forced vital capacity (FVC) and 18 forced expiratory volume in one second (FEV₁); mild bronchoconstriction; rapid, shallow 19 breathing patterns during exercise; symptoms of cough and pain on deep inspiration (PDI); 20 increased airway responsiveness; and pulmonary inflammation. Such studies provide direct 21 evidence of relationships between short-term O₃ exposure and an array of respiratory-related 22 effects, however, there are only sufficient exposure-response data at different concentrations to 23 develop quantitative risk estimates for O_3 -related decrements in FEV₁. Other responses to O_3 24 which may be equally or more important then FEV_1 decrements (e.g., inflammation) do not 25 necessarily correlate with FEV_1 responses (ISA, section 6.2.3.1) and this risk assessment is not 26 able to address these other responses. 27 As stated in the 2006 Criteria Document (Table 8-3, p.8-68) for adults with lung disease, 28 even moderate functional responses (e.g., FEV₁ decrements \geq 10% but < 20%) would likely

29 interfere with normal activities for many individuals, and would likely result in more frequent

30 medication use. In a recent letter to the Administrator, the CASAC O₃ Panel stated that

31 "Clinically relevant' effects are decrements > 10%, a decrease in lung function considered

32 clinically relevant by the American Thoracic Society" (Samet, 2011, p.2). The CASAC O₃ Panel

33 also stated that:

a 10% decrement in FEV₁ can lead to respiratory symptoms, especially in
 individuals with pre-existing pulmonary or cardiac disease. For example,
 people with chronic obstructive pulmonary disease have decreased ventilatory

1 reserve (i.e., decreased baseline FEV₁) such that $a \ge 10\%$ decrement could lead 2 to moderate to severe respiratory symptoms (Samet, 2011, p.7). 3 4 This is consistent with the most recent official statement of the American Thoracic Society on 5 what constitutes an adverse lung function health effect of air pollution: 6 The committee recommends that a small, transient loss of lung function, by 7 itself, should not automatically be designated as adverse. In drawing the 8 distinction between adverse and nonadverse reversible effects, this committee 9 recommended that reversible loss of lung function in combination with the 10 presence of symptoms should be considered adverse (ATS, 2000, p.672). 11 12 For this lung function risk assessment, a focus on the mid- to upper-end of the range of 13 moderate levels of functional responses and higher (FEV₁ decrements \geq 15%) is appropriate for 14 estimating potentially adverse lung function decrements in active healthy adults, while for people 15 with asthma or lung disease, a focus on moderate functional responses (FEV₁ decrements down 16 to 10%) may be appropriate.

17 6.2.2 Approach for Estimating Health Risk Based on Controlled Human Exposure 18 Studies

19 The major components of the health risk assessment based on data from controlled 20 human exposure studies are illustrated in Figure 3-3 in Chapter 3. As shown in this figure, under 21 this portion of the risk assessment, exposure estimates for a number of different air quality 22 scenarios (i.e., recent year of air quality, just meeting the existing 8-hour and alternative 23 standards) are combined with probabilistic exposure-response relationships derived from the 24 controlled human exposure studies to develop risk estimates associated with recent air quality 25 and after simulating just meeting the existing and alternative standards. The health effect 26 included in this portion of the risk assessment is lung function decrement, as measured by 27 changes in FEV₁. The population risk estimates for a given lung function decrement (e.g., $\geq 15\%$ 28 reduction in FEV₁) are estimates of the expected number of people who will experience that lung 29 function decrement, the number of times that people experience repeated occurrences of given 30 lung function decrements, and the number of occurrences (person-days) of the given lung 31 function decrement. The air quality and exposure analysis components that are integral to this 32 portion of the risk assessment are discussed in Chapters 4 and 5. 33 We used two approaches to estimate health risk. As done for the risk assessment 34 conducted during the previous O₃ NAAQS review, a Bayesian Markov Chain Monte Carlo

35 approach was used to develop probabilistic exposure-response functions. These functions were

- 36 then applied to the APEX estimated population distribution of 8-hour maximum exposures for
- 37 persons at or above moderate exertion ($\geq 13 \text{ L/min-m}^2$ body surface area) to estimate the number

1 of persons expected to experience lung function decrements. The primary approach, based on the

- 2 McDonnell-Stewart-Smith FEV_1 model, uses the time-series of O_3 exposure and corresponding
- 3 ventilation rates for each APEX simulated individual to estimate their personal time-series of
- 4 FEV₁ reductions, selecting the daily maximum reduction for each person. A key difference
- 5 between these approaches is that the previous method estimates a population distribution of
- $6 \quad FEV_1$ reductions, where the MSS model estimates FEV_1 reductions at the individual level. Each
- 7 of these approaches is discussed in detail below.

8 6.2.3 Controlled Human Exposure Studies

Modeling of risks of lung function decrements as a function of exposures to O₃ is based
on application of results from controlled human exposure studies. As discussed in Chapter 6 of
the ISA (EPA, 2013a), there is a significant body of controlled human exposure studies reporting
lung function decrements and respiratory symptoms in adults associated with 1- to 8-hour
exposures to O₃. In the ISA sections on controlled human exposure (Sections 6.2.1.1, 6.2.2.1,
6.2.3.1, and 6.3.1) over 140 references to human clinical studies are reported.

15 6.2.3.1 Life Stages

16 Consistent with the approach used in the previous O_3 NAAQS review and lacking a 17 significant body of controlled human exposure studies on children, we judge that it is reasonable 18 to estimate exposure-response relationships for lung function decrements associated with O₃ 19 exposures in children 5-18 years old based on data from young adult subjects (18-35 years old). 20 As discussed in the ISA (EPA, 2013a), findings from clinical studies for children and summer 21 camp field studies of children 7-17 years old in at least six different locations in the U.S. and 22 Canada found lung function decrements in healthy children similar to those observed in healthy 23 young adults exposed to O₃ under controlled chamber conditions. There are fewer studies of 24 young children than adolescents to draw upon, which may add to uncertainties in the modeling. 25 Additional uncertainties are likely introduced since the lungs and airways of children are 26 developing, while development is complete in adults (Dietert et al., 2000). The primary period of 27 alveolar development is from birth to around eight years of age, but there is evidence for 28 continued development through adolescence. The adult number of alveoli is reached by 2-3 29 years of age and the size and surface area of the alveoli increase until after adolescence (Hislop, 30 2002; Narayanan et al., 2012).

Lung function responses to O_3 exposure for adults older than 18 decrease with age until around age 55, when responses are minimal. "Children, adolescents, and young adults appear, on average, to have nearly equivalent spirometric responses to O_3 , but have greater responses than middle-aged and older adults when similarly exposed to O_3 " (ISA p. 6-21). "In healthy individuals, the fastest rate of decline in O_3 responsiveness appears between the ages of 18 and 35 years (Passannante et al., 1998; Seal et al., 1996), more so for females then males (Hazucha et
al., 2003). During the middle age period (35-55 years), O₃ sensitivity continues to decline, but at
a much lower rate. Beyond this age (>55 years), acute O₃ exposure elicits minimal spirometric
changes" (ISA p. 6-23).

5 6.2.3.2 Asthma

6 There have been several controlled human exposure studies of the effects of O_3 on 7 asthmatic subjects, going back to 1978 (Linn et al., 1978). In reference to these studies, the ISA 8 states that "[b]ased on studies reviewed in the 1996 and 2006 O_3 AQCDs, asthmatic subjects 9 appear to be at least as sensitive to acute effects of O_3 as healthy nonasthmatic subjects" (ISA p. 6-20). Studies published since the 2006 O_3 AQCD do not alter this conclusion (ISA, p. 6-20 to 6-21). In the 2010 O_3 NAAQS proposal (75 FR 2969-2972), EPA describes the evidence that 12 people with asthma are as sensitive as, if not more sensitive than, normal subjects in manifesting

13 O₃-induced pulmonary function decrements.

14 In reference to epidemiologic studies, the ISA states that "[t]he evidence supporting

15 associations between short-term increases in ambient O₃ concentration and increases in

16 respiratory symptoms in children with asthma is derived mostly from examination of 1-h max,

17 8-h max, or 8-h avg O₃ concentrations and a large body of single-region or single-city studies.

18 The few available U.S. multicity studies produced less consistent associations." (ISA, p. 6-101 to

19 6-102). "Although recent studies contributed mixed evidence, the collective body of evidence

20 supports associations between increases in ambient O₃ concentration and increased asthma

21 medication use in children" (ISA, p. 6-109).

22 6.2.3.3 **Ethnicity**

There are two controlled human exposure studies that have assessed differences in lung function responses comparing ethnic groups (ISA, p. 6-23 to 6-24). Both of these studies show greater FEV_1 decrements in blacks than whites, however, epidemiologic studies were less supportive of this difference in response. The data available are insufficient to quantify any differences that might exist due to the limited number of studies and a lack of consistency between disciplines.

29 6.2.3.4 Body Mass Index

30 Some studies have found greater FEV_1 decrements to be associated with increasing BMI.

BMI was included in some of the models of McDonnell et al. (2012); however, the BMI terms

- 32 were found to be statistically insignificant, indicating that the effect of BMI on FEV_1 in the
- 33 presence of O_3 is likely to be small, within the range of BMIs of the subjects studied.

1 6.2.3.5 **Outdoor Workers**

Although there are no controlled human exposure studies that have had specifically
outdoor workers as subjects, the studies are applicable to outdoor workers: the 6.6-hour
experimental protocol was intended to simulate the performance of heavy physical labor for a
full workday (ISA, p. 6-9).

6 6.2.3.6 Variability of Responses

7 Responses to O_3 exposure are variable within the population, even within cohorts of 8 similar people (e.g., healthy young adult white males) (ISA, p. 6-16 to 6-20). Factors which 9 contribute to interindividual variability include health status, body mass index, age, sex, 10 race/ethnicity, and the intrinsic responsiveness of individuals. Other factors which contribute to 11 the variability of responses include the duration and concentration of O_3 exposure, the level of 12 exercise and breathing rate, attenuation due to repeated exposures, and co-exposures with other 13 pollutants. For specific individuals, lung function responses tend to be reproducible over a period 14 of several months.

15 6.2.4 The McDonnell-Stewart-Smith (MSS) Model

16 In this review, EPA is investigating the use of a new model that estimates FEV_1 17 responses for individuals associated with short-term exposures to O_3 (McDonnell, Stewart, and 18 Smith, 2007; McDonnell, Stewart, and Smith, 2010). This is a fundamentally different approach 19 than the previous approach, for which the exposure-response function is at a population level, not 20 an individual level. This model was developed using the controlled human exposure data 21 described in Section 6.2.5 as well as incorporating several additional data sets from studies using 22 shorter exposure durations and different exertion levels and breathing rates. These data were 23 from 15 controlled human O_3 exposure studies that included exposure of 541 volunteers (ages 18) 24 to 35^{1}) on a total of 864 occasions. These data are described in McDonnell et al. (1997). 25 Schelegle et al. (2009) found that there appears to be a delay in response when modeling FEV_1 26 decrements as a function of cumulative dose and estimated a threshold associated with the delay. 27 McDonnell et al. (2012) refit their 2010 model using data from eight additional studies with 201 28 subjects and incorporating a threshold parameter into the model. Their threshold parameter 29 allows for modeling a delay in response until cumulative dose rate (taking into account decreases 30 over time according to first order reaction kinetics) reaches a threshold value and is found by

31 McDonnell et al. (2012) to slightly improve model fit. That latest model is the model described

¹ The ages in these studies range from 18 years 1 month to 35 years 1 month.

here and is the model used in this risk assessment. The threshold is not a concentration thresholdand does not preclude responses at low concentration exposures.

3 Schelegle et al. (2012) have also developed a 2-compartment model for predicting FEV_1 4 decrements (ISA, p. 6-15,16). Their model is similar to the MSS model in that it accounts for the 5 effects of cumulative dose coupled with an exponential decay and also has a threshold, below 6 which response is delayed. The primary difference between this model and the MSS model is 7 that in the Schelegle et al. model the net cumulative dose is multiplied by an individual's 8 responsiveness coefficient to obtain a predicted FEV_1 decrement, whereas in the MSS model the 9 FEV₁ decrement increases as a sigmoid-shaped function of the net cumulative dose rate. Also, 10 the Schelegle et al. model's threshold is based on cumulative intake dose (the integral of 11 concentration x volume inhaled), where the MSS model's threshold is based on net cumulative 12 dose rate (taking into account the first order decay). A direct comparison of the results of these 13 two models has not been performed.

14 The MSS model is conceptually a two-compartment model. The cumulative amount of

15 exposure to O_3 (exposure concentration times ventilation rate, loosely speaking a measure of

16 dose rate) is modeled in the first compartment and modified by an exponential decay factor to

17 yield an intermediate quantity **X**. The response (lung function decrement) of the individual to **X**

18 is modeled in the second compartment (Figure 6-1). The threshold parameter imposes the

19 constraint that there is no response while the value of **X** is below the threshold value.



20 21

22 23

Figure 6-1. Two-Compartment Model

C is exposure concentration, V is ventilation rate, t is time, X is an intermediate quantity, a is a decay constant. Adapted from Figure 1 in McDonnell et al. (1999).

25

 \mathbf{X} is given by the solution of the differential equation (6-1):

2

1

$$\frac{dX}{dt} = C(t)V(t)^{\beta 6} - \beta_5 X(t)$$
 (Equation 6-1)

3

4 $\mathbf{X}(t)$ increases with "dose" $(C \cdot V^{\beta 6})$ over time for an individual and allows for removal of 5 O_3 with a half-life of $1/\beta_5$ through the 2nd term in equation (6-1). In APEX, because the exposure 6 concentration, exertion level, and ventilation rate are constant over an event, this equation has an 7 analytic solution for each event ("events" in APEX are intervals of constant activity and 8 concentration, where an individual is in one microenvironment, and range in duration from one 9 to 60 minutes):

10
$$X(t) = X(t_o) e^{-\beta 5(t-t_0)} + \frac{C(t)}{\beta_5} V(t)^{\beta 6} (1 - e^{-\beta 5(t-t_0)})$$
(Equation 6-2)

11

12

This model calculates the FEV_1 decrement due to O_3 exposure (compartment 2) as:

$$\mathscr{A}FEV1_{ijk} = e^{Ui}[\beta_1 + \beta_2(Age_{ik} - \bar{A})] \left\{ \frac{1}{1 + \beta_4 e^{-\beta_3 Tijk}} - \frac{1}{1 + \beta_4} \right\} + \varepsilon_{ijk}$$
(Equation 6-3)

13 where $T_{ijk} = \max\{0, \mathbf{X}_{ijk} - \beta_9\}$. β_9 is a threshold parameter which allows **X** to increase up to the 14 threshold before the median response is allowed to exceed zero.

- 15 The variables in the above equations are defined as:
- 16 The indices *i*,*j*,*k* refer to the *i*th subject at the *j*th time for the *k*th experiment for that subject,

17 C(t) is the O₃ exposure concentration at time t (ppm) during the event,

- 18 V(t) = VE(t)/BSA is the ventilation rate normalized by body surface area at time t 19 $(L/min-m^2)$,
- 20 VE(t) is the expired minute volume at time t (L min⁻¹),
- 21 BSA is the body surface area (m^2) ,
- 22 t is the time (minutes), t_0 is the time at the start of the event,
- 23 Age_{ik} is age in years of the *i*th subject in the *k*th study,
- \bar{A} is an age parameter (taken to be the approximate mean age of the clinical study subjects in the McDonnell, Stewart, and Smith 2007 (\bar{A} =25), 2010 (\bar{A} =25), and 2012 (\bar{A} =23.8) papers),
- 26 U_i is a subject-level random effect (between-individual variability not otherwise captured by 27 the model), and
- 28 ε_{ijk} is a variability term, which includes measurement error and intra-individual variability
- 29 not otherwise captured by the model.

1	The β s and the variances of the {U _i } and { ε_{ijk} } are fitted model parameters (see
2	McDonnell, et al. (2007, 2010, and 2012) for details). In APEX, values of U_i and ε_{ijk} are drawn
3	from Gaussian distributions with mean zero and variances $var(U)$ and $var(\mathcal{E})$, constrained to be
4	within ± 2 standard deviations from the means. The values of U _i are chosen once for each
5	individual and remain constant for individuals throughout the simulation. The ε_{ijk} are sampled
6	daily for each individual. The best fit values (based on maximum likelihood) for these
7	parameters are listed in Table 6-1. The values in parentheses are standard errors of the estimates
8	(given here to two significant digits; the values in the papers are given to up to five significant
9	digits). Although some of the parameters are quite different in the three models in Table 6-1, the
10	predictions of these three models are similar. The relative influences of the parameters are
11	discussed in Section 6.5.1.

	-
1	\mathbf{n}
	4

 Table 6-1. Estimated Parameters in the MSS Models

Model	β1	β2	β3	β4	β5	β6	β9	var(U)	var(E)
2007 ¹ ,	9.9047	-0.4106	0.0164	46.9397	0.003748	0.9123		0.835	13.8279
2010 ²	(0.61)	(0.11)	(0.0030)	(7.3)	(0.00027)	(0.054)		(0.080)	(0.36)
2012 ³	9.8057 (0.74)	-0.1907 (0.28)	0.01839 (0.0051)	65.826 (12)	0.003191 (0.00021)	0.8753 (0.086)	0	0.9449 (0.083)	17.120 (1.2)
2012T ⁴	10.916	-0.2104	0.01506	13.497	0.003221	0.8839	59.284	0.9373	17.0816
	(0.84)	(0.31)	(0.0033)	(4.7)	(0.00021)	(0.065)	(10)	(0.082)	(1.2)

14 ¹ McDonnell, Stewart, and Smith (2007). $\overline{A} = 25$.

15 ² McDonnell, Stewart, and Smith (2010). $\overline{A} = 25$.

16 ³ McDonnell, et al. (2012). $\overline{A} = 23.8$. No-threshold.

⁴ McDonnell, et al. (2012). $\overline{A} = 23.8$. Threshold.

18 We are using this model to estimate lung function decrements for people ages 5 and 19 older. However this model was developed using only data from individuals aged 18 to 35 and the 20 age adjustment term $[\beta_1 + \beta_2 (Age_{ijk} - \bar{A})]$ in the model is not appropriate for all ages. In addition 21 to this age term, the effects of age are also taken into account through the dependence of

22 ventilation rate and body surface area on age. The APEX estimates of lung function risk for

23 different age groups are also influenced by the time spent outdoors and the activities engaged in

by those groups, which vary by age (see Appendix 6-E).

Clinical studies data for children which could be used to fit the model for children are not
available at this time. In the absence of data, we are extending the model to ages 5 to 18 by

holding the age term constant at the age 18 level. Since the response increases as age decreases

in the range 18 to 35, this trend may extend into ages of children, in which case the responses of

1 children could be underestimated. However, the slope of the age term in the MSS model is

- 2 estimated based on data for ages 18 to 35 and does not capture differences in age trend within
- 3 this range; in particular, we don't know at what age the response peaks, which could be above or
- 4 below 18. The evidence from clinical studies indicates that the responsiveness of children to O_3

5 is about the same as for young adults (ISA, 2012, p. 6-21). This suggests that the age term for

6 children should not be higher than the age term for young adults.

7 Because the responses to O_3 decline from age 18 until around age 55 and for ages older 8 than 55 the response are minimal, we let the age term for ages 35 to 55 linearly decrease to zero 9 and set it to zero for ages > 55.

"In healthy individuals, the fastest rate of decline in O₃ responsiveness
appears between the ages of 18 and 35 years During the middle age period
(35-55 years), O₃ sensitivity continues to decline, but at a much lower rate.
Beyond this age (>55 years), acute O₃ exposure elicits minimal spirometric
changes." (ISA, 2012, p. 6-23)

In order to extend the age term to ages outside the range of ages the MSS model is based on (ages 18-35), we parameterize the age term by $[\beta_1 + \beta_2(\alpha_1 \operatorname{Age} + \alpha_2)]$, for different ranges of ages (α_1 and α_2 depend on age), requiring that these terms match at each boundary to form a piecewise linear continuous function of age. The foregoing assumptions result in the following values of α_1 and α_2 for four age ranges (Table 6-2).

NIO	vioder to All Ages							
	Age Range	β1	β2	α_1	α_2			
	5 – 17	10.916	-0.2104	0	-5.8			
	18 – 35	10.916	-0.2104	1	-23.8			
	36 – 55	10.916	-0.2104	2.0341	-59.994			
	> 55	0	0	0	0			

Table 6-2. Age Term Parameters for Application of the 2012 MSS Threshold Model to All Ages

22

23 The lung function decrements estimated by the MSS (2010) model for a particular case 24 are illustrated in Figure 6-2 and Figure 6-3. Figure 6-2 shows the predictions of the MSS model for 20-year old individuals with a (typical) body surface area (BSA) of 2 m^2 and a target 25 ventilation rate of 40 L/min (moderate exertion) and an O₃ exposure level of 100 ppb, under the 26 27 conditions of a typical 6.6-hour clinical study. Subjects alternated 50 minutes of moderate 28 exercise with 10 minutes of rest for the first three hours, with the exercise occurring first. For the 29 next 35 minutes (lunch), subjects continued exposure at rest. For the remaining three hours of the 30 exposure period, subjects again alternated 50 minutes of exercise with 10 minutes of rest. The

- 1 inter-individual variability predicted by this model is depicted by the boxplots in this figure. The
- 2 predictions for the median individual over time are given by the line. Minute-by-minute
- 3 predictions for the median individual for an exposure level of 100 ppb are shown in Figure 6-3.
- 4 The stairstep response results from the pattern of exercise and rest during the experiment.
- 5



Figure 6-2. Distribution of Responses (Lung Function Decrements in FEV₁)
 Predicted by the MSS Model for 20-Year Old Individuals. Exposure to 100 ppb O₃ at
 Moderate Exercise (40 L/min, BSA=2 m²) Under the Conditions of a Typical 6.6-hour

- 10 Clinical Study.
- 11



13 drawn at the 50th percentile (median). The whiskers are at the 1st and 99th percentiles.



Figure 6-3. Median Response (Lung Function Decrements in FEV₁) Predicted by
 the MSS Model for 20-Year Old Individuals. Exposure to 100 ppb O₃ at Moderate Exercise
 (40 L/min, BSA=2 m²) Under the Conditions of a Typical 6.6-hour Clinical Study.

Figure 6-4 and Figure 6-5 illustrate the threshold effect based on McDonnell et al.

7 (2012). Figure 6-4 is a graph of the median response for a population of 20-year old individuals

8 over a 6.6-hour time period. The exposure concentration is a constant 100 ppb over this time

- 9 period, while the individuals are exercising from hour 1 to hour 3 and at rest otherwise. There is
- 10 a 30-minute delay in response due to the threshold; without the threshold, the response starts
- 11 increasing when exercise starts. Figure 6-5 shows the corresponding probability of a response
- 12 (FEV₁ decrement) \geq 10% over the time period for the two models. There is very little difference
- 13 in response between the threshold and non-threshold models.



Figure 6-4. Median Response (FEV₁ Decrements) Predicted by the MSS Threshold and Non-Threshold Models for 20-Year Old Individuals, Constant 100 ppb O₃ Exposure, 2 Hours Heavy Exercise (30 L/min-m² BSA).



Figure 6-5. Probability of Response $\geq 10\%$ Predicted by the MSS Threshold and Non-Threshold Models for 20-Year Old Individuals, Constant 100 ppb O₃ Exposure, 2 Hours Heavy Exercise (30 L/min-m² BSA).

1 6.2.5 The Exposure-Response Function Approach Used in Prior Reviews

2 As described in section 3.1.2 of the 2007 Risk Assessment Technical Support Document 3 (EPA, 2007b), a Bayesian Markov Chain Monte Carlo approach (Lunn et al., 2012) was used to 4 estimate probabilistic exposure-response relationships for lung function decrements associated 5 with 8-hour O_3 exposures occurring at moderate exertion. In the previous review, summary data 6 from the Folinsbee et al. (1988), Horstman et al. (1990), McDonnell et al. (1991), and Adams 7 (2002, 2003, 2006) studies were combined to estimate exposure-response relationships for 8hour exposures at moderate exertion for each of the three measures of lung function decrement 8 9 $(\geq 10, \geq 15, \geq 20\%$ decrements in FEV₁). In this second draft REA we have updated this 10 exposure-response function with the results from two additional studies (Kim et al., 2011; 11 Schelegle et al., 2009). The controlled human exposure study data were corrected for the effect 12 of exercise in clean filtered air on an individual basis to remove any systematic bias that might 13 be present in the data attributable to an exercise effect (ISA, Section 6.2.1.1). This is done by 14 subtracting the FEV_1 decrement in filtered air from the FEV_1 decrement (at the same time point) 15 during exposure to O_3 . An example of this calculation is given in Appendix 6-D. 16 Table 6-3 presents a summary of the study-specific results based on correcting all 17 individual responses for the effect on lung function decrements of exercise in clean air.

Table 6-3. Study-specific O₃ Exposure-response Data for Lung Function Decrements Based on Correcting Individual Responses for the Effect on Lung Function of Exercise in Clean Air, Ages 18-35

Study Grouped by		Number	Number of Responses ^a				
Average O ₃ Exposure	Protocol	Exposed	$\frac{\Delta FEV_1 \geq}{10\%}$	ΔFEV ₁ ≥ 15%	ΔFEV ₁ ≥ 20%		
0.04 ppm O ₃							
Adams et al. (2002)	Square-wave, face mask	30	2 (2)	0 (0)	0 (0)		
Adams et al. (2006)	Triangular	30	0 (0)	0 (0)	0 (0)		
0.06 ppm O ₃							
A dama at al. (2000)	Square-wave	30	2 (2)	0 (0)	0 (0)		
Adams et al. (2006)	Triangular	30	2 (2)	2 (2)	0 (0)		
Kim et al. (2011)	Square-wave	59	3 (6)	1 (3)	0 (0)		
Schelegle et al. (2009) Variable levels (0.06 ppm avg)		31	4 (8)	2 (3)	1 (1)		
0.07 ppm O ₃	0.07 ppm O ₃						
Schelegle et al. (2009)	Variable levels (0.07 ppm avg)	31	6 (12)	3 (7)	2 (3)		
0.08 ppm O ₃	0.08 ppm O ₃						
Adams et al. (2002)	Square-wave, face mask	30	6 (6)	5 (5)	2 (2)		

Table 6-3. Study-specific O₃ Exposure-response Data for Lung Function Decrements Based on Correcting Individual Responses for the Effect on Lung Function of Exercise in Clean Air, Ages 18-35

Study Grouped by		Numbor	Nu	Number of Responses ^a			
Average O ₃ Exposure Protocol		Exposed	$\begin{array}{c} \Delta FEV_1 \geq \\ 10\% \end{array}$	ΔFEV₁≥ 15%	$\frac{\Delta FEV_1}{20\%} \ge$		
	Square-wave, chamber	30	6 (6)	2 (2)	1 (1)		
	Square-wave, face mask	30	5 (5)	2 (2)	2 (2)		
Adams et al. (2003)	Variable levels (0.08 ppm avg), chamber	30	6 (6)	1 (1)	1 (1)		
avg), chamber Variable levels (0.08 ppm avg), face mask Adams et al. (2006) Square-wave F-H-M ^b Square-wave Kim et al. (2011) Square-wave	30	5 (5)	1 (1)	1 (1)			
A dame at al. (2006)	Square-wave	30	7 (7)	2 (2)	1 (1)		
Adams et al. (2000)	Triangular	30	9 (9)	3 (3)	1 (1)		
F-H-M ^b	Square-wave	60	17 (19)	11 (14)	8 (8)		
Kim et al. (2011)	Square-wave	30	4 (6)	1 (1)	0 (0)		
Schelegle et al. (2009)	Variable levels (0.08 ppm avg)	31	10 (15)	5 (8)	4 (6)		
0.087 ppm O ₃							
Schelegle et al. (2009)	Variable levels (0.087 ppm avg)	31	14 (17)	10 (12)	7 (9)		
0.1 ppm O ₃							
F-H-M ^b	Square-wave	32	13 (13)	11 (12)	6 (9)		
0.12 ppm O ₃							
A dome at al. (2002)	Square-wave, chamber	30	17 (17)	12 (12)	10 (10)		
Adams et al. (2002)	Square-wave, face mask	30	21 (21)	13 (13)	7 (7)		
F-H-M ^b	Square-wave	30	18 (19)	15 (15)	10 (10)		

a. The first number in each cell is the number of responses based on post-exposure decrements in FEV_1 (i.e., we used only the last FEV_1 measurement and the pre-exposure FEV_1 to obtain a single percentage change in FEV_1 for each subject in each experiment). The numbers in parentheses are the numbers of responses based on maximum FEV_1 decrements. Specifically, when there were multiple FEV_1 measurements after the beginning of the exposure, we calculated multiple FEV_1 percentage changes for each subject in each experiment and used the maximum change when calculating the numbers of responses greater than 10%, 15%, and 20%.

- b. F-H-M combines data from Folinsbee et al. (1988), Horstman et al. (1990), and McDonnell et al. (1991).
- 1 2
- For the risk assessment conducted during the 2007 O_3 NAAQS review (EPA, 2007b),
- 3 EPA considered both linear and logistic functional forms in estimating the exposure-response
- 4 relationship and chose a 90 percent logistic/10 percent piecewise-linear split using a Bayesian
- 5 Markov Chain Monte Carlo approach. This Bayesian estimation approach incorporates both
- 6 model uncertainty and uncertainty due to sampling variability.

For each of the three measures of lung function decrement, EPA assumed a 90 percent
 probability that the exposure-response function has the following 3-parameter logistic form:²

$$y(x; \alpha, \beta, \gamma) = \frac{\alpha^* e^{\gamma} (1 - e^{\beta x})}{(1 + e^{\gamma})(1 + e^{\beta x + \gamma})},$$
 (Equation 6-4)

4 where *x* denotes the O₃ concentration (in ppm) to which the individual is exposed, y 5 denotes the corresponding response (decrement in FEV₁ \ge 10%, \ge 15% or \ge 20%), and α , β , and 6 γ are the three parameters whose values are estimated.

7 We assumed a 10 percent probability that the exposure-response function has the8 following linear with threshold (hockey stick) form:

3

9
$$y(x; \alpha, \beta) = \begin{cases} \alpha + \beta x, & \text{for } \alpha + \beta x > 0\\ 0, & \text{for } \alpha + \beta x < 0 \end{cases}$$
 (Equation 6-5)

10 We assumed that the number of responses, S, out of N subjects exposed to a given 11 concentration, x, has a binomial distribution with response probability given by Eq (6-4) with 90 12 percent probability and response probability given by Eq (6-5) with 10 percent probability. In the 13 2007 review, we also considered 80/20 and 50/50 probabilities for the logistic and hockey stick 14 forms, and ran those as sensitivity analyses. We performed those analyses with the updated data 15 and found that for each of the three exposure-response curves, the 90/10 mix has smaller error in 16 fit (weighted RMSE) than the other two combinations of probabilities, and we are using only that 17 function in this review.

18 In some of the controlled human exposure studies, subjects were exposed to a given O_3 19 concentration more than once – for example, using a constant (square-wave) exposure pattern in 20 one protocol and a variable (triangular) exposure pattern in another protocol. However, because 21 there were insufficient data to estimate subject-specific response probabilities, we assumed a 22 single response probability (for a given definition of response) for all individuals and treated the 23 repeated exposures for a single subject as independent exposures in the binomial distribution. 24 For each of the two functional forms (logistic and linear), we derived a Bayesian 25 posterior distribution using this binomial likelihood function in combination with prior 26 distributions for each of the unknown parameters (Box and Tiao, 1973). We assumed lognormal 27 priors with maximum likelihood estimates of the means and variances for the parameters of the 28 logistic function, and normal priors, similarly with maximum likelihood estimates for the means 29 and variances, for the parameters of the linear function. For each of the two functional forms

² The 3-parameter logistic function is a special case of the 4-parameter logistic, in which the function is forced to go through the origin, so that the probability of response to 0.0 ppm is 0.

considered, we used 1,000 iterations as the "burn-in" period³ followed by 9,000 iterations for the 1 2 estimation. Each iteration corresponds to a set of values for the parameters of the (logistic or 3 linear) exposure-response function. We combined the 9,000 sets of values from the logistic 4 model runs with the last 1,000 sets of values from the linear model runs to get a single combined 5 distribution of 10,000 sets of values reflecting the 90 percent/10 percent assumption stated 6 above. WinBUGS version 1.4.3 was used for these analyses (WinBUGS; Lunn et al., 2012). For any O_3 concentration, x, we can derive the nth percentile response value, for any n, by 7 8 evaluating the exposure-response function at x using each of the 10,000 sets of parameter values 9 (9,000 of which were for a logistic model and 1,000 of which were for a linear model). The resulting 2.5th percentile, median (50th percentile), and 97.5th percentile exposure-response 10 11 functions for changes in $FEV_1 \ge 10\%$ are shown in Figure 6-6, along with the response data to which they were fit. The corresponding exposure-response functions for changes in $FEV_1 >$ 12 15% and > 20% are shown in Appendix 6-A. The values of the functions are also provided in 13 14 Appendix 6-A.

³ Markov chain Monte Carlo (MCMC) simulations require an initial adaptive "burn-in" set of iterations, which are not used. This allows the MCMC sampling to stabilize.



Figure 6-6. Probabilistic Exposure-Response Relationships for FEV_1 Decrements \geq 10% for 8-Hour Exposures At Moderate Exertion, Ages 18-35. Values associated with data points are the number of subject-exposures at each exposure concentration.

The population risk is estimated by multiplying the expected risk by the number of people in the relevant population, as shown in Equation 6-6 below. The risk (i.e., expected fractional response rate) for the k^{th} fractile, R_k is estimated as:

Ν

10
$$R_k = \sum_{j=1}^{k} P_j x(RR_k \mid e_j)$$

(Equation 6-6)

11 where:

 $e_j = (\text{the midpoint of}) \text{ the } j\text{th category of personal exposure to } O_3;$

 P_j = the fraction of the population having personal exposures to O₃ concentration 14 of e_j ppm;

15
$$RR_k | e_j = k$$
-fractile response rate at O₃ exposure concentration e_j;

N = number of intervals (categories) of O₃ personal exposure concentration.

1 Exposure estimates used in this portion of the risk assessment were obtained from APEX 2 for each of the fifteen urban areas and the five air quality scenarios. Chapter 5 provides 3 additional details about the inputs and methodology used to estimate population exposure in the 4 urban areas. Exposure estimates for all and asthmatic school-aged children (ages 5 to 18) were 5 combined with probabilistic exposure-response relationships for lung function decrements 6 associated with 8-hour exposure while engaged in moderate exertion. Individuals engaged in activities that resulted in an average equivalent (BSA-normalized) ventilation rate (EVR) for the 7 8-hour period at or above 13 L/min-m² BSA were included in the exposure estimates for 8-hour 8 9 moderate or greater exertion. This range was selected based on the EVRs for the group of 10 subjects in the controlled human exposure studies that were the basis for the exposure-response 11 relationships used in this portion of the risk assessment.

12 6.3 O₃ RISK ESTIMATES

13 This section provides lung function risk estimates associated with several air quality 14 scenarios: five recent years of air quality as represented by 2006 to 2010 monitoring data, and air 15 quality in those years after simulating just meeting the existing O_3 standard and alternative 16 standard levels of 0.070, 0.065, and 0.060 ppm. The risk measures presented here are the 17 percents of the population estimated to experience lung function responses greater then 10, 15, 18 and 20%, one or more times or six or more times during an O_3 season, for three age groups: 19 school-aged children (ages 5-18), young adults (ages 19-35) and adults ages 36-55. Results for 20 adults older than 55 are not presented since the responses for this age group are estimated to be 21 minimal. People with multiple events with large lung function decreases are more at risk than 22 those with only one such event during the O₃ season. Although six events is less than once per 23 month, we see dramatic decreases in population risk in going from one or more to six or more 24 events during a season, which is why we report on six or more events rather than a higher 25 number.

In the figures and tables that follow, "base" indicates the base case scenario of recent air 26 27 quality for the indicated year. "75," "70," "65," and "60" respectively indicate the existing O₃ 28 standard and alternative standard levels of 0.070, 0.065, and 0.060 ppm. "75 6-8" indicates the 29 0.075 ppm existing 8-hour standard based on rollback for the 2006-2008 period, while "75 8-10" 30 indicates the existing standard scenario based on rollback for the 2008-2010 period. There are 31 two estimates of results for the 2008 existing and alternative standard scenarios (because 2008) 32 overlaps the two rollback periods) and one for each of the other four years. These two estimates 33 for 2008 can be quite different because of the relationship between the design value over the 34 three-year period and the amount of adjustment to the air quality distribution in 2008 that can 35 result.

6-21

6.3.1 Lung Function Risk Estimates Based on the McDonnell-Stewart-Smith Model

2 Results based on the McDonnell-Stewart-Smith (2012) threshold model are summarized
3 in this section; detailed results can be found in Appendix 6-B.

Figure 6-7 shows the results for school-aged children in the same format used in exposure results, explained in Section 5.3.1. Figure 6-7 depicts results for all cities, year, and scenarios for ages 5 to 18 with \geq 1 occurrences of FEV₁ decrements \geq 10, 15, 20% and illustrates the variation of results across cities, year, and scenarios.

8 Figure 6-8 shows the variation across cities (horizontally) and years (vertically) for the 9 percent of school-aged children with ≥ 1 occurrences of FEV₁ decrements $\geq 10\%$ with air quality 10 just meeting the potential alternative standard of 0.07 ppm. The points above each study area on 11 this graph represent the risk for the six years for the study area (2008 has two points,

the study area (2008 has two points,

12 corresponding to the different 2006-2008 and 2008-2010 design values used to adjust the air

13 quality to meet 0.07 ppm). There is substantial variability both across years and across cities.

14 Denver has the highest overall risks, while Cleveland and New York have the lowest. Los

15 Angeles has the smallest variation across years, with a range of 2.3% (from 14.3% to 16.6%).

16 The other cities have a range of around 4% to 7.5% across years.

17Table 6-4 and Table 6-5 present summary results (ranges over cities and years) of FEV_1 18decrements ≥ 10 and 15% estimated by the MSS model for the different age groups. The results19for asthmatic school-aged children are very similar to the results for all school-aged children and20are not presented here.

21 Figure 6-9 illustrates the distribution of responses (FEV₁ decrements > 10%) across

22 ranges of ambient concentrations of O_3 for school-aged children for one city and scenario (Los

23 Angeles, 2006 recent air quality). The concentrations are daily 8-hour average ambient

concentrations during the 8-hour period with maximum 8-hour average exposure for that day.



Figure 6-7. Risk results for all school-aged children with ≥ 1 occurrences of FEV₁ decrements ≥ 10 , 15, 20% for all cities, year, and scenarios (y-axis is percent of children affected).



3Figure 6-8. Risk results for all school-aged children with ≥ 1 occurrences of FEV14decrements $\geq 10\%$ under the 0.07 ppm alternative standard showing variability across5cities (horizontally) and years (vertically).

Table 6-4. Ranges of percents of population experiencing one or more days during the O_3 season with lung function decrement (ΔFEV_1) more than 10%. The numbers in this table are the minimum and maximum percents estimated over all cities and years.

		percent experienci ∆FEV₁ ≥	$mg \ge 1$ day with $\ge 10\%$	$\begin{array}{c} \mbox{percent experiencing} \geq 6 \mbox{ days with} \\ \Delta FEV_1 \geq 10\% \end{array}$		
Age group	Scenario	minimum	maximum	minimum	maximum	
5 to 18	base	11%	31%	1%	9%	
5 to 18	75	11%	22%	1%	6%	
5 to 18	70	8%	20%	1%	5%	
5 to 18	65	2%	18%	0%	4%	
5 to 18	60	4%	13%	0%	3%	
19 to 35	base	3%	13%	0%	1%	
19 to 35	75	3%	9%	0%	1%	
19 to 35	70	2%	8%	0%	1%	
19 to 35	65	1%	6%	0%	1%	
19 to 35	60	1%	5%	0%	0%	
36 to 55	base	1%	4%	0%	0%	
36 to 55	75	1%	2%	0%	0%	
36 to 55	70	0%	2%	0%	0%	
36 to 55	65	0%	2%	0%	0%	
36 to 55	60	0%	1%	0%	0%	
> 55	All	0%	0%	0%	0%	

Table 6-5. Ranges of percents of population experiencing one or more days during the O₃ season with lung function decrement (Δ FEV₁) more than 15%. The numbers in this table are the minimum and maximum percents estimated over all cities and years.

3 4

		percent experie AFEV	ncing \geq 1 day with $T_1 \geq$ 15%	percent experiencing \geq 6 days with $\Delta FEV_1 \geq 15\%$		
Age group	Scenario	minimum	maximum	minimum	maximum	
5 to 18	base	2%	12%	0%	3%	
5 to 18	75	2%	6%	0%	1%	
5 to 18	70	2%	5%	0%	1%	
5 to 18	65	0%	4%	0%	1%	
5 to 18	60	0%	3%	0%	0%	
19 to 35	base	0%	3%	0%	0%	
19 to 35	75	0%	2%	0%	0%	
19 to 35	70	0%	1%	0%	0%	
19 to 35	65	0%	1%	0%	0%	
19 to 35	60	0%	1%	0%	0%	
36 to 55	base	0%	1%	0%	0%	
36 to 55	75	0%	0%	0%	0%	
36 to 55	70	0%	0%	0%	0%	
36 to 55	65	0%	0%	0%	0%	
36 to 55	60	0%	0%	0%	0%	
> 55	All	0%	0%	0%	0%	

These concentrations are less than but close to daily maximum 8-hour average ambient concentrations and are greater than daily maximum 8-hour average exposure concentrations. The 9 percents in this chart reflect the frequencies of person-days with FEV₁ decrements $\geq 10\%$ within a concentration bin as percents of all person-days with FEV₁ decrements $\geq 10\%$. Figure 6-9 10 shows that more than 90% of daily instances of FEV_1 decrements $\geq 10\%$ occur when 8-hour 11 average ambient concentrations are above 40 ppb for this modeled scenario. This distribution 12 13 will be different for different cities, years, and air quality scenarios.



1 8-hour average concentration (ppb) 2 Figure 6-9. Distribution of Daily FEV₁ Decrements ≥ 10% Across Ranges of 8-hour 3 Average Ambient O₃ Concentrations (Los Angeles, 2006 recent air quality). 4

5 Outdoor workers spend more time outdoors than the general population and therefore are 6 at higher risk for health effects due to O₃. We conducted simulations of outdoor workers ages 19-7 35 for Atlanta (2006) for the current and alternative standards to estimate the risk of this group 8 for experiencing FEV₁ decrements \geq 15%. The methodology for simulating outdoor workers 9 involves modifying activity diaries to represent outdoor workers and is described in Section 10 5.3.3 in Chapter 5. Table 6-6 shows the results of these simulations and compares them with the results for the general population, ages 19-35. The percents of people experiencing one or more 11 12 FEV₁ decrements \geq 15% during the 2006 O₃ season in Atlanta are 3.6 times higher for outdoor 13 workers than for the general population (ages 19-35) under the current standard, and range up to 14 5.3 times higher for the alternative standards. The percents of people experiencing six or more FEV₁ decrements \geq 15% during the 2006 O₃ season in Atlanta are 20 times higher for outdoor 15 16 workers than for the general population under the current standard, and range up to 150 times 17 higher for the alternative standards. As expected, we see that the risk of repeated occurrences of 18 FEV_1 decrements $\geq 15\%$ is much greater for outdoor workers than for the general population.

- 1 Part of the reason for this is that APEX tends to underestimate the number of individuals who
- 2 have very repetitive activity patterns (e.g., 9 to 5 weekdays office workers) when using the
- 3 CHAD activity database and the method selected for generating longitudinal diary profiles (see
- 4 Section 5.3.1).
- 5
- 6
- 7 8

Table 6-6. Percents of the General Population and Outdoor
Workers (ages 19-35) Experiencing 1 or More and 6 or More FEV ₁
Decrements $\geq 15\%$ (based on Atlanta 2006 APEX simulations)

	General population ages 19-35	Outdoor workers ages 19-35
1 or more		
Current standard	1.2%	4.3%
70 ppb alt. std.	0.84%	3.2%
65 ppb alt. std.	0.55%	2.5%
60 ppb alt. std.	0.32%	1.7%
6 or more		
Current standard	0.06%	1.2%
70 ppb alt. std.	0.018%	0.93%
65 ppb alt. std.	0.005%	0.74%
60 ppb alt. std.	0.005%	0.55%

6.3.2 Lung Function Risk Estimates Based on the Exposure-Response Functions Approach Used in Prior Reviews

In this section we present lung function risk estimates for all school-aged children following the methodology used in previous reviews, based on updated exposure-response (E-R) functions. In Appendix 6-C we compare these estimates with those from the previous review.
Table 6-7 provides an overall summary of results for each air quality scenario by tabulating the minimum and maximum estimates over all cities and years of percents of all

- 18 school-aged children (ages 5 to 18) experiencing one or more days (during the O_3 season) with
- 19 FEV_1 decrement more than 10 and 15%. This table can be compared with Table 6-4 and Table 6-
- 20 5, which have analogous results for the MSS model. These results are much lower than the MSS
- 21 model results. The reasons for this are described in Section 6.3.3 below.

- 1
- 2 Table 6-7. Ranges of percents of school-aged children experiencing one or more 3 days during the O₃ season with lung function decrement (Δ FEV₁) more than 10 and 15%.
- days during the O₃ season with lung function decrement (ΔFEV₁) more than 10 and 15%
 The numbers in this table are the minimum and maximum percents estimated over all
- 5 cities and years.

Scenario	$\begin{array}{l} \mbox{minimum}\\ \mbox{percent}\\ \mbox{experiencing}\\ \geq 1 \mbox{ day with}\\ \mbox{\Delta FEV}_1 \geq 10\% \end{array}$	$\begin{array}{l} \text{maximum} \\ \text{percent} \\ \text{experiencing} \\ \geq 1 \text{ day with} \\ \Delta \text{ FEV}_1 \geq 10\% \end{array}$	$\begin{array}{l} \mbox{minimum}\\ \mbox{percent}\\ \mbox{experiencing}\\ \geq 1 \mbox{ day with}\\ \mbox{\Delta FEV}_1 \geq 15\% \end{array}$	$\begin{array}{l} maximum \\ percent \\ experiencing \\ \geq 1 \ day \ with \\ \Delta \ FEV_1 \geq 15\% \end{array}$
base	2%	11%	0%	5%
75	2%	6%	1%	2%
70	2%	6%	0%	2%
65	1%	5%	0%	1%
60	2%	3%	0%	1%

8 6.3.3 Comparison of the MSS Model with the Exposure-Response Function Approach

9 There are two key differences between the MSS and E-R models. The E-R model 10 estimates the distribution of FEV_1 decrements across the population or study group, whereas the 11 MSS model estimates FEV_1 decrements at the individual level and then these are aggregated to 12 obtain the population distribution. Thus the MSS model allows for detailed analyses of 13 conditions that influence risk. Second, the E-R model estimates FEV_1 decrements only for 8-14 hour average exposures when the 8-hour average exertion level is moderate or greater. The MSS 15 model estimates FEV₁ decrements for any averaging time and therefore accounts for a wider 16 range of activities that might result in FEV₁ decrements. 17 A comparison of the MSS model with the exposure-response function approach for the

17 A comparison of the MSS model with the exposure-response function approach for the 18 2006 existing standard scenarios is summarized in Table 6-8, which lists estimates of the

19 percents of school-aged children estimated to experience lung function responses greater then 10,

20 15, and 20%. The MSS model estimates are significantly higher than the exposure-response

21 function approach estimates. In most cases, the MSS model gives results about a factor of three

higher than the exposure-response function model for school-aged children. This is expected,

23 since, as discussed above, the MSS model includes responses for a wider range of exposure

24 protocols (under different levels of exertion, lengths of exposures, and patterns of exposure

25 concentrations) than the exposure-response model of previous reviews.

T	
2	
2	

Table 6-8. Comparison of responses from the MSS model with responses from the population exposure-response (E-R) method. 2006 existing standard, ages 5 to 18

Urban area	\geq 10% FEV ₁ decrement		\geq 15% FEV ₁ decrement		\geq 20% FEV ₁ decrement	
	MSS model	E-R method	MSS model	E-R method	MSS model	E-R method
Atlanta	19.2%	5.6%	5.3%	1.7%	2.1%	0.7%
Baltimore	18.6%	5.4%	5.2%	1.6%	2.1%	0.7%
Boston	13.6%	4.5%	3.7%	1.2%	1.4%	0.5%
Chicago	14.4%	4.7%	3.9%	1.3%	1.6%	0.5%
Cleveland	13.5%	4.2%	3.3%	1.1%	1.2%	0.4%
Dallas	21.6%	6.0%	6.4%	1.8%	2.7%	0.8%
Denver	20.2%	5.8%	5.6%	1.7%	2.2%	0.7%
Detroit	13.6%	4.4%	3.5%	1.2%	1.3%	0.4%
Houston	16.2%	4.7%	4.2%	1.3%	1.6%	0.5%
Los Angeles	18.2%	4.8%	4.2%	1.2%	1.5%	0.5%
New York	12.7%	4.2%	3.2%	1.1%	1.2%	0.4%
Philadelphia	16.4%	4.8%	4.2%	1.3%	1.6%	0.5%
Sacramento	17.9%	5.1%	4.8%	1.4%	2.8%	0.6%
St. Louis	18.6%	5.4%	5.1%	1.6%	2.0%	0.6%
Washington	15.9%	4.6%	4.0%	1.3%	1.5%	0.5%

12

4 Since the E-R method of the previous reviews only looks at 8-hour exposures 5 concomitant with EVR ≥ 13 L/min-m² BSA (hereafter, EVR ≥ 13), it is of interest to compare 6 the E-R method results with the corresponding MSS model results (instances of $\Delta FEV_1 \ge 10, 15$,

7 20% concomitant with EVR \geq 13).

8 We performed this comparison for four APEX simulations: the Atlanta March 1-October

9 30, 2006 base case, ages 18-35; the Los Angeles May 29-July 28, 2006 base case, age 25; the

10 Los Angeles May 29-July 28, 2006 base case, ages 18-35; and the Los Angeles Jan 1-Dec 31,

11 2006 base case, ages 18-35.

For the Atlanta simulation, the E-R function approach gives 5.0, 1.8, and 0.9%

13 responding for $\Delta FEV_1 \ge 10$, 15, 20%. The MSS model approach gives 11.54, 3.26, and 1.28%

14 responding for $\Delta FEV_1 \ge 10$, 15, 20%. The percents of the population for $\Delta FEV_1 \ge 10$, 15, 20% at

15 the end of the daily max 8-hour average exposure period where the concomitant 8-hour average

16 EVR is \geq 13 are 6.67%, 2.09%, and 0.84%. 15.17% of the population never have any instances

17 of EVR \geq 13 and 0.41% have at least one occurrence of $\Delta FEV_1 \geq$ 10% while never having EVR

18 \geq 13 for any 8-hour period. 4.46% (not among the 15.17%) have instances of $\Delta FEV_1 \geq$ 10% but

- 1 none of those instances with concomitant EVR \geq 13. The 11.54% responding is made up of
- 2 6.67% of the population with instances of $\Delta FEV_1 \ge 10\%$ concomitant with EVR ≥ 13 and 4.87%
- 3 with instances of $\Delta FEV_1 \ge 10\%$ not concomitant with EVR ≥ 13 .
- 4 Table 6-9 and Table 6-10 summarize the pertinent results for the Atlanta and three Los
- 5 Angeles simulations. Looking at the first rows of these tables shows that these models have
- 6 similar corresponding results. The broader scope of activity/exposure patterns encompassed by
- 7 the MSS model, beyond the 8-hour average EVR \geq 13 restriction of the E-R model, contributes
- 8 from a third to a half to the total MSS model risk and to a large part explains the differences
- 9 between the models for ages 18-35. The difference between the MSS and E-R models is larger
- 10 for school-aged children than for adults ages 18-35 due to the increased EVR and time spent
- 11 outdoors in children compared to adults.
| L | | | | / | / / U | |
|---|-----------------------------------|-----------------------------------|-----------------------------------|-----------------------------------|-----------------------------------|-----------------------------------|
| Component of results | MSS model $\Delta FEV_1 \ge 10\%$ | E-R model $\Delta FEV_1 \ge 10\%$ | MSS model $\Delta FEV_1 \ge 15\%$ | E-R model $\Delta FEV_1 \ge 15\%$ | MSS model $\Delta FEV_1 \ge 20\%$ | E-R model $\Delta FEV_1 \ge 20\%$ |
| profiles with instances of $\Delta FEV1 \ge cutoff$
concomitant with 8-hour EVR ≥ 13 | 6.7% | 5.0% | 2.1% | 1.8% | 0.8% | 0.9% |
| profiles with instances of $\Delta FEV_1 \ge$ cutoff never concomitant with 8-hour EVR ≥ 13 | 4.8% | | 1.2% | | 0.5% | |
| Final result of each model | 11.5% | 5.0% | 3.3% | 1.8% | 1.3% | 0.9% |

Table 6-9. Comparison of MSS Model and E-R Model of Previous Reviews for Atlanta, Mar 1-Oct 30, 2006, ages 18-35

Table 6-10. Comparison of MSS Model and E-R Model of Previous Reviews for Los Angeles, Jan 1-Dec 31, 2006, ages 18-35

Component of results	$\begin{array}{l} MSS \ model \\ \Delta FEV_1 \geq 10\% \end{array}$	E-R model $\Delta FEV_1 \ge 10\%$	$\begin{array}{c} MSS \ model \\ \Delta FEV_1 \geq 15\% \end{array}$	E-R model $\Delta FEV_1 \ge 15\%$	$\begin{array}{c} MSS \ model \\ \Delta FEV_1 \geq 20\% \end{array}$	E-R model $\Delta FEV_1 \ge 20\%$
profiles with instances of $\Delta FEV1 \ge \text{cutoff}$ concomitant with 8-hour EVR ≥ 13	7.9%	6.2%	2.6%	2.6%	1.2%	1.4%
profiles with instances of $\Delta FEV_1 \ge$ cutoff never concomitant with 8-hour EVR ≥ 13	6.5%		1.8%		0.8%	
Final result of each model	14.4%	6.2%	4.4%	2.6%	2.0%	1.4%

1 Figure 6-10 compares the E-R function to the response curve of the MSS model restricted

2 to 8-hour average EVR \geq 13 and shows that these curves are very close. The MSS model has a

- 3 higher response for the low and high ranges of exposure concentrations, while the E-R model is
- 4 higher in the mid-range of exposures.





7 Another element of the difference between the models derives from the distribution of 8 EVR in the clinical studies the E-R approach is based on and how this compares to the 9 distribution of EVR in the APEX simulations. Most of the clinical studies are conducted with a target EVR of 20 L/min-m² BSA and the actual EVRs vary somewhat around this value. The 10 rationale for the cutpoint of 13 L/min-m² BSA is described in EPA's responses to comments on 11 12 the 1996 proposed rule on the NAAQS for O₃ (Federal Register, 1996) as "for the 8-hr health 13 risk assessment the range (based on being within 2 standard deviations of the mean) of EVRs 14 observed in the subjects who participated in the study [McDonnell et al., 1991] was 13-27 liters 15 per minute per meter squared [BSA] (L/min-m²)." Figure 6-11 shows the distribution of EVRs \geq 1 13 for the Atlanta simulation and is clearly shifted much lower than the distribution of EVR in

2 the clinical studies. This could lead to an overestimation of the percent of responders by the E-R

- 3 method, since higher EVRs lead to higher lung function decrements and it is applying an E-R
- 4 function based on EVRs around 20 to a population with median EVRs around 14.5.
- 5



1 6.4 EVALUATION OF THE MSS MODEL

2 6.4.1 Summary of Published Evaluations

McDonnell et al. (2010) performed a detailed evaluation of their model using two
methods: (1) cross-validation and (2) comparison of an independent data set against the
predictions of the model.

6 The cross-validation was based on the data set of 15 EPA studies from which their 7 original model was developed (McDonnell et al., 2007). This data set has 541 subjects, each with 8 multiple measurements during single experiments. Subjects were omitted from the data set, one 9 at a time, the model refit to the reduced data set, and the resulting parameters used to predict the 10 FEV₁ decrements for the omitted subject. The authors then compare the mean predictions and 11 mean observed values for each subject and presented these results in a scatter plot (Figure 1b, 12 McDonnell et al., 2010). The observations exhibit much more variability than the predictions; for 13 observed values of 20%, predicted values range from around 2 to 19%; and all observed values 14 above 20% are underpredicted (the observed values range from -20 to 60%, while the predicted 15 values range from 0 to 20%). These features result from the omission of the inter- and intra-16 individual variability terms (U_i and ε_{iik}) in the MSS model (equation 6-3), which are accounted

17 for in the risk estimates in this chapter.

Model predictions were compared against an independent data set of seven clinical studies with a total of 204 subjects (McDonnell et al., 2010). Graphs of predicted and observed study means vs. time show fair to good model fit. The authors do not present overall fit statistics that are directly commensurate with the statistics of interest in this risk assessment: the proportions of people with FEV₁ decrements greater than 10, 15, and 20%.

McDonnell et al. (2012) do compare observed and predicted proportions of people with FEV₁ decrements greater than 10, 15, and 20% and provide the corresponding scatter plots (Figure 4). They find the model to be unbiased, with the slopes of the observed vs. predicted lines for 10, 15, and 20% to be around 1.0 and the R^2 respectively 0.78, 0.73, and 0.67. The higher observed proportions of people with FEV₁ decrements greater than 10, 15, and 20% tended to be substantially underpredicted.

29 **6.4.2** Children

A clinical study with children (ages 8-11; mean, 10 years; n=22), exposed to 120 ppb O_3 over 2.5 hours at heavy exertion levels was done by McDonnell et al. (1985). This study could be used to fit the model for children if all of the measurements of FEV₁ and ventilation rates were available. The paper lists the end-of exposure FEV₁ responses for each individual (but not ventilation rates), which we use to compare with the MSS model with the age term extension described in Section 6.2.4. The numbers of subjects with clean-air adjusted responses greater

- 1 than 10%, 15%, and 20% are respectively 4, 2, and 1, corresponding to 18.2%, 9.1%, and 4.5%
- 2 of the number of subjects. We ran the MSS 2010 model using the mean and standard deviation of
- 3 the ventilation rates reported in the paper. Resting ventilation rates were assumed to be 10.4
- 4 L/min (Avol et al., 1985) and BSA to be 1.08 m^2 (EPA, 2011). Details of this comparison can be
- 5 found in Appendix 6-D.
- 6 Table 6-11 compares the results of this simulation with the results of McDonnell et al.
- 7 (1985). The agreement is fairly good. Due to the limited sample size of 22 subjects from only
- 8 one study and the assumptions made in running the MSS model, this does not provide
- 9 confirmation that the age term extension is correct; on the other hand, this comparison does not
- 10 indicate that there is a problem with the age term extension. Information is not available that
- 11 would allow us to provide respectable confidence intervals for these estimates.
- 12

13Table 6-11. Comparison of Responses from the MSS 2010 Model with Responses14from McDonnell et al. (1985)

	≥ 10% FEV1 decrement		≥15% FEV	/1 decrement	≥ 20% FEV1 decrement	
	MSS model	McDonnell et al. (1985)	MSS model	McDonnell et al. (1985)	MSS model	McDonnell et al. (1985)
Percent responding	18.4%	18.2% (4 subjects)	6.8%	9.1% (2 subjects)	2.3%	4.5% (1 subject)

15

16 6.4.3 Threshold vs. Non-Threshold Models

17 The difference between the results of the MSS threshold and non-threshold models is 18 minor, with the threshold version estimates of lung function decrements almost identical to the 19 no-threshold version for the Atlanta 2006 recent air quality base case, as can be seen by 20 comparing Table 6-12 with Table 6-13. This is consistent with the logistic form of the model, 21 where the impact of exposures to low concentrations on risk is small.

22

1Table 6-12. Percents of the population by age group with one or more days during2the O3 season with lung function (FEV1) decrements more than 10, 15, and 20% (Atlanta32006 base case). MSS Threshold model, monitors air quality.4

4

Age Group	ΔFEV ₁ ≥ 10%	ΔFEV ₁ ≥ 15%	ΔFEV ₁ ≥ 20%
5 to 18	31%	13%	6.4%
19 to 35	11%	3.1%	1.3%
36 to 55	3.7%	0.60%	0.14%

5

Table 6-13. Percents of the population by age group with one or more days during
the O₃ season with lung function (FEV₁) decrements more than 10, 15, and 20% (Atlanta
2006 base case). MSS No-Threshold model, monitors air quality.

9

Age Group	$\frac{\Delta FEV_1 \geq}{10\%}$	ΔFEV ₁ ≥ 15%	$\frac{\Delta FEV_1 \geq}{20\%}$
5 to 18	31%	13%	6.6%
19 to 35	11%	3.1%	1.3%
36 to 55	3.8%	0.60%	0.15%

10

11

12 6.5 CHARACTERIZATION OF UNCERTAINTY

In the controlled human exposure study based risk assessment, there are two broad sources of uncertainty to the risk estimates. One of the most important sources of uncertainty is the estimation by APEX of the population distribution of individual time series of O₃ exposures and ventilation rates. The uncertainty regarding these estimated exposures is discussed in

17 Chapter 5; they are not discussed further here.

⁴ In the first draft REA, monitor-level air quality was provided as input to the APEX model. As discussed in Chapter 5, tract-level air quality was used in APEX for this second draft REA. Monitor-level air quality is used for the APEX simulations here, since these simulations take less time to run. This does not affect the analyses here, since the two air quality formats yield very similar results (see Appendix 6-F).

In this section, uncertainties associated with the second broad source of uncertainty in the
 risk calculation are discussed, namely, uncertainties in the lung function risk model. The specific
 sources of uncertainty covered are:

- Statistical model form
- 5 6

7

8

- Convergence of APEX resultsApplication of model for all lifestages
- Application of model for asthmatic children
- Interaction between O₃ and other pollutants
- 9 6.5.1 Statistical Model Form

10 The MSS model is a 2-compartment model, the form of which is based on physical 11 considerations. It accomodates these key features of human exposure studies: (1) FEV_1 12 responses increase with increasing O_3 concentration, ventilation rate, and duration of exposure, 13 (2) the effect of each of these three variables depends on the levels of the other two variables, (3) 14 FEV_1 responses depend on age, (4) certain individuals are consistently more responsive to O_3 15 exposure, and (5) O_3 –induced FEV₁ decrements improve within a few hours of cessation of 16 exposure (McDonnell et al, 2007). These considerations support the form of the model, as do model evaluation that have been performed (Section 6.4.1). Although the model does not have 17 good predictive ability for individuals (psuedo- $R^2 0.28$), it does better at predicting the 18 proportion of individuals with FEV₁ decrements \geq 10, 15, and 20% (psuedo-R²s of 0.78, 0.74, 19 20 0.68) (McDonnell et al, 2012). 21 The clinical studies that these models' estimates are based on were conducted with young 22 adult volunteers rather than randomly selected individuals, so it may be that selection bias has 23 influenced the model parameter estimates.

24 The parameter estimates are not very precise, as the result of the likelihood surface being 25 somewhat flat in the neighborhood of the maximum likelihood estimates. Table 6-14 gives 95 26 percent confidence intervals for each of the parameter estimates as percents of the estimates, 27 based on the standard errors reported by McDonnell et al. (2012). Figure 6-10 shows how much 28 the modeled number of children with one or more FEV₁ decrements $\geq 10\%$ changes when each 29 parameter is increased by five percent (keeping the other parameters fixed at their estimates). 30 The scenario modeled is the Los Angeles 2006 recent conditions base case. The physiological 31 parameter MET, a measure of the level of exertion for a given activity (see Appendix 6-E), is 32 also included here for comparison. MET is a key variable in calculating ventilation rates and is 33 specified by a distribution for each activity. Here we have shifted all MET distributions by +5% 34 of their means.

	β1	β2	β3	β4	β5	β6	β9	var(U)	var(E)
parameter estimate	10.916	-0.2104	0.01506	13.497	0.003221	0.8839	59.284	0.9373	17.0816
standard error	0.8446	0.31	0.00333	4.734	0.000207	0.0647	10.192	0.0824	1.1506
95% conf. interval	±15%	±289%	±43%	±69%	±13%	±14%	±34%	±17%	±13%

Table 6-14. MSS threshold model estimated parameters with confidence intervals

2

from McDonnell et al. (2012).

3 The most influential parameter in Figure 6-12 is β_6 , the power to which ventilation rate is raised in the MSS model. An increase of five percent in β_6 leads to 27, 40, and 47 percent 4 5 increases respectively in the modeled number of children with FEV₁ decrements $\geq 10, 15$ and 6 20%. The next most influential parameter is the variance of E, the intra-individual variability 7 term. The least influential parameter is β_2 , the slope of the age term. These changes of five 8 percent are much less than the 95 percent confidence intervals of the parameter estimates, so the 9 uncertainty in the risk estimates resulting from parameter uncertainty is likely to be more than is 10 indicated in Figure 6-12.

11 Age Term Significance

12 As discussed in Section 6.5.3 below, there are uncertainties in extrapolating the MSS model down to age 5 from the age range of 18 to 35 to which the model was fit. Further 13 14 considerations indicating that the uncertainty of the extension to children of the MSS model could be substantial are that the age coefficient $\beta_2 = -0.21$ (s.e. 0.31) in the MSS model is not 15 16 statistically significantly different from zero; and when the MSS model is fit to the U.C. Davis 17 clinical data the age term is <u>positive</u>, $\beta_2 = +0.19$ (0.60), although also not statistically 18 significantly different from zero (McDonnell et al., 2012). Note that, in the previous section, β_2 19 was found to be the least influential model parameter.



Figure 6-12. Sensitivity (Percent Change) of Population With One or More FEV_1 Decrements $\geq 10\%$ to a 5% Increase in Individual MSS Model Parameter Estimates.

1 The Variability Term ε

2 The variability term ε in equation 6-3 is assumed by the MSS model to have a Gaussian 3 distribution with mean zero and estimated standard deviation 4.135 (in the threshold model). 4 Since the actual values are bounded, we truncate the variability term distribution at ± 2 standard deviations (± 8.27), a convention we use for the distributions of several physiological variables 5 6 input to APEX in the physiology input file. To look at the effect of truncating the variability 7 distribution, we conducted simulations with the variability term truncated at ± 20 , the range of the 8 actual values of the variability term. We find that this constraint has a very large effect on 9 estimates of percents of the population with FEV_1 decrements ≥ 10 and 15% and less of an effect 10 for 20%. The percent of children with FEV₁ decrements \geq 10% increases from 31% to 92% 11 when increasing the truncation point from 8.27 to 20. Details of this comparison and additional results are presented in Appendix 6-F. The assumption that the distribution of the variability term 12 13 ε is Gaussian is convenient for fitting the model, but is not accurate. The extent to which this 14 mis-specification affects the estimates of the parameters of the MSS model is not clear.

15

6.5.2 Convergence of APEX Results

16 APEX accounts for several sources of variability by drawing random variables from 17 specified distributions. Some variables are drawn once for each simulated individual (e.g., age, 18 location of residence), some are drawn every day or every hour for each simulated individual, 19 and others are drawn more frequently, at the event level (e.g., activity). Increasing the number of 20 individuals simulated in an APEX run increases the accuracy of the modeled variability and the 21 results of the APEX runs are more reproducible. In order to assess the number of individuals to 22 simulate to achieve convergence of APEX results, we perform multiple APEX runs with 23 identical inputs except for the random number seed, and look at the variability of the results of 24 these model runs. Table 6-15 summarizes the results of 40 APEX simulations of the Atlanta 25 2006 base case with 200,000 simulated individuals. For each of these measures, the range of 26 results over the 40 APEX runs is less than one percent. This analysis of the convergence of 27 APEX results shows that modeling 200,000 simulated individuals is adequate for reasonable 28 convergence of the FEV_1 risk measures.

1Table 6-15. Convergence results for the Atlanta 2006 base case with 200,0002simulated individuals. Percents of the population by age group with one or more days (and3six or more days) during the O3 season with lung function (FEV1) decrements more than

4 10, 15, and 20%. Minimum and maximum values and ranges over 40 APEX runs.

	ΔF	$EV_1 \ge 10$)%	ΔF	$\mathrm{EV}_1 \ge 1$	5%	ΔF	$\mathrm{TEV}_1 \geq 2$	0%
Age group	min	max	range	min	max	range	min	max	range
1 or more da	ys in the	season							
5 to 18	31.3%	32.1%	0.88%	12.4%	12.9%	0.49%	6.21%	6.71%	0.50%
19 to 35	11.1%	11.5%	0.39%	3.00%	3.26%	0.26%	1.11%	1.32%	0.22%
36 to 55	3.54%	3.79%	0.25%	0.55%	0.68%	0.13%	0.13%	0.20%	0.07%
6 or more da	ys in the	season							
5 to 18	9.28%	9.73%	0.45%	2.80%	3.18%	0.38%	1.11%	1.37%	0.27%
19 to 35	1.09%	1.25%	0.16%	0.15%	0.21%	0.06%	0.03%	0.06%	0.03%
36 to 55	0.22%	0.30%	0.08%	0.01%	0.03%	0.02%	0.00%	0.01%	0.01%

5

6 6.5.3 Application of Model for All Lifestages

7 The exposure-response functions derived from controlled human exposure studies 8 involving 18-35 year old subjects were used to estimate responses for school-aged children (ages 9 5-18). This was in part justified by the findings of McDonnell et al. (1985) who reported that 10 children 8-11 years old experienced FEV₁ responses similar to those observed in adults 18-35 11 years old when both groups were exposed to 120 ppb O_3 at an EVR of 32-35 L/min/m². In 12 addition, a number of summer camp studies of school-aged children exposed in outdoor environments in the Northeast also showed O₃-induced lung function changes similar in 13 14 magnitude to those observed in controlled human exposure studies using adults, although the 15 studies may not directly comparable. The MSS model predicts increasing responsiveness with 16 younger participants in the age range of 18-35 years, as shown in Figure 6E-4 (Appendix 6-E), 17 which might indicate that responsiveness would continue to increase as age decreases from 18. 18 In extending the MSS model to children, we fixed the age term in the model at its highest value, 19 the value for age 18. If continuing the MSS model trend were to accurately describe continued 20 increased response in children, then the fixed age term for children may have underestimated the 21 effects on children, and particularly younger children. On the other hand, if FEV₁ responses for 22 children are similar to those observed in adults 18-35 years old, as the evidence suggests, then 23 our approach to extending the age term would overestimate the response to children (see Table 24 6E-3 in Appendix 6-E).

In considering extending the MSS model to ages older than 36, we note that, in general, O₃ responsiveness steadily declines for persons aged 35-55, with persons >55 eliciting minimal responsiveness (ISA, section 6.2.1.1). As described in Section 6.2.4, we extended the age term from the value at 36 linearly to zero at age 55, and set it to zero for ages above 55 (see **Error! Reference source not found.**). The uncertainty of this extrapolation may be substantial, but these age groups are not the primary focus in the clinical risk assessment.

7 6.5.4 Application of Model for Asthmatic Children

8 The risk assessment used the same exposure-response relationship, developed from data 9 collected from healthy study subjects, and applied it to all persons, children, and asthmatic 10 children. Based on limited evidence from a few human exposure studies, it is likely that subjects 11 having asthma are at least as sensitive to acute effects of O_3 as other subjects not having this 12 health condition (ISA, page 6-20 to 6-21). An analysis by Romieu et al. (2002) indicated a larger 13 O₃-associated decrement in FEV₁ among children with moderate to severe asthma than 14 among all children with asthma (ISA, page 6-54). This suggests that the lung function 15 decrements presented in this assessment for asthmatic children may be underestimated. The 16 magnitude of influence this element might have on our risk estimates remains unknown at this 17 time. In addition, asthmatic children may have less reserve lung capacity to draw upon when 18 faced with decrements, and therefore a $\geq 10\%$ decrement in lung function may be a more adverse 19 event in an asthmatic child than a healthy child.

20 6.5.5 Interaction Between O₃ and Other Pollutants

Because the controlled human exposure studies used in the risk assessment involved only
 O₃ exposures, it was assumed that estimates of O₃-induced health responses would not be
 affected by the presence of other pollutants (e.g., SO₂, PM_{2.5}, etc). The magnitude of influence
 that potential interactions might have on our risk estimates remains unknown at this time.

25 6.5.6 Qualitative Assessment of Uncertainty

EPA staff have identified key sources of uncertainty with respect to the lung function risk estimates. These are: the physiological model in APEX for ventilation rates, the O₃ exposures estimated by APEX, the MSS model applied to ages 18 to 35, and extrapolation of the MSS model to children ages 5 to 18. The first two of these are discussed in Chapter 5. At this time we do not have quantitative estimates of uncertainty for any of these. Table 6-16 provides a qualitative assessment of the uncertainty resulting from each of these key sources. The primary source of uncertainty is the MSS model, applied to ages 18 to 35.

Table 6-16. Summary of Qualitative Uncertainties of Key Modeling Elements in the O₃ Lung Function Risk Assessment

		Potential influence of uncertainty on risk		Knowledge-	
Source	Description	Direction	Magnitude	Base uncertainty*	Comments
The physiological model in APEX for ventilation rates	The physiological model in APEX takes into account the population distribution of individual physiological characteristics and activities and models minute-by-minute ventilation rates for each simulated individual using a series of physiological relationships known with varying degrees of certainty.	Over	Low- Medium	Low- Medium	Ventilation rates are a key input to the MSS model.Figure 6E-3 in Appendix 6-E gives an overview of the physiological model in APEX for ventilation rates.Comparisons with ventilation rates reported in the literature show fairly good agreement with APEX ventilation rates (Section 5.4.4).
O ₃ exposures	The O_3 exposures estimated by APEX and their uncertainties are discussed in Chapter 5.	Both	Low- Medium	Low	O_3 exposures are a key input to the MSS model.
The McDonnell- Stewart-Smith (MSS) FEV ₁ model for ages 18 to 35	The MSS model is integrated into APEX and predicts FEV ₁ decrements for each simulated individual.	Both	Medium- High	Low	There is a good conceptual foundation for the structure of this model, but the variability in measurements of FEV ₁ and estimated parameters of the model introduce uncertainty into the model predictions of large FEV ₁ decrements. The estimated parameters have fairly wide confidence intervals (Table 6-1) and the risk results are sensitive to varying the parameters (Figure 6-12). The most influential parameter is $\beta 6$, the power to which ventilation rate is raised in the MSS model. An increase of five percent in $\beta 6$ leads to a 27 percent increase in the modeled number of children with FEV ₁ decrements $\geq 10\%$. (The 95 percent confidence interval of this parameter estimate is $\pm 14\%$.) The variability term ϵ [in equation 6-3] is assumed by the MSS model to have a Gaussian distribution with mean zero and estimated standard deviation 4.135. Since the actual values are bounded, we truncate the variability term distribution at ± 2 standard deviations (± 8.27), a convention we use for the distributions of several physiological

		Potential i uncertair	influence of ity on risk	Knowledge-	
		estir	estimates		
Source	Description	Direction	Magnitude	uncertainty*	Comments
					variables input to APEX in the physiology input file. To look at the effect of truncating the variability distribution, we conducted simulations with the variability term truncated at ± 20 , the range of the actual values of the variability term. We find that this constraint has a large effect on estimates of percents of the population with FEV ₁ decrements ≥ 10 and 15% and less of an effect for 20%. The percent of children with FEV ₁ decrements $\geq 10\%$ increases from 31% to 92% when increasing the truncation point from 8.27 to 20.
Extrapolation of the MSS model to children	The MSS model is based on studies with subjects ranging in age from 18 to 35 years; therefore prediction for individuals outside this age range involves assumptions for extrapolation of the MSS model for individuals <18 and >35 years of age	Both	Medium	Low	Summer camp studies and one clinical study of children indicate that FEV_1 responses for children are similar to those observed in adults 18-35 years old. See discussion in Section 6.5.3.

* Refers to the degree of uncertainty associated with our understanding of the phenomenon, in the context of assessing and characterizing its uncertainty

1 6.6 DISCUSSION

2 The second draft lung function risk assessment evaluated risks of lung function 3 decrements due to O_3 exposure for all three groups: school-age children ages 5 to 18, young 4 adults ages 19 to 35, and adults ages 36 to 55. Adults older than 55 have minimal O_3 -induced 5 lung function risk. Two models were used, one based on application of an individual level 6 exposure-response function, the MSS model introduced in this review, and one based on 7 application of a population level E-R function consistent with the model used in the previous O_3 8 review which applies probabilistic population-level exposure-response relationships for lung 9 function decrements (measured as percent reductions in FEV₁) associated with 8-hour moderate 10 exertion exposures. The MSS model is preferred, due to its ability to model individual exposures 11 for a wide range of exposure times and levels of exercise (Section 6.2.4; ISA pages 6-15 to 6-12 16). Both models provide estimates of the percent of the groups experiencing a reduction in lung 13 function for three different levels of impact, 10, 15, and 20% decrements in FEV₁. These levels 14 of impact were selected based on the literature discussing the adversity associated with these 15 types of lung function decrements (US EPA, 2012, Section 6.2.1.1; Henderson, 2006). For the 16 second draft assessment, lung function risks were estimated for 15 cities: Atlanta, Baltimore, 17 Boston, Chicago, Cleveland, Dallas, Denver, Detroit, Houston, Los Angeles, New York, 18 Philadelphia, Sacramento, St. Louis, and Washington, DC. 19 Based on the MSS model, the percents of population estimated to experience lung

20 function responses greater then 10, 15, and 20%, associated with O₃ exposure while engaged in 21 various levels of exertion, vary considerably for different years and cities under the recent air 22 quality scenarios and also for the existing and alternative standard scenarios (Figure 6-7 and 23 Figure 6-8, Table 6-4 and Table 6-5). The estimates for $\geq 10\%$ FEV₁ decrement for school-age children for recent O₃ concentrations range across cities and years from 11 to 31 percent, and 24 25 range from 11 to 22 percent after simulating just meeting the existing standard. The estimates for 26 \geq 15% FEV₁ decrement for school-age children for recent O₃ concentrations range across cities 27 and years from 2 to 12 percent, and range from 2 to 6 percent after simulating just meeting the 28 existing standards. The estimates for $\geq 20\%$ FEV₁ decrement for recent O₃ concentrations range 29 across cities and years from 1 to 6 percent, and range from 1 to 3 percent after simulating just 30 meeting the existing standards.

Figure 6-13 displays the risks and the incremental increases in risk for increasing standard levels, where risk is taken to be the highest value for each study area (over years) of the percent of school-aged children with FEV₁ decrement \geq 10%. The risks in this figure for Washington, DC, for example, are about 9.6% for the alternative standard level of 60 ppb and 13.4% for the alternative standard level of 65 ppb. The length of the orange bar is the 1 incremental risk (3.8%) in going from the 60 ppb to the 65 ppb alternative standards. This figure

- 2 shows that there are significant increases in incremental risk for all 15 cities in the progression of
- 3 alternative standard levels from 60 ppb to the level of the existing standard, 75 ppb. The pattern
- 4 of reductions for lung function decrements larger than 15 and 20% are similar. As discussed in
- 5 Section 4.3.1, the New York 60 ppb alternative standard was not modeled and the risk for NY for
- 6 that scenario would not necessarily be zero. Figure 6-14 displays the risks and the incremental
- 7 increases in risk for increasing standard levels, where risk is taken to be the mean value for each
- 8 study area (over years) of the percent of school-aged children with FEV₁ decrement $\geq 10\%$.

Similar to the MSS model results, the percents of school-age children estimated to
experience lung function responses greater then 10, 15, and 20% based on the population level
E-R function exhibit variation across years and cities. However, the MSS model estimates are
significantly higher than the E-R approach estimates. For lung function responses greater than
10, 15, and 20% the MSS model gives results typically a factor of three higher than the E-R

14 model for school-aged children. Both models give higher responses for higher concentrations,

15 compared to lower concentrations, as can be seen in Figures 6-6, 6-9, and 6-10.

16 The MSS model was applied to estimate lung function risk for outdoor workers (ages 19-

17 35) in Atlanta for one year (2006). The proportion of outdoor workers with FEV₁ decrements \geq

18 15% ranges from 3.6 to 5.3 times the proportion of the general population (ages 19-35) with

19 FEV_1 decrements $\geq 15\%$ across the different standards simulated. The proportion of outdoor

20 workers with multiple occurrences of FEV₁ decrements $\geq 15\%$ is much greater than for the

- 21 general population.
- 22



Figure 6-13. Lung Function Risk Results, Incremental Increases In Risk For Increasing Standard Levels: Percent of All School-aged Children With FEV₁ Decrement $\geq 10\%$, Highest Value For Each Study area Over Years⁵

⁵ New York level 60 was not modeled . We do not know what the percent risk would be for NY under the 60 ppb alternative standard, but it would not necessarily be zero.



Figure 6-14. Lung Function Risk Results, Incremental Increases In Risk For Increasing Standard Levels: Percent of All School-aged Children With FEV₁ Decrement \geq 10%, Mean Value For Each Study Area Over Years⁶

⁶ New York level 60 was not modeled . We do not know what the percent risk would be for NY under the 60 ppb alternative standard, but it would not necessarily be zero.

1 6.7 REFERENCES

3	Adams, W.C. 2002. "Comparison of Chamber and Face-mask 6.6-hour Exposures to Ozone on
4	Pulmonary Function and Symptoms Responses." Inhalation Toxicology, 14:745-764.
5	Adams, W.C. 2003. "Comparison of Chamber and Face Mask 6.6-hour Exposure to 0.08 ppm
6	Ozone via Square-wave and Triangular Profiles on Pulmonary Responses." Inhalation
7	<i>Toxicology</i> , 15: 265-281.
8	Adams, W.C. 2006, "Comparison of Chamber 6.6-h Exposures to 0.04-0.08 ppm Ozone via

- Adams, w.C. 2000. Comparison of Chamber 6.0-n Exposures to 0.04-0.08 ppm Ozone via
 Square-wave and Triangular Profiles on Pulmonary Responses." *Inhalation Toxicology*, 18:
 127-136.
- American Thoracic Society. 2000. "What Constitutes an Adverse Effect of Air Pollution?"
 American Journal of Respiratory and Critical Care Medicine, 161: pp. 665-673.
- Box, G. E. P. and G. C. Tiao. 1973. *Bayesian Inference in Statistical Analysis*. Wesley
 Publishing Co.: Wiley Classic Library, ISBN 0-471-57428-7.
- Burmaster, D.E. 1998. "Lognormal Distributions for Skin Area as a Function of Body Weight."
 Risk Analysis, 18(1):27-32.
- Dietert, R.R.; R. A. Etzel; D. Chen, et al. 2000. "Workshop to Identify Critical Windows of
 Exposure for Children's Health: Immune and Respiratory Systems Work Group Summary."
 Environ Health Perspect, 108(suppl 3): 483-490.
- Esmail, S.; Y. Bhambhani and S. Brintnell. 1995. "Gender Differences in Work Performance on
 the Baltimore Therapeutic Equipment Work Simulator." *American Journal of Occupational Therapy*, 49: 405 411.
- Federal Register. 1996. Responses to Significant Comments on the 1996 Proposed Rule on the
 National Ambient Air Quality Standards for Ozone (December 13, 1996; Fed. Reg. 61 FR
 65716).
- Folinsbee, L.J., et al. 1988. "Pulmonary Function and Symptom Responses after 6.6-hour
 Exposure to 0.12 ppm Ozone with Moderate Exercise." *Journal of the Air Pollution Control Association*, 38: 28-35.
- Graham, S. and T. McCurdy. 2005. *Revised Ventilation Rate (Ve) Equations for Use in Inhalation-Oriented Exposure Models: A NERL Internal Research Report.* Research
 Triangle Park, NC: EPA National Exposure Research Laboratory.
- Hazucha, M.J.; L. J. Folinsbee and P. A. Bromberg. 2003. "Distribution and Reproducibility of
 Spirometric Response to Ozone by Gender and Age." *Journal of Applied Physiology*, vol.
 95, no. 5, pp. 1917-1925.
- Hislop, A.A. 2002. "Airway and Blood Vessel Interaction During Lung Development." *Journal of Anatomy*, 201:325–334. Available at:
- 37 <<u>www.ncbi.nlm.nih.gov/pmc/articles/PMC1570917/</u>>.

- Horstman, D.H. et al. 1990. "Ozone Concentration and Pulmonary Response Relationships for
 6.6-hour Exposures with Five Hours of Moderate Exercise to 0.08, 0.10, and 0.12 ppm."
 American Review of Respiratory Disease, 142:1158-1163.
- Isaacs, K.; G. Glen; T. Mccurdy and L. Smith. 2008. "Modeling Energy Expenditure and Oxygen
 Consumption in Human Exposure Models: Accounting for Fatigue and EPOC." *Journal of Exposure Science and Environmental Epidemiology*, 18, 289–298.
- Isaacs, K. and L. Smith. 2005. New Values for Physiological Parameters for the Exposure Model
 Input File 'Physiology.txt.' Technical memorandum to Tom McCurdy, EPA, December 20.
- 9 Kim, C.S.; N. E. Alexis; A. G. Rappold; H. Kehrl; M. J. Hazucha; J. C. Lay; M. T. Schmitt; M.
- Case; R. B. Devlin; D. B. Peden; D. Diaz-Sanchez. 2011. "Lung Function and Inflammatory
 Responses in Healthy Young Adults Exposed to 0.06 ppm Ozone for 6.6 hours." *American Journal of Respiratory and Critical Care Medicine*, 183: 1215-1221.
- Linn, W.S.; R. D. Buckley; C. E. Spier; R. L. Blessey; M. P. Jones; D. A. Fischer and J. D.
 Hackney. 1978. "Health Effects of Ozone Exposure in Asthmatics." *The American Review of Respiratory Disease*, vol. 117, no. 5, pp. 835-843.
- Lunn, D.; C. Jackson; N. Best; A. Thomas and D. Spiegelhalter. 2012. *The BUGS Book A Practical Introduction to Bayesian Analysis*. CRC Press: Chapman and Hall.
- McCurdy, T.; G. Glen; L. Smith; Y. Lakkadi. 2000. "The National Exposure Research
 Laboratory's Consolidated Human Activity Database." *Journal of Exposure Analysis and Environmental Epidemiology*, 10: 566-578.
- McDonnell, W.F.; R. S. Chapman; M. W. Leigh; G. L. Strope; A. M. Collier. 1985. "Respiratory
 Responses of Vigorously Exercising Children to 0.12 ppm Ozone Exposure." *American Review of Respiratory Disease*, 132: 875-879.
- McDonnell, W. F.; H. R. Kehrl; S. Abdul-Salaam; P. J. Ives; L. J. Folinsbee; R. B. Devlin; J. J.
 O'Neil; D. H. Horstman. 1991. "Respiratory Response of Humans Exposed to Low Levels
 of Ozone for 6.6 hours." *Archives of Environmental Health*, 46(3):145-150.
- McDonnell, W.F. et al. 1993. "Predictors of Individual Differences in Acute Response to Ozone
 Exposure." *American Review of Respiratory Disease*, 147:818-825.
- McDonnell, W.F. et al. 1997. "Prediction of ozone-induced FEV₁ changes." *American Journal of Respiratory and Critical Care Medicine*, 156:715:722.
- McDonnell, W.F.; P. W. Stewart and M. V. Smith. 2007. "The Temporal Dynamics of Ozone induced FEV₁ Changes in Humans: An Exposure-response Model." *Inhalation Toxicology*,
 19:483-494.
- McDonnell, W.F.; P. W. Stewart and M. V. Smith. 2010. "Prediction of Ozone-induced Lung
 Function Responses in Humans." *Inhalation Toxicology*, 22(2):160-8.
- 36 McDonnell, W.F.; P. W. Stewart; M. V. Smith; C. S. Kim and E. S. Schelegle. 2012. "Prediction
- of Lung Function Response for Populations Exposed to a Wide Range of Ozone
- 38 Conditions." *Inhalation Toxicology*, 24:619-633.
- 39

1 2 3 4 5 6	 Narayanan, M.; J. Owers-Bradley; C. S. Beardsmore; M. Mada; I. Ball; R. Garipov; K. S. Panesar; C. E. Kuehni; B. D. Spycher; S. E. Williams; M. Silverman. 2012. "Alveolarization Continues During Childhood and Adolescence: New Evidence from Helium-3 Magnetic Resonance." <i>American Journal of Respiratory and Critical Care Medicine</i>, January 15; 185(2): 86–191. Available at: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC3410735/>.
7	Passannante, A. N.; M. J. Hazucha; P. A. Bromberg; E. Seal; L. Folinsbee; G. Koch. 1998.
8	"Nociceptive Mechanisms Modulate Ozone-induced Human Lung Function Decrements."
9	<i>Journal of Applied Physiology</i> , 85:1863-1870.
10 11 12 13 14	 Romieu, I; J. J. Sienra-Monge; M. Ramirez-Aguilar; M. M. Tellez-Rojo; H. Moreno-Macias; N. I. Reyes-Ruiz; B. E. Del Rio-Navarro; M. X. Ruiz-Navarro; G. Hatch; R. Slade; M. Hernandez-Avila. 2002. "Antioxidant Supplementation and Lung Functions Among Children with Asthma Exposed to High Levels of Air Pollutants." <i>American Journal of Respiratory and Critical Care Medicine</i>, 166: 703-709.
15	Samet, J.M. 2011. "Clean Air Scientific Advisory Committee (CASAC) Response to Charge
16	Questions on the Reconsideration of the 2008 Ozone National Ambient Air Quality
17	Standards." (Report number EPA-CASAC-11-004), dated March 30, 2011. Available at:
18	http://yosemite.epa.gov/sab/sabproduct.nsf/0/F08BEB48C1139E2A8525785E006909AC/
19	§File/EPA-CASAC-11-004-unsigned+.pdf>.
20	Schelegle, E.S.; C. A. Morales; W. F. Walby; S. Marion; R. P. Allen. 2009. "6.6-Hour Inhalation
21	of Ozone Concentrations from 60 to 87 ppb in Healthy Humans." <i>American Journal of</i>
22	<i>Respiratory and Critical Care Medicine</i> , 180:265-272.
23	Schelegle, E.S.; W. C. Adams; W. F. Walby and S. Marion. (2012). "Modeling of Individual
24	Subject Ozone Exposure Response Kinetics." <i>Inhalation Toxicology</i> , 24(7):401-415.
25	Schofield, W.N. 1985. "Predicting Basal Metabolic Rate, New Standards and Review of
26	Previous Work." <i>Clinical Nutrition</i> , 1985; 39C (suppl): 5–41.
27	Seal, Jr., E.; W. F. McDonnell; D. E. House. 1996. "Effects of Age, Socioeconomic Status, and
28	Menstrual Cycle on Pulmonary Response to Ozone." Archives of Environmental and
29	Occupational Health, 1996;51:132-137.
30	U.S. Environmental Protection Agency. 1996. <i>Review of National Ambient Air Quality</i>
31	<i>Standards for Ozone: Assessment of Scientific and Technical Information - OAQPS Staff</i>
32	<i>Paper</i> . Research Triangle Park, NC: EPA Office of Air Quality Planning and Standards.
33	(EPA document number EPA/452/R-96-007). Available at:
34	< <u>http://www.epa.gov/ttn/naaqs/standards/ozone/s_o3_pr.html</u> >.
35	U.S. EPA. 2007a. <i>Review of National Ambient Air Quality Standards for Ozone: Policy</i>
36	<i>Assessment of Scientific and Technical Information - OAQPS Staff Paper</i> . Research Triangle
37	Park, NC: EPA Office of Air Quality Planning and Standards. (EPA document number
38	EPA-452/R-07-007). Available at:
39	< <u>http://www.epa.gov/ttn/naaqs/standards/ozone/s_o3_cr_sp.html</u> >.
10	

- U.S. EPA. 2007b. Ozone Population Health Risk Assessment for Selected Urban Areas.
 Research Triangle Park, NC: EPA Office of Air Quality Planning and Standards. (EPA document number EPA-452/R-07-009). Available at:
- 4 <<u>http://www.epa.gov/ttn/naaqs/standards/ozone/s_o3_cr_td.html</u>>.
- 5 U.S. EPA. 2009. *Metabolically Derived Human Ventilation Rates: A Revised Approach Based* 6 Upon Oxygen Consumption Rates. Washington, DC: EPA National Center for
- 7 Environmental Assessment. (EPA document number EPA/600/R-06/129F). Available at:
- 8 <<u>http://cfpub.epa.gov/ncea/cfm/recordisplay.cfm?deid=202543</u>>.
- 9 U.S. EPA. 2011. O₃ National Ambient Air Quality Standards: Scope and Methods Plan for
 10 Health Risk and Exposure Assessment. Research Triangle Park, NC: EPA Office of Air
 11 Quality Planning and Standards. (EPA document number EPA-452/P-11-001). Available at:
 12 http://www.epa.gov/ttn/naags/standards/ozone/s o3 2008 pd.html>.
- U.S. EPA. 2012b. Total Risk Integrated Methodology (TRIM) Air Pollutants Exposure Model
 Documentation (TRIM.Expo / APEX, Version 4.4) Volume I: User's Guide. Research
 Triangle Park, NC: EPA Office of Air Quality Planning and Standards(EPA document
- 16 number EPA-452/B-12-001a). Available at:
- 17 <<u>http://www.epa.gov/ttn/fera/human_apex.html</u>>.
- 18 U.S. EPA. 2012c. Total Risk Integrated Methodology (TRIM) Air Pollutants Exposure Model
- 19 Documentation (TRIM.Expo / APEX, Version 4.4) Volume II: Technical Support
- 20 Document. Research Triangle Park, NC. (EPA document number EPA-452/B-12-001b).
 21 Available at: <<u>http://www.epa.gov/ttn/fera/human_apex.html</u>>.
- U.S. EPA. 2013a. Integrated Science Assessment of Ozone and Related Photochemical Oxidants.
 EPA National Center for Environmental Assessment. (EPA document number EPA/600/R-
- 24 10/076F, 2013). Available at:
- 25 <<u>http://www.epa.gov/ttn/naaqs/standards/ozone/s_o3_2008_isa.html</u>>.
- 26 WinBUGS, version 1.4.3. Available at: <<u>http://www.mrc-</u>
 27 bsu.cam.ac.uk/bugs/winbugs/contents.shtml>.

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7 CHARACTERIZATION OF HEALTH RISK BASED ON EPIDEMIOLOGICAL STUDIES

4 This chapter provides an overview of the methods used to estimate health risks in 5 selected urban areas based on application of results of epidemiology studies. Section 7.1.1 6 discusses the basic structure of the risk assessment, identifying the modeling elements and 7 related sources of input data needed for the analysis and presenting an overview of the approach 8 used in calculating health effect incidence using concentration-response (C-R) functions based 9 on epidemiological studies. Section 7.2 discusses air quality considerations. Section 7.3 10 discusses the selection of model inputs including: (a) selection of urban study areas, (b) selection 11 of epidemiological studies and specification of C-R functions, (c) specification of baseline health 12 effect incidence and prevalence rates, and (d) estimation of population (demographic) counts. 13 Section 7.4 describes how uncertainty and variability are addressed in the risk assessment, 14 including specification of the sensitivity analyses completed for the risk assessment and how 15 these differ from the core risk estimates. Section 7.5 summarizes the risk estimates that are 16 generated, including both the core estimates and sensitivity analyses. Finally, Section 7.6 17 provides an assessment of overall confidence in the risk assessment together with a set of key 18 observations regarding the risk estimates generated.

19 7.1 GENERAL APPROACH

20 7.1.1 Basic Structure of the Risk Assessment

This risk assessment involves the estimation of the incidence of specific health effect endpoints associated with exposure to ambient O_3 for defined populations located within a set of urban study areas. Because the risk assessment focuses on health effect incidence experienced by defined populations, it represents a form of population-level risk assessment and does not estimate risks to individuals within the population. Furthermore, because it models risk for residents in a set of urban study areas, it is not intended to provide an estimate of national-level risk¹.

The general approach used in both the prior and current O_3 risk assessments relies on C-R functions based on effect estimates and model specifications obtained from epidemiological studies. Since these studies derive effect estimates and model specifications using averages of ambient air quality data from fixed-site, population-oriented monitors, uncertainty arising from

¹ Chapter 8 provides a limited assessment of national risk focused on the mortality burden associated with recent O₃ levels. This risk and exposure assessment does not provide an analysis of the risk reductions that would be expected for the entire U.S. after meeting either the existing or alternative standards.

- 1 the application of these functions in an O₃ risk assessment is decreased if, in modeling risk, we
- 2 also use ambient air quality data at fixed-site, population-oriented monitors to characterize
- 3 exposure. Therefore, we developed a composite monitor for each urban study area to represent a
- 4 surrogate population exposure by averaging O₃ concentrations across the monitors in that study
- 5 area to produce a single composite hourly time series of values. The O_3 metrics used in
- 6 evaluating risk are derived from the composite monitor hourly time series distribution (see
- 7 sections 7.2 and Chapter 4 for additional detail on the characterization of ambient O_3 levels).²
- 8 The general O_3 health risk model, illustrated in Figure 7-1, combines O_3 air quality data, 9 C-R functions, baseline health incidence and prevalence data, and population data (all specific to 10 a given urban study area) to derive estimates of the annual incidence of specified health effects for that urban study area attributable to O_3 exposure. This risk assessment models risk for 12 11 12 urban study areas we selected to provide coverage for the types of urban O₃ scenarios likely to 13 exist across the U.S. (see section 7.3.1). Chapter 8 provides an assessment of the degree to which 14 the 12 selected urban areas are representative of other urban areas in the U.S. that are likely to 15 experience elevated risks from exposure to ambient O₃ under recent conditions.
- This risk assessment provides an updated set of estimates for risk under recent O_3 conditions and just meeting the existing standard, and additional estimates of risk if alternative standards are just met, with an emphasis on reductions in risk between just meeting the existing standard and just meeting alternative standards (the full set of risk estimates, including simulation of risk under current conditions is presented in Appendix 7-B). The alternative standard levels evaluated are 70, 65 and 60 ppb (expressed using the current form of the O_3
- standard).
- We simulated just meeting the existing and alternative O₃ standards by adjusting hourly O₃ concentrations measured over the O₃ season using a model-based adjustment methodology that estimates O₃ sensitivities to precursor emissions changes.³ These sensitivities, which estimate the response of O₃ concentrations to reductions in anthropogenic NOx and VOC emissions, are developed using the Higher-order Decoupled Direct Method (HDDM) capabilities in the Community Multi-scale Air Quality (CMAQ) model. More details on the HDDM-
- adjustment approach is presented in Chapter 4 of this REA and in Simon et al. (2013).
- 30 As discussed in Chapters 2 and 3, in modeling risk we employ continuous non-threshold
- 31 C-R functions relating O₃ exposure to health effect incidence. The use of non-threshold

 $^{^2}$ This holds for all air quality metrics used in modeling short-term mortality and morbidity endpoints. However, the air metric used in modeling long-term mortality is based on a seasonal average of maximum hourly values derived for each O₃ monitor within an urban study area with those individual averages then combined to generate a single seasonal average composite monitor value for each study area (see section 7.2 for more detail).

³ In the first draft of this REA, we used a statistical quadratic rollback approach to simulate just meeting the existing O₃ standards. In that first draft, we proposed using the model based approach used in this draft, and received support for the model based approach from CASAC (Frey, H.D., 2012).

1 functions reflects the discussion of the relevant studies in the O₃ ISA (see O₃ ISA, section

- 2 2.5.4.4, U.S. EPA 2013a). However, also consistent with the conclusions of the O₃ ISA, we
- 3 recognize that the evidence from the studies indicates less confidence in specifying the shape of
- 4 the C-R function at O₃ concentrations towards the lower end of the distribution of data used in
- 5 fitting the curve due to the reduction in the number of data points available. The O_3 ISA noted
- 6 that the studies indicate reduced certainty in specifying the shape of the C-R function
- 7 specifically for short-term O₃-attributable respiratory morbidity and mortality, in the range
- 8 generally below 20 ppb (for both 8hr-maximum and 24hr metrics) (O₃ ISA, section 2.5.4.4).
- 9 However, care needs to be taken in interpreting this range of reduced confidence indicated in the
- 10 studies and applying it to the interpretation of risk estimates generated for a specific urban study
- 11 area. This is because there is considerable heterogeneity in the effect of O_3 on mortality across
- 12 urban study areas (O₃ ISA section 6.6.2.3). Additionally, it is likely that levels of confidence
- 13 associated with C-R functions (including ranges of reduced confidence in specifying the
- 14 function) also vary across urban study areas reflecting underlying differences in factors
- 15 impacting the exposure-response relationship for O_3 , such as demographic differences and
- 16 exposure measurement error. For these reasons, the ≤ 20 ppb range discussed in the O₃ ISA
- 17 should be viewed as a more generalized range to be considered qualitatively or semi-
- 18 quantitatively, along with many other factors, when interpreting the risk estimates rather than as
- 19 a fixed, bright-line.⁴

Based on comments we received from CASAC on the 1st draft REA, we are no longer including estimates of risk down to the lowest measured level (LML).⁵ Instead, through the use of heat map tables, we focus on providing estimates of total risk, and the distribution of risk over concentrations of O_3 .⁶ Coupled with information about what the studies indicate about the C-R function at lower O_3 concentrations, this provides for a more complete understanding of confidence in estimated risk than simply truncating risk at the LML.

- 26
- 27

In modeling risk for all health endpoints included in the analysis, for recent O_3 conditions and just meeting the existing standard, we estimated <u>total risk (down to zero)</u>. For meeting the

 $^{^4}$ This range of reduced confidence in the shape of the C-R function is most appropriately applied to area-wide averages (i.e., composite monitor values) of the type often used in epidemiological studies rather than to the range of O₃ associated with a particular monitor. This reflects the fact that the observations presented in the O₃ ISA are themselves based on consideration for epidemiological studies which use composite monitor values.

⁵ Based on their November 19, 2012 letter commenting on the 1st draft REA, CASACrecommended against inclusion of risk estimates based on the LML in the core analysis due to the fact that there is little difference between these estimates and risk estimates based on total O₃ exposure and that LML information is not available for many of the epidemiological studies used in the REA (Frey and Samet, 2012). However, they recommend a more limited exploration of the LML and its implications for risk for one or more areas. In response, we have included coverage for LML as part of our discussion of the heat maps results presented in section 7.5.1.

⁶ Heat map tables illustrate the distribution of estimated O_3 -related deaths across daily O_3 levels for each urban study and allow a quick visual comparison of trends (in the distribution of total O_3 risk as we as risk reductions) across ambient O_3 ranges both within and across study areas (see section 7.5).

1 existing and alternative standards, we estimated both <u>total risk</u> as well as the difference in risk,

- 2 representing the degree of <u>risk reduction</u> associated with just meeting the existing and alternative
- 3 standard levels. When calculating risk differences, we focus on comparing total risk after just
- 4 meeting each alternative standard with total risk after just meeting the existing standards. We
- 5 also evaluate the incremental change in risk from meeting increasingly lower alternative standard
- 6 levels. Risk results are presented in terms of absolute numbers and changes in the O₃ attributable
- 7 incidence of mortality and morbidity, and in terms of the percent of baseline mortality and
- 8 morbidity attributable to O_3 . We also provide risks per 100,000 population (to normalize risks
- 9 across urban areas with different size populations to facilitate comparisons).
- 10 As with previous NAAQS-related risk assessments, for this analysis we have generated 11 two categories of risk estimates, including a set of core (or primary) estimates and an additional 12 set of sensitivity analyses. The core risk estimates utilize C-R functions based on
- 13 epidemiological studies for which we have relatively greater overall confidence and which
- 14 provide the best coverage for the broader O_3 monitoring period (rather than focusing only on the
- 15 summer season). Although it is not strictly possible to assign quantitative levels of confidence to
- 16 these core risk estimates due to data limitations, they are generally based on inputs having higher
- 17 overall levels of confidence relative to risk estimates that are generated using other C-R
- 18 functions. Therefore, emphasis is placed on the core risk estimates in making observations
- 19 regarding total risk and risk reductions associated with recent conditions and after just meeting
- 20 the existing and alternative standard levels. By contrast, the sensitivity analysis results typically
- 21 reflect application of C-R functions covering a wider array of design elements which can impact
- 22 risk (e.g., length of season, copollutants models, lag structures, statistical modeling methods etc).
- 23 The sensitivity analysis results provide insights into the potential impact of these design elements
- 24 on the core risk estimates, thereby informing our characterization of overall confidence in the
- 25 core risk estimates.⁷ We have significantly expanded our sensitivity analysis relative to that
- 26 completed for the 1st draft REA to address a wider range of modeling elements which can impact
- 27 the core risk estimates. Details of the design of the core and sensitivity analyses (including
- 28 modeling element composition) for each of the health effect endpoints categories covered in this
- risk assessment are presented in section 7.4.3 and briefly summarized below.
- 30

For short-term exposure related mortality, our core analysis is based on application of C-

31 R functions obtained from the Smith et al., 2009 epidemiological study (see section 7.3.2). In

⁷ In presenting both the core and sensitivity analysis, we include both point estimates and 95th percentile confidence intervals (CIs). The 95th percentile CIs reflect the statistical fit of the underlying effect estimates and therefore reflect the statistical power of the epidemiological studies supplying the effect estimates. Often in comparing sensitivity analysis with core risk estimates, we focus not only on the point estimates, but also on the confidence intervals since these inform our understanding of confidence in the respective risk estimates.

- 1 addition, we have completed an expanded array of sensitivity analyses which provide coverage
- 2 for a number of modeling elements including: (a) time period reflected in risk modeling (summer
- 3 season versus full monitoring period), (b) peak O₃ metric (8hr maximum versus 8hr mean) (c)
- 4 use of regional versus national-based Bayesian adjustment in deriving effect estimates,⁸ (d) use
- 5 of single (O₃-only) versus copollutant (O₃ and PM_{10}) models, ⁹ (e) application of alternative C-R
- 6 functions based on Zanobetti and Schwartz, 2008 (see section 7.3.2) and (f) size of the urban
- 7 study area (CBSA versus smaller multi-county study area)¹⁰ (see sections 7.4.3 and 7.5.3 for
- 8 additional detail on the sensitivity analyses completed). In addition to these sensitivity analyses,
- 9 we have considered alternative methods for adjusting air quality to attain existing and alternative
- 10 standards (NOx-only versus combination of VOC and NOx reductions). Additional sensitivity
- 11 analyses exploring lag structure may also provide useful information, but are not possible due to
- 12 the lack of availability of Bayes adjusted estimates for alternative lag structures.
- 13 For short-term exposure morbidity, we have effect estimates covering a wide range of
- 14 design elements including co-/single-pollutant models and lag structure. However, we were not
- 15 in a position to differentiate between these alternative model forms in terms of overall
- 16 confidence and have therefore included all of these estimates in the core analysis. This range of
- 17 risk estimates can also be viewed as a sensitivity analysis where there is no clear "core" estimate
- 18 and instead, the full range of risk estimates is considered to provide the best overall picture of
- 19 risk for a specific endpoint (see section 7.3.2 and 7.4.3).
- 20 Our analysis also includes estimates of long-term exposure related respiratory mortality,
- 21 including a core estimate based on a co-pollutant model (with PM_{2.5}) together with sensitivity
- 22 analyses exploring regional heterogeneity in the effect estimate and application of a national-

⁸ Short-term O₃-attributable mortality in this analysis is modeled using Bayesian-adjusted effect estimates. This approach involves adjustment of each city's effect estimate using a prior distribution reflecting the O₃-mortality relationship seen across the broader set of cities considered in the epidemiological study. For the sensitivity analysis, we compare the use of a national prior distribution (the core approach) for the Bayesian adjustment with use of a regional prior.

⁹ The copollutants model results are limited by the reduced number of days with copollutants sampling (either 1 in 3 or 1 in 6) which makes it difficult to evaluate the statistical significance of these results in view of the large posterior standard deviations (Smith et al., 2009). This increased uncertainty associated with the estimates prevents these results from being treated as part of the core analysis. Never the less, they provide perspective on the potential magnitude of risk associated with copollutants modeling and as such make an important contribution as a sensitivity analysis for short-term O₃-attributable mortality.

¹⁰ Core based statistical areas (CBSAs) are U.S. geographic areas defined by the Office of Management and Budget (OMB). They include an urban center of at least 10,000 people combined with adjacent urban and surburban areas that are socioeconomically tied to the urban center by commuting. CBSAs tend to be significantly larger than the study areas used in the epidemiological studies providing effect estimates. We have used risk estimates based on CBSAs in the core analysis in order to better represent the changes in risk that could be experienced in the broader urban areas and to avoid the introduction of known bias into the risk assessment. We have included risk estimates based on the smaller study areas from the original epidemiological studies as sensitivity analyses (see discussion later in this section for additional detail).

- 1 level estimates focusing only on O_3 (see section 7.5.3).¹¹ The decision to model this endpoint is
- 2 based on our evaluation of the evidence as summarized in the O₃ ISA and comments received
- 3 from CASAC based on the 1st draft risk assessment (Frey H.D., 2012 p.).
- 4 As noted earlier, for this draft, we have modeled all core risk estimates using study areas 5 based on the core-based statistical area (CBSA) regardless of whether the epidemiological 6 studies providing the effect estimates used the CBSA spatial definition or a different spatial 7 study area definition. The decision to use CBSA-based study areas in all core simulations for this 8 draft reflects our desire to better represent the changes in risk that could be experienced in the urban 9 areas and avoid introducing substantial known bias into the risk estimates. As discussed in 10 Chapter 4 (section 4.3.1.2), most nonattaining O_3 monitors are not located in the center of the urban 11 area, but instead in the surrounding areas, reflecting the transport and atmospheric chemistry 12 governing O_3 formation. The monitors in the urban core areas are usually most affected by local 13 sources of NOx and experience lower concentrations of O_3 since the NO is titrating the O_3 in these 14 areas. For these monitors, simulating attainment of the existing and alternative standard levels can 15 result in an increase in O₃ concentrations, while areas further out from the core experience the 16 expected reduction in O₃ level. Had we focused risk estimates on the smaller urban core areas 17 used in some of the epidemiological studies, we would not have fully captured the changes in 18 risk estimated to be experienced by the broader urban area since we would have been focusing 19 only on those areas experiencing net increases in O_3 (when simulating attainment of the existing 20 and alternative standard levels). By modeling risk for the core analysis using the more inclusive 21 CBSA study areas, we insure that risk estimates will include consideration both for the relatively 22 smaller core urban areas experiencing increases in O₃ as well as the broader urban and suburban 23 area experiencing risk reductions. We will also insure that, to a greater extent, the analysis 24 includes the county with the design value monitor in the assessment of risk (see section 7.2). 25 There is a degree of uncertainty introduced through application of effect estimates to 26 study areas (i.e., CBSAs) that do not match those used in the underlying epidemiological studies. 27 This uncertainty should be viewed within the context of the overall larger uncertainty associated with transferring effect estimates from the context of the epidemiological studies to the context 28 29 of the risk assessment. The epidemiological studies used in modeling short-term exposure-related 30 endpoints generate effect estimates based on day to day variation in O_3 and health effects, using
- 31 the area wide average O_3 concentrations. Area wide O_3 averaging masks the specific population
- 32 distribution of O_3 exposures which reflects the times and durations of exposures to O_3 measured
- 33 at individual monitors in an urban area. We apply those effect estimates to the air quality

¹¹ The seasonal average metrics used in the long-term mortality estimate are not very sensitive to the reduced number of days with co-pollutant monitoring, and as such it is appropriate to use the co-pollutant model in generating the core risk estimates.

1 scenarios of just meeting existing and alternative standards, where we are shifting the entire

- 2 distribution of daily O_3 concentrations, and altering the relationships between O_3 concentrations
- 3 at different monitors, and thus likely altering the relationship between area wide average O_3 and
- 4 the population distribution of O_3 exposures. By doing so, we introduce an additional source of
- 5 exposure measurement error, which goes beyond the impact that measurement error has on the
- 6 effect estimate, and introduces additional uncertainty into the estimates of risk associated with
- 7 simulating meeting existing and alternative standards.

8 Our decision to use the CBSA to define the spatial extent of each urban study area 9 reflects the greater weight we place on minimizing biases relative to minimizing uncertainty, 10 although we strive to minimize both where possible. The sensitivity analysis related to using 11 study-based spatial definitions for urban areas shows clearly that using the smaller urban areas 12 biases downward the risk reductions across an urban area. Thus, to avoid this bias in risk 13 estimates we accept a measure of increased uncertainty associated with the application of effect 14 estimates to study areas that are larger than those used in some of the original epidemiological 15 studies providing those effect estimates.

16 Using the CBSA definitions of urban areas can partially address the bias caused by 17 focusing only on urban core areas. However, it does not address this bias fully in some areas 18 because of the unevenness in monitoring throughout urban areas. In some urban areas the 19 monitors are more evenly distributed across the CBSA, while in other areas they are not. For 20 example, in some urban areas, there is a high density of monitors in the urban core counties, with 21 less density of monitors in surrounding counties also in the CBSA. Because we use a simple 22 average (to match the averaging used in the epidemiology studies) of monitors across the CBSA, 23 this means that O_3 concentrations in areas where there are more monitors (e.g. in urban core 24 counties) will get a higher weight in the average O_3 concentrations relative to O_3 concentrations 25 in other parts of the CBSA. To the extent that the area with the higher density of monitors experiences increases in O₃ while the remaining area experiences decreases in O₃, the overall 26 27 average O₃ concentrations applied to populations in the entire CBSA will be weighted more 28 towards O₃ increases, which will attenuate the overall risk reduction that may be associated with 29 meeting alternative O_3 standards. We are not able to determine the magnitude of this remaining 30 bias; however, it is expected to be higher in locations with a high percentage of total CBSA 31 monitors concentrated in urban core counties.

The risk assessment reflects consideration for five years of recent air quality data from 2006 through 2010, with these five years reflecting two three-year attainment simulation periods that share a common overlapping year (i.e., 2006-2008 and 2008-2010 - see section 7.2). We selected these two attainment simulation periods to provide coverage for a more recent time period with relatively elevated O₃ levels (2006-2008) and recent time period with relatively

1 lower O₃ levels (2008-2010). For the REA, we model risk for the middle year of each three-year 2 attainment simulation period in order to provide estimates of risk for a year with generally higher 3 O_3 levels (2007) and a year with generally lower O_3 levels (2009). In modeling risk, we matched 4 the population data used in the risk assessment to the year of the air quality data. For example, 5 when we used 2007 air quality data, we used 2007 population estimates. For baseline incidence 6 and prevalence, rather than interpolating rates for the two specific years modeled in the risk 7 assessment, we selected the closest year for which we had existing incidence/prevalence data 8 (i.e., for simulation year 2007, we used available data for 2005 and for simulation year 2009, we 9 used data from 2010). The calculation of baseline incidence and prevalence rates is described in 10 section 7.3.4.

11 The risk assessment procedures described in more detail below are diagramed in Figure 12 7-1. To estimate the change in incidence of a given health effect resulting from a given change in 13 ambient O_3 concentrations in an assessment location, the following analysis inputs are necessary:

- Air quality information including: (1) O₃ air quality data from each of the
 simulation years included in the analysis (2007 and 2009) from population-oriented
 monitors in the assessment location (these are aggregated to form composite monitor
 values used to represent population exposure), and (2) a method for adjusting the air
 quality data to simulate just meeting the current or alternative suite of O₃ standards.
 (These air quality inputs are discussed in more detail in Chapter 4).
- 20 • **C-R function**(s): which provide an estimate of the relationship between the health 21 endpoint of interest and O₃ concentrations (for this analysis, C-R functions used were 22 applied to urban study areas matching the assessment locations from the 23 epidemiological studies used in deriving the functions, in order to increase overall 24 confidence in the risk estimates generated - see section 7.3.2). For O_3 , 25 epidemiological studies providing information necessary to specify C-R functions are 26 readily available for O_3 -related health effects associated with short-term exposures 27 (Section 7.1.2 describes the role of C-R functions in estimating health risks associated 28 with O₃). In addition, the Jerrett et al. (2009) study provided a C-R function for 29 modeling mortality risks associated with longer-term exposures to O₃.
- Population information (*baseline health affects incidence and prevalence rates and population*): The baseline incidence provides an estimate of the incidence rate
 (number of cases of the health effect per year or day, depending on endpoint, usually
 per 10,000 or 100,000 general population) in the assessment location corresponding
 to recent ambient O₃ levels in that location. The baseline prevalence rate describes the
 prevalence of a given disease state or conditions (e.g., asthma) within the population

1 (number of individuals with the disease state/condition, usually per 10,000 or 100,000 2 general population). To derive the total baseline incidence or prevalence per year, this 3 rate must be multiplied by the corresponding population number (e.g., if the baseline 4 incidence rate is number of cases per year per 100,000 population, it must be 5 multiplied by the number of 100,000s in the population) (Section 7.3.4 summarizes 6 considerations related to the baseline incidence and prevalence rates and population 7 data inputs to the risk assessment). 8 9 In addition to the inputs described above, it is also necessary to specify the spatial extent 10 of the study areas that will be modeled. These study areas definitions determine the composition 11 of (a) the composite monitor values (which specific set of monitors are used in constructing the 12 composite monitor, reflecting the area-wide average across monitors for each study area), (b) the 13 specific set of effect estimates that will be used (matching the study areas to the specific set of

14 effect estimates in the epidemiological studies being used to support modeling of endpoints), (c)

15 the baseline incidence data and (d) the population demographic (count) data for each study area.

16 As mentioned earlier, for this REA we have modeled 12 urban study areas and have used the

17 CBSA spatial definition to specify the extent of each of these urban areas (see section 7.3.1 for

18 additional details on study area selection).



Figure 7-1 Flow Diagram of Risk Assessment for Short-term Exposure Studies

1 This risk assessment was implemented using the EPA's Environmental Benefits Mapping 2 and Analysis Program—Community Edition, Version 0.63 (BenMAP-CE) (U.S. EPA, 2013b). 3 This GIS-based computer program draws upon a database of population, baseline 4 incidence/prevalence rates and effect coefficients to automate the calculation of health impacts. 5 For this analysis, the standard set of effect coefficients and health effect incidence data available 6 in BenMAP has been augmented to reflect the latest studies and data available for modeling O_3 7 risk. EPA has traditionally relied upon the BenMAP program to estimate the health impacts 8 avoided and economic benefits associated with adopting new air quality rules. For this analysis, 9 EPA used the model to estimate O₃-related risk for the suite of health effects endpoints described 10 in section 3.2. There are three primary advantages to using BenMAP for this analysis, as 11 compared to the procedure for estimating population risk followed in the last review. First, once 12 we have configured the BenMAP software for this particular O₃ analysis, the program can 13 produce risk estimates for an array of modeling scenarios across a large number of urban areas. 14 Second, the program can more easily accommodate a variety of sensitivity analyses. Third, BenMAP allowed us to complete the national assessment of O₃ mortality described in Chapter 8, 15 16 which plays in important role in assessing the representativeness of the urban study area analysis.

17 7.1.2 Calculating O₃-Related Health Effects Incidence

18 The C-R functions used in the risk assessment are empirically estimated associations 19 between average ambient concentrations of O₃ and the health endpoints of interest (e.g., 20 mortality, hospital admissions, emergency department visits). This section describes the basic 21 method used to estimate changes in the incidence of a health endpoint associated with changes in 22 O₃, using a "generic" C-R function of the most common functional form.

Although some epidemiological studies have estimated linear C-R functions and some have estimated logistic functions, most of the studies used a method referred to as "Poisson regression" to estimate exponential (or log-linear) C-R functions in which the natural logarithm of the health endpoint is a linear function of O₃:

- 27
- 28

 $y = Be^{\beta x} \tag{1}$

where x is the ambient O_3 level, y is the incidence of the health endpoint of interest at O_3 level x, β is the coefficient relating ambient O_3 concentration to the health endpoint, and B is the incidence at x=0, i.e., when there is no ambient O_3 . The relationship between a specified ambient O_3 level, x₀, for example, and the incidence of a given health endpoint associated with that level (denoted as y₀) is then

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- 35

 $y_0 = Be^{\beta x_0} \tag{2}$

1 2 3

4

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Because the log-linear form of a C-R function (equation (1) is by far the most common form, we use this form to illustrate the "health impact function" used in the O₃ risk assessment. If we let x₀ denote the baseline (upper) O₃ level, and x₁ denote the lower O₃ level, and y₀ and y₁ denote the corresponding incidences of the health effect, we can derive the following relationship between the change in x, Δx= (x₀- x₁), and the corresponding change in y, Δy, from

7 8

9

equation (1).¹²

$$\Delta y = (y_0 - y_1) = y_0 [1 - e^{-\beta \Delta x}].$$
(3)

- 10 Alternatively, the difference in health effects incidence can be calculated indirectly using 11 relative risk. Relative risk (RR) is a measure commonly used by epidemiologists to characterize 12 the comparative health effects associated with a particular air quality comparison. The risk of mortality at ambient O_3 level x_0 relative to the risk of mortality at ambient O_3 level x_1 , for 13 14 example, may be characterized by the ratio of the two mortality rates: the mortality rate among 15 individuals when the ambient O_3 level is x_0 and the mortality rate among (otherwise identical) 16 individuals when the ambient O_3 level is x_1 . This is the RR for mortality associated with the 17 difference between the two ambient O_3 levels, x_0 and x_1 . Given a C-R function of the form shown in equation (1) and a particular difference in ambient O_3 levels, Δx , the RR associated 18 with that difference in ambient O₃, denoted as RR Δx , is equal to $e^{\beta \Delta x}$. The difference in health 19 effects incidence, Δy , corresponding to a given difference in ambient O₃ levels, Δx , can then be 20 21 calculated based on this RR Δx as: 22
- 23
- Ζ.

 $\Delta y = (y_0 - y_1) = y_0 [1 - (1/RR_{\Delta x})].$ (4)

24

Equations (3) and (4) are simply alternative ways of expressing the relationship between a given difference in ambient O_3 levels, $\Delta x > 0$, and the corresponding difference in health effects incidence, Δy .¹³ These health impact equations are the key equations that combine air quality information, C-R function information, and baseline health effects incidence information to estimate ambient O_3 health risk.

¹² If $\Delta x < 0 - i.e.$, if $\Delta x = (x_1 - x_0) - then the relationship between <math>\Delta x$ and Δy can be shown to be $\Delta y = (y_1 - y_0) = y_0 [e^{\beta \Delta x} - 1]$. If $\Delta x < 0$, Δy will similarly be negative. However, the *magnitude* of Δy will be

the same whether $\Delta x > 0$ or $\Delta x < 0 - i.e.$, the absolute value of Δy does not depend on which equation is used. ¹³ When calculating total risk associated with a specific air quality scenario, Δx is the total O₃ concentration

associated with a given study area (as noted earlier in section 7.1.1, we are not incorporating thresholds, such as LMLs into this analysis).

1 7.2 AIR QUALITY CONSIDERATIONS

2 Air quality data are discussed in detail in Chapter 4 of this report. Here we describe those 3 air quality considerations that are directly relevant to the estimation of health risks in the 4 epidemiology based portion of the risk assessment. As described in section 7.1.1, the risk 5 assessment uses composite (area-wide average) monitor values derived for each urban study area 6 as the basis for characterizing population exposure in modeling risk. The use of composite 7 monitors reflects consideration for the way ambient O₃ data are used in the epidemiological 8 studies providing the C-R functions (see section 7.1.1). For the short-term exposure related 9 health endpoints, the composite monitor values derived for this analysis include hourly time 10 series for each study area (where the O_3 value for each hour is the average of measurements 11 across the monitors in that study area reporting values for that hour). Once these composite 12 monitor hourly time series are constructed, we can then extract short-term peak O₃ metrics 13 needed to model specific health effects endpoints. For short-term O₃-attributable endpoints, 14 reflecting consideration for available evidence in the published literature (see section 7.3.2), we 15 have focused the analysis on short-term peak O₃ metrics including 1hr maximum, 8hr mean and 16 8hr maximum. The 24 hour average has been deemphasized for this analysis, although it is still 17 used in risk modeling when use of C-R functions based on this metric allow us to cover a specific health effect endpoint/location of particular interest 14 (see section 7.3.2). 18 19 For modeling mortality risk associated with long-term O₃-attributable we construct 20 seasonally-averaged maximum hourly O_3 values (see section 7.3.2). The derivation of composite 21 monitor distributions used in modeling this health effect endpoint is different than that used for 22 short-term O₃-attributable endpoints. Specifically, for the long-term O₃-attributable endpoint we

first construct the seasonally-averaged peak O_3 metric for each monitor within a given study area and then average those monitor-specific metric values together to generate a single composite value to use in generating risk estimates for that study area.

26 In applying effect estimates obtained from epidemiological studies we attempted to 27 match the modeling period (e.g. O₃ monitoring season) associated with each epidemiology study. 28 This increases overall confidence in the risk compared with using a single more generalized 29 specification of the modeling period. As discussed earlier, we modeled all health effect endpoints 30 for the core analysis using a CBSA-based study area. The use of the CBSA-based study areas 31 addresses potential bias that would have occurred had we focused the risk assessment on the 32 smaller core urban study areas. (see section 7.1.1). Table 7-1 identifies (a) the counties 33 associated with the CBSA definition for each of the 12 urban study areas, (b) the number of O_3 34 monitors associated with each CBSA (and a flag for whether the design value monitor is

¹⁴ In order to provide estimates of respiratory-related HA for LA, we did include a C-R function based on Linn et al., 2000, which utilizes a 24 hour average exposure metric.

1 contained within the CBSA), (c) the number of monitors associated with the smaller Smith et al.,

2 2009-based study areas, and (d) the specific O₃ modeling period for each study area. A map

3 showing the counties and monitors for these 12 urban areas can be found in Chapter 4 (figure 4-

4 5, Section 4.3.2.1).

- 5
- 6

Table 7-1 Information on the 12 Urban Case Study Areas in the Risk Assessment

Study Area	Counties associated with the CBSA definition	# of O ₃ Monitors within the CBSA ^a	Required O_3 Monitoring Season
Atlanta	Barrow, Bartow, Butts, Carroll, Cherokee Clayton, Cobb, Coweta, Dawson, DeKalb, Douglas, Fayette, Forsyth, Fulton, Gwinnett, Haralson, Heard, Henry, Jasper, Lamar, Meriwether, Newton, Paulding, Pickens, Pike, Rockdale, Spalding, Walton	13 (3)	March - October
Baltimore	Anne Arundel, Baltimore, Carroll, Harford, Howard, Queen Anne's, Baltimore	7 (1)	April - October
Boston	Essex, Middlesex, Norfolk, Plymouth, Suffolk, Rockingham, Strafford	11* (2)	April - September
Cleveland	Cuyahoga, Geauga, Lake, Lorain, Medina	10* (4)	April - October
Denver	Adams, Arapahoe, Broomfield, Clear Creek, Denver, Douglas, Elbert, Gilpin, Jefferson, Park	16 (6)	March - September
Detroit	Lapeer, Livingston, Macomb, Oakland, St. Clair, Wayne	8 (4)	April - September
Houston	Austin, Brazoria, Chambers, Fort Bend, Galveston, Harris, Liberty, Montgomery, San Jacinto, Waller	22 (17)	January - December
Los Angeles	Los Angeles, Orange	21* (17)	January - December
New York	Bergen, Essex, Hudson, Hunterdon, Middlesex, Monmouth, Morris, Ocean, Passaic, Somerset, Sussex, Union, Bronx, Kings, Nassau, New York, Putnam, Queens, Richmond, Rockland, Suffolk, Westchester, Pike	22 (7)	April - October
Philadelphia	New Castle, Cecil, Burlington, Camden, Gloucester, Salem, Bucks, Chester, Delaware, Montgomery, Philadelphia	15 (4)	April - October
Sacramento	El Dorado, Placer, Sacramento, Yolo	17 (8)	January - December
St. Louis	Bond, Calhoun, Clinton, Jersey, Macoupin, Madison, Monroe, St. Clair, Franklin, Jefferson, Lincoln, St. Charles, St. Louis, Warren, Washington, St. Louis	17 (2)	April - October

7 8 9 a - This column presents the number of monitors within each CBSA, whether the design value falls outside of the CBSA (denoted with an "*") and the number of monitors within the smaller Smith et al., 2009-based study area (in parenthesis).

10

11

We estimate risk associated with recent O₃ conditions as well as risk associated with

12 simulating just meeting the existing and alternative standards. While the derivation of composite
1 monitor hourly O_3 distributions (and associated peak exposure metrics) for recent conditions is 2 relatively straightforward, the generation of these estimates for the scenarios of just meeting the 3 existing and alternative standards is more complex. The procedures for simulating attainment of 4 both existing and alternative O_3 standards are presented in Chapter 4 and Chapter 4 appendices. 5 Summary statistics for the air metrics used in modeling risk for each of the 12 urban 6 study areas under recent conditions and simulated attainment of the existing and alternative 7 standard levels are presented in Chapter 4 (see section 4.3.3.2, Figures 4-10 (2007) and 4-11 8 (2009)).

9 7.3 SELECTION OF MODEL INPUTS AND ASSUMPTIONS

10 7.3.1 Selection of Urban Study Areas

11 This analysis focuses on modeling risk for a set of urban study areas, reflecting the goal 12 of providing risk estimates that have greater overall confidence due to the use of location-13 specific data when available for these urban locations. In addition, given the greater availability 14 of location-specific data, a more rigorous evaluation of the impact of uncertainty and variability 15 can be conducted for a set of selected urban study areas than would be possible for a broader 16 regional or national-scale analysis. We considered the following factors in selecting the 12 urban 17 study areas included in this analysis:

- Air Quality Data: An urban area has reasonably comprehensive monitoring data for the period of interest (2006-2010) to support the risk assessment. This criterion was
 evaluated qualitatively by considering the number of monitors within the CBSA of the prospective urban areas. Locations with one or two monitors would be excluded since
 they had relatively limited spatial coverage in characterizing O₃ levels.
- Elevated Ambient O₃ Levels: Because we are interested in evaluating the potential
 magnitude of risk reductions associated with just meeting the existing and alternative O₃
 standard levels, we focus on study areas with elevated ambient O₃ levels at or above the
 existing standard, such that just meeting alternative O₃ standard levels would result in
 some degree of risk reduction.
- Location-specific C-R Functions: Given the health endpoints selected for inclusion in
 the analysis (see section 7.3.2), there are epidemiological studies of sufficient quality
 available for these urban study areas to provide the C-R functions necessary for modeling
 risk. This criterion primarily applies to short-term epidemiological studies since the
 associated health effect endpoints are the primary focus of the REA. Short-term O₃-

attributable epidemiological studies often include city-specific effect estimates, and in
 some cases are multi-city studies that provide estimates for multiple cities.

Baseline Incidence Rates and Demographic Data: The required urban area-specific
 baseline incidence rates and population data are available for a recent year for at least one
 of the health endpoints.

6 **Geographic Heterogeneity**: Because O₃ distributions and population characteristics vary • 7 geographically across the U.S., we selected urban study areas to provide coverage for 8 regional variability in factors related to O_3 risk including variability in the spatial pattern 9 of O_3 in the urban area, population exposure (differences in residential housing density, 10 air conditioning use and commuting patterns), demographic characteristics (baseline 11 incidence rates, SES) and variability in effect estimates. The degree to which the set of 12 urban study areas provided coverage for regional differences across the U.S. in many of 13 these O_3 risk-related factors was evaluated as part of the representativeness analysis 14 presented in Chapter 8.

- 15 Application of the above criteria resulted in the selection of 12 urban study areas for 16 inclusion in the risk assessment including:
- 17 Atlanta, GA
- 18 Baltimore, MD
- 19 Boston, MA
- 20• Cleveland, OH
- Denver, CO
- Detroit, MI
- Houston, TX
- Los Angeles, CA
 - New York, NY
- 26• Philadelphia, PA
- Sacramento, CA
 - St. Louis, MO
- 28 29

25

The specific set of counties used in defining each of the 12 urban study areas based onthe CBSA is presented in Table 7-1.

7.3.2 Selection of Epidemiological Studies and Specification of Concentration-Response Functions

Once the set of health effect endpoints to be included in the risk assessment has been specified, the next step was to select the set of epidemiological studies that will provide the effect estimates and model specifications used in the C-R functions. This section describes the approach used in completing these tasks and presents a summary of the epidemiological studies and associated C-R functions specified for use in the risk assessment.

8 In Chapter 2, section 2.5 we identified the set of health effect categories and associated 9 endpoints to be included in this assessment, based on review of the evidence provided in the O_3 10 ISA (U.S. EPA, 2013a). The selection of specific health effect endpoints to model within a given

11 health effect endpoint category is an iterative process involving review of both the strength of

12 evidence (for a given endpoint) as summarized in the O_3 ISA together with consideration for the

13 available epidemiological studies supporting a given endpoint and the ability to specific key

14 inputs needed for risk modeling, including effect estimates and model forms. Ultimately,

15 endpoints are only selected if (a) they are associated with an overarching effect endpoint

16 category selected for inclusion in the risk assessment and (b) they have sufficient

17 epidemiological study support to allow their modeling in the risk assessment. Health effect

18 endpoints selected for inclusion in the second draft REA include:

19

20

Short-term O₃-attributable:

- 21 Mortality (likely to be a casual relationship) 22 o All-cause (non-accidental) 23 o Cardiovascular 24 o Respiratory 25 Respiratory effects (causal relationship) • ED (asthma, wheeze, all respiratory symptoms) 26 HA (COPD, asthma, all respiratory)¹⁵ 27 0 28 • Respiratory symptoms 29
- 30 **Long-term** O₃-attributable:

¹⁵ Regarding COPD-related HA, the O₃ ISA states that "Although limited in number, both single- and multi-city studies consistently found positive associations between short-term O₃ exposures and asthma and COPD hospital admissions." (U.S. EPA 2013a, p. 6-128). It is also important to point out that when modeling of COPD-related HA is limited to the summer months (as was done for the REA), available effect estimates have tighter confidence intervals and are generally positive, which increases overall confidence in the resulting risk estimate (see U.S. EPA 2013a, Figure 6-19).

1	• Respiratory effects, focusing on respiratory-related mortality (likely causal
2	relationship). ¹⁰
3	
4 5	We selected epidemiological studies to support modeling of the health effect endpoints listed above by applying a number of criteria including ¹⁷ :
6	• The study was peer-reviewed, evaluated in the O ₃ ISA, and judged adequate by EPA
7	staff for purposes of inclusion in the risk assessment. We considered the following
8	criteria: whether the study provides C-R relationships for locations in the U.S.,
9	whether the study has sufficient sample size to provide effect estimates with a
10	sufficient degree of precision and power, and whether adequate information is
11	provided to characterize statistical uncertainty.
12	• Preference for multicity studies given that they typically have greater power and
13	reflect patterns of O_3 related health effects over a range of urban areas (and regions)
14	which can display variability in key risk-related factors such as exposure
15	measurement error. In the case of short-term O_3 -attributable mortality, we also
16	favored those multi-city studies for which we could obtain Bayesian-adjusted city-
17	specific estimates from the study authors, since these incorporate both city-specific
18	effect information with information from the broader array of cities included in the
19	study. In those instances where we did not have multi-city studies (e.g., with many of
20	the short-term respiratory-related morbidity endpoints) we use single-city studies.
21	• The study design is considered robust and scientifically defensible, particularly in
22	relation to methods for covariate adjustment, including treatment of confounders, as
23	well as treatment of effect modifiers. For example, if a given study used ecological-
24	defined variables (e.g., smoking rates) as the basis for controlling for confounding,
25	concerns may be raised as to the effectiveness of that control.
26	• The study is not superseded by another study (e.g., if a later study is an extension or
27	replication of a former study, the later study would effectively replace the former
28	study), unless the earlier study has characteristics that are clearly preferable (e.g.,
29	inclusion of copollutants models, or use of a peak exposure metric of interest).

¹⁶ The O₃ ISA classifies long-term O₃-attributable respiratory health effects, including respiratory-related mortality, as having a likely causal classification. By contrast, it classifies long-term O₃-attributable total mortality as having a suggestive of a causal relationship classification (O₃ ISA, 2012, Chapter 1). We have focused on modeling long-term O₃-attributable respiratory-related mortality given the greater support for this health endpoint relative to total mortality.

¹⁷ In addition to the criteria listed here, we also attempted to include studies that provide coverage for populations considered particularly at-risk for a particular health (e.g., children, individuals with preexisting disease). However, a study would have to meet the criteria listed here (in addition to providing coverage for an at-risk population) in order for that study to be used to derive C-R functions.

- We applied the above criteria and selected the set of epidemiological studies presented in
 Table 7-2 for use in specifying C-R functions (Table 7-2 also describes elements of the C-R
 functions specified using each epidemiological study, as discussed below).
- 4 As part of methods refinement for this risk assessment, we considered studies that 5 utilized more sophisticated and potentially representative exposure surrogates in characterizing 6 population exposure (e.g., using population-weighted O_3 monitor values instead of equally-7 weighted monitors, linking exposures in individual counties or U.S. Census tracts to the nearest 8 monitor, rather than using a composite monitor value to represent the entire study area). 9 However, analysis conducted by EPA demonstrated that use of the simpler composite monitor 10 approach (as used for other short-term O₃-attributable morbidity endpoints) generated risk 11 estimates that were very close to those generated using the population-weighted O₃ metric (see 12 REFERENCE- Karen Wesson???). Therefore, in order to conserve time and resources, we 13 modeled this endpoint using the more generalized composite monitor-based metric. And finally, 14 a number of the long-term O_3 -attributable morbidity studies originally considered for modeling 15 this endpoint category did involve more complex O₃ metrics (e.g., Atkinbami et al., 2010, Meng 16 et al., 2010, and Moore et al., 2008). However, limitations in the study-level data required to 17 support risk assessment prevents us at this point from completing a quantitative risk assessment for this category of health endpoints with a reasonable degree of confidence.¹⁸ 18

19 Based on additional evaluation of the literature, we have substituted Smith et al., 2009 for 20 Bell et al., 2004 as a source of Bayes-adjusted city-specific effect estimates to support modeling 21 short-term O₃-attributable mortality. This decision reflects a number of factors. The Smith et al., 22 2009 study includes a wider range of simulations exploring sensitivity of the mortality effect to 23 different model specifications including (a) regional versus national Bayes-based adjustment, (b) 24 copollutants models considering PM_{10} , and (c) all - year versus O_3 -season based estimates. This 25 is contrasted with the Bell et al., 2004 study which does not provide this degree of model 26 exploration. In obtaining the city-specific Bayes-adjusted effect estimates for the Smith et al., 27 2009 study from the study authors, we were provided with estimates reflecting this range of 28 alternative model specifications which allowed us to incorporate them into both the core and 29 sensitivity analysis portions of the REA (see section 7.4.3). In addition, the Smith et al., 2009 30 study does not use the trimmed mean approach employed in the Bell et al., 2004 study in 31 preparing O_3 monitor data. We have a number of concerns regarding the trimmed mean approach 32 including (1) the potential loss of temporal variation in the data when the approach is used (this 33 could impact the size of the effect estimate) and (2) a lack of complete documentation for the

¹⁸ However, as noted in section 7.7.3 of the first draft REA, these limitations do not prevent the use of this evidence from informing consideration of the levels of exposure at which specific types of health effects may occur (i.e., the evidence analysis, which is an important aspect of the O₃ NAAQS review). Rather, these limitations only prevent the quantitative estimation of risk with a reasonable degree of confidence.

1 approach which prevents us from fully reviewing the technique and using it in preparing O_3

2 metrics for the REA. Given these concerns, we view it as advantageous that the Smith et al.,

3 2009 study does not use the trimmed mean approach.

4 With the exception of the trimmed mean approach, the Smith et al., 2009 study was 5 intended to reproduce the results of the Bell et al., 2004 analysis. Thus, the core risk results based on Smith et al 2009 are comparable to the 1st draft REA estimates based on Bell et al 2004, 6 7 while the alternative models provided in Smith et al 2009 allow for an expanded set of sensitivity 8 analyses. The comparability of the Smith et al 2009 and Bell et al 2004 estimates is confirmed by 9 the graphical comparison in Smith et al 2009 of mortality effect estimates (for the 24hr O_3) 10 metric) with matching effect estimates from Bell et al., 2004. This comparison demonstrates the 11 close match of the two studies (for this particular scenario).

12 Reflecting the points made above, in modeling short-term O₃-attributable mortality, we 13 have included a core analysis based on the national-Bayesian adjusted city-specific effect 14 estimates (reflecting the full O_3 monitoring period in each city) obtained from Smith et al., 2009. 15 As sensitivity analyses, we have included effect estimates obtained from Smith et al., 2009 16 which reflect application of copollutants models (including PM_{10}), Bayes adjustment using a regional prior, ¹⁹ and a shorter fixed O_3 measurement period (April-October). In the 1st draft 17 18 REA, we had also included national Bayes-adjusted effect estimates (reflecting a fixed June-19 August period) obtained from Zanobetti and Schwartz, 2008 as part of the core analysis. However, we have decided to instead include these as part of the sensitivity analysis in this 2nd 20 21 draft of the REA since these effect estimates cover a more limited warm-weather period and 22 consequently will generate only partial characterizations of mortality risk (since they exclude 23 risk occurring during the non-summer months).

24 We have also included estimates of respiratory-related mortality associated with long-25 term O_3 exposures based on effect estimates obtained from Jerrett et al., 2009. The decision to 26 model long-term O₃-attributable mortality reflects consideration for evidence supporting a likely 27 to be a causal relationship for long-term O_3 -attributable respiratory effects, including mortality 28 (O₃ ISA, section 2.5.2, U.S. EPA, 2013a). After considering its strengths and weaknesses, we 29 consider the Jerrett et al. (2009) study to be an appropriate basis for estimating long-term O_3 -30 related respiratory mortality risk. Key strengths of this study are that it (a) included 1.2 million 31 participants in the American Cancer Society cohort from all 50 states, DC, and Puerto Rico; 32 included O_3 data from 1977 (5 years before enrollment in the cohort began) to 2000; (b) 33 considered co-pollutant models that controlled for $PM_{2.5}$; and (c) explored the potential for a 34 threshold concentration associated with the long-term mortality endpoint. Importantly, this study

¹⁹ With application of a regional prior within Bayesian adjustment, city-specific effect estimates are adjusted towards the regional value rather than a national value as is the case with the application of a national prior.

1 was also the first to explore the relationship between long-term O₃ exposure and respiratory

- 2 mortality (rather than focusing on cardiopulmonary mortality). Key limitations are possible
- 3 exposure misclassification and uncontrolled confounding by temperature, which are endemic to

4 most long-term epidemiological studies. While Jerrett et al. (2009) found negative associations

5 between O_3 exposure and cardiovascular mortality when controlling for $PM_{2.5}$, null or negative

- 6 associations for O_3 are consistent with the evidence that $PM_{2.5}$ is the pollutant most strongly
- 7 associated with cardiovascular disease (EPA 2009 PM ISA).
- 8 Our analysis includes a core estimate based on a co-pollutant model (with PM_{2.5}). The 9 seasonal average metrics used in the long-term exposure mortality estimate are not very sensitive 10 to the reduced number of days with co-pollutant monitoring, and as such it is appropriate to 11 include the copollutant model as the core estimate. We also include two sensitivity analyses for
- 12 long-term O₃-attributable respiratory mortality including: (a) application of regionally-

13 differentiated effect estimates (although these do not include a copollutants model specification)

14 and (b) application of a single pollutant (O₃-only) national-based effect estimate.

15 The effect estimates used in modeling long-term O₃-attributable mortality (see Table 7-2) 16 utilize a seasonal average of peak (1hr maximum) measurements. These long-term exposure 17 metrics can be viewed as long-term exposures to daily peak O₃ over the warmer months, as 18 compared with annual average levels such as are used in long-term PM exposure calculations. 19 This increases the need for care in interpreting these long-term O₃-attributable mortality

20 estimates together with the short-term O₃-attributable mortality estimates, in order to avoid

21 double counting. It is also important to keep in mind that our estimates of short-term O_3 -

22 attributable mortality are for all-causes, while estimates of long-term O_3 -attributable mortality 23 are focused on respiratory-related mortality. This further limits the ability to compare estimates

24 of long-term and short-term exposure related mortality.

25 Once the set of epidemiology studies described above was selected, the next step was to 26 specify C-R functions for use in the risk assessment. Several factors were considered in 27 identifying the effect estimates and model forms used in specifying C-R functions for each 28 endpoint. These factors are described below:

29

O₃ Exposure Metric: In the risk assessment supporting the previous O₃ NAAQS review,
 for short-term exposure, we included C-R functions based on 24hr averages as well as a
 number of peak O₃ measurements. However, given that the the current O₃ NAAQS
 standard uses an 8hr form and given that many of the clinical studies involving O₃ also
 utilize shorter exposures (on the order of 2 to 8 hrs – see O₃ ISA, section 6.2.1.1), we
 wanted to see if the latest epidemiological studies for O₃ also supported use of an 8hr
 averaging time in modeling risk. Several epidemiological studies completed since the last

review provide limited support for stronger associations between health endpoints and
peak O_3 metrics (i.e., 1hr maximum, 8hr maximum and 8hr means) relative to 24hr
averages. Specifically, a study of respiratory ED visits in Atlanta (Darrow et al., 2011)
found stronger associations with peak metrics (including 1hr and 8hr maximum
measurements) compared with 24hr averages (see O ₃ ISA section 6.2.7.3 and Figure 6-
17, U.S. EPA, 2013a). Similarly, for short-term exposure-related mortality, there are also
a limited number of epidemiologic studies that have compared mortality associations
with peak O_3 metrics and the 24hr average metric. Although the O_3 ISA recognizes that
24hr exposure metrics when used in time series studies may result in smaller risk
estimates, ultimately it concludes that "Overall, the evidence from time-series and panel
epidemiologic studies does not indicate that one exposure metric is more consistently or
strongly associated with mortality or respiratory-related health effects" (U.S. EPA,
2013a, section 2.5.4.2). Based on consideration for the evidence summarized in the O_3
ISA, we have decided to focus on peak exposure metrics because of the limited evidence
that these metrics may be associated with higher risk estimates relative to the 24 hr
exposure metric. However, we recognize that, as summarized in the O_3 ISA, there is only
weak support for differentiating between these two categories of short-term exposure
metric.

Epidemiological study		Location					
(stratified by O_3 -		(urban study	Exposure metric				
attributable health	Health	area(s)	(and modeling	Additional study design details	Notes recording application in the analysis		
enupoinis)	enupoints	covereu)	Sh ant tanu (Additional study design details	Notes regarding appreation in the analysis		
Smith at al. 2000	Non	05 lange unber	Snort-term	A directing for time remains	Ecourad on the the may based matrix C.D.		
Smith et al., 2009	accidental, respiratory, cardiovascul ar	95 large urban communities (provides coverage for all 12 urban study areas)	24nr avg, snr max, 1hr max. April through October and all year	Adjusting for time-varying confounders (PM, weather, seasonality). Lag structure included 0, 1, 2 and day 3 lag as well as 0-6 day distributed lag. Age range: all ages.	Focused on the Shr max-based metric C-R functions for the REA (see text discussion later in this section). Obtained Bayes-adjusted city-specific effect estimates for non-accidental all-cause mortality from Dr. Smith (personal communication, Dr. Richard L. Smith, January 15, 2013) reflecting consideration for the following modeling elements: (a) regional- versus national-prior Bayes model adjustment, (b) single pollutant versus copollutants (PM_{10}) models, and (c) full monitoring period versus summer only (April-October). For the core analysis, we focused on the single pollutant (O_3 - only) model covering the full monitoring period		
					The copollutants model (with PM_{10}) was included as a sensitivity analysis (see section 7.4.3).		
Zanobetti and Schwartz (2008)	Non- accidental, respiratory, cardiovascul ar	48 U.S. cities (provides coverage for the 12 urban study areas)	8hr max. June- August	Effect controlled for season, day of week, and temperature. Lag structure included 0-3d, 0-20 and 4-20 day). Age range: all ages	Obtained Bayes-adjusted city-specific effect estimates for non-accidental, respiratory and cardiovascular from Dr. Zanobetti (personal communication, Dr. Antonella Zanobetti, January 5, 2012). These effect estimates reflect a 0-3 day distributed lag and are based on 8hr mean O_3 levels measured between June and August. Estimates were generated for each study area using this constrained warm-season period.		
	Short-term O_3 -attributable morbidity - HA for respiratory effect)						
Medina-Ramon et al., 2006.	HA: COPD, pneumonia	36 cities (provides coverage for all 12 urban study areas)	8hr mean. warm (May-September), cool (October- April), all year	Distributed lag (0-1 day). Age range: \geq 65yrs. Controlled for day of the week and weather (including temperature).	Generated risk estimates based on warm season for COPD only (May-September).		
Linn et al., 2000	HA: unscheduled	LA only	24hr mean, LA O_3 season (all year),	Lag 0. Age range: all ages. Used subgroup analysis to explore the	Included effect estimate based on 24hr avg metric (for summer) since this provided additional		

Table 7-2 Overview of Epidemiological Studies Used in Specifying C-R Functions

Epidemiological study (stratified by O ₃ - attributable health endpoints)	Health endpoints	Location (urban study area(s) covered)	Exposure metric (and modeling period)	Additional study design details	Notes regarding application in the analysis
	for pulmonary illness		winter, spring, summer and autumn	effect of temporal variation, weather and autocorrelation on O_3 effect.	coverage for HA in L.A. Modeled using air quality for June-August.
Lin et al., 2008	HA: respiratory disease	NY State (used to cover NYC)	1hr max (for 10am-6pm interval), warm season (April- October)	Lag 0, 1, 2, 3. Age range: <18yrs. Models adjusted for the confounding effects of demographic characteristics, particulate matter(PM_{10}), meteorological conditions, day of the week, seasonality, long- term trends, and different lag periods of exposure.	Used 1hr max metric applied to the city-specific O_3 season for NYC (April-October).
Katsouyanni et al 2009	HA: cardiovascul ar disease, chronic obstructive pulmonary disease, pneumonia, all respiratory	14 cities (provides coverage for Detroit only)	lhr max. Summer only and all year	Lag 0-1day. Age range: ≥ 65 yrs. Models accounted for seasonal patterns, but also, for weekend and vacation effects, and for epidemics of respiratory disease. The data were also analyzed to detect potential thresholds in the concentration–response relationships.	C-R function applied only for all respiratory endpoint. Used June-August-based composite monitor.
Silverman et al., 2010	HA: asthma (ICU and non-ICU)	NYC	8hr max. Warm season (April- August)	Includes control for $PM_{2.5}$. Lag 0-1 day. Age range: children 6-18yrs. The model adjusted for temporal trends, weather, and day of the week.	Applied C-R function (for O_3 and O_3 with control for PM _{2.5}) to the city-specific O_3 season for NYC (slightly longer than the modeling period used in the study).
		Short-to	erm O_3 -attributable n	norbidity– ED and ER visits (respirato	ry)
Ito et al., 2007	ED: asthma	NYC	8hr max. Warm season (April- September)	Includes models controlling for SO ₂ , NO ₂ , CO and PM _{2.5} . Lag: 0, 1, and distributed lag (0-1 day). Age range: all ages. Model adjusts for temporal trends, weather terms, day-of-week and other pollutants.	Applied C-R functions (for O_3 alone and $\overline{O_3}$ with control for listed pollutants) to the city-specific O_3 season for NYC (slightly longer than the modeling period used in the study).
Tolbert et al., 2007	ED: all respiratory	Atlanta	8hr max. Summer (March-October)	Includes models controlling for NO ₂ , CO, PM ₁₀ and NO ₂ / NO ₂ . Age range: all ages. Model controls for temporal trends, temperature.	Applied C-R functions (for O_3 alone and O_3 with control for listed pollutants) to the city-specific O_3 season for Atlanta.

Epidemiological study (stratified by O ₃ - attributable health endpoints)	Health endpoints	Location (urban study area(s) covered)	Exposure metric (and modeling period)	Additional study design details	Notes regarding application in the analysis	
				other pollutants.		
Strickland et al., 2010	ER: respiratory	Atlanta	8hr max (based on population weighted average across monitors). Warm season (May to October) and cool (November to April)	Lag: average of 0-2 day, distributed lag 0-7 day. Age range: 5-17yrs. Model controls for seasonal trends and meteorology.	Included effect estimates based on both lag structures and used composite monitor values for city-specific O_3 season.	
Darrow etl al., 2011	ED: all respiratory	Atlanta	8hr max, 1hr max, 24hr avg for summer (March- October).	Lag: 1day. Age range: all ages. The study used a time series analysis similar to case-crossover with crossover matching based on daily temperature (rather than day of the week) to provide control for this key risk-related factor.	Used city-specific O_3 season-based composite monitor values.	
		She	ort-term O_3 -attributal	ble morbidity – respiratory symptoms		
Gent et al., 2003	Respiratory symptoms: wheeze, persistent cough, chest tightness, shortness of breath	Springfield MA (study used to cover Boston)	1hr max, 8hr max	Lag: 0 and 1 day. Age range: asthmatic children <12 yrs. Model adjusted for temperature.	Included effect estimates for different symptoms based on both 8hr max and 1hr max metrics (for city-specific O_3 season composite monitor values for Boston). The study area (which focuses on Springfield and the northern portion of Connecticut) does not encompass Boston. However, we are willing to accept uncertainty associated with using effect estimates from this study to provide coverage for Boston given the goal of providing coverage for this morbidity endpoint. However, there is increased uncertainty associated with modeling for this endpoint.	
Long-term O_3 -attributable respiratory mortality						
Jerrett et al., 2009	Respiratory, cardiovascul ar, cardiopulmo nary	96 metropolitan statistical areas (provides coverage for all 12 study areas)	Seasonal average (i.e. Apr-Sep) of the peak daily 1hr max values.	>30 yrs of age, includes national- level and regional effect estimates (only national-level estimate has copollutants modeling considering PM2.5 along with O ₃). Modeling included consideration for a range of potential confounders evaluated	Included national copollutants model-based effect estimates in core analysis and single-pollutant model regional effect estimates and national effect estimates as sensitivity analyses.	

Epidemiological study (stratified by O ₃ - attributable health endpoints)	Health endpoints	Location (urban study area(s) covered)	Exposure metric (and modeling period)	Additional study design details	Notes regarding application in the analysis
1	•		• • •	at both the ecological level and	
				personal level.	

• Single-and Multi-pollutant Models (pertains to both short-term and long-term

2 *exposure studies*): Epidemiological studies often consider health effects associated with 3 ambient O₃ using both single-pollutant and co-pollutant models. To the extent that any of 4 the co-pollutants present in the ambient air may have contributed to health effects 5 attributed to O_3 in single pollutant models, risks attributed to O_3 may be overestimated or 6 underestimated if C-R functions are based on single pollutant models. This would argue 7 for inclusion of models reflecting consideration of co-pollutants. Conversely, in those 8 instances where co-pollutants are highly correlated with O_3 , inclusion of those pollutants 9 in the health impact model can produce unstable and statistically insignificant effect 10 estimates for both O₃ and the co-pollutants. Furthermore, there are often significant 11 differences in sampling frequencies for each pollutant included in copollutants models, 12 which can lead to a loss of statistical power in copollutants models (relative to single 13 pollutant models). These last points could argue for inclusion of a model based 14 exclusively on O₃. Given that single and multi-pollutant models each have potential 15 advantages and disadvantages, to the extent possible, given available information we 16 have included both types of C-R functions in the risk assessment.

17 Multiple Effect Estimates within a Given CBSA-based Study Area: As noted earlier 18 in section 7.1.1, for this analysis, all health endpoints, including short-term O_{3} -19 attributable mortality are modeled using CBSA-based study areas. In the case of both 20 Smith et al., 2009 and Zanobetti and Schwartz 2008, these CBSA-based study areas are 21 larger than the study areas used in these epidemiological studies to derive effect 22 estimates. Furthermore, for some of the CBSA-based urban study areas, several of the 23 smaller study areas evaluated in the epidemiological study fall within a single larger 24 CBSA-based study area. For example, with the Smith et al., 2009 study, multiple effect 25 estimates are available for the CBSA-defined study areas of Los Angeles and New York 26 City. Specifically, the Smith et al., 2009 study provides separate effect estimates for (a) 27 Santa Anna/Anaheim and Los Angeles study areas, both of which fall within the larger 28 CBSA-based Los Angeles study area and (b) New York, Jersey City and Newark study 29 areas, all of which fall within the larger CBSA-defined New York study area (see Table 30 7-3). This raises the question of how to specify the effect estimate for these larger CBSA-31 based study areas when there are multiple effect estimates available from the 32 epidemiological study. For this analysis, in those instances where there are multiple effect 33 estimates, we have decided to use the effect estimate that represents the largest number of 34 residents within each CBSA-based study area. There is uncertainty associated with this 35 decision which is discussed both in section 7.4.2 and section 7.5.3 (as part of the air 36 quality-related sensitivity analysis discussion).

Table 7-3 CBSA-based Study Areas with Multiple Effect Estimates from the Smith et al., 2009 Study*

CBSA	Smith et al., 2009 (smaller) study areas with CBSA-based	Population	Mortality effect	
Study Area	study area	totals	estimate	Comments
	New York, NY	9,100,000	0.0009	New York study area dominates from
	Jersey City, NJ	630,000	0.0001	a population standpoint, so that effect
New York City	Newark, NJ	780,000	0.0005	estimate was chosen to represent the entire CBSA. An additional 8.3 million people live in portions of the New York CBSA not covered by the Smith et al., 2009 study areas.
Los Angeles	Santa Ana/Anaheim, CA	3,000,000	0.0002	Los Angeles dominates from a population standpoint, so that effect estimate was chosen to represent the
Los Aligeles	Los Angeles, CA	9,800,000	0.0001	entire CBSA. In this case, the full CBSA-based study area is covered by the Smith et al., 2009-based subareas.

3

1

2

* Source: obtained from Dr. Smith (personal communication, Dr. Richard L. Smith, January 15, 2013)

4 Single-city Versus Multi-city Studies: All else being equal, we judge C-R functions 5 estimated in the assessment location as preferable to a function estimated in some other location, to avoid uncertainties that may exist due to differences associated with 6 7 geographic location. There are several advantages, however, to using estimates from 8 multi-city studies versus studies carried out in single cities. Multi-city studies are 9 applicable to a variety of settings, since they estimate a central tendency across multiple 10 locations. Multi-city studies also tend to have more statistical power and provide effect 11 estimates with relatively greater precision than single-city studies due to larger sample 12 sizes, reducing the uncertainty around the estimated health coefficient. By contrast, 13 single-city studies, while often having lower statistical power and varying study designs 14 which can make comparison across cities challenging, reflect location-specific factors 15 such as differences in underlying health status, and differences in O₃ exposure-related 16 factors such as air conditioner use and patterns of urban residential density. There is a 17 third type of study design that generates Bayes-adjusted city-specific effect estimates, 18 thereby combining the advantages of both city-specific and multi-city studies. Bayes-19 adjusted city-specific estimates begin with a city-specific effect estimate and shrink that 20 towards a multi-city mean effect estimate based on consideration for the degree of 21 variance in both estimates. We have elected to place greater confidence on these types of 22 Bayesian-adjusted effect estimates when they are available. Otherwise, given the 23 advantages for both city-specific and multi-city effect estimates, we have used both types 24 when available.

7-28

Multiple Lag Models: Based on our review of evidence provided in the O_3 ISA, we 1 2 believe there is increased confidence in modeling both short-term O₃-attributable 3 mortality and respiratory morbidity risk based on exposures occurring up to a few days 4 prior to the health effect, with less support for associations over longer exposure periods 5 or effects lagged more than a few days from the exposure (see O₃ ISA section 2.5.4.3, U.S. EPA, 2013a). Consequently, we have favored C-R functions reflecting shorter lag 6 7 periods (e.g., 0, 1 or 1-2 days). With regard to the specific lag structure (e.g., single day 8 versus distributed lags), the O_3 ISA notes that epidemiological studies involving 9 respiratory morbidity have suggested that both single day and multi-day average 10 exposures are associated with adverse health effects (see O₃ ISA section 2.5.4.3). 11 Therefore, when available both types of lag structures where considered in specifying C-12 R functions for short-term O₃-attributable mortality and morbidity. 13 Seasonally-differentiated Effects Estimates: The previous O₃ Air Quality Criteria 14 Document (AQCD) (published in 2006) concluded that aggregate population time-series studies demonstrate a positive and robust association between ambient O₃ concentrations 15 16 and respiratory-related hospitalizations and asthma ED visits during the warm season (see 17 O₃ ISA section 2.5.3.1 U.S. EPA, 2013a). The current O₃ ISA notes that recent studies of 18 short-term O₃-attributable respiratory mortality in the U.S. suggest that the effect is 19 strengthened in the summer season (O₃ ISA section 2.5.3.1, U.S. EPA, 2013a). In addition, many of the key epidemiological studies discussed in the current O₃ ISA 20

exploring both short-term exposure related mortality and morbidity have larger (and more statistically significant) effect estimates when evaluated for the summer (O_3) season,

21

22

- relative to the full year (see O₃ ISA Figures 6-20 and 6-27, U.S. EPA, 2013a). However,
 if we focus the assessment of risk on the warm season, we bias our estimate by excluding
 potential effects associated with cooler (non-summer) months. Given our desire to
 provide a more complete picture of overall risk in each of the study areas, we have
 favored (for the core analysis) effect estimates that cover the full O₃ monitoring period
 specific to each study area, rather than the more limited warm (summer) period.
- Shape of the Functional Form of the Risk Model (including threshold): The current
 O₃ ISA concludes that there is little support in the literature for a population threshold for
 short-term O₃-attributable effects. However, specifically in relation to mortality, the O₃
 ISA concludes that a national or combined analysis may not be appropriate to identify
 whether a threshold exists (see O₃ ISA, section 2.5.4.4, U.S. EPA, 2013a).²⁰ Given the

 $^{^{20}}$ Specifically, given the multi-city nature of these mortality studies combined with the variability in O₃ and other factors related to exposure and risk, the O₃ ISA concludes that these studies are not well positioned to evaluate the potential for a threshold in the mortality effect.

1	above general observation from the O ₃ ISA regarding the low potential for thresholds, we
2	did not include C-R functions for any of the short-term O ₃ -attributable health endpoints
3	modeled that incorporated a threshold. ²¹
4	Application of the above criteria resulted in an array of C-R functions specified for the
5	risk assessment (see Table 7-2), including functions covering short-term O ₃ -attributable
6	mortality and morbidity and long-term O3-attributable mortality. In presenting the C-R functions
7	in Table 7-2, we have focused on describing key attributes of each C-R function (and associated
8	source epidemiological study) relevant to a review of their use in the risk assessment. More
9	detailed technical information including effect estimates and model specification is provided in
10	Appendix 7A. Specific summary information provided in Table 7-2 includes:
11	• <i>Health endpoints:</i> identifies the specific endpoints evaluated in the study. Generally
12	we included all of these in our risk modeling, however, when a subset was modeled,
13	we reference that in the "Notes" column (last column in the table).
14	• Location: identifies the specific urban areas included in the study and maps those to
15	the set of 12 urban study areas included in the risk assessment.
16	• <i>Exposure metric</i> : describes the exposure metric used in the study, including the
17	specific modeling period (e.g., O3 season, warm season, full year). We developed two
18	categories of composite monitor values to match the modeling periods used in the two
19	short-term O ₃ -attributable mortality studies providing C-R functions for the analysis.
20	For the remaining morbidity endpoints, we mapped specific C-R functions to
21	whichever of these two composite monitor categories most closely matched the
22	modeling period used in the underlying epidemiological study. This mapping (for
23	morbidity endpoint C-R functions) is described in the "Notes" column (the seasons
24	reflecting in modeling for each C-R function are also presented in Appendix 7A).
25	• Additional study design details: this column provides additional information primarily
26	covering the lag structure and age ranges used in the study.
27	• Notes regarding application in second draft analysis: as the name implies, this
28	column provides notes particular to the application of a particular epidemiological
29	study and associated C-R functions in the risk assessment.

²¹ While clinical studies have suggested the presence of a threshold for respiratory effects, these should not be used to support specification of population-level thresholds for use in the epidemiological-based risk assessment. The clinical studies focus on relatively small and clearly defined populations of healthy adults which are not representative of the broader residential populations typically associated with epidemiological studies, including older individuals and individuals with existing health conditions which place them at greater risk for O₃-related effects. Therefore, the clinical studies are unlikely to have the power to capture population thresholds in a broader and more diverse urban residential population, should those thresholds exist.

1 7.3.3 Baseline Health Effect Incidence and Prevalence Data

2 As discussed earlier (section 7.1.2), the most common epidemiological-based health risk 3 model expresses the change in health risk (Δy) associated with a given change in O₃ 4 concentrations (Δx) as a percentage of the baseline incidence (y). To accurately assess the impact 5 of O₃ air quality on health risk in the selected urban areas, information on the baseline incidence 6 of health effects (i.e., the incidence under recent air quality conditions) in each location is 7 needed. In some instances, health endpoints are modeled for a population with an existing health 8 condition, necessitating the use of a prevalence rate. Where at all possible, we use county-9 specific incidences or incidence rates (in combination with county-specific populations). In some 10 instances, when county-level incidence rates were not available, BenMAP can employ more 11 generalized regional rates (see BenMAP Guidance Manual for additional detail, Abt Associates, 12 Inc. 2010). For prevalence rates (which were only necessary for modeling respiratory symptoms 13 among asthmatic children using Gent et al., (2008) - see Table 7-2), we utilized a national-level 14 prevalence rate appropriate for the age group being modeled. A summary of available baseline 15 incidence data for specific categories of effects (and prevalence rates for asthma) is presented 16 below: 17 • Baseline incidence data on mortality: County-specific (and, if desired, age- and race-18 specific) baseline incidence data are available for all-cause and cause-specific mortality from CDC Wonder.²² The most recent year for which data are available 19 online is 2005 and this was the source of incidence data for the risk assessment.²³ 20 21 • Baseline incidence data for hospital admissions and emergency room (ER) visits: 22 Cause-specific hospital admissions baseline incidence data are available for each of 23 40 states from the State Inpatient Databases (SID). Cause-specific ER visit baseline incidence data are available for 26 states from the State Emergency Department 24 25 Databases (SEDD). SID and SEDD are both developed through the Healthcare Cost 26 and Utilization Project (HCUP), sponsored by the Agency for Healthcare Research and Quality (AHRQ). In addition to being able to estimate State-level rates, SID and 27 28 SEDD can also be used to obtain county-level hospital admission and ER visit counts 29 by aggregating the discharge records by county. 30 • Asthma prevalence rates: state-level prevalence rates that are age group stratified are 31 available from the Centers for Disease Control and Prevention (CDC) Behavioral 32 Risk Factor Surveillance System (BRFSS) (U.S. CDC, 2010).

²² <u>http://wonder.cdc.gov/mortsql.html</u>

²³ Note: For years 1999 – 2005, CDC Wonder uses ICD-10 codes; for years prior to 1999, it uses ICD-9 codes. Since most of the studies use ICD-9 codes, this means that EPA will have to create or find a mapping from ICD-9 codes to ICD-10 codes if the most recent data available are to be used.

Incidence and prevalence rates are presented as part of the full set of model inputs
 documented in Appendix 7A. The incidence rates and prevalence rates provided in Table 7A-1
 are weighted average values for the age group associated with each of the C-R functions. These
 weighted averages are calculated within BenMAP using more refined age-differentiated
 incidence and prevalence rates originally obtained from the data sources listed in the bullets
 above.

7 7.3.4 Population (demographic) Data

8 To calculate baseline incidence rates, in addition to the health baseline incidence data we 9 also need the corresponding population. We obtained population data from the 2010 U.S. Census 10 (<u>http://www.census.gov/popest/counties/asrh/</u>). These data are then used as the basis for back-11 casting estimates for simulation years (in this case, 2007 and 2009) (see Appendix J of the 12 BenMAP User's Manual for additional detail, U.S. EPA, 2012b). Total population counts used in 13 modeling each of the health endpoints evaluated in the analysis (differentiated by urban study 14 area and simulation year) are provided as part model inputs presented in Appendix 7A.

15 7.4 ADDRESSING VARIABILITY AND UNCERTAINTY

16 An important component of a population risk assessment is the characterization of both 17 uncertainty and variability. Variability refers to the heterogeneity of a variable of interest within 18 a population or across different populations. For example, populations in different regions of the 19 country may have different behavior and activity patterns (e.g., air conditioning use, time spent 20 indoors) that affect their exposure to ambient O_3 and thus the population health response. The 21 composition of populations in different regions of the country may vary in ways that can affect 22 the population response to exposure to $O_3 - e.g.$, two populations exposed to the same levels of 23 O_3 might respond differently if one population is older than the other. Variability is inherent and 24 cannot be reduced through further research. Refinements in the design of a population risk 25 assessment are often focused on more completely characterizing variability in key factors 26 affecting population risk - e.g., factors affecting population exposure or response - in order to 27 produce risk estimates whose distribution adequately characterizes the distribution in the 28 underlying population(s). 29 Uncertainty refers to the lack of knowledge regarding the actual values of inputs to an 30 analysis. Models are typically used in analyses, and there is uncertainty about the true values of

31 the parameters of the model (parameter uncertainty) – e.g., the value of the coefficient for O_3 in a

32 C-R function. There is also uncertainty about the extent to which the model is an accurate

33 representation of the underlying physical systems or relationships being modeled (model

34 uncertainty) – e.g., the shapes of C-R functions. In addition, there may be some uncertainty

surrounding other inputs to an analysis due to possible measurement error—e.g., the values of daily O_3 concentrations in a risk assessment location, or the value of the baseline incidence rate for a health effect in a population.²⁴ In any risk assessment, uncertainty is, ideally, reduced to the maximum extent possible through improved measurement of key variables and ongoing model refinement. However, significant uncertainty often remains, and emphasis is then placed on characterizing the nature of that uncertainty and its impact on risk estimates. The characterization of uncertainty can be both qualitative and, if a sufficient knowledgebase is available,

8 quantitative.

9 The selection of urban study areas for the O_3 risk assessment was designed to cover the 10 range of O_3 -related risk experienced by the U.S. population and, in general, to adequately reflect 11 the inherent variability in those factors affecting the public health impact of O_3 exposure. 12 Sources of variability reflected in the risk assessment design are discussed in section 7.4.1, along 13 with a discussion of those sources of variability which are not fully reflected in the risk

14 assessment and consequently introduce uncertainty into the analysis.

15 The characterization of uncertainty associated with risk assessment is often addressed in 16 the regulatory context using a tiered approach in which progressively more sophisticated 17 methods are used to evaluate and characterize sources of uncertainty depending on the overall 18 complexity of the risk assessment (WHO, 2008). Guidance documents developed by EPA for 19 assessing air toxics-related risk and Superfund Site risks (U.S.EPA, 2004 and 2001, respectively) 20 as well as recent guidance from the World Health Organization (WHO, 2008) specify multi-

21 tiered approaches for addressing uncertainty.

22 The WHO guidance, in particular, presents a four-tiered approach for characterizing 23 uncertainty (see Chapter 3, section 3.2.6 for additional detail on the four tiers included in the 24 WHO's guidance document). With this four-tiered approach, the WHO framework provides a 25 means for systematically linking the characterization of uncertainty to the sophistication of the 26 underlying risk assessment. Ultimately, the decision as to which tier of uncertainty 27 characterization to include in a risk assessment will depend both on the overall sophistication of 28 the risk assessment and the availability of information for characterizing the various sources of 29 uncertainty. We used the WHO guidance as a framework for developing the approach used for 30 characterizing uncertainty in this risk assessment. 31

- The overall analysis in the O_3 NAAQS risk assessment is relatively complex, thereby
- 32 warranting consideration of a full probabilistic (WHO Tier 3) uncertainty analysis. However,
- 33 limitations in available information prevent this level of analysis from being completed at this

²⁴ It is also important to point out that failure to characterize variability in an input used in modeling can also introduce uncertainty into the analysis. This reflects the important link between uncertainty and variability with the effort to accurately characterize variability in key model inputs actually reflecting an effort to reduce uncertainty about population means and population variability.

1 time. In particular, the incorporation of uncertainty related to key elements of C-R functions 2 (e.g., competing lag structures, alternative functional forms, etc.) into a full probabilistic WHO 3 Tier 3 analysis would require that probabilities be assigned to each competing specification of a 4 given model element (with each probability reflecting a subjective assessment of the probability 5 that the given specification is the "correct" description of reality). However, for many model 6 elements there is insufficient information on which to base these probabilities. One approach that 7 has been taken in such cases is expert elicitation; however, this approach is resource- and time-8 intensive and consequently, it was not feasible to use this technique in the current O_3 NAAQS 9 review to support a WHO Tier 3 analysis.²⁵

10 For most elements of this risk assessment, rather than conducting a full probabilistic 11 uncertainty analysis, we have included qualitative discussions of the potential impact of 12 uncertainty on risk results (WHO Tier1). As discussed in section 7.1.1, for this draft of the risk 13 assessment, we have also expanded the sensitivity analysis considerably to cover a range of 14 model elements (this represents a WHO Tier 2 analysis). The specific modeling elements 15 covered in the sensitivity analysis for each health effects endpoint together with the specification 16 of the core analysis is presented in section 7.4.3. As part of the sensitivity analysis, we have also completed an influence analysis using estimated elasticities of response²⁶ designed to determine 17 18 which of the input factors used in calculating risk are primarily responsible for inter-city 19 variability in risk. This influence analysis focuses on the response of core short-term exposure-20 related mortality risk to inputs since this is one of the key risk metrics completed for the REA 21 (see section 7.4.3).

22 In addition to the qualitative and quantitative treatment of uncertainty and variability 23 which are described here, we have also completed an analysis to evaluate the representativeness 24 of the selected urban study areas against national distributions for key O₃ risk-related attributes 25 to determine whether they are nationally representative or more focused on a particular portion 26 of the distribution for a given attribute (see Chapter 8, section 8.2.1). In addition, we have 27 completed a second analysis addressing the representativeness issue, which identified where the 12 urban study areas included in this risk assessment fall along a distribution of national-level 28 29 short-term and long-term exposure-related mortality risk (see Chapter 8, section 8.2.2). This 30 analysis allowed us to assess the degree of which the 12 urban study areas capture locations 31 within the U.S. likely to experience elevated levels of risk related to O_3 exposure (for both short-32 term and long-term O₃-attributable mortality).

²⁵ While a full probabilistic uncertainty analysis was not completed for this risk assessment, we were able to use confidence intervals associated with effects estimates (obtained from epidemiological studies) to incorporate statistical uncertainty associated with sample size considerations in the presentation of risk estimates.

²⁶ Elasticities are a measure of sensitivity calculated as the percent change in the response variable for a one percent change in the input variable.

1 The remainder of this section is organized as follows. Key sources of variability which 2 are reflected in the design of the risk assessment, along with sources excluded from the design, 3 are discussed in section 7.4.1. A qualitative discussion of key sources of uncertainty associated 4 with the risk assessment (including the potential direction, magnitude and degree of confidence 5 associated with our understanding of the source of uncertainty – the knowledge base) is 6 presented in section 7.4.2. The design of the core analysis and sensitivity analysis completed for 7 each of the health effect endpoint categories modeled in the risk assessment is discussed in 8 section 7.4.3.

9 7.4.1 Treatment of Key Sources of Variability

10 The risk assessment was designed to cover the key sources of variability related to 11 population exposure and exposure response, to the extent supported by available data. Here, the 12 term key sources of variability refers to those sources that we believe have the potential to play 13 an important role in impacting population incidence estimates generated for this risk assessment. 14 Specifically, have have concluded that these sources of variability, if fully addressed and 15 integrated into the analysis, could result in adjustments to the core risk estimates which might be 16 relevant from the standpoint of interpreting the risk estimates in the context of the O₃ NAAQS 17 review. The identification of sources of variability as "key" reflects consideration for sensitivity 18 analyses conducted for previous O₃ NAAQS risk assessments, which have provided insights into 19 which sources of variability can influence risk estimates, as well as information presented in the 20 O₃ ISA.

21 As with all risk assessments, there are sources of variability which have not been fully 22 reflected in the design of the risk assessment and consequently introduce a degree of uncertainty 23 into the risk estimates. While different sources of variability were captured in the risk 24 assessment, it was generally not possible to separate out the impact of each factor on population 25 risk estimates, since many of the sources of variability are reflected collectively in a specific 26 aspect of the risk model. For example, inclusion of urban study areas from different regions of 27 the country likely provides some degree of coverage for a variety of factors associated with O_3 28 risk (e.g., air conditioner use, differences in population commuting and exercise patterns, 29 weather). However, the model is not sufficiently precise or disaggregated to allow the individual 30 impacts of any one of these sources of variability on the risk estimates to be characterized. 31 Key sources of potential variability that are likely to affect population risks are discussed 32 below, including the degree to which they are captured in the design of the risk assessment:

- 33
- 34
- 35

• Heterogeneity in the Effect of O₃ on Health Across Different Urban Areas: A number of studies cited in the O₃ ISA have found evidence for regional heterogeneity in the short-term O₃-attributable mortality effect (Smith et al., 2009 and Bell and

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1	Dominici, 2008, Bell et al., 2004, Zanobetti an Schwartz 2008 – see O ₃ ISA section
2	6.6.2.2, U.S. EPA, 2013a). These studies have demonstrated that differences in effect
3	estimates between cities can be quite substantial (see O_3 ISA Figures 6-32 and 6-33).
4	Therefore, for the short-term O3-attributable mortality endpoint modeled using Smith
5	et al., 2009-based effect estimates, we have included Bayes-adjusted city-specific
6	effect estimates reflecting application of both a regional- and national-prior, both of
7	which are intended to capture cross-city differences in effect estimates for the
8	mortality endpoint, while still reflecting input from the more stable regional, or
9	national-level signal. The national-prior based estimates are included in the core
10	analysis since they have greater overall power, while the regional-prior based
11	estimates are included as sensitivity analyses to explore the impact of using regional
12	prior in developing the Bayes-adjusted estimates (see section 7.4.3). ²⁷ For short-term
13	morbidity endpoints, typically we have used city-specific effect estimates; however,
14	for most endpoints, we only have estimates for a subset of the urban study areas
15	(typically NYC, Atlanta and/or LA). Therefore, although our risk estimates do reflect
16	the application of city-specific effect estimates, because we do not have estimates for
17	all 12 urban study areas, we do not provide comprehensive coverage for
18	heterogeneity in modeling the respiratory morbidity endpoint category. Long-term
19	O ₃ -attributable mortality has been shown to demonstrate regional heterogeneity.
20	Specifically, Jerrett et al., 2009 presented regional effect estimates that demonstrated
21	considerable heterogeneity ranging from essentially a no-effect (for the Northeast and
22	Industrial Midwest) to effects substantially larger than the national effect (Southeast,
23	Southwest and Upper Midwest) (see Table 4 in Jerrett et al., 2009). There are many
24	potential explanations for regional heterogeneity including differences in O ₃ -
25	attributable factors and potential confounding, potential for the presence of (and
26	regional differences in) averting behavior, and variation in sample sizes which can
27	impact stability of effect estimates. For the core analysis, we use a national effect
28	estimate in modeling long-term exposure related mortality. Consideration of regional
29	effect estimates are included as a sensitivity analysis (see section 7.4.3 and 7.5.3).
30	• Exposure Measurement Error Associated with O ₃ Effect Estimates: Exposure
31	measurement error refers to uncertainty associated with using ambient monitor based
32	exposure surrogate metrics to represent the actual exposure of an individual or
33	population. As such, this factor can be an important contributor to variability in
34	epidemiological study results across locations, and uncertainty in results for any

²⁷ Note, that in some instances, there may be insufficient variance between cities to generate city-specific estimates using a regional prior, which compromises their use in the core analysis.

- 1 specific city (O_3 ISA, p. 1xii). Exposure measurement error can result from a number 2 of factors (e.g., central site monitors not representing actual patterns of personal 3 exposure including activity patterns, presence of non-ambient sources of exposure for 4 the pollutant of interest) (O_3 ISA, 1xii). These factors can vary across urban study 5 areas (and even within urban study areas), thereby contributing to differences in the 6 nature and magnitude of exposure measurement error across locations and ultimately 7 to differences in effect estimates and associated confidence levels. Exposure 8 measurement error is related to heterogeneity in effect estimates, since regional 9 differences in effect estimates can result in part, from differences in exposure 10 measurement error as noted here.
- 11 • Intra-urban Variability in Ambient O₃ Levels: The picture with regard to within 12 city variability in ambient O_3 levels and the potential impact on epidemiologic-based 13 effect estimates is somewhat more complicated. The O₃ ISA notes that spatial 14 variability in O₃ levels is dependent on spatial scale with O₃ levels being more 15 homogeneous over a few kilometers due to the secondary formation nature of O_3 , 16 while levels can vary substantially over tens of kilometers. Community exposure may 17 not be well represented when monitors cover large areas with several subcommunities 18 having different sources and topographies as exemplified by Los Angeles which 19 displays significantly greater variation in inter-monitor correlations than does, for 20 example, Atlanta or Boston (see O₃ ISA section 4.6.2.1 U.S. EPA 2013a). Despite the 21 potential for substantial variability across monitors the O₃ ISA notes that studies have 22 tended to demonstrate that monitor selection has only a limited effect on the 23 association of short-term O_3 exposure with health effects. The likely explanation for 24 this is that, while absolute values for a fixed point in time can vary across monitors in 25 an urban area, the temporal patterns of O_3 variability across those same monitors 26 tends to be well correlated. Given that most of the short-term O_3 -attributable O_3 27 epidemiological studies are time series in nature, the O_3 ISA notes that the stability of 28 temporal profiles across monitors within most urban areas means that monitor 29 selection will have little effect on the outcomes of an epidemiological study 30 examining short-term O₃-attributable mortality or morbidity (see O₃ ISA section 4.6.2.1 U.S. EPA 2013a). For this reason, we conclude that generally intra-city 31 32 heterogeneity in O₃ levels is not a significant factor likely to impact estimates of 33 short-term O₃-attributable risk. One exception is LA which, due to its size and 34 variation in O_3 sources and other factors impacting O_3 patterns such as topography, 35 displays significant variation in ambient O₃ levels with a subsequent impact on risk. 36 However, in the case of LA (as with the other urban study areas), we model risk using

- composite monitors which do not provide spatially-differentiated representations of 1 2 exposure and consequently, we do not address this source of variability in the risk 3 assessment. As discussed in the uncertainty section, short-term exposure mortality 4 effect estimates for the New York CBSA (Smith et al., 2009) display significant 5 variability. However, it is not clear which factors are primarily responsible for this 6 heterogeneity (e.g., differences in the urban structure, residential behavior, or ambient 7 O_3 levels within the CBSA). The potential for intra-city heterogeneity in O_3 levels to 8 affect risk is more pronounced with long-term O₃-attributable mortality where the 9 relationship between annual trends in ambient O_3 (as represented using composite 10 monitor values) and annual mortality is compared between urban study areas in order 11 to derive effect estimates. Here, pronounced heterogeneity in O₃ levels within a given 12 city can result in exposure misclassification, if that heterogeneity is not well 13 represented by the composite monitor for that city. Different degrees of exposure 14 misclassification across urban study areas can introduce uncertainty into the overall 15 national-level effect estimate for long-term exposure-related mortality. Furthermore, 16 if that exposure measurement error has a regional trend, then measurement error can 17 potentially result in apparent regional heterogeneity in the effect estimates. The 18 degree to which there is true regional heterogeneity is made uncertain by the presence 19 of differential measurement error across regions. 20 Variability in the Patterns of Ambient O₃ Reduction Across Urban Areas: The •
- 21 simulated patterns of ambient O₃ concentrations across an urban area can vary based 22 on the methodology used to adjust ambient O₃ concentrations to represent just 23 meeting the current or alternative suites of standards. For the 1st draft REA, we used 24 a statistical approach called the "quadratic rollback" method for simulating just 25 meeting the current O₃ standard. Although the quadratic rollback method replicates 26 historical patterns of air quality changes better than some alternative methods, its 27 implementation relies on a statistical relationship instead of on a mechanistic 28 characterization of physical and chemical processes in the atmosphere. Because of its 29 construct as a statistical fit to measured O_3 values, the quadratic rollback technique 30 cannot capture spatial and temporal heterogeneity in O₃ response and also cannot 31 account for nonlinear atmospheric chemistry that causes increases in O_3 as a result of 32 emissions reductions of certain O_3 precursors under some circumstances. As noted in 33 section 7.1.1, for this draft of the REA, we have employed a model-based O₃ 34 adjustment methodology in the risk assessment for simulating O_3 concentrations 35 under current and alternate standard levels. Use of this model-based approach allows 36 the risk assessment results to more fully account for non-linearities in O_3 formation

1 and to reflect spatial and temporal heterogeneity in O_3 response, including NOx 2 titration conditions under which a reduction in NOx causes an increase in O_3 3 concentrations, in some core urban locations. 4 Demographics and Socioeconomic-status (SES)-related Factors: Variability in • 5 population density, particularly in relation to elevated levels of O₃ has the potential to 6 influence population risk, although the significance of this factor also depends on the 7 degree of intra-urban variation in O₃ levels (as discussed above). In addition, 8 community characteristics such as pre-existing health status, ethnic composition, SES 9 and the age of housing stock (which can influence rates of air conditioner use thereby 10 impacting rates of infiltration of O₃ indoors) can contribute to observed differences in 11 O₃-related risk (discussed in O₃ ISA – section 2.5.4.5, U.S. EPA, 2013a). Some of the 12 heterogeneity observed in effect estimates between cities in the multicity studies may 13 be due to these community characteristics, and while we cannot determine how much 14 of that heterogeneity is attributable to these factors, the degree of variability in effect

16 variability in these community characteristics. 17 Baseline Incidence of Disease: We collected baseline health effects incidence data • 18 (for mortality and morbidity endpoints) from a number of different sources (see 19 section 7.3.4). Often the data were available at the county-level, providing a relatively 20 high degree of spatial refinement in characterizing baseline incidence given the 21 overall level of spatial refinement reflected in the risk assessment as a whole. 22 Otherwise, for urban study areas without county-level data, either (a) a surrogate 23 urban study area (with its baseline incidence rates) was used, or (b) less refined state-

estimates between cities in our analysis should help to capture some of the latent

- 24 level or national incidence rate data were used.
- 25

15

7.4.2 Qualitative Assessment of Uncertainty

26 As noted in section 7.4, we have based the design of the uncertainty analysis carried out 27 for this risk assessment on the framework outlined in the WHO guidance document (WHO, 28 2008). That guidance calls for the completion of a Tier 1 qualitative uncertainty analysis, 29 provided the initial Tier 0 screening analysis suggests there is concern that uncertainty associated 30 with the analysis is sufficient to significantly impact risk results (i.e., to potentially affect 31 decision making based on those risk results). Given previous sensitivity analyses completed for 32 prior O₃ NAAQS reviews, which have shown various sources of uncertainty to have a potentially 33 significant impact on risk results, we believe that there is justification for conducting a Tier 1 34 analysis.

1 For the qualitative uncertainty analysis, we have described each key source of uncertainty 2 and qualitatively assessed its potential impact (including both the magnitude and direction of the 3 impact) on risk results, as specified in the WHO guidance. Similar to our discussion of 4 variability in the last section, the term key sources of uncertainty refers to those sources that the 5 we believe have the potential to play an important role in impacting population incidence 6 estimates generated for this risk assessment (i.e., these sources of uncertainty, if fully addressed 7 could result in adjustments to the core risk estimates which might impact the interpretation of 8 those risk estimates in the context of the O_3 NAAQS review). These key sources of uncertainty 9 have been identified through consideration for sensitivity analyses conducted for previous O₃ 10 NAAQS risk assessments, together with information provided in the final O₃ ISA and comments 11 provided by CASAC on the analytical plan for the risk assessment.

Table 7-4 includes the key sources of uncertainty identified for the O_3 REA. For each source of uncertainty, we have (a) provided a description, (b) estimated the direction of influence (over, under, both, or unknown) and magnitude (low, medium, high) of the potential impact of each source of uncertainty on the risk estimates, (c) assessed the degree of uncertainty (low, medium, or high) associated with the knowledge-base (i.e., assessed how well we understand each source of uncertainty), and (d) provided comments further clarifying the qualitative assessment presented.

19 The categories used in describing the potential magnitude of impact for specific sources 20 of uncertainty on risk estimates (i.e., low, medium, or high) reflect our consensus on the degree 21 to which a particular source could produce a sufficient impact on risk estimates to influence the interpretation of those estimates in the context of the O₃ NAAQS review.²⁸ Sources classified as 22 23 having a "low" impact would not be expected to impact the interpretation of risk estimates in the 24 context of the O₃ NAAQS review; sources classified as having a "medium" impact have the 25 potential to change the interpretation; and sources classified as "high" are likely to influence the 26 interpretation of risk in the context of the O₃ NAAQS review. Because this classification of the 27 potential magnitude of impact of sources of uncertainty is not based on our direct quantitative 28 assessments, we use qualitative judgments, in some cases informed by other relevant quantitative 29 analyses. Therefore, the results of the qualitative analysis of uncertainty are not useful for 30 making quantitative estimates of confidence, e.g. probabilistic statements about risk. However, 31 they can be used to support the interpretation of the risk estimates, including the assessment of 32 overall confidence in the risk estimates. In addition, they can also be used in guiding future 33 research to reduce uncertainty related to O₃ risk assessment. As with the qualitative discussion of

²⁸ For example, if a particular source of uncertainty were more fully characterized (or if that source was resolved, potentially reducing bias in a core risk estimate), could the estimate of incremental risk reduction in going from the current to an alternative standard level change sufficiently to produce a different conclusion regarding the magnitude of that risk reduction in the context of the O₃ NAAQS review?

- 1 sources of variability included in the last section, the characterization and relative ranking of
- 2 sources of uncertainty addressed here is based on our consideration of information provided in
- 3 previous O₃ NAAQS risk assessments (particularly past sensitivity analyses), the results of risk
- 4 modeling completed for the current O₃ NAAQS risk assessment and information provided in the
- 5 third draft O_3 ISA as well as earlier O_3 Criteria Documents. Where appropriate, in Table 7-4, we
- 6 have included references to specific sources of information considered in arriving at a ranking
- 7 and classification for a particular source of uncertainty.

		Potential influence of			
		uncertainty on risk		Knowledge-	Comments
		estima	ates	Base	(KB: knowledge base, INF: influence of uncertainty on risk
Source	Description	Direction	Magnitude	uncertainty*	estimates)
A. Adjustment of recent air quality measurements of O_3 to simulate attainment of both existing and alternative standard levels	See Chapter 4 for details	Both	Low- Medium	Low-medium	See Chapter 4 for more details (specific call-outs to be added)
B. Use of CBSA-based study areas in modeling risk (i.e., potential mismatch between study areas used in the REA and study areas used in the epidemiological studies providing the effect estimates used in modeling health effect endpoints)	If the set of monitors used in a particular urban study area to characterize population exposure as part of an ongoing risk assessment do not match the ambient monitoring data used in the original epidemiological study, then uncertainty can be introduced into the risk estimates. This uncertainty is balanced in part by the reduction in bias that results from using the expanded CBSA definition. (See section 7.1.1 for more details.) However, it should be noted that because these epidemiological studies occurred in the past, sometimes it can be impossible to exactly match the monitors used in the study using recent air quality data given that monitors may have moved to a different location or there may not be measurements available at specific monitors in the more recent time period.	Both	Low- medium	Low-medium	KB and INF: In modeling risk for the current draft of the REA, we used CBSA-based study areas for all health effect endpoints. As discussed in section 7.1.1, the use of the larger CBSA study areas allows us to better reflect how the change in air quality affects risk across the entire urban area and to avoid introducing known bias into the REA by focusing risk estimates on that subpopulation living in areas likely to experience potential increases in O_3 (and excluding the larger population of urban and suburban areas likely to experience reductions in ambient O_3 levels). While the use of the larger CBSA-based study areas addresses this source of known bias, it also introduces uncertainty into the REA since we are no longer matching the REA study areas to the study areas in the epidemiological studies providing the effect estimates used in modeling health effects endpoints. Given available data, it is not possible at this point to reliably characterize the degree of uncertainty introduced into the REA by having this mismatch in study areas. However, the potential bias avoided through the use of the larger CBSA study areas (with its acknowledged uncertainty) is substantial, as illustrated in the sensitivity analyses exploring spatial study area (see section 7.5.3).
C. Application of C-R functions based on a specific temporal and spatial pattern of correlations between O_3 monitors in an urban area (as reflected in the	The effect estimates used in this risk assessment reflect a specific spatial and temporal pattern of ambient O_3 (as represented by the particular monitoring network providing data for the underlying epidemiological study). However, if the spatial and	Both	Low- medium	Low-medium	KB and INF: With application of the HDDM adjustment approach, we simulate potential changes in the spatial and temporal pattern of O_3 for a study areas when just meeting the existing and alternative standards relative to patterns under recent conditions. This introduces uncertainty into the application of the original effect estimates, since the exposure surrogate represented by the composite monitor values may no longer match that of the underlying epidemiological study. However, it is not

Table 7-4 Summary of Qualitative Uncertainty Analysis of Key Modeling Elements in the O3 NAAQS Risk Assessment

		Potential influence of uncertainty on risk estimates		Knowledge- Base	Comments (KB: knowledge base, INF: influence of uncertainty on risk
Source	Description	Direction	Magnitude	uncertainty*	estimates)
epidemiological study providing the effect estimates) to a simulated change in the patterns of those correlations when we estimate risk in the REA.	temporal pattern of O_3 levels in the study areas being modeled differ significantly from the patterns in the original epidemiological study (for those same study areas), then uncertainty can be introduced into the risk estimates.				possible, given available data, to characterize quantitatively the magnitude of this uncertainty. This is probably most true in the urban areas of New York and Los Angeles where simulation meeting the existing and alternative standards using the HDDM-adjustment approach relied on large NOx reductions and there is very little day-to-day variability in the resulting O_3 concentrations.
D. Characterizing intra- urban population exposure in the context of epidemiology studies linking O_3 to specific health effects	Exposure misclassification within communities that is associated with the use of generalized population monitors (which may miss important patterns of exposure within urban study areas) introduces uncertainty into the effect estimates obtained from epidemiology studies.	Under (generally)	Low- medium	Medium	KB and INF: Despite the potential for substantial variability in O_3 levels across monitors (particularly in larger urban areas with greater variation in sources and topography such as L.A.), the O_3 ISA notes that studies have tended to demonstrate that monitor selection has only a limited effect on the association of short-term O_3 exposure with health effects (see O_3 ISA section 4.6.2.1, US EPA, 2013a). However, this issue could be more of a concern in larger urban areas which may exhibit greater variation in O_3 levels due to diverse sources, topography and patterns of commuting.
E. Statistical fit of the C- R functions	Exposure measurement error combined with other factors (e.g., size of the effect itself, sample size, control for confounders) can effect the overall level of confidence associated with the fitting of statistical effect-response models in epidemiological studies.	Both	Medium (short-term health endpoints)	Medium	INF: For short-term mortality and morbidity health endpoints, there is greater uncertainty associated with the fit of models given the smaller sample sizes often involved, difficulty in identifying the etiologically relevant time period for short-term O_3 exposure, and the fact that models tend to be fitted to individual counties or urban areas (which introduces the potential for varying degrees of confounding and effects modification across the locations). These studies can also have effects estimates that are not statistically significant. For this risk assessment, in modeling short-term mortality, we are not relying on location-specific models. Instead, we are using city-specific effects estimates derived using Bayesian techniques (these combine national-scale models with local-scale models). Exposure measurement error (uncertainty associated with the exposure metrics used to represent exposure of an individual or population) can also be an important contributor to uncertainty in effect estimates (O_3 ISA, p. 1xii). Together with other factors (e.g., low data density), exposure measurement error can result in the smoothing of epidemiologically-derived response functions and the obscuring of thresholds should they exist (O_3 ISA, p. Ixix). In addition, exposure measurement error can vary across different populations even within the same urban study area. For

		Potential influence of uncertainty on risk		Knowledge-	Comments	
		estimates		Base	(KB: knowledge base, INF: influence of uncertainty on risk	
Source	Description	Direction	Magnitude	uncertainty*	estimates)	
					example a particular group could have an activity pattern that results in central site monitors (in that urban study area) being particularly poor	
					at representing that group's exposure to ambient O_3 . In this example,	
					an effect estimate derived for that specific population based on O_3 exposure characterized using central site monitors would have increased uncertainty relative to effect estimates generated for other population with different activity patterns and lower levels of exposure measurement error.	
F. Shape of the C-R functions	Uncertainty in predicting the shape of the C-R function, particularly in the lower exposure regions which are often the focus in O_3 NAAQS regulatory reviews.	Both	Medium	Low-medium	KB and INF: Studies reviewed in the O_3 ISA that attempt to characterize the shape of the O_3 C-R curve along with possible "thresholds" (i.e., O_3 concentrations which must be exceeded in order to elicit an observable health response) have indicated a generally linear C-R function with no indication of a threshold (for analyses that have examined 8-h max and 24-h avg O_3 concentrations). However, the ISA notes that the studies from which the C-R functions are derived indicate there is less certainty in the shape of the C-R curve at the lower end of the distribution of O_3 concentrations (in the range below 20 ppb) due to the low density of data in the studies in this range.	
G. Addressing co- pollutants	The inclusion or exclusion of co- pollutants which may confound, or in other ways, affect the O_3 effect, introduces uncertainty into the analysis.	Both	Low- medium	Medium	KB and INF: The O_3 ISA notes that across studies, the potential impact of PM indices on O_3 -mortality risk estimates tended to be much smaller than the variation in O_3 -mortality risk estimates across cities. This suggests that O_3 effects are independent of the relationship between O_3 and mortality. However, interpretation of the potential confounding effects of PM on O_3 -mortality risk estimates requires caution. This is because the PM- O_3 correlation varies across regions, due to the difference in PM components, complicating the interpretation of the combined effect of PM on the relationship between O_3 and mortality. Additionally, the limited PM or PM component datasets used as a result of the every-3rd- and 6th-day PM sampling schedule instituted in most cities limits (in most cases) the overall sample size employed to examine whether PM or one of its components confounds the O_3 -mortality relationship (O_3 ISA section 2.5.4.5, US EPA, 2013a).	
H. Specifying lag structure (short-term exposure studies)	There is uncertainty associated with specifying the exact lag structure to use in modeling short-term O ₃ -	Both	Low- Medium	Low	KB and INF: The majority of studies examining different lag models suggest that O_3 effects on mortality occur within a few days of	

		Potential in	fluence of		
		uncertainty on risk		Knowledge-	Comments
		estimates		Base	(KB: knowledge base, INF: influence of uncertainty on risk
Source	Description	Direction	Magnitude	uncertainty*	estimates)
	attributable mortality and respiratory-related morbidity.				exposure. Similar, studies examining the impact of O_3 exposure on respiratory-related morbidity endpoints suggests a rather immediate response, within the first few days of O_3 exposure (see O_3 ISA section 2.5.4.3, US EPA, 2013a). Consequently, while the exact nature of the ideal lag models remains uncertain, generally, we are fairly confident that they would be on the order of a day to a few days following exposure.
I. Using studies from one geographic area to cover urban areas outside of the study area	In the case of Gent et al., 2003 (used in modeling asthma exacerbations in Boston), we are using C-R functions based on an epidemiological study of a region (northern Connecticut and Springfield) that does not encompass the actual urban study area assessed for risk (Boston).	Both	Medium	Low	INF: Factors related to O_3 exposure including commuting patterns, exercise levels etc may differ between the region reflected in the epidemiological study and Boston. If these differences are great, then applying the effect estimate from the epidemiological study to Boston could be subject to considerable uncertainty and potential bias.
J. Characterizing baseline incidence rates	Uncertainty can be introduced into the characterization of baseline incidence in a number of different ways (e.g., error in reporting incidence for specific endpoints, mismatch between the spatial scale in which the baseline data were captured and the level of the risk assessment).	Both	Low- medium	Low	INF: The degree of influence of this source of uncertainty on the risk estimates likely varies with the health endpoint category under consideration. There is no reason to believe that there are any systematic biases in estimates of the baseline incidence data. The influence on risk estimates that are expressed as incremental risk reductions between alternative standards should be relatively unaffected by this source of uncertainty. KB: The county level baseline incidence and population estimates at the county level were obtained from data bases where the relative degree of uncertainty is low.

* Refers to the degree of uncertainty associated with our understanding of the phenomenon, in the context of assessing and characterizing its uncertainty (specifically in the context of modeling PM risk)

1 7.4.3 Description of Core and Sensitivity Analyses

2 As discussed in section 7.1.1, this risk assessment includes a set of core (higher 3 confidence) risk estimates which are supplemented by sensitivity analyses. The sensitivity 4 analyses explore the potential impact that variation in specific model design elements can have 5 on the core risk estimates. This section specifies which design elements are included in both the 6 core and sensitivity analyses completed for each of the health effect endpoint categories included 7 in the risk assessment. We divided the sensitivity analyses into two categories: (a) those 8 involving air quality characterization and (b) those associated directly with the specification of 9 the C-R functions used in estimating risk. We recognize that there can be overlap between these 10 categories with some modeling elements (e.g., modeling period) affecting both the composite 11 monitor distribution as well as representing an element of C-R function specification. However, 12 we have retained these two categories to aid in the presentation and discussion of sensitivity analysis results.²⁹ The sensitivity analyses also included an initial influence analysis designed to 13 14 evaluate which of the model inputs are primarily responsible for inter-city variability 15 (heterogeneity) in risk. The influence analysis uses estimated elasticities of risk with respect to 16 the risk function input variables, focusing on the short-term exposure-related mortality endpoint 17 and associated input parameters since this is one of the key risk estimates generated for the REA 18 (additional detail on how the influence analysis was conducted is presented in section 7.5.3). 19 Table 7-5 presents the alternative approaches for adjusting the O_3 distributions used in 20 the sensitivity analysis and also identifies the approaches used in the core analysis for each of the 21 study areas. The alternative air quality adjustment approaches examine the differences in 22 changes in air quality and risk when applying NOx-only versus NOx and VOC reductions in the 23 HDDM-adjustment approach. It should be noted that when NOx and VOC reductions were used 24 in the HDDM-adjustment approach in this sensitivity analysis, the same percent reduction for 25 both pollutants was used in the air quality adjustment for meeting the existing and alternative 26 standard in each urban area. More details on these alternative air quality adjustment approaches

are discussed in Chapter 4 and appendices.

Besides the approach used to adjust the distributions of O_3 , another fact which has a direct impact on composite monitor composition is the specification of the study area (since this determines the mix of monitors that will be included in constructing the composite monitor). As discussed in section 7.1.1, for the core analysis, we modeled all endpoints (for all study areas)

²⁹ As noted in 7.1.1, in presenting both the core and sensitivity analyses, we include both point estimates and 95th CIs, with the latter reflecting the statistical fit of the effect estimates (and hence the power of the underlying epidemiological study). In comparing core and sensitivity analyses, we not only focus on point estimates, but also on the CIs since they provide insights into differences in the degree of statistical support for the effect estimates underlying the risk estimates and therefore, overall confidence in those estimates.

- 1 using CBSA-based study areas. For the sensitivity analysis (for the short-term O₃-attributable
- 2 mortality endpoint), we included a smaller study area based on the original study area definition
- 3 used in the Smith et al., 2009 study.³⁰
- 4 Table 7-6 presents the model elements included in sensitivity analyses exploring 5 alternative C-R function specifications. These sensitivity analyses were applied both to short-6 term O₃-attributable mortality and long-term O₃-attributable mortality. As discussed in section 7.1.1, we were not able to differentiate between alternative C-R function specifications for short-7 8 term O₃-attributable morbidity endpoints and therefore included the full set of alternative C-R 9 function specifications in the core analysis. This results in a distribution of core risk estimates for 10 each endpoint which can be used to gain insights into the impact of different C-R function 11 specifications on risk. Because separate sensitivity analyses were not completed for short-term
- 12 O_3 -attributable morbidity endpoints, this category is not included in Table 7-6.
- 13
- 14

 Table 7-5
 Specification of the Core and Sensitivity Analyses (air quality simulation)

Study Area	Core simulation (type of precursor reduced to adjust O_3 distribution)	Sensitivity analysis	
Atlanta, GA	NOx-only		
Baltimore, MD	NOx-only	Alternative modeling	
Boston, MA	NOx-only	approach not evaluated	
Cleveland, OH	NOx-only		
Denver, CO	NOx & VOC	NOx-only	
Detroit, MI	NOx-only	NOx & VOC	
Houston, TX	NOx-only	NOx & VOC	
Los Angeles, CA	NOx-lower bound*	NOx & VOC-lower bound*	
New York, NY	NOx-lower bound* (exclude 60)	NOx& VOC-lower bound*	
Philadelphia, PA	NOx-only	NOx & VOC	
Sacramento, CA	NOx-only	NOx & VOC	

³⁰ We did not include an alternative study area simulation as a sensitivity analysis for long-term exposure related mortality, since we are using a single (national) effect estimate in modeling this endpoint, and consequently, the use of an effect estimate from a smaller study area to represent a somewhat larger area (as is the case with short-term O₃-attributable mortality) is likely to introduce less uncertainty.

Study Area	$\begin{array}{c} \textbf{Core simulation} \\ \textbf{(type of precursor reduced to} \\ \textbf{adjust } O_3 \textbf{ distribution} \end{array}$	Sensitivity analysis
St. Louis, MO	NOx-only	Alternative modeling approach not evaluated

A lower-bound fit of the HDDM-based O_3 sensitivities (reflecting a greater increment of O_3 reduction per unit of VOC and/or NOx reduction) was required in simulation of the alternative standard levels.

Table 7-6 Specification of the Core and Sensitivity Analyses (alternative C-R function specification)

Health effect	Modeling elements included			
endpoint category	Core analysis	Sensitivity analysis		
Short-term O ₃ - attributable mortality	 Full monitoring period (specific to each study area), 8hr max metric, national-Bayes adjusted, single pollutant model. effect estimates obtained from: Smith et al., 2009 study 	 summer (warm month), 8hr mean, regional-bayes adjusted, multi-pollutant (with PM₁₀). effect estimates obtained from Zanobetti and Schwartz, 2008 and Smith et al., 2009 		
Long-term O ₃ - attributable mortality	 Single national estimate, two-pollutant model (PM2.5), long-term peak trend metric (based on daily 1hr max values), CBSA-based study area. effect obtained from Jerrett et al., 2009 study 	 Regional-differentiated effect estimates, single pollutant model. National-level effect estimate, single pollutant model. effect estimates also obtained from Jerrett et al., 2009 study 		

6

7 7.5 URBAN STUDY AREA RESULTS

8 This section discusses risk estimates generated for the set of 12 urban study areas, 9 including both the core risk estimates and accompanying sensitivity analyses. In summarizing 10 risk estimates, this discussion focuses on results most relevant to two policy-related questions: 11 (a) to what extent is the existing O_3 standard protective of public health , and, (b) what is the 12 nature and magnitude of additional public health protection provided by the suite of alternative 13 standards under consideration? Consequently, we focus on two types of risk estimates including 14 the magnitude of O_3 -attributable risk after simulation of just meeting the existing standard and the degree of risk reduction potentially provided by each of the alternative standards relative to
 just meeting the existing standard.³¹

3 This section is organized as follows. We begin by presenting the core risk estimates in 4 both tabular and graphical format at the end of this section. We then present key observations 5 about the risk estimates for just meeting the existing standard (for core risk) in section 7.5.1. Key 6 observations related to risk estimates for just meeting alternative standard levels, and for 7 estimates of risk changes comparing alternative standards to just meeting the existing standard 8 (again, for core risk) are presented in section 7.5.2. After presenting key observations related to 9 the core risk estimates, we then present key observations resulting from the sensitivity analyses 10 (section 7.5.3).

11 A number of details regarding the design of the core risk assessment should be kept in 12 mind when reviewing the core risk estimates presented in this section (see section 7.1.1 for 13 additional detail on these design elements):

14 • All risk estimates reflect application of a CBSA-based study area. 15 • Estimates are presented for two simulation years (2007 and 2009): 16 • Short-term O₃-attributable mortality estimates are generated for all 12 urban study 17 areas, while most short-term O₃-attributable morbidity estimates (depending on the 18 specific health endpoint) are generated for only a subset of urban study areas. Long-19 term O₃-attributable mortality is modeled for all 12 urban study areas. 20 • For all health effect endpoints, we model risk down to zero O_3 and do not include either consideration for LML or alternative threshold levels. 21 22 There are several categories of risk metrics generated for the core mortality and

morbidity endpoints modeled in this analysis. Below we describe both the types of risk metrics
generated for the core analysis and the specific types of tables and figures used in presenting
those metrics.

26	
27	<u>I. Core short-term O₃-attributable mortality estimates</u>
28	• Table presenting estimates of O ₃ -attributable mortality incidence after just
29	meeting the existing standard and the estimated change in incidence associated
30	with meeting each of the alternative standard levels relative to the existing
31	standard (Table 7-7): These estimates include point estimates and 95 th percentile
32	confidence intervals representing uncertainty associated with the statistical fit of the
33	effect estimates.

³¹ As part of this draft of the risk assessment, we have also generated estimates of risk under recent conditions as well as estimates of the degree of risk reduction (relative to risk under recent conditions) associated with the simulated attainment of the existing standard. See Appendix 7B.

1 Table presenting estimates of the percent of total mortality attributable to O_3 • 2 after just meeting the existing standard and the percent reduction in O₃-3 attributable risk associated with each alternative standard (Table 7-8). 4 Heat maps for mortality illustrating distribution across daily O₃ levels of total 5 O₃-attributable risk after just meeting the existing standard and risk reductions 6 after meeting alternative standards (Figures 7-2 and 7-3): Heat maps are provided 7 for each of the 12 urban areas. The color gradient in each figure reflects the 8 distribution of mortality (or the change in mortality) across the range of daily 8-hour 9 O₃ levels and provides a visual tool to explore trends in mortality counts across daily 10 O₃ levels and between cities. Visual patterns in the figures presenting total risk and 11 risk reduction are interpreted differently: 12 • For figures depicting total O₃-attributable risk, colors range from blue (lower mortality count) to red (higher mortality count). As an example, with Figure 7-2, 13 14 top heat map (which presents total O₃-attributable risk for the existing standard in 15 2007, based on Smith et al., 2009 C-R functions), if we focus on the first row 16 (Atlanta, GA), we see a value of 38 under the column 55-60 ppb. This value 17 reflects the fact that 38 of the 270 O₃-attributable deaths estimated for Atlanta 18 after just meeting the existing 75 ppb standard in 2007 occurred on days with 19 composite monitor O_3 levels between 55 and 60 ppb. Similarly, in the same row, 20 we see that only 3 O₃ attributable deaths occurred on days when the composite 21 monitor value was between 20 and 25 ppb. We also include the total O₃-22 attributable incidence (for each study area) in the final column marked "Total". 23 • For figures depicting changes in risk associated with simulation of existing and 24 alternative standard levels, we see that the pattern is more complex since we can 25 have a combination of increases and decreases in risk in the heat maps, with increases in risk identified as red to yellow and decreases in risk identified as 26 27 yellow to blue. Increases in risk are negative numbers, decreases are positive. In 28 addition, in the final three columns of each map, we provide estimates of the total 29 O₃-attributable incidence, as well as the total broken down into the subtotals 30 across days with increases (negative) and days with decreases (positive) in that 31 incidence. The increase and decrease for a given study area should sum 32 (accounting for rounding in these subtotals) to the overall total for O_3 -attributable 33 deaths for that study area. Several factors can contribute to the patterns of changes 34 in O₃-attributable risk reflected in these maps. For example, non-linearities in O₃
- 1 formation can result in increases in O₃ on some days, even when simulating 2 attainment of a lower alternative standard (see section 7.1.1). In addition, 3 simulation of alternative standard levels can result in a change in the overall 4 distribution of the composite monitor ambient O₃ distribution. Often, this change 5 will take the form of a shift in the upper tail of the distribution towards the mean, 6 given that simulated attainment of alternative standard levels targets higher O_3 7 days. If we look at figure 7-2 at the second map (Decrease 75 to 70) and 8 specifically at the row for Houston, we see that there is a -4 increase in deaths 9 distributed across 20-35 ppb days and a decrease in deaths of 9, primarily 10 distributed across 40-60 ppb days.
- 11

• Graphic plots of O₃-attributable deaths per 100,000 population for just meeting 12 the existing and alternative standards (Figures 7-4): O₃This plot provides 13 estimates that are adjusted for the size of the underlying urban population, thereby 14 allowing the mortality estimates and associated trends to be more readily compared 15 across urban study areas (consideration of absolute O_3 mortality is complicated by the 16 role that underlying urban population plays in driving total O_3 -attributable mortality – 17 larger study areas like Los Angeles and New York having substantially larger 18 mortality estimates primarily due to their higher underlying populations). These 19 figures allow us to evaluate the overall magnitude of risk reductions across standard 20 levels and determine the degree to which those trends differ for different study areas.

21 Tables summarizing incidence, percent of baseline incidence and percent reduction in O₃-22 attributable risk for short-term O₃-attributable morbidity (Tables 7-9 through 7-11): Three 23 categories of short-term O₃-attributable mortality effects were modeled for the analysis 24 (respiratory related HA, respiratory-related ER visits and asthma exacerbations). As discussed in 25 section 7.1.1, these morbidity effects were modeled for a combination of all 12 urban study areas 26 and a subset of those study areas depending on the endpoint (see below). The C-R functions 27 available for modeling many of these morbidity endpoints included consideration for a number 28 of design elements (e.g., copollutants and lag structure). However, as noted earlier in section 29 7.1.1, for short-term exposure morbidity endpoints with multiple C-R functions, we were not 30 able to differentiate between C-R functions in terms of overall confidence and consequently we 31 could not identify a single core model. Therefore, when we have multiple C-R functions 32 reflecting different treatments of key design elements such as lag structure, we consider the risk 33 estimates that result from the full set of C-R functions to represent a core range of risk. Each of 34 the tables summarizing short-term O₃-attributable morbidity risk present several risk metrics

1	including: (a) total O ₃ -attributable incidence (after just meeting the existing standard), (b)
2	reductions in O ₃ -attributable incidence (for each of the alternative standard levels relative to just
3	meeting the existing standard), (c) percent of baseline incidence attributable to O ₃ (after just
4	meeting the existing standard) and (d) percent reductions in O ₃ -attributable risk (for each of the
5	alternative standard levels). In presenting these morbidity risk estimates, we do not include 95 th
6	percentile confidence intervals in order to conserve space. Specific tables summarizing these
7	morbidity incidence estimates include:
8	• HA visits (for respiratory symptoms including asthma): Table 7-9 presents
9	estimates of the incidence of HA (for respiratory symptoms, chronic lung disease
10	and asthma). Risk estimates are generated for a subset of the urban study areas for
11	some of the health endpoints (e.g., New York City for HA [chronic lung disease
12	and asthma]), while HA (respiratory-related) estimates cover all 12 urban study
13	areas.
14	• ER visits (for respiratory symptoms including asthma): Table 7-10 presents
15	estimates of the incidence of ER visits for respiratory symptoms and asthma)
16	specifically for New York City and Atlanta based on C-R functions obtained from
17	several epidemiological studies.
18	• Asthma exacerbations: Table 7-11 presents estimates of the incidence of asthma
19	exacerbations (including estimates for a range of symptoms) for Boston, the only
20	urban study area with C-R functions supporting modeling for this endpoint.
21	• Graphic plots of O ₃ -attributable respiratory-related HA per 100,000 residents
22	for the existing and alternative standard levels (Figures 7-5): This figure is
23	intended to complement Figure 7-4 which presents the same type of risk information
24	for short-term O ₃ -attributable mortality. By plotting respiratory HA per 100,000, we
25	adjust for the underlying population which makes trends in risk more comparable
26	across urban study areas. We have only created this graphic for respiratory HA (based
27	on application of Medina-Ramon et al., 2006) since that is the only morbidity
28	endpoint modeled for all 12 urban study areas. As with the mortality figure, this
29	figure allows us to evaluate the overall magnitude of risk reductions across standard
30	levels and determine the degree to which those trends differ for different study areas.
31	III. Core long-term O ₃ -attributable mortality estimates
32	

1	•	Table presenting estimates of long-term O_3 -attributable mortality incidence
2		including total risk after just meeting the existing standard and risk reductions
3		based on comparing risks after meeting alternative standards to risks after
4		meeting the existing standard (Table 7-12): Estimates presented in Table 7-12
5		reflect respiratory mortality and include 95 th percentile confidence intervals
6		representing uncertainty associated with the statistical fit of the effect estimates used.
7		Estimates presented in these tables allow for consideration for the magnitude of risk
8		associated with just meeting the existing standard and the pattern of risk reduction (in
9		incidence) in meeting alternative standards relative to the existing standard.
10	•	Table presenting estimates of the percent of respiratory mortality attributable to
11		O_3 and percent reductions in O_3 -attributable risk for long-term O_3 -attributable
12		mortality (Table 7-13).
13	•	Graphic plots of O_3 -attributable deaths per 100,000 population for just meeting
14		the existing and alternative standards (Figures 7-6): This plot provides estimates
15		that are adjusted for the size of the underlying urban population, thereby allowing the
16		mortality estimates and associated trends to be more readily compared across urban
17		study areas (consideration of absolute O_3 mortality is complicated by the role that
18		underlying urban population plays in driving total O3-attributable mortality – larger
19		study areas like Los Angeles and New York having substantially larger mortality
20		estimates primarily due to their higher underlying populations). These figures allow
21		us to evaluate the overall magnitude of risk reductions across standard levels and
<u> </u>		

Table 7-7 Short-Term O₃-attributable All Cause Mortality Incidence (2007 and 2009) (Smith et al., 2009 C-R Functions) (O₃ season, CBSA-based study area, no threshold)

		Air Ouslain Ca		-
		Air Quality Sc		
	Absolute Incidence		inge in incidei	nce
Study Area	75ppb	/5-/U	/5-65	/5-60
	2007 SIMU	12	21	24
Atlanta, GA	(270, 800)	(16 20)	(20, 72)	54
	(-370-890)	(-10-39)	(-30 - 72)	(-47 - 110)
Baltimore, MD	440	13	2/	45
	(-250 - 1100)	(-/-33)	(-15 - 68)	(-25 - 110)
Boston, MA	350	/	20	32
	(-500 - 1200)	(-10 - 24)	(-28-67)	(-45 - 110)
Cleveland, OH	430	14	32	64
	(-41 - 890)	(-1 - 28)	(-3-67)	(-6 - 130)
Denver, CO	80	2	4	8
	(-280 - 440)	(-6 - 10)	(-14 - 23)	(-25 - 40)
Detroit, MI	660	23	42	69
	(32 - 1300)	(1 - 44)	(2 - 81)	(3 - 130)
Houston, TX	680	5	11	24
	(130 - 1200)	(1-9)	(2 - 20)	(4 - 43)
Los Angeles, CA	1300	43	8/	160
	(-530 - 3000)	(-18 - 100)	(-36 - 210)	(-66 - 380)
New York, NY	2800	130	(280, 800)	NA
	(1700 - 3900)	(80 - 190)	(380-890)	120
Philadelphia, PA	(270, 2200)	33 (9.62)	/0	(25 210)
	(270-2200)	(8-62)	(17 - 140)	(25-210)
Sacramento, CA	(200, 1100)	(7.20)	(12 20)	(24 70)
	(-590-1100)	(-7-20)	20	(-24 - 70)
St. Louis, MO	(-110 - 950)	(-5, 41)	(-10 - 86)	(-15 - 120)
	2009 Simu	lation Vear	(-10-00)	(-15 - 150)
	240	9	16	23
Atlanta, GA	(-340 - 800)	(-12 - 28)	(-22 - 54)	(-32 - 77)
	400	7	17	28
Baltimore, MD	(-220 - 1000)	(-4 - 19)	(-10 - 44)	(-15 - 71)
	320	-1	5	14
Boston, MA	(-450 - 1100)	(24)	(-7 - 17)	(-19 - 47)
	400	12	29	49
Cleveland, OH	(-38 - 830)	(-1 - 24)	(-3 - 60)	(-5 - 100)
	83	0	2	7
Denver, CO	(-270 - 420)	(-1 - 2)	(-6 - 10)	(-23 - 37)
Detect M	580	-21	-6	15
Detroit, ivii	(28 - 1100)	(-141)	(012)	(1 - 30)
Houston TV	700	-1	4	14
nousion, TX	(130 - 1200)	(01)	(1-7)	(3 - 26)
	1300	41	89	160
LUS Aligeles, CA	(-540 - 3100)	(-17 - 100)	(-37 - 210)	(-68 - 390)
New York NV	2600	84	440	NA
	(1600 - 3700)	(50 - 120)	(260 - 610)	NA
Philadelphia PA	1100	19	44	69
	(240 - 2000)	(4 - 34)	(10 - 78)	(15 - 120)
Sacramento, CA	370	6	12	21
	(-390 - 1100)	(-7 - 19)	(-13 - 38)	(-22 - 64)
St. Louis, MO	380	8	21	37
	(-96 - 840)	(-2 - 18)	(-5 - 46)	(-9 - 83)

3 4 5

NA: for NYC, the model-based adjustment methodology was unable to adjust O_3 distributions such that they would meet the

5 lower alternative standard level of 60 ppb.

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Table 7-8 Percent of Total All-Cause Mortality Attributable to 03 and Percent Change in
03-Attributable Risk (2007 and 2009) (Smith et al., 2009 C-R functions) (03 season,

CBSA-based study area, no threshold)

	Ai	ir Quality S	cenario	
	% of Baseline	% Change	e in O ₃ -Att	ributable
	Incidence		Risk	
Study Area	75ppb	75-70	75-65	75-60
	2007 Simulat	ion Year		
Atlanta, GA	1.1	4	8	12
Baltimore, MD	1.9	3	6	10
Boston, MA	1.2	2	5	9
Cleveland, OH	2.4	3	7	14
Denver, CO	0.8	2	5	9
Detroit, MI	3.0	3	6	10
Houston, TX	1.9	1	2	3
Los Angeles, CA	1.0	3	7	13
New York, NY	4.1	5	22	NA
Philadelphia, PA	3.2	3	6	9
Sacramento, CA	1.2	2	3	6
St. Louis, MO	2.5	4	9	14
	2009 Simulat	ion Year		
Atlanta, GA	1.0	3	7	9
Baltimore, MD	1.8	2	4	7
Boston, MA	1.1	-0.3	2	4
Cleveland, OH	2.3	3	7	12
Denver, CO	0.8	0.3	2	8
Detroit, MI	2.7	-4	-1	3
Houston, TX	1.9	-0.1	0.5	2
Los Angeles, CA	1.1	3	7	13
New York, NY	3.9	3	16	NA
Philadelphia, PA	3.0	2	4	6
Sacramento, CA	1.2	2	3	6
St. Louis, MO	2.3	2	5	9

NA: for NYC, the model-based adjustment methodology was unable to adjust O_3 distributions such that they would meet the

11 lower alternative standard level of 60 ppb.

Figure 7-2 Heat Maps for Short Term O₃-attributable Mortality (Just meeting existing standard and risk reductions from just meeting alternative standards) (2007) (Smith et al., 2009 C-R functions) (see Key at bottom of figure)

Current Standard (75)

Study area	Daily 8hr f	Max Ozone	Level (pp	b)													Total
	0-5	5-10	10-15	15-20	20-25	25-30	30-35	35-40	40-45	45-50	50-55	55-60	60-65	65-70	70-75	>75	
Atlanta, GA	0	0	0	0	3	5	18	24	41	52	63	38	15	6	3	0	267
Baltimore, MD	0	0	0	1	1	11	22	43	84	71	69	73	44	12	9	3	443
Boston, MA	0	0	0	0	4	20	45	50	58	57	35	20	30	9	13	11	353
Cleveland, OH	0	0	0	1	5	14	40	65	89	81	43	40	31	12	10	0	431
Denver, CO	0	0	0	0	0	0	1	5	6	13	17	23	15	4	2	0	86
Detroit, MI	0	0	0	0	2	7	42	72	123	147	75	52	56	20	43	17	655
Houston, TX	0	0	0	0	17	49	126	146	148	95	50	49	3	0	0	0	683
Los Angeles, CA	0	0	0	0	0	0	0	17	340	445	388	44	13	5	0	0	1,253
New York, NY	0	0	0	0	21	98	297	544	741	475	364	233	39	0	0	0	2,812
Philadelphia, PA	0	0	0	2	0	34	62	156	213	236	209	165	101	42	9	10	1,238
Sacramento, CA	0	0	0	0	1	18	53	98	67	65	40	20	5	2	0	0	367
St. Louis, MO	0	0	0	1	3	7	18	65	66	76	74	47	29	29	12	3	430

Decrease 75 to 70

Study area	Daily 8hr	Max Ozone	e Level (pp	b)													Total	Change	e in risk
	0-5	5-10	10-15	15-20	20-25	25-30	30-35	35-40	40-45	45-50	50-55	55-60	60-65	65-70	70-75	>75		Inc.	Dec.
Atlanta, GA	0	0	0	0	0	0	0	1	1	2	3	2	1	0	0	0	12	0	12
Baltimore, MD	0	0	0	0	0	0	0	0	1	2	3	3	2	1	1	0	13	-1	14
Boston, MA	0	0	0	0	0	0	0	0	1	1	1	1	1	0	1	1	7	-1	8
Cleveland, OH	0	0	0	0	0	0	0	0	2	3	2	2	2	1	1	0	14	-2	15
Denver, CO	0	0	0	0	0	0	0	0	0	0	0	1	1	0	0	0	2	0	2
Detroit, MI	0	0	0	0	0	0	-1	0	2	5	3	3	4	2	3	1	23	-2	24
Houston, TX	0	0	0	0	-1	-1	-2	0	2	2	2	2	0	0	0	0	5	-4	9
Los Angeles, CA	0	0	0	0	0	0	0	0	6	16	17	2	1	0	0	0	43	0	43
New York, NY	0	0	0	0	-1	-2	0	12	27	32	35	25	5	0	0	0	134	-11	146
Philadelphia, PA	0	0	0	0	0	-1	0	0	3	7	8	9	6	3	1	1	35	-3	38
Sacramento, CA	0	0	0	0	0	-1	-1	1	2	2	2	1	0	0	0	0	7	-1	8
St. Louis, MO	0	0	0	0	0	0	0	1	2	3	4	3	2	2	1	0	18	0	19

Decrease 75 to 65

Study area	Daily 8hr	Max Ozone	e Level (pp	b)													Total	Change	e in risk
	0-5	5-10	10-15	15-20	20-25	25-30	30-35	35-40	40-45	45-50	50-55	55-60	60-65	65-70	70-75	>75		Inc.	Dec.
Atlanta, GA	0	0	0	0	0	0	1	1	3	4	6	4	2	1	1	0	21	0	22
Baltimore, MD	0	0	0	0	0	0	0	0	3	4	5	7	5	1	1	0	27	-2	28
Boston, MA	0	0	0	0	0	-1	0	1	2	3	3	2	3	1	2	2	20	-2	22
Cleveland, OH	0	0	0	0	0	0	0	2	6	7	5	5	4	2	2	0	32	-3	34
Denver, CO	0	0	0	0	0	0	0	0	0	0	1	2	1	0	0	0	4	-1	5
Detroit, MI	0	0	0	0	0	0	-1	1	4	8	6	6	7	3	6	3	42	-3	45
Houston, TX	0	0	0	0	-2	-3	-3	0	4	5	4	5	0	0	0	0	11	-9	20
Los Angeles, CA	0	0	0	0	0	0	0	0	13	33	35	4	1	1	0	0	87	0	87
New York, NY	0	0	0	0	-1	2	24	85	149	136	136	90	19	0	0	0	640	-6	646
Philadelphia, PA	0	0	0	0	0	-1	-1	0	7	15	17	18	12	6	1	2	76	-5	81
Sacramento, CA	0	0	0	0	0	-1	-1	2	3	4	3	2	0	0	0	0	13	-3	15
St. Louis, MO	0	0	0	0	0	0	0	3	4	7	8	6	4	4	2	1	39	-1	39

Decrease 75 to 60

Study area	Daily 8hr	Max Ozone	e Level (pp	b)													Total	Change	e in risk
	0-5 5-10 10-15 15-20 20-25 25-30 30-35 35-40 40-45 45-50 50-55 55-60 60-65 65-70 70-75 5																		
	0-5	5-10	10-15	15-20	20-25	25-30	30-35	35-40	40-45	45-50	50-55	55-60	60-65	65-70	70-75	>75		Inc.	Dec.
Atlanta, GA	0	0	0	0	0	0	1	2	4	7	9	6	3	1	1	0	34	0	34
Baltimore, MD	0	0	0	0	0	0	0	1	5	7	9	11	8	2	2	1	45	-2	47
Boston, MA	0	0	0	0	0	-1	1	2	4	5	5	3	5	2	3	2	32	-2	34
Cleveland, OH	0	0	0	0	-1	0	0	5	12	14	10	10	8	4	3	0	64	-3	67
Denver, CO	0	0	0	0	0	0	0	0	0	1	2	3	2	1	0	0	8	-1	8
Detroit, MI	0	0	0	0	0	0	-1	2	8	14	10	9	11	5	9	4	69	-4	73
Houston, TX	0	0	0	0	-3	-5	-4	2	8	10	7	8	1	0	0	0	24	-13	37
Los Angeles, CA	0	0	0	0	0	0	0	2	41	59	48	6	2	1	0	0	159	0	159
New York, NY										NA									
Philadelphia, PA	0	0	0	0	0	-2	-1	1	11	23	26	27	17	8	2	2	116	-7	123
Sacramento, CA	0	0	0	0	0	-2	-2	4	6	7	5	3	1	0	0	0	23	-4	27
St. Louis, MO	0	0	0	0	0	0	0	5	7	11	12	9	6	7	3	1	60	-1	61

NA: for NYC, the model-based adjustment methodology was unable to adjust O_3 distributions such that they would meet the lower alternative standard level of 60 ppb.

Key: For current standard (75) which is an absolute risk metric expressed as incidence of mortality, color gradient ranges from

blue (smallest O3-related mortality count) to red (highest O3-related mortality count). For estimates of decreases in risk, color

gradient ranges from red (increase in risk – negative cell values) to blue (reduction in risk – positive cell values).

Figure 7-3 Heat Maps for Short Term 0₃-attributable Mortality (Just meeting existing standard and risk reductions from just meeting alternative standards) (2009) (Smith et al., 2009 C-R functions) (see Key at bottom of figure)

Current Standard (75)

Study area	Daily 8hr 1	Max Ozone	e Level (ppl	b)													Total
	0-5	5-10	10-15	15-20	20-25	25-30	30-35	35-40	40-45	45-50	50-55	55-60	60-65	65-70	70-75	>75	
Atlanta, GA	0	0	1	2	8	16	18	33	49	44	29	30	9	2	0	0	241
Baltimore, MD	0	0	0	1	3	13	41	71	64	92	64	45	11	0	0	0	404
Boston, MA	0	0	1	0	11	25	45	57	50	53	48	7	3	6	9	3	319
Cleveland, OH	0	0	0	0	5	25	46	68	75	81	57	28	11	6	0	0	401
Denver, CO	0	0	0	0	0	1	2	4	9	17	22	20	6	2	1	0	83
Detroit, MI	0	0	1	9	7	26	46	67	114	148	38	51	46	0	21	6	579
Houston, TX	0	0	0	6	28	51	122	123	114	90	84	36	27	7	4	4	695
Los Angeles, CA	0	0	0	0	0	0	2	17	281	328	496	152	9	0	0	0	1,285
New York, NY	0	0	0	6	36	215	427	356	632	469	274	175	56	0	0	0	2,645
Philadelphia, PA	0	0	0	2	16	51	159	126	219	175	198	97	68	0	0	0	1,112
Sacramento, CA	0	0	0	0	2	23	64	69	73	56	42	33	6	0	0	0	367
St. Louis, MO	0	0	1	6	7	17	29	54	52	77	66	53	13	8	0	0	383

Decrease 75 to 70

Study area	Daily 8hr I	Max Ozone	e Level (pp	b)													Total	Change	in risk
	0-5	5-10	10-15	15-20	20-25	25-30	30-35	35-40	40-45	45-50	50-55	55-60	60-65	65-70	70-75	>75		Inc.	Dec.
Atlanta, GA	0	0	0	0	0	0	0	1	2	3	2	2	1	0	0	0	8	-2	10
Baltimore, MD	0	0	0	0	0	0	-1	0	1	3	2	2	1	0	0	0	7	-2	9
Boston, MA	0	0	0	0	-1	-1	-1	0	0	1	1	0	0	0	0	0	-1	-6	5
Cleveland, OH	0	0	0	0	0	-1	0	1	2	3	3	2	1	0	0	0	12	-2	14
Denver, CO	0	0	0	0	0	0	0	0	0	0	0	1	0	0	0	0	0	-1	1
Detroit, MI	0	0	-1	-3	-2	-6	-5	-5	-5	-2	0	2	2	0	2	1	-21	-29	8
Houston, TX	0	0	0	-1	-2	-3	-3	-1	1	2	3	2	2	0	0	0	-1	-10	10
Los Angeles, CA	0	0	0	0	0	0	0	0	4	11	20	6	0	0	0	0	41	0	41
New York, NY	0	0	0	-1	-3	-14	-8	8	22	31	22	19	6	0	0	0	84	-38	121
Philadelphia, PA	0	0	0	0	-1	-2	-2	-1	5	5	8	5	4	0	0	0	19	-9	28
Sacramento, CA	0	0	0	0	0	-1	-1	1	2	2	2	2	0	0	0	0	6	-2	8
St Louis MO	0	0	0	-1	-1	-1	0	1	1	3	3	3	1	1	0	0	8	-4	12

Decrease 75 to 65

Study area	Daily 8hr	Max Ozone	e Level (pp	b)													Total	Change	e in risk
	0-5	5-10	10-15	15-20	20-25	25-30	30-35	35-40	40-45	45-50	50-55	55-60	60-65	65-70	70-75	>75		Inc.	Dec.
Atlanta, GA	0	0	0	0	-1	-1	0	1	3	5	3	4	2	0	0	0	16	-3	19
Baltimore, MD	0	0	0	0	0	-1	-1	0	3	6	5	4	1	0	0	0	17	-3	21
Boston, MA	0	0	0	0	-1	-1	-1	0	1	2	3	1	0	0	1	0	5	-6	11
Cleveland, OH	0	0	0	0	-1	-1	0	4	5	8	7	4	2	1	0	0	29	-3	32
Denver, CO	0	0	0	0	0	0	0	0	0	0	1	2	1	0	0	0	2	-1	3
Detroit, MI	0	0	-1	-4	-2	-7	-5	-4	-2	3	2	4	5	0	3	1	-6	-27	21
Houston, TX	0	0	0	-2	-4	-5	-5	-1	2	4	6	3	3	1	1	1	4	-18	21
Los Angeles, CA	0	0	0	0	0	0	0	0	10	23	42	14	1	0	0	0	89	0	89
New York, NY	0	0	0	-1	-5	-17	16	52	107	120	81	63	21	0	0	0	437	-43	479
Philadelphia, PA	0	0	0	-1	-2	-4	-4	-1	10	11	17	10	8	0	0	0	44	-15	59
Sacramento, CA	0	0	0	0	0	-2	-1	2	3	4	3	3	1	0	0	0	12	-4	16
St. Louis, MO	0	0	-1	-2	-1	-1	0	2	3	6	6	6	2	1	0	0	21	-5	26

Decrease 75 to 60

Study area	Daily 8hr	Max Ozon	e Level (pp	b)													Total	Change	in risk
	0-5	5-10	10-15	15-20	20-25	25-30	30-35	35-40	40-45	45-50	50-55	55-60	60-65	65-70	70-75	>75		Inc.	Dec.
Atlanta, GA	0	0	-1	0	-1	-1	0	2	5	6	5	5	2	0	0	0	23	-3	26
Baltimore, MD	0	0	0	0	-1	-1	-1	1	5	10	8	6	2	0	0	0	28	-4	32
Boston, MA	0	0	0	0	-2	-1	-1	2	2	5	5	1	1	1	2	1	14	-6	20
Cleveland, OH	0	0	0	0	-1	-1	1	7	9	13	11	6	2	1	0	0	49	-3	53
Denver, CO	0	0	0	0	0	0	0	0	0	1	2	3	1	0	0	0	7	-1	8
Detroit, MI	0	0	-1	-4	-2	-7	-5	-2	2	10	4	8	8	0	4	2	15	-26	41
Houston, TX	0	0	0	-2	-7	-7	-7	0	5	8	11	6	5	1	1	1	14	-25	40
Los Angeles, CA	0	0	0	0	0	0	0	2	32	44	63	22	1	0	0	0	164	0	164
New York, NY										NA									
Philadelphia, PA	0	0	0	-1	-3	-6	-4	0	16	17	25	14	11	0	0	0	69	-20	88
Sacramento, CA	0	0	0	0	-1	-3	-1	3	6	6	5	5	1	0	0	0	21	-6	27
St. Louis. MO	0	0	-1	-2	-1	-2	0	3	5	10	9	9	3	2	0	0	37	-6	43

4 5

NA: for NYC, the model-based adjustment methodology was unable to adjust O_3 distributions such that they would meet the lower alternative standard level of 60 ppb.

Key: For current standard (75) which is an absolute risk metric, color gradient ranges from blue (smallest O_3 -related mortality count) to red (highest O_3 -related mortality count). For *estimates of decreases in risk*, color gradient ranges from red (increase in risk – negative cell values) to blue (reduction in risk – positive cell values).

Figure 7-4 Plots of Short-Term 0₃-attributable All-Cause Mortality for Meeting Existing standard and Alternative Standards (Smith et al., 2009) (Simulation year 2007 and 2009)



Table 7-9 Short-Term O₃-attributable Morbidity Incidence, Percent of Baseline and Reduction in O₃-attributable Risk – Respiratory-Related Hospital Admissions (2007 and 2009)

		Air Quality Scenario							
		Absolute				Percent of	% Chang	e in Ozone	-Related
		Incidence	Chan	ge in Incid	ence	Baseline		Risk	
En	dpoint/Study Area/Descriptor	75ppb	75-70	75-65	75-60	75ppb	75-70	75-65	75-60
			2007 Sir	nulation Ye	ear				
HA (respiratory); Detroit (Katsouyan	ni et al., 2009	9)						-
	1hr max, penalized splines	230	13	23	37	2.8	5	10	15
	1hr max, natural splines	230	12	22	36	2.7	5	10	15
HA (respiratory); NYC (Silverman ar	nd Ito, 2010; Li	n et al., 20	08)					-
	HA Chronic Lung Disease (Lin)	120	6.7	29		3.3	5	23	
	HA Asthma (Silverman)	420	28	120	NA	27.6	5	21	NA
	HA Asthma, PM2.5 (Silverman)	310	20	84		20.1	5	22	
HA (respiratory); LA (Linn et al., 2000)							
	1hr max penalized splines	790	19	38	60	2.4	2	5	7
HA (COPD less asthma); all 12 study	areas (Medina	a-Ramon, e	et al., 2006)					
	Atlanta, GA	67	4	6	10	2.5	5	9	15
	Baltimore, MD	77	3	6	9	2.6	4	7	12
	Boston, MA	100	2	6	10	2.2	2	6	9
	Cleveland, OH	61	2	5	10	2.4	4	8	17
	Denver, CO	27	1	2	3	2.9	3	6	11
	Detroit, MI	90	3	6	9	2.5	3	6	10
	Houston, TX	68	1	2	4	2.1	1	3	6
	Los Angeles, CA	180	8	16	25	2.7	4	9	13
	New York, NY	180	11	50	NA	2.2	6	28	NA
	Philadelphia, PA	130	4	10	15	2.5	3	7	11
	Sacramento, CA	34	1	2	4	2.5	3	7	11
	St. Louis, MO	53	3	5	8	2.6	5	10	15
			2009 Sir	nulation Ye	ear				
HA (respiratory); Detroit (Katsouyan	ini et al., 2009	9)						
	1hr max, penalized splines	220	3.6	13	25	2.5	2	6	11
	1hr max, natural splines	210	3.4	12	24	2.4	2	6	11
HA (respiratory); NYC (Silverman ar	nd Ito, 2010; Li	n et al., 20	08)					-
	HA Chronic Lung Disease (Lin)	120	5.1	21		3.2	4	17	
	HA Asthma (Silverman)	410	24	96	NA	27.2	4	17	NA
	HA Asthma, PM2.5 (Silverman)	310	17	68		19.8	4	18	
HA (respiratory); LA (Linn et al., 2000)							-
	1hr max penalized splines	640	18	39	62	2.4	2	4	7
HA (COPD less asthma); all 12 study	areas (Medina	a-Ramon, e	et al., 2006)				1	-
	Atlanta, GA	65	3	5	8	2.2	5	8	12
	Baltimore, MD	74	2	4	6	2.3	2	5	8
	Boston, MA	92	0	1	4	2.0	0	1	4
	Cleveland, OH	58	2	5	8	2.2	3	8	14
	Denver, CO	27	0	1	3	2.7	1	4	11
	Detroit, MI	81	-3	-2	1	2.2	-4	-2	1
	Houston, TX	71	1	2	4	2.2	1	2	5
	Los Angeles, CA	200	8	16	26	2.7	4	8	13
	New York, NY	170	7	35	NA	2.1	4	20	NA
	Philadelphia, PA	120	2	6	9	2.3	2	4	7
	Sacramento, CA	41	1	3	4	2.4	3	7	11
	St. Louis, MO	51	2	4	6	2.4	3	8	12

NA: for NYC, the model-based adjustment methodology was unable to adjust O_3 distributions such that they would meet the lower alternative standard level of 60 ppb.

Table 7-10 Short-Term O₃-attributable Morbidity Incidence, Percent of Baseline and Reduction in O₃-attributable Risk – Emergency Room Visits (2007 and 2009)

		Air Quality Scenario							
		Absolute				Percent of	% Chang	e in Ozone	-Related
		Incidence	Change in Incidence			Baseline		Risk	
Er	ndpoint/Study Area/Descriptor	75ppb	75-70	75-65	75-60	75ppb	75-70	75-65	75-60
			2007 Sir	nulation Y	ear				
ER۱	/isits (repiratory); Atlanta (Strick	dand et al., 2	.007)	r	r	1	n	1	
	Distributed lag 0-7 days	7,500	410	740	1,200	19.6	4	8	13
	Average day lag 0-2	4,500	230	420	670	11.6	5	8	13
ER-	visits (respiratory); Atlanta (Tolb	ert et al., 20	07, Darrow	et al., 201	1)			1	1
	Tolbert	8,100	360	670	1,100	5.8	4	8	12
	Tolbert-CO	7,200	320	590	940	5.1	4	8	12
	Tolbert-NO2	6,500	290	530	840	4.6	4	8	12
	Tolbert-PM10	5,100	230	420	660	3.6	4	8	12
	Tolbert-PM10, NO2	4,900	220	400	640	3.5	4	8	12
	Darrow	4,400	190	360	560	3.1	4	8	12
ER-	visits (asthma); NYC (Ito et al, 20	007)		r	r			n	-
	single pollutant model	9,000	530	2,300		19.9	5	22	
	PM2.5	7,100	410	1,800		15.5	5	22	
	NO2	5,800	330	1,500	NA	12.8	5	23	NA
	со	9,500	570	2,500		21.0	5	22	
	SO2	7,300	420	1,900		16.0	5	22	
			2009 Sir	nulation Y	ear				
ER \	/isits (repiratory); Atlanta (Strick	dand et al., 2	.007)	r	r		T	1	-
	Distributed lag 0-7 days	6,800	310	570	800	17.2	4	7	10
	Average day lag 0-2	4,000	170	320	460	10.1	4	7	10
ER-	visits (respiratory); Atlanta (Tolb	ert et al., 20	07, Darrow	et al., 201	1)				-
	Tolbert (single pollutant	7,400	270	500	720	5.1	3	6	9
	Tolbert-CO	6,600	240	450	640	4.5	3	6	9
	Tolbert-NO2	6,000	210	400	580	4.1	3	6	9
	Tolbert-PM10	4,700	170	310	450	3.2	3	6	9
	Tolbert-PM10, NO2	4,500	160	300	430	3.1	3	6	9
	Darrow (single pollutant	4,000	140	270	380	2.8	3	6	9
ER-	visits (asthma); NYC (Ito et al, 20	007)		-	-				
	single pollutant model	8,800	400	1,800	ļ	19.3	4	17	
	PM2.5	6,900	310	1,400		15.0	4	17	
	NO2	5,700	250	1,100	NA	12.4	4	18	NA
	со	9,300	430	1,900	ļ	20.4	4	17	
	SO2	7,100	320	1,400		15.5	4	17	

NA: for NYC, the model-based adjustment methodology was unable to adjust O_3 distributions such that they would meet the lower alternative standard level of 60 ppb.

Table 7-11 Short-Term O₃-attributable Morbidity Incidence, Percent of Baseline and Reduction in O₃-attributable Risk – Asthma Exacerbations (2007 and 2009)

		Air Quality Scenario							
		Absolute	te		Percent of	% Chang	e in Ozone	-Related	
		Incidence	Chan	ge in Incid	ence	Baseline		Risk	
E	indpoint/Study Area/Descriptor	75ppb	75-70	75-65	75-60	75ppb	75-70	75-65	75-60
			2007 Sir	nulation Y	ear				
A	Asthma exacerbation (wheeze); Boston (Gent et al., 2003, 2004)								
	Chest Tightness	69,000	2,100	5,600	8,600	28.8	2	5	9
	Shortness of Breath	49,000	1,400	3,700	5,700	16.2	2	6	10
	Chest Tightness (1hr max)	51,000	1,200	3,200	5,000	21.2	2	5	8
	Shortness of Breath (1hr max)	59,000	1,300	3,600	5,800	19.6	2	5	8
	Chest Tightness (PM2.5)	69,000	2,100	5,600	8,700	29.1	2	5	9
	Chest Tightness (PM2.5)	64,000	1,900	5,100	8,000	26.8	2	5	9
	Wheeze (PM2.5)	130,000	3,800	10,000	16,000	23.2	2	6	9
			2009 Sir	nulation Y	ear				
A	sthma exacerbation (wheeze); Bo	ston (Gent et	t al., 2003,	2004)					
	Chest Tightness	63,000	490	2,400	4,800	27.0	0.4	2	5
	Shortness of Breath	45,000	330	1,600	3,200	15.1	1	3	6
	Chest Tightness (1hr max)	47,000	-180	790	2,200	19.8	-0.4	1	3
	Shortness of Breath (1hr max)	54,000	-210	910	2,500	18.3	-0.4	1	4
	Chest Tightness (PM2.5)	64,000	500	2,400	4,800	27.2	0.4	2	5
	Chest Tightness (PM2.5)	59,000	450	2,200	4,400	25.1	0.4	3	5
	Wheeze (PM2.5)	120,000	900	4,300	8,700	21.7	0.5	3	6

Figure 7-5 Plots of Short-Term 0₃-attributable Respiratory HA for Meeting Existing standard and Alternative Standards (Medina-Ramon, et al., 2006) (Simulation year 2007 and 2009)



	Air Qualtiy Scenario					
	Absolute Incidence	Cha	nge in Incide	nce		
Study Area	75ppb	75-70	75-65	75-60		
	2007 Simu	lation Year				
	710	43	78	120		
Atlanta, GA	(260 - 1100)	(15 - 71)	(26 - 130)	(41 - 200)		
Poltimore MD	750	33	67	110		
balumore, wid	(270 - 1200)	(11 - 55)	(23 - 110)	(37 - 180)		
Poston MA	1,100	35	93	140		
boston, MA	(400 - 1800)	(12 - 58)	(32 - 150)	(49 - 240)		
Clausiand Oli	530	26	56	100		
Cleveland, OH	(190 - 820)	(9 - 43)	(19 - 93)	(35 - 170)		
	480	19	39	64		
Deriver, CO	(170 - 740)	(6 - 31)	(13 - 64)	(22 - 100)		
Dotroit MI	760	35	63	99		
Detroit, wi	(270 - 1200)	(12 - 59)	(21 - 100)	(33 - 160)		
Houston TV	550	9.5	19	32		
nousion, 1x	(190 - 860)	(3 - 16)	(6 - 31)	(11 - 53)		
	2,600	140	260	410		
Los Angeles, CA	(930 - 4000)	(46 - 230)	(89 - 430)	(140 - 670)		
Now York, NV	1,800	120	480	NIA		
New York, NY	(660 - 2900)	(41 - 200)	(160 - 790)	NA NA		
Philadelphia, PA	1,300	56	120	170		
	(450 - 1900)	(19 - 93)	(40 - 190)	(59 - 290)		
Sacramonto CA	680	31	60	100		
Sacramento, CA	(250 - 1100)	(10 - 51)	(20 - 98)	(34 - 170)		
St Louis MO	600	34	69	100		
St. Louis, NIO	(210 - 930)	(230 - 1000)	(23 - 110)	(35 - 170)		
	2009 Simu	lation Year				
Atlanta GA	700	41	76	100		
Atlanta, GA	(250 - 1100)	(14 - 68)	(26 - 120)	(36 - 170)		
Atlanta, GA Baltimore, MD	730	25	54	84		
	(260 - 1100)	(8 - 41)	(18 - 90)	(28 - 140)		
Baltimore, MD Boston, MA	1,100	6.8	42	85		
	(380 - 1700)	(2 - 11)	(14 - 69)	(29 - 140)		
Cleveland, OH	510	24	54	84		
	(180 - 800)	(8 - 41)	(18 - 89)	(29 - 140)		
Denver, CO	490	9.0	28	70		
	(180 - 770)	(3 - 15)	(10 - 47)	(24 - 120)		
Detroit. MI	720	-8.9	18	51		
	(260 - 1100)	(-315)	(6 - 30)	(17 - 85)		
Houston, TX	610	14	30	49		
	(220 - 950)	(5 - 23)	(10 - 49)	(17 - 82)		
Los Angeles, CA	2,800	130	280	430		
	(1000 - 4300)	(45 - 220)	(94 - 460)	(150 - 710)		
New York, NY	1,900	110	390	NA		
	(670 - 2900)	(37 - 180)	(130 - 630)			
Philadelphia, PA	1,200	44	94	140		
	(430 - 1900)	(15 - 73)	(32 - 160)	(47 - 230)		
Sacramento, CA	730	34	66	110		
	(260 - 1100)	(12 - 57)	(22 - 110)	(36 - 170)		
St. Louis, MO	580	24	53	86		
	(210 - 900)	(210 - 910)	(18 - 87)	(29 - 140)		

Table 7-12 Long-Term 0₃-attributable Respiratory Mortality Incidence (2007 and 2009) (Jerrett et al., 2009 C-R Functions) (CBSA-based study area, no threshold)

NA: for NYC, the model-based adjustment methodology was unable to adjust O_3 distributions such that they would meet the lower alternative standard level of 60 ppb.

Table 7-13 Long-Term 03-attributable Respiratory Mortality Percent of BaselineIncidence and Percent Reduction in 03-attributable Risk (simulation years 2007 and2009) (Jerrett et al., 2009 C-R Functions) (CBSA-based study area, no threshold)

	Air Quality Scenario					
	% of Baseline					
	Attributable to					
	Ozone	Change	in O3-Attributa	able Risk		
Study Area	70ppb	75-70 75-65 75-6				
	2007 Sir	nulation Year				
Atlanta, GA	17.7	5	9	15		
Baltimore, MD	18.1	4	8	12		
Boston, MA	16.7	3	7	11		
Cleveland, OH	16.9	4	9	17		
Denver, CO	20.1	3	7	11		
Detroit, MI	17.7	4	7	11		
Houston, TX	16.1	1	3	5		
Los Angeles, CA	19.6	4	9	13		
New York, NY	15.9	6	24	NA		
Philadelphia, PA	17.7	4	8	12		
Sacramento, CA	17.1	4	7	13		
St. Louis, MO	17.9	5	10	15		
	2009 Sir	nulation Year				
Atlanta, GA	16.1	5	9	13		
Baltimore, MD	16.9	3	6	10		
Boston, MA	15.9	1	3	7		
Cleveland, OH	16.1	4	9	15		
Denver, CO	19.7	1	5	12		
Detroit, MI	17.1	-1	2	6		
Houston, TX	16.6	2	4	7		
Los Angeles, CA	19.9	4	8	13		
New York, NY	15.9	5	18	NA		
Philadelphia, PA	16.7	3	7	10		
Sacramento, CA	17.3	4	8	12		
St. Louis, MO	17.1	4	8	13		

NA: for NYC, the model-based adjustment methodology was unable to adjust O_3 distributions such that they would meet the lower alternative standard level of 60 ppb.

Figure 7-6 Plots of Long-Term O₃-attributable Respiratory Mortality for Meeting Existing standard and Alternative Standards (Jerrett et al., 2009) (Simulation year 2007 and 2009)



1	The presentation of key observations drawn from review of the core risk estimates is							
2	divided into two sections: 1) the assessment of health risks associated with just meeting the							
3	existing standard (section 7.5.1) and 2) the assessment of risk changes from meeting alternative							
4	standards relative to meeting the existing standard (section 7.5.2). The presentation of key							
5	observations in each of these two sections is further separated into those associated with (a)							
6	short-term O ₃ -attributable mortality, (b) short-term O ₃ -attributable morbidity and (c) long-term							
7	O3-attributable mortality. Unless otherwise noted, all risk estimates discussed in these three							
8	sections are core risk estimates. In some cases we refer to the confidence intervals around risk							
9	estimates. When an effect estimate is drawn from a study with low statistical power, confidence							
10	intervals can be wide, and can include negative values because of the assumptions of normality							
11	in the distribution of the effect estimate. Negative lower-confidence bounds do not imply that							
12	additional exposure to O_3 has a beneficial effect, but rather that the estimated O_3 effect estimate							
13	in the C-R function was not statistically significantly different from zero, and thus has a higher							
14	degree of uncertainty as to the magnitude of the estimated risk. As noted earlier, presentation of							
15	sensitivity analysis results and their use in interpreting the core risk estimates is covered in							
16	section 7.5.3.							
17	7.5.1 Assessment of Health Risk After Just Meeting the Existing 75 ppb standard							
18	The analysis of risk after simulating just meeting the existing standard focuses on							
19	absolute risk, since this is of greatest relevance in evaluating the adequacy of the existing							
20	standard.							
21								
22	Short-term O ₃ -attributable mortality							
23								
29								
23 24	• After meeting the existing standard, estimates of O ₃ -related all-cause mortality range							
23 24 25	• After meeting the existing standard, estimates of O ₃ -related all-cause mortality range across urban areas from 86 to 2,800 deaths (for simulation year 2007) and from 83 to							
23 24 25 26	• After meeting the existing standard, estimates of O ₃ -related all-cause mortality range across urban areas from 86 to 2,800 deaths (for simulation year 2007) and from 83 to 2,600 deaths (for simulation year 2009) (see Table 7-7). This translates into from 0.8							
23 24 25 26 27	• After meeting the existing standard, estimates of O ₃ -related all-cause mortality range across urban areas from 86 to 2,800 deaths (for simulation year 2007) and from 83 to 2,600 deaths (for simulation year 2009) (see Table 7-7). This translates into from 0.8 to 4.1% of baseline all-cause mortality (for simulation year 2007) and from 0.8 to							
23 24 25 26 27 28	• After meeting the existing standard, estimates of O ₃ -related all-cause mortality range across urban areas from 86 to 2,800 deaths (for simulation year 2007) and from 83 to 2,600 deaths (for simulation year 2009) (see Table 7-7). This translates into from 0.8 to 4.1% of baseline all-cause mortality (for simulation year 2007) and from 0.8 to 3.9% (for simulation year 2009) (see Table 7-8) in these study areas. Generally, O ₃ -							
23 24 25 26 27 28 29	• After meeting the existing standard, estimates of O ₃ -related all-cause mortality range across urban areas from 86 to 2,800 deaths (for simulation year 2007) and from 83 to 2,600 deaths (for simulation year 2009) (see Table 7-7). This translates into from 0.8 to 4.1% of baseline all-cause mortality (for simulation year 2007) and from 0.8 to 3.9% (for simulation year 2009) (see Table 7-8) in these study areas. Generally, O ₃ -attributable all-cause mortality risks continue to be lower for the 2009 simulation year							
23 24 25 26 27 28 29 30	• After meeting the existing standard, estimates of O ₃ -related all-cause mortality range across urban areas from 86 to 2,800 deaths (for simulation year 2007) and from 83 to 2,600 deaths (for simulation year 2009) (see Table 7-7). This translates into from 0.8 to 4.1% of baseline all-cause mortality (for simulation year 2007) and from 0.8 to 3.9% (for simulation year 2009) (see Table 7-8) in these study areas. Generally, O ₃ -attributable all-cause mortality risks continue to be lower for the 2009 simulation year as compared with the 2007 simulation year (with the exception of Houston),							
23 24 25 26 27 28 29 30 31	• After meeting the existing standard, estimates of O ₃ -related all-cause mortality range across urban areas from 86 to 2,800 deaths (for simulation year 2007) and from 83 to 2,600 deaths (for simulation year 2009) (see Table 7-7). This translates into from 0.8 to 4.1% of baseline all-cause mortality (for simulation year 2007) and from 0.8 to 3.9% (for simulation year 2009) (see Table 7-8) in these study areas. Generally, O ₃ -attributable all-cause mortality risks continue to be lower for the 2009 simulation year as compared with the 2007 simulation year (with the exception of Houston), reflecting the generally lower ambient O ₃ levels associated with 2009 for most of the							
23 24 25 26 27 28 29 30 31 32	• After meeting the existing standard, estimates of O ₃ -related all-cause mortality range across urban areas from 86 to 2,800 deaths (for simulation year 2007) and from 83 to 2,600 deaths (for simulation year 2009) (see Table 7-7). This translates into from 0.8 to 4.1% of baseline all-cause mortality (for simulation year 2007) and from 0.8 to 3.9% (for simulation year 2009) (see Table 7-8) in these study areas. Generally, O ₃ -attributable all-cause mortality risks continue to be lower for the 2009 simulation year as compared with the 2007 simulation year (with the exception of Houston), reflecting the generally lower ambient O ₃ levels associated with 2009 for most of the study areas (see Tables 7-7 and 7-8).							
24 25 26 27 28 29 30 31 32 33	 After meeting the existing standard, estimates of O₃-related all-cause mortality range across urban areas from 86 to 2,800 deaths (for simulation year 2007) and from 83 to 2,600 deaths (for simulation year 2009) (see Table 7-7). This translates into from 0.8 to 4.1% of baseline all-cause mortality (for simulation year 2007) and from 0.8 to 3.9% (for simulation year 2009) (see Table 7-8) in these study areas. Generally, O₃-attributable all-cause mortality risks continue to be lower for the 2009 simulation year as compared with the 2007 simulation year (with the exception of Houston), reflecting the generally lower ambient O₃ levels associated with 2009 for most of the study areas (see Tables 7-7 and 7-8). Confidence intervals (CIs) reflecting the statistical fit of the effect estimates used in 							

- general, the upper 95th percentile CI tends to be from 2-3 times larger than the point 1 2 estimate for the 12 urban study areas (see Table 7-7). However, some cities have 3 markedly wider confidence intervals (e.g., Denver where the upper CI is ~5 times the 4 point estimate), while others have tighter relative CIs (e.g., New York, where the 5 upper CI is ~ 1.4 times larger than the point estimate). This variation in the CIs 6 associated with risk estimates can reflect a number of factors including the statistical power of the underlying epidemiological study, which is based on the population size, 7 8 and differences in the magnitude of such factors as exposure measurement error and 9 correlations between O₃ and other pollutants.
- 10 After just meeting the existing O₃ standard, all-cause mortality estimates based on C-11 R functions from Smith et al., 2009 (for simulation year 2007) continue to be driven largely by days with total O₃ levels falling in the range of 30 to 70 ppb, with 87 to 12 99% of the mortality estimate across the 12 urban study areas associated with days in 13 14 this range. A smaller, but still significant fraction (9 to 24%) of the mortality risk is associated with days above 60 ppb (see Figure 7-2, "Existing standard (75)" plot).³² 15 For 2009, this trend continues although risk distributions are shifted down somewhat 16 17 (reflecting the lower ambient O_3 levels generally seen in this simulation year compared with 2007) (see Figure 7-3, "current standard (75)" plot). For 2009, a 18 19 substantial portion (2% to 24%) of O₃-attributable mortality risk is now associated 20 with days having O₃ measurements 55-60 ppb or higher. A relatively smaller fraction 21 (~0% to 2%) of total mortality estimates for the existing standard are associated with days having ambient O₃ levels of 20 ppb or less.³³ 22
- Estimates of O₃- attributable respiratory-related HA range from 10's to 100's of cases
 (after just meeting the existing standard) depending on the type of respiratory HA
 endpoint modeled and the specific urban study areas evaluated (see Table 7-9). All 12
 urban study areas were modeled for one of more respiratory-related HA endpoints.

 ³² Houston has a significantly smaller percentage (<1) of its mortality signal associated with days above 60ppb.
 ³³ In the first draft O₃ REA, we included consideration for surrogate LMLs (based on the lowest composite monitor values used in modeling short-term exposure-related mortality for each urban study area – see Table 7-5, First Draft REA, U.S. EPA, 2012). For the 8hr max monitoring season LML (applicable to the Smith et al.., 2009-based core risk estimate generated for this second draft REA), we have values ranging from 4 to 17 ppb and from 5 to 16 ppb (across the 12 urban study areas for 2007 and 2009, respectively). If we look at heat maps characterizing the distribution of short-term exposure-related mortality for recent conditions (see "recent conditions" heat maps in Figures 7B-1 and 7B-2 in Appendix B) we see that the vast majority of ozone-related mortality falls above these surrogate LML ranges. Consequently we see, that had we integrated consideration for the surrogate LMLs into modeling of short-term exposure-related mortality, there would have been very little change in estimates of risk.

1	•	O ₃ - attributable ER (for respiratory symptoms) ranged into the thousands for both
2		New York and Atlanta under simulated attainment of the existing standard (these
3		were the only two study areas modeled for this health endpoint) (see Table 7-10).
4	•	Estimates of O ₃ -attributable asthma exacerbation (wheeze) in Boston are in the tens
5		of thousands to over 100,000 (see Table 7-11). The percent of baseline for this
6		endpoint (after just meeting the existing standard) is generally in the 20-30% range
7		which is markedly higher than other short-term morbidity endpoints modeled for this
8		analysis (see Table 7-11 and compare to values in 7-9 and 7-10).
9		
10	Long-ter	<u>mO₃-attributable mortality</u>
11	•	After simulating just meeting the existing standard, estimates of O ₃ -related respiratory
12		mortality range across urban areas from 480 to 2,600 deaths (for 2007) and from 490
13		to 2,800 deaths (for 2009) (see Table 7-12). This translates into from 16.3 to 20.8% of
14		baseline across the 12 urban study areas (for 2007) and from 15.9 to 20.7% (for 2009)
15		using the single Jerrett et al., 2009 C-R national-scale function applied to each urban
16		area (see Table 7-13). As discussed in section 7.3.2, because of the long-term
17		exposure metric (seasonal mean of daily 8-hour maximum) employed in risk
18		modeling, there is the potential for some degree of overlap between short-term and
19		long-term exposure-related mortality estimates. For that reason, these two categories
20		of mortality estimates cannot be considered distinct and should not be added to
21		estimate total mortality.
22	•	95 th percentile CIs for long-term O ₃ -attributable respiratory mortality suggest greater
23		power (and potential less heterogeneity) associated with modeling this health
24		endpoint, compared with short-term O3-attributable mortality. None of the CIs for
25		long-term O ₃ -attributable mortality include negative estimates as lower bounds (see
26		Table 7-12).
27	7.5.2 A	Assessment of Health Risk Associated with Simulating Meeting Potential
28	A	Alternative Standards of 70, 65, and 60 ppb
29	A	As discussed earlier, we have considered three alternative standard levels (70, 65 and 60
30	ppb), ead	ch evaluated using the form and averaging time of the existing standard. In presenting
31	risk estir	nates associated with the simulated attainment of each of these alternative standard

levels, we focus on the change in risk associated with a comparison of O₃ levels after simulation
 of the existing standard with levels after simulation of each of the alternative standard levels.
 This is of greatest relevance in comparing the potential public health benefit associated with each
 of the alternative standards relative to the level of protection afforded by just meeting the
 existing standard.

6 In reviewing these risk estimates, it is important to keep in mind that simulation of 7 alternative standard levels is based on a reaching a peak-based attainment metric. Based on the 8 simulated air quality information for the 12 urban study areas, there is a tendency for O_3 to increase on lower concentration days and decrease on higher concentration days.³⁴ Therefore, it 9 10 is not immediately clear that we would expect risk reductions when applying C-R functions that 11 are based on the full distribution of daily 8-hour max values. Specifically, risk reductions are 12 only expected to the extent that the composite monitor daily 8-hour max values decrease as 13 lower alternative standards are simulated. As discussed in Chapter 4 (section ???), after 14 adjustment to alternative standard levels, decreases in O_3 typically occur on higher O_3 days 15 which tend to occur during warmer (summer) months and are concentrated in suburban areas. 16 Conversely, increases in O_3 , typically occur lower O_3 days which tend to occur in the cooler 17 portions of the year and are focused in core urban areas. In general, variability in predicted daily

- 18 O₃ concentrations decreases when meeting lower standard levels.
- 19

20 <u>Short-term O₃-attributable mortality</u>

²² In our analysis, the mortality risk metric is generally not responsive to meeting the • 23 existing and alternative standard levels. This reflects a number of factors all related to 24 1) how O_3 concentrations respond to reductions in NOx emissions used to meet the 25 standards, and 2) how the risk metrics are associated with temporal and spatial 26 patterns of O₃. As discussed in section 7.1.1, mortality risk is modeled using 27 composite monitor values (i.e., averages of O_3 measurements across monitors in an 28 urban study area) which removes spatial variability in measured O₃ within an urban 29 study area (also removing variability in changes in O_3 across an urban area resulting 30 from NOx reductions). Furthermore, in modeling total mortality risk for the core 31 analysis, we add the risk changes occurring across all days within the monitored O_3 32 season, including days with low values of O_3 as well as days with high values of O_3 . 33 This means that we include both decreases in risk on those days when O₃ is estimated 34 to decrease (generally occurring on days with higher values of O_3) and increases in

³⁴ This relationship is also observed in ambient air quality measurements as discussed in Chapter 4 and appendices.

1		risk when O_3 is simulated to increase (generally associated with lower values of O_3).
2		The dampened response of short-term mortality risk can be contrasted with clinical
3		study-based risk estimates. The clinical study-based estimates primarily reflect
4		changes in the upper end of the O ₃ distribution where we tend to see more uniform
5		reductions under simulation of alternative standard levels. In addition, clinical-based
6		estimates of risk are based on detailed micro-environmental exposure modeling which
7		uses individual monitor values instead of composite monitor values, thereby resulting
8		is less dampening of spatial variability in O_3 within a given urban study area.
9	٠	Generally, the magnitude of risk reduction increases as lower alternative standard
10		levels are simulated. For example, for the lowest alternative standard we evaluated,
11		60 ppb, across the 12 urban study areas, we predict from 8 to 160 fewer O_3 -
12		attributable deaths for simulation year 2007 (relative to risk after just meeting the
13		existing standard) (see Table 7-7). This range is from 7 to 160 deaths for simulation
14		year 2009. These ranges (for the 60 ppb standard level) translate into a 3 to 14%
15		reduction in O ₃ -attributable risk relative to risk after just meeting the existing
16		standard (see Table 7-8).
17	•	As noted in section 7.1.1, some of the urban study areas are projected to experience
18		increases in O_3 (and hence risk) when attainment with the existing standard and some
19		of the alternative standard levels is simulated. Focusing specifically on the alternative
20		standard levels, we see that, for the core analysis, this potential increase in risk only
21		occurs for the 2009 simulation year and specifically for three of the urban study areas
22		(Boston, Detroit and Houston – see Table 7-7). For example, Detroit is predicted to
23		have an increase of 21 O_3 -attributable deaths after meeting the 70 ppb standard (when
24		compared to risk remaining after meeting the existing standard). However, we
25		estimate a net reduction of 15 O_3 -attributable deaths after meeting the 60 ppb level
26		(again based on comparison to risk after meeting the existing standard). Furthermore,
27		for all three urban study areas with initial risk increases (based on comparing meeting
28		the existing standard to meeting alternative standards), we see that these increases are
29		offset after meeting the lowest alternative standard simulated (60 ppb) (see Table 7-
30		7). The potential for risk increases is increased somewhat for several of the urban
31		study areas when we simulate how the O ₃ distribution shifts from recent conditions to
32		just meeting the existing standard (see Appendix 7B, Tables 7B-1 and 7B-2).
33		Specifically, in simulating estimated risk from moving from recent conditions to
34		attaining the existing standard, we see that for the 2007 simulation year, two of the

1 study areas (Houston and Los Angeles) have risk increases after meeting the existing 2 standard compared to recent conditions while half of the twelve urban study areas 3 have risk increases for the 2009 simulation year in adjusting air quality to meetthe 4 existing standard relative to recent conditions. It is also important to keep in mind 5 that, for the urban areas of New York and Los Angeles, there are additional 6 uncertainties in the simulation of existing and alternative standards given the 7 limitations in the application of the adjustment methodology to very large emissions 8 perturbations and the fact that the 95th percent confidence interval lower bound 9 estimate of hourly O₃ concentrations was used to capture a scenario in which these 10 cities could meet lower standard levels (65 ppb for New York and 60 ppb for Los Angeles). In five of these eight cases, the initial risk increases (including the increase 11 12 in going from recent conditions to the existing standard) is fully offset after meeting the lowest alternative standard level (60 ppb).³⁵ 13

14 Figure 7-4 provides plots of short-term mortality risk for the existing and alternative • standards adjusting for total exposed population (i.e., O₃-attributable deaths per 15 16 100,000 exposed). From this figure it can be seen that total O₃-attributable risk, even 17 when adjusted for population, varies substantially across the 12 urban study areas, 18 with New York and Philadelphia having the highest risk and Boston and Denver the 19 lowest. This spread in risk (adjusted for population) reflects, to a great extent, 20 differences in the effect estimates used in modeling this endpoint for each study area, 21 which can in turn reflect a number of factors (e.g., differences in behavior such as 22 outdoor activity across cities and differences in exposure measurement error). 23 However, despite considerable variability in absolute O₃-attributable risk, Figure 7-4 24 also suggests that most of the study areas display relatively limited reduction in O₃-25 attributable risk across the three alternative standards (with the exception of New York, which has a notable decrease in risk for the 70 to 65 ppb standard level).³⁶ This 26 27 suggests that a substantial fraction of O₃-attributable risk would still remain, even 28 after simulated attainment of the lowest alternative standard considered.

29 30 • Heat map plots of risk reductions for 2007 suggest that most of the risk reductions associated with simulation of all three alternative standards occur on days with

³⁵ For both LA and Houston (in 2007) and Houston (2009) a modest net risk increase still persists (compared to risk under recent conditions), even when we have simulated the lowest alternative standard considered (60 ppb) (see Tables 7B-1 and 7B-2).

³⁶ With the New York City study area, we recognize however that there is significant uncertainty associated with the use of the CBSA-based study area due to significant heterogeneity in short-term O₃-attributable mortality effect estimates (from Smith et al., 2009) falling within that larger urban study area (see discussion in section 7.6.1).

1	composite O_3 level between 35 and 60 ppb (see Figure 7-4). By contrast, most of the
2	risk increases occur on days with composite O3 levels between 20 and 35ppb (see
3	Figure 7-4). This is expected given that most of the increases in urban core O_3 are
4	associated with lower O ₃ days where NOx titration is prevalent (see Appendix D,
5	section 4.6 Figures 40-54). Very little of the projected change in risk (increases, or
6	decreases) for any of the alternative standards considered occurred on days with O_3
7	levels below 20 ppb O ₃ Similar observations hold for risk results generated for
8	simulation year 2009.

9 <u>Short-term O₃-attributable morbidity</u>

10 Generally, because the short-term O_3 exposure-related morbidity endpoints use the • 11 same air metrics as used in modeling short-term O₃-attributable mortality (i.e., 8hr 12 maximum and 8hr mean) the pattern of risk reduction seen for these morbidity 13 endpoints are similar to those seen with short-term mortality (see Tables 7-9 though 14 7-11 and Figure 7-5). However, New York, as mentioned with regard to short-term 15 O₃-attributable mortality, has substantially higher percent reductions (for O₃-16 attributable risk) compared with the other study areas. For example, with ER visits 17 (asthma), under the lowest alternative standard in simulation year 2007, New York is 18 estimated to have a 22 to 23% reduction in the number of ER-visits associated with 19 O_3 exposure (see Table 7-10).

20 Long-term O₃-attributable mortality

21 Although long-term O₃-attributable mortality is modeled using a different O₃ metric • 22 (essentially a long-term trend in the 1hr maximum for the hottest two seasons - see 23 section 7.3.2) the overall magnitude and pattern of reduction in O_3 -related risk is 24 similar to that seen with short-term exposure related mortality. Specifically, for the 25 2007 simulation year, for most urban study areas risk reductions range from 11 to 26 15% (for the 60 ppb standard) (see Table 7-13). Risk reductions are generally slightly 27 smaller across alternative standard levels for simulation year 2009. For the 2009 28 simulation year, for Detroit, we see a relatively small risk increase for the 70 ppb 29 alternative standard (compared to risk under the existing standard). However that 30 initial increase is offset by risk reductions for the other (lower) alternative standard 31 levels simulated (see Table 7-13).

1 7.5.3 Sensitivity Analyses Designed to Enhance Understanding of the Core Risk Estimates

2 We have completed a number of sensitivity analyses intended to support interpretation of 3 the core risk estimates. These sensitivity analyses, which are described in section 7.4.3, can be 4 divided into two categories: (a) sensitivity analyses exploring factors impacting air quality 5 characterization (specifically composite monitor composition) and (b) sensitivity analyses 6 exploring the impact of alternative C-R function specifications. As noted in section 7.4.3, we 7 also completed an initial influence analysis designed to identify which of the input factors to the 8 risk model (for short-term exposure-related mortality) are primarily responsible for inter-city 9 variability in that risk metric. This section summarizes the results of these sensitivity analyses 10 and presents key observations related to those analyses, beginning with the influence analysis 11 and then proceeding to sensitivity analyses focused on air quality characterization and alternative 12 C-R function specification.

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14 <u>Influence analysis</u>

15 The influence analysis considered three factors involved in modeling risk for the short-16 term exposure-related mortality endpoint including: baseline incidence, composite monitor O_3 17 levels and Bayes-adjusted city-specific effect estimates (recall that the core risk estimate is based 18 on effect estimates derived as part of analyses published in Smith et al., 2009). Each of these 19 input factors displays inter-study area variation and are responsible, collectively, for heterogeneity in risk estimates.³⁷ In completing the analysis, we first calculated a central 20 21 tendency estimate of risk based on the mean of each input factor across the 12 urban study areas 22 for the 2009 simulation year (i.e., using the average of the city-specific values for each of the 23 input factors). We then systematically varied each of the three heterogeneity-related factors 24 (effect estimate, composite monitor-based O_3 level and baseline incidence) to one standard 25 deviation (SD) above its mean value (reflecting variance across the 12 urban study area values) 26 and noted the percent increase produced by that perturbation over the initial mean risk estimate. 27 This influence analysis allowed us to explore the impact of both model form – specifically, 28 potential non-linearities in the model – as well as the relative magnitude of variability in each of 29 the three heterogeneity-related input factors on risk. The influence analysis generated the 30 following results: baseline incidence (23%), composite monitor-based O₃ level (8%), and effect 31 estimate (58%). In other words, the 58% result for *effect estimate* means that use of a value 1 SD

32 over the mean (for the effect estimate) in generating risk, resulted in a risk estimate that was 58%

³⁷ Note, that the demographic count input factor also varies across the study areas and is an important factor in determining total incidence. However, for the influence analysis, we used deaths per 100,000 as the risk metric which standardizes on demographic count and therefore allowed us to exclude this input parameter in conducting the influence analysis.

1 larger than the risk estimate based on the mean of all input factors. These results clearly show

- 2 that, of the three input factors considered, the *effect estimate* is primarily responsible for inter-
- 3 city variability in short-term exposure-related risk.

4 Interestingly, when we look at the coefficient of variation (CV) for these three 5 heterogeneity-related input factors we see values almost identical to the influence analysis results 6 in terms of relative magnitude to each other (i.e., 0.232, 0.084, and 0.527 for baseline incidence, 7 composite monitor-based O₃ levels and effect estimate, respectively). Given that the CV values 8 only reflect variability in each input factor and not model form (i.e., do not reflect potential non-9 linearities in the model), the fact that the CV values almost exactly match the influence analysis 10 results in terms of relative magnitude suggest that there is very little if any non-linearity in the 11 model calculations involving these three input factors. Had non-linearity existed to a significant 12 extent, then the influence anlaysis results would have differed substantially from the CV results. 13 The fact that both analyses suggest a primary role for the effect estimate in driving inter-city 14 variability in risk emphasizes the importance of the sensitivity analyses exploring alternative C-R 15 functions specifications that were completed for the REA (see below). 16 17 Air quality-related sensitivity analyses

18

19 This category of sensitivity analysis covers (a) the use of a smaller study area (the Smith 20 et al., 2009 study areas) as contrasted with the CBSA-based study areas used in the core analysis, 21 and (b) the use of alternative approaches to simulate attainment of the existing and alternative 22 standards (for a subset of the study areas) (see section 7.4.3 for additional detail). This category 23 of sensitivity analysis was applied to short-term O₃-attributable mortality given the importance of 24 the endpoint in the policy-context.³⁸

25 To allow for easier visual comparisons, we have presented the results of this sensitivity 26 analysis category in graphical form (see Figure 7-7, numerical results are presented in Appendix 7C). This figure presents point estimates and 95th percentile confidence ranges for the core model 27 28 and for two sensitivity analyses: (a) SA1 (use of the smaller Smith et al., 2009 based study area) 29 and (b) SA2 (use of the alternative approach to simulating attainment). SA2 is not presented for 30 all of the study areas, only for the subset included in these alternative simulations (see section 31 7.4.3). The sensitivity analyses results presented in Figure 7-7 are the changes in O_3 -related risk 32 that result from meeting the three alternative standards relative to meeting the existing standard. 33 Furthermore, these changes reflect deaths per 100,000, which standardizes the estimates on

 $^{^{38}}$ Observations regarding the sensitivity of core short-term O₃-attributable mortality risk to these sensitivity analyses can be applied with care to the core short-term O₃-attributable morbidity endpoints, since many of these used similar air quality metrics in modeling risk.

population. This removes variation in the size of the underlying exposed population as a factor to
 consider in interpreting these results.

For the sensitivity analysis examining use of the smaller Smith et al., 2009 study area, we have also included heat maps similar to those used in conveying core estimates for short-term exposure related mortality (see section 7.5 for a description of the heat maps used in the core analysis). These heat maps (included in Appendix C – see Figure 7C-1) allow us to consider how changes in risk, including both reductions in risk and increases in risk are distributed across the O₃ air quality distributions for each study area.

9 10 Key observations related to the air quality-related sensitivity analyses include:

11 Use of smaller study area reduces magnitude of risk reduction: For most of the • 12 study areas, use of the smaller Smith et al., 2009-based study area resulted in smaller 13 risk reductions (again expressed in terms of changes in deaths per 100,000). For 14 example, in Figure 7-7 (Baltimore plot), we see that estimated change in risk for SA1 15 (the smaller study area) are lower than estimated change in risk for the core scenario. 16 This likely reflects the mix of monitors in the smaller study areas which results in a 17 smaller change in the composite monitor value (for the existing standard versus 18 alternative standard levels) as compared with composite monitor values based on the 19 larger CBSA study area. However, it is important to keep the relative small 20 magnitude of these risk reductions in mind when considering these sensitivity 21 analysis results. Most of these differences in risk reductions are less than 1 individual 22 per 100,000 which reflects the fact that total risk reduction (for short-term O₃-23 attributable mortality) across the urban study areas is relatively small (see Table 7-7).

24 \circ Reductions in risk are focused on higher O₃ days while increases 25 are focused on lower O₃ days: Figure 7C-1 allows us to consider patterns 26 in risk reductions and increases when using the smaller Smith et al., 2009-27 based study areas in modeling risk. Figure 7C-1 (particularly the plots of 28 risk decreases) suggests that decreases in risk tend to occur on days with 29 composite monitor O₃ concentrations ranging from 40-70ppb, while 30 increases in risk tend to occur on days with composite monitor values in the range at or below 30-40 ppb (with most risk increases falling in the 31 32 range of 15ppb to 40ppb). As noted in 7.1.1, there is less confidence in 33 specifying the nature of the C-R function (and therefore less confidence in 34 specifying risk) in the range below 20 ppb.

- 1 Application of effect estimates derived for smaller study areas to larger CBSA-• 2 **based study areas:** As noted in section 7.3.2, in those instances where an 3 epidemiological study provides effect estimates for multiple subareas within a larger 4 CBSA-based study area, we are selecting the effect estimates that represent the 5 largest number of individuals to model that CBSA-based study areas. There is 6 uncertainty associated with this approach. Specifically, as illustrated in Table 7-3, 7 effect estimates within some of the CBSA-based study areas can display considerable 8 heterogeneity. For example, consider the Smith et al., 2009-based effect estimates 9 that fall within the CBSA-based New York study areas (these vary from 0.0001 to 10 0.0009 – almost a 10 fold factor, see Table 7-3). Furthermore, with the CBSA-based 11 New York study area, Smith et al., 2009-based effect estimates only cover about half 12 of the total population, with 8.3 million residents living within portions of the CBSA 13 not covered by Smith et al., 2009-based effect estimates. As noted in section 7.3.2, in 14 these types of situations, we have decided to use the single effect estimates 15 representing the largest number of residents in modeling the larger CBSA-study area. 16 This reflects the observation that, in the case of the New York CBSA, one of the 17 available effect estimates (for the New York study area), represents ~7 times the 18 population of the other effect estimates (see Table 7-3). In the case of the Los 19 Angeles CBSA, there is significantly less difference between the available effect 20 estimates, making the issue of heterogeneity (and the specification of a single effect 21 estimate for this study area) less important. Never the less, we recognize that the issue 22 of heterogeneity does complicate extrapolation of effect estimates for smaller study 23 areas to the larger CBSA study areas modeled in this analysis and does introduce a 24 degree of uncertainty that is difficult to characterize. 25 • Use of alternative approach for simulating attainment of existing and alternative 26 standard levels: Use of an alternative approach to simulate attainment of the existing 27 and alternative standard levels did not produce a consistent trend in terms of changes 28 in risk between existing and alternative standards relative to the core analysis. For 29 example, if we look at Figure 7-7 (plot for Houston), we see that SA2 (reflecting
- 30 application of the alternative simulation approach) has a larger risk reduction than the 31 core estimate. By contrast, if we look at the plot for Los Angeles, we see that the SA2 32 risk change is lower than the core estimate. Again, as with the sensitivity analysis 33 results looking at study area size, it is important to keep in mind that the magnitude of 34 these differences is relatively small, reflecting the small magnitude of mortality risk 35 associated with these analyses in general (see Table 7-7). It is also important to note 36 that in the alternative simulation approach, the HDDM-adjustment approach assumed

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Sensitivity analyses related to specification of C-R functions

9 10 This category of sensitivity analysis covers a number of factors related to the 11 specification of C-R functions for both short-term O₃-attributable and long-term O₃-attributable 12 mortality. In the case of short-term O_3 -attributable mortality, we consider (a) the use of Bayes 13 adjusted effect estimates using regional priors (as contrasted with the Bayes adjusted values 14 using a national prior applied in the core analysis), (b) the use of a copollutants model 15 considering PM_{10} (as contrasted with the single pollutant model used in the core analysis) and (c) 16 application of effect estimates from Zanobetti and Schwartz 2008 reflecting a summer focused 17 analysis (as contrasted with the Smith et al., 2009-based analysis reflecting the entire monitoring 18 period in each study area, which is used in the core analysis). For long-term O₃-attributable 19 mortality, we consider the use of regionally-differentiated single pollutant effect estimates 20 obtained from Jerrett et al., 2009, as contrasted with the single national copollutants model used 21 in the core analysis (see section 7.1.1). We also present estimates for long-term O₃ attributable 22 mortality based on application of results from a national level single pollutant model. 23 For sensitivity analyses examining alternative specification of the C-R function for short-24 term O_3 -attributable mortality, we have used the same graphical approach as used in presenting 25 results of the sensitivity analyses examining air quality characterization (i.e., plots of point estimates with 95th percentile C.I.s for the core and sensitivity analyses for each of the study 26 27 areas – see Figure 7-8). Here we also plot estimates of risk changes using deaths per 100,000 to 28 standardize in terms of total exposed population. For the sensitivity analysis considering

the same percent reductions of NOx and VOC and did not examine if a different air

quality distribution could have been obtained with a different combination of NOx

reductions were very similar to the NOx-only percent reductions. The similarity in the

NOx reductions between the two approaches could be the reason for there being little

versus VOC reductions. For most of the urban areas, the percent NOx and VOC

difference in the risk estimates between the core and the alternative approach.

- 29 alternative C-R functions for long-term O₃-attributable mortality, we present results in tabular
- 30 form (Table 7-14). Specifically, for both the core and sensitivity analysis, we present (a) the
- 31 percent of baseline mortality attributable to O_3 (under simulated attainment of the existing
- 32 standard) and (b) the percent reduction in O₃-attributablerisk for each of the alternative standard
- 33 levels. Key observations related to sensitivity analyses examining alternative C-R functions
- 34 specifications include:
- 35 36
- Use of regional Bayes-adjusted effect estimates in modeling short term O₃attributable mortality: The use of Bayes-adjusted effect estimates with regional

- 1 priors in modeling short-term O₃-attributable mortality, had a mixed impact across the 2 urban study areas, with some study areas having increased changes in risk and others 3 having smaller changes, relative to the core analysis. For example, in Figure 7-8 (plot 4 for Baltimore), SA1 had a larger change in risk compared with the core analysis. 5 However, as with the sensitivity analyses examining air quality-related factors 6 (discussed above), it is important to keep in mind that the overall magnitude of the 7 O₃-attributable mortality risk is relatively small and that these differences in changes 8 in risk (comparing SA1 to the core analysis) are generally in the fraction of a person 9 per 100,000 exposed population.
- 10 • Use of a copollutants model (with PM₁₀) in modeling short-term O₃-attributable 11 **mortality:** The use of the PM₁₀ copollutant model in modeling short-term O₃-12 attributable mortality (as contrasted with the single pollutant model used in the core 13 analysis) tended to have a relatively small effect on estimates of risk changes for the 14 alternative standards considered. For example, in Figure 7-8 (plot for Boston), we see 15 that the estimates of risk changes for SA2 (reflecting application of the PM_{10}) 16 copollutant model) is essentially the same as the core risk estimate. It is important to 17 keep in mind that the PM_{10} copollutant model suffers from significantly reduced 18 power due to the 1/3 to 1/6 day sampling frequency used in measuring PM₁₀ (this 19 reduces the number of observations available to support epidemiological analysis). 20 This has the impact of greatly increasing the confidence intervals on the SA2 risk 21 estimates relative to the core estimates. 22
- Use of Zanobetti and Schwartz 2008 effect estimates in modeling short-term O₃-23 attributable mortality: The use of Zanobetti and Schwartz, 2008 effect estimates 24 (reflecting a focus on the warmer summer months) produces a mixed set of results 25 when compared to the core risk estimates. If we look at Figure 7-8 we see that, for 26 Boston, estimates of risk changes for SA3 (reflecting application of the Zanobetti and 27 Schwartz 2008 effect estimates) are significantly larger than core estimates. By 28 contrast, SA3 estimates of risk changes for Houston are significantly smaller than the 29 core estimates. It is important, however to keep in mind that the Zanobetti and 30 Schwartz 2008 effect estimates will tend to under-estimate total risk since they only 31 model impacts during the summer months (while the Smith et al., 2009 effect estimates allow us to model impacts for the entire O₃ monitoring season in each study 32 33 area). Note that if the O_3 effect were only occurring during the summer months, then 34 the total risk estimated using effect estimates from the two studies would be similar. 35 However, because the risks in many locations are smaller (using the Zanobetti and

1	Schwartz, 2008 based effect estimates), this suggests that the O ₃ effect occurs outside
2	of the summer months evaluated in this study.
3 •	Use of regional-differentiated effect estimates in modeling long-term O_3 -
4	attributable mortality: Risk estimates generated using regional-specific effect
5	estimates for long-term O_3 -attributable mortality differ substantially from the core
6	estimates based on a single national-level effect estimate (see Table 7-14).
7	Furthermore, the risk estimates generated using the regional effect estimates display
8	considerable variability (see Table 7-14) reflecting the significant variability in the
9	underlying effect estimates (see Jerrett et al., 2009, Table 4). The regional effect
10	estimates range from 0.99 (for the Northeast) to 1.21 (for the Southwest) and include
11	1.00 (no O_3 effect for the Industrial Midwest). As noted earlier in section 7.5,
12	negative risk estimates should not be interpreted as suggesting that O ₃ exposure is
13	beneficial. Rather, these suggest that there may be instability in the underlying
14	estimates or that potential confounding has not been fully addressed. Regional effect
15	estimates used in this analysis have considerably larger confidence intervals than the
16	national estimate (compare values in Jerrett et al., 2009 Table 3 with values in Table
17	4). This suggests that the regional estimates are less stable than the national estimates
18	and are subject to considerably greater uncertainty. For this reason, while the results
19	of this sensitivity analysis point to the potential for regional heterogeneity in the long-
20	term O ₃ -attributable mortality effect estimate, we do not have significant confidence
21	in the regionally-based risk estimates themselves given the relatively large confidence
22	intervals associated with those estimates.
23 •	Use of national-based single pollutant model in modeling long-term O_3 -
24	attributable mortality: Risk estimates generated using the national-level O ₃ -only
25	effect estimate were significantly lower (~30%) than the core risk estimates which
26	utilize a copollutants model (which includes $PM_{2.5}$) (see Table 7-15). In this case,
27	control for another pollutant results in a stronger O_3 signal, possibly due to an
28	association between $PM_{2.5}$ and a confounder or effect modifier associated with the
29	O ₃ -related effect.

Figure 7-7 Sensitivity Analysis: Short-Term O₃-attributable Mortality (air quality-related factors including study area size and method used to simulate attainment of existing and alternative standard levels) (2009) SA1-smaller (Smith-based) study area, SA2-alternative method for simulating standards.



Figure 7-8 Sensitivity Analysis: Short-Term O₃-attributable Mortality (C-R function specification) (2009) SA1-regional Bayesbased adjustment; SA2-copollutant model (PM₁₀); SA3-Zanobetti and Schwartz-based effect estimates.



Table 7-14 Sensitivity Analysis for Long-Term 0₃-attributable Respiratory Mortality – Alternative C-R Function Specification (regional effect estimates) % of baseline all *cause mortality and change in 0₃-attribuable risk* (2009) (Smith et al., 2009, 0₃ season))

	Air Quality Scenario					
	Baseline					
	Incidence					
	Attributable to					
	Ozone	Change i	n O₃-Attribut	able Risk		
Study Area	75ppb	75-70	75-65	75-60		
	Core Si	mulation				
Atlanta, GA	16.6	5	10	14		
Baltimore, MD	17.4	3	6	10		
Boston, MA	15.9	1	3	7		
Cleveland, OH	16.8	4	9	15		
Denver, CO	20.0	1	5	12		
Detroit, MI	17.0	-1	2	6		
Houston, TX	16.9	2	4	7		
Los Angeles, CA	20.7	4	8	13		
New York, NY	16.7	5	18	18		
Philadelphia, PA	17.2	3	7	10		
Sacramento, CA	18.0	4	8	12		
St. Louis, MO	17.7	4	8	13		
	Sensitivi	ty analysis				
Atlanta, GA	41.21	4	8	11		
Baltimore, MD	-7.01	4	9	13		
Boston, MA	-6.19	1	5	9		
Cleveland, OH	0.00	0	0	0		
Denver, CO	27.38	1	4	11		
Detroit, MI	0.00	0	0	0		
Houston, TX	41.15	2	3	6		
Los Angeles, CA	4.46	3	6	10		
New York, NY	-6.61	7	24	NA		
Philadelphia, PA	-6.89	4	9	13		
Sacramento, CA	24.79	4	7	11		
St. Louis, MO	0.00	0	0	0		

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NA: for NYC, the model-based adjustment methodology was unable to adjust O_3 distributions such that they would meet the lower alternative standard level of 60 ppb.

Table 7-15 Sensitivity Analysis for Long-Term O₃-attributable Respiratory Mortality – Alternative C-R Function Specification (national O₃-only effect estimates) % of baseline all-cause mortality and change in O₃-attribuable risk (2009) (Smith et al., 2009, O₃ season))

	Air Quality Scenario					
	Percent of Baseline Incidence	Change in	O3-Attribu	ıtable Risk		
Study Area	75ppb	75-70	75-65	75-60		
	Core Sim	ulation				
Atlanta, GA	16.6	5	10	14		
Baltimore, MD	17.4	3	6	10		
Boston, MA	15.9	1	3	7		
Cleveland, OH	16.8	4	9	15		
Denver, CO	20.0	1	5	12		
Detroit, MI	17.0	-1	2	6		
Houston, TX	16.9	2	4	7		
Los Angeles, CA	20.7	4	8	13		
New York, NY	16.7	5	18	18		
Philadelphia, PA	17.2	3	7	10		
Sacramento, CA	18.0	4	8	12		
St. Louis, MO	17.7	4	8	13		
	Sensitivity	analysis				
Atlanta, GA	11.9	5	10	14		
Baltimore, MD	12.2	3	7	10		
Boston, MA	11.1	1	3	7		
Cleveland, OH	11.8	4	10	15		
Denver, CO	14.1	2	5	12		
Detroit, MI	11.9	-1	2	6		
Houston, TX	11.9	2	4	7		
Los Angeles, CA	14.6	4	9	14		
New York, NY	11.7	5	19	19		
Philadelphia, PA	12.0	3	7	10		
Sacramento, CA	12.6	4	8	13		
St. Louis, MO	12.4	4	8	13		

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NA: for NYC, the model-based adjustment methodology was unable to adjust O_3 distributions such that they would meet the lower alternative standard level of 60 ppb.

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7.6 KEY OBSERVATIONS REGARDING OVERALL CONFIDENCE IN THE RISK ASSESSMENT AND RISK ESTIMATES

This section discusses our overall confidence associated with risk estimates presented in 3 4 this draft of the REA. We begin by presenting a set of key observations related to overall 5 confidence in the risk assessment. These observations are drawn largely from (a) consideration 6 for the systematic approach used in designing the risk assessment, (b) our assessment of the 7 degree to which we have captured key sources of variability in the analysis (section 7.4.1) (c) our 8 qualitative assessment of uncertainty in the risk assessment (section 7.4.2), and (d) the results of 9 the sensitivity analyses completed (section 7.5.3). Once we present these observations, we 10 provide a synthesis statement reflecting our overall degree of confidence in the risk estimates (at 11 the end of this section). Key observations addressing overall confidence in the analysis include:

- 12 • A deliberative process was used in specifying each of the analytical elements 13 comprising the risk model. This is in line with recommendations made by the 14 National Research Council in Science and Decisions, Advancing Risk Assessment 15 (NRC, 2009. P. 89-90) for improving risk assessment as applied in the regulatory 16 context. This deliberative process included first identifying specific goals for the 17 analysis, and then designing the analysis to meet those goals, given available 18 information and methods. Specific analytical elements reflected in the design include: 19 selection of urban study areas, characterization of ambient air O_3 levels, selection of 20 health endpoints to model and selection of epidemiological studies (and specification 21 of C-R functions) (see sections 7.1.1 and 7.3). In addition, the design of this draft of 22 the REA reflects consideration for comments provided by the public and by CASAC in their review of the 1st draft REA in letter form (Frey, H. C. 2012.). 23
- 24 • Review of available literature (as specified in the O₃ ISA, U.S. EPA. 2013a), resulted 25 in a decision not to incorporate a true (no effect) threshold into our risk modeling. 26 Conversely, the studies used to develop the C-R functions indicate a range of ambient 27 O₃ (area-wide daily levels, based on averaging across monitors in locations with 28 multiple monitors, of \leq 20ppb) below which there is reduced confidence in specifying 29 the nature of the concentration-response relationship, based on less data in the studies, 30 specifically for short-term O_3 -attributable respiratory mortality and morbidity (see 31 section 7.1.1). In any case, only a relatively small fraction of short-term O₃-32 attributable mortality reflected in the risk estimates is associated with days in this 33 range with the vast majority of the risk estimates reflecting days with peak O_3 34 measurements well above this level (see section 7.5.1 and 7.5.2). O₃ 35 Modeling of short-term O₃-attributable mortality utilized Bayes-adjusted city-specific
- 36

effect estimates (see section 7.1.1 and section 7.3.2). These effect estimates are

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considered to have increased overall confidence since they combine elements of the local city-specific signal with a broader scale (national) signal.

- 3 Use of CBSA-based study areas in modeling all health endpoints in order to address 4 known bias associated with using smaller study areas. As discussed in 7.1.1, we have 5 used larger CBSA-based study areas to avoid focusing the risk assessment only on 6 core urban areas (often used in the epidemiological studies providing effect estimates) 7 which can experiences increases in O_3 based on simulated attainment of both existing 8 and alternative standard levels. There is uncertainty in using effect estimates based on 9 smaller study areas to represent larger CBSA-based study areas (see section 7.4.2 and 10 7.5.3). A key concern is heterogeneity in the effect estimates which may suggest 11 increased uncertainty in applying effect estimates to larger study areas (since larger 12 study areas may display heterogeneity in the nature of the relationship between O_3 13 exposure and risk). It is possible also that this heterogeneity varies across urban areas, 14 or regionally. For both categories of mortality endpoints (short-term and long-term 15 O₃-attributable), potential heterogeneity in the mortality effect even within larger 16 urban areas remains a potentially important source of uncertainty.
- 17 Specifically in relation to short-term exposure-related mortality and morbidity which • 18 depend on time-series studies, there is uncertainty in applying effect estimates derived 19 based on evaluating the longitudinal (in terms of time) relationship between ambient 20 O_3 and a particular health effect to the modeling of a discrete shift in the entire 21 distribution that occurs when you simulate an alternative standard. Specifically, the 22 time-series studies relate unit changes in day to day O_3 with a degree of impact on 23 baseline health effect rates. In the risk assessment, we use this effect estimate to 24 predict risk for a unit shift in daily composite monitor value. There is uncertainty in 25 this application of the effect estimates, although it is not possible at this time to 26 characterize either qualitatively, or quantitatively the magnitude of this uncertainty 27 and the degree of any potential bias that could be introduced into the simulation of 28 risk.
- 29 Use of HDDM-adjustment approach to simulate attainment of both the existing and • 30 alternative standard levels provides more refined estimates of ambient O₃ 31 distributions given its ability to characterize the physical and chemical processes of 32 O₃ formation in the atmosphere However, in the case of both the New York and Los 33 Angeles study areas, given the limitations in the application of the adjustment 34 methodology to very large emissions perturbations and the need to use the 95th 35 percent confidence interval lower bound estimate to simulate attainment of these 36 standard levels, we have reduced overall confidence in the simulation of the O₃

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1	concentrations for these study areas and consequently all health endpoints modeled
2	for risk for these two study areas (see section 7.4.2 and 7.5.3).
3	• Sensitivity analyses exploring alternative C-R functions for modeling <u>short-term O₃-</u>
4	attributable mortality (e.g., Bayes regional prior based estimates, copollutants
5	models) suggested that alternative models can have a moderate impact on risk (see
6	section 7.5.3). This modest impact reflects primarily the relatively small magnitude of
7	short-term O ₃ -attributable mortality reductions simulated for the alternative standard
8	levels.
9	• The use of alternative C-R functions for modeling <u>long-term</u> O ₃ -attributable mortality
10	(specifically the regional-based estimates referenced earlier) was shown to have a
11	significant impact on risk (see section 7.5.3). However, concerns over the power and
12	hence stability of the regional effect estimates used in this simulation limit our ability
13	to draw firm conclusions regarding the potential magnitude of that regional
14	heterogeneity.
15	
16	Based on the key observations regarding confidence presented above, we draw the
17	following conclusions regarding overall confidence in the risk estimates generated for this draft
18	of the REA. We have a reasonable degree of confidence in <u>short-term</u> O ₃ -attributable mortality
19	and morbidity estimates for ten of the twelve study areas. This confidence is tempered somewhat
20	by concerns over potential heterogeneity in effect estimates for mortality which can impact the
21	risk assessment given our use of larger CBSA-based study areas. Our confidence in risk
22	estimates generated for both New York and Los Angeles is considerably lower than for the
23	remaining ten study areas due to (a) concerns over air quality modeling (specifically the use of
24	lower-bound fits to the DDM model) and (b) specifically in the case of New York, evidence for
25	significant heterogeneity in the mortality effect estimates for subareas within the CBSA. For
26	long-term O ₃ -attributable mortality, we also have a reasonable degree of confidence in our risk
27	estimates. However, as with short-term O3-attributable mortality, this confidence is also
28	tempered by concerns over regional heterogeneity in the O_3 effect. If we had regionally-
29	differentiated effect estimates for this endpoint that had sufficient power and stability, we would
30	consider using these as the basis for generating core risk estimates (rather than the national-level
31	effect estimate used in the current analysis).
32	
1 7.7 REFERENCES

3	Abt Associates Inc. 1996. A Particulate Matter Risk Assessment for Philadelphia and Los Angeles.
4	Prepared for Office of Air Quality Planning and Standards. Research Triangle Park, NC: EPA
5	Office of Air and Radiation, OAQPS, July 1996. Available at:
6	http://www.epa.gov/ttnnaqs/standards/pm/data/jly3amd.pdf >.
7 8 9 10	Abt Associates, Inc. 2010. <i>Environmental Benefits and Mapping Program (Version 4.0)</i> . Prepared for U.S. Environmental Protection Agency, Bethesda, MD. Research Triangle Park, NC: EPA Office of Air Quality Planning and Standards. Available on the Internet at: http://www.epa.gov/air/benmap >.
11	Akinbami, LJ; C. D. Lynch; J. D. Parker and T. J. Woodruff. 2010. "The Association Between Childhood
12	Asthma Prevalence and Monitored Air Pollutants in Metropolitan Areas, United States, 2001-
13	2004." <i>Environmental Research</i> , 110: 294-301.
14	Bell, M. L. and F. Dominici. 2008. "Effect Modification by Community Characteristics on the Short-term
15	Effects of O ₃ Exposure and Mortality in 98 U.S. Communities." <i>American Journal of</i>
16	<i>Epidemiology</i> , 167: 986-997.
17 18	Bell, M. L.; A. McDermott; S. L. Zeger; J. M. Samet; F. Dominici. 2004. "O ₃ and Short-term Mortality in 95 U.S. Urban Communities, 1987-2000." <i>JAMA</i> , 292: 2372-2378.
19	Darrow, L. A.; M. Klein; J. A. Sarnat; J. A. Mulholland; M. J. Strickland; S. E. Sarnat, et al. 2011. "The
20	Use of Alternative Pollutant Metrics in Time-series Studies of Ambient Air Pollution and
21	Respiratory Eemergency Department Visits." <i>Journal of Exposure Science and Environmental</i>
22	<i>Epidemiology</i> , 21, 10-19.
23 24 25 26 27	 Frey, H. C. 2012. "Letter from Clean Air Scientific Advisory Committee to the Honorable Lisa P. Jackson, Administrator, US EPA. CASAC Review of the EPA's Health Risk and Exposure Assessment for O₃ (First External Review Draft - Updated August 2012) and Welfare Risk and Exposure Assessment for O₃ (First External Review Draft - Updated August 2012)," November, 19, 2012.
28 29 30	 Gent, J. F.; E. W. Triche; T. R. Holford; K. Belanger; M. B. Bracken; W. S. Beckett, et al. 2003. "Association of Low-level O₃ and Fine Particles with Respiratory Symptoms in Children with Asthma." <i>Journal of the American Medical Association</i>, 290(14), 1859-1867.
31	Ito, K.; G. D. Thurston and R. A. Silverman. 2007. "Characterization of PM _{2.5} , Gaseous Pollutants, and
32	Meteorological Interactions in the Context of Time-series Health Effects Models." <i>Journal of</i>
33	<i>Exposure Science and Environmental Epidemiology</i> , 17 Suppl 2, S45-60.
34	Jerrett, M.; R. T. Burnett; C. A. Pope, III; K. Ito; G. Thurston; D. Krewski; Y. Shi and E. Calle; M. Thun.
35	2009. "Long-term O ₃ Exposure and Mortality." <i>New England Journal of Medicine</i> , 360:1085-
36	1095.
37	Katsouyanni, K.; J. M. Samet; H. R. Anderson; R. Atkinson; A. L. Tertre; S. Medina, et al. 2009. "Air
38	Pollution and Health: A European and North American Approach (APHENA)," Health Effects
39	Institute.
40	Lin, S.; E. M. Bell; W. Liu; R. J. Walker; N. K. Kim and S. A. Hwang. 2008a. "Ambient O ₃
41	Concentration and Hospital Admissions Due to Childhood Respiratory Diseases in New York
42	State, 1991–2001." <i>Environmental Research</i> , 108, 42-47.
43 44	Lin, S; X. Liu; L. H. Le; S. A. Hwang. 2008. "Chronic Exposure to Ambient O ₃ and Asthma Hospital Admissions Among Children." <i>Environonmental Health Perspective</i> , 116: 1725-1730.

- Linn, W. S.; Y. Szlachcic; H. Gong, Jr.; P. L. Kinney and K. T. Berhane. 2000. "Air Pollution and Daily Hospital Admissions in Metropolitan Los Angeles." *Environmental Health Perspective*, 108(5), 427-434.
 Medice Demon Mark. Zenchetti and L. Schwartz. 2006. "The Effort of O. and DM. on Heavital."
- Medina-Ramon, M.; A. Zanobetti and J. Schwartz. 2006. "The Effect of O₃ and PM₁₀ on Hospital
 Admissions for Pneumonia and Chronic Obstructive Pulmonary Disease: A National Multicity
 Study." *American Journal of Epidemiology*, 163(6), 579-588.
- Meng, Y. Y.; R. P. Rull; M. Wilhelm; C. Lombardi; J. Balmes and B. Ritz. 2010. "Outdoor Air Pollution
 and Uncontrolled Asthma in the San Joaquin Valley, California." *Journal of Epidemiology Community Health*, 64: 142-147.
- Moore, K; R. Neugebauer; F. Lurmann; J. Hall; V. Brajer; S. Alcorn; I. Tager. 2008. "Ambient O₃
 Concentrations Cause Increased Hospitalizations for Asthma in Children: An 18-year Study in
 Southern California." *Environmental Health Perspective*, 116: 1063-1070.
- National Research Council (NRC). 2009. Science and Decisions, Advancing Risk Assessment. Committee
 on Improving Risk Analysis Approaches. Washington, DC: The National Academies Press.
- Silverman, R. A.; and K. Ito. 2010. "Age-related Association of Fine Particles and O₃ with Severe Acute
 Asthma in New York City." *Journal of Allergy Clinical Immunology*, 125(2), 367-373 e365.
- Smith, R.L.; B. Xu and P. Switzer. 2009. "Reassessing the Relationship Between O₃ and Short- term
 Mortality in U.S. Urban Communities." *Inhalation Toxicology*, 21: 37-61.
- Strickland, M. J.; L. A. Darrow; M. Klein; W. D. Flanders; J. A. Sarnat; L. A. Waller, et al. 2010. "Short-term Associations between Ambient Air Pollutants and Pediatric Asthma Emergency Department Visits." *American Journal of Respiratory Critical Care Medicine*, 182, 307-316.
- Tolbert, P. E.; M. Klein; J. L. Peel; S. E. Sarnat and J. A. Sarnat. 2007. "Multipollutant Modeling Issues
 in a Study of Ambient Air Quality and Emergency Department Visits in Atlanta." *Journal of Exposure Science and Environmental Epidemiology*, 17 Suppl 2, S29-35.
- U.S. Center for Disease Control. 2010. "Centers for Disease Control and Prevention, Behavioral Risk
 Factor Surveillance System (BRFSS), 2010, Table "Table C1 Adult Self-Reported Current
 Asthma Prevalence Rate (Percent) and Prevalence (Number) by State or Territory." Available at:
 http://www.cdc.gov/asthma/brfss/2010/current/tableC1.htm
- U.S. Environmental Protection Agency. 2001. *Risk Assessment Guidance for Superfund. Vol. III, Part A. Process for Conducting Probabilistic Risk Assessment (RAGS 3A).* Washington, DC: EPA. (EPA document number EPA 540-R-02-002; OSWER 9285.7-45; PB2002 963302). Available at:
 .
- U.S. EPA. 2004. *EPA's Risk Assessment Process for Air Toxics: History and Overview*. In: Air Toxics
 Risk Assessment Reference Library, Technical Resource Manual, Vol. 1., pp. 3-1 3-30. (EPA
 document number EPA-453-K-04-001A). Washington, DC: EPA. Available at:
 http://www.epa.gov/ttn/fera/data/risk/vol_1/chapter_03.pdf)
- U.S. EPA. 2007. O₃ *Health Risk Assessment for Selected Urban Areas*. Research Triangle Park, NC: EPA
 Office of Air Quality Planning and Standards. (EPA document number EPA 452/R-07-009).
 Available at: http://www.epa.gov/ttn/naaqs/standards/O3/s_O3_cr.html.
- 40 U.S. EPA. 2011. O₃ National Ambient Air Quality Standards: Scope and Methods Plan for Health Risk
 41 and Exposure Assessment. Research Triangle Park, NC: EPA. (EPA document number EPA 42 452/P-11-001).

1 2 U.S. EPA. 2012. Health Risk and Exposure Assessment for O₃ First External Review Draft. Research Triangle Park, NC: U.S. Environmental Protection Agency, Research Triangle Park, NC. (EPA 3 document number EPA 452/P-12-001). 4 U.S. EPA. 2013a. Integrated Science Assessment for O₃ and Related Photochemical Oxidants: Final. 5 Research Triangle Park, NC: U.S. Environmental Protection Agency. (EPA document number 6 EPA/600/R-10/076F). 7 U.S. EPA. 2013b. Environmental Benefits Mapping and Analysis Program—Community Edition (Version 8 0.63), 2013a. Research Triangle Park, NC: U. S. Environmental Protection Agency. Available at: 9 <hhtp://www.epa.gov/air/benmap/beta.html>. 10 World Health Organization. 2008. Part 1: Guidance Document on Characterizing and Communicating 11 Uncertainty in Exposure Assessment, Harmonization Project Document No. 6. Published under 12 joint sponsorship of the World Health Organization, the International Labour Organization and 13 the United Nations Environment Programme. WHO Press, World Health Organization, 20 14 Avenue Appia, 1211 Geneva 27, Switzerland (tel.: +41 22 791 2476). 15 Zanobetti, A; J. Schwartz. 2008. "Mortality Displacement in the Association of O₃ with Mortality: An 16 Analysis of 48 Cities in the United States." American Journal of Respiratory and Critical Care 17 Medicine, 177: 184-189.

7-89

18NATIONAL SCALE MORTALITY RISK BURDEN BASED ON2APPLICATION OF RESULTS FROM EPIDEMIOLOGICAL STUDIES

3 As described in Chapter 2, the O_3 ISA (U.S. EPA 2013) concluded that there is likely to 4 be a causal relationship between short-term O_3 exposure and all-cause mortality and that there is 5 likely to be a causal relationship between long-term O₃ exposure and respiratory effects, 6 including respiratory mortality. Chapter 7 estimated health risks associated with recent O₃ 7 concentrations and meeting the current and alternative O₃ standards in 12 selected urban study 8 areas. In this chapter we estimate nationwide premature mortality attributable to recent short-9 term and long-term exposures to ambient O_3 (Section 8.1); and assess the degree to which the 10 selected urban case study areas represent the full national distribution of risk-related attributes 11 and air quality dynamics (Section 8.2). Compared with the urban scale analysis in Chapter 7, this 12 analysis includes full spatial coverage across the U.S. but has less geographic specificity in the 13 concentration-response functions that are used to calculate O₃.attributable mortality. The national 14 scale analysis is therefore intended as a complement to the urban scale analysis, providing both a 15 broader assessment of O_3 -related health risks across the U.S. as well as an evaluation of how 16 well the urban study areas examined in Chapter 7 represent the full distribution of O₃-related 17 health risks and air quality dynamics in the U.S.

18

19 8.1 NATIONAL-SCALE ASSESSMENT OF MORTALITY RELATED TO O₃ 20 EXPOSURE

21 This section estimates the total annual deaths for 2007 populations associated with 22 average 2006-2008 O₃ levels across the continental U.S. We first describe the methods and 23 inputs used to estimate O_3 -attributable risk across the continental U.S., including O_3 exposure 24 estimates, population and baseline mortality rate estimates, and epidemiologically derived O₃-25 mortality effect estimates. Results for the estimation of O₃-attributable risk are then discussed in 26 terms of the magnitude and percent of total mortality attributable to O₃ exposure. We provide 27 two analyses to give perspective on the confidence in the estimates of O_3 -related mortality: (1) 28 risk estimated only within the urban areas for which O₃ mortality effect estimates are available; 29 and (2) the distribution of O_3 -related deaths across the range of observed 2006-2008 average O_3 30 concentrations fused with modeled 2007 concentrations. These results are then synthesized and 31 compared with previous estimates of the burden of O₃ exposure on mortality in the U.S. from the 32 literature in a discussion section.



2 Figure 8.1 Conceptual Diagram for National-scale Mortality Risk Assessment

1

4 **8.1.1 Methods**

5 This section describes the inputs and datasets used to conduct the national-scale 6 assessment of O₃-attributable risk. As shown in the conceptual diagram in Figure 8-1, we 7 conduct this analysis using the BenMAP software, which uses projections of the size and 8 geographic distribution of the potentially exposed population along with estimates of the ambient 9 O_3 concentrations to estimate O_3 -attributable health risks. In general, this analysis uses the same 10 analytical structure and many of the same inputs as are used in the epidemiology-based 11 assessment of O₃-attributable risk in the selected urban case study areas in Chapter 7. We refer 12 back to Chapter 7 for details on these shared inputs, and describe where the urban-scale and 13 national-scale analyses use divergent methods. 14

15

16

8.1.1.1 Ambient O₃ Concentrations

1	Air quality inputs to this analysis are described in detail in Chapter 4. In contrast to the
2	urban study areas analysis in Chapter 7, the national-scale analysis employs a data fusion
3	approach that takes advantage of the accuracy of monitor observations and the comprehensive
4	spatial information of the CMAQ modeling system to create national-scale "fused" spatial
5	surfaces of seasonal average O ₃ . Measured O ₃ concentrations from 2006-2008 were fused with
6	modeled concentrations from a 2007 CMAQ model simulation, run for a 12 km domain covering
7	the contiguous U.S. In the first draft of the REA, the spatial surfaces were created using the
8	enhanced Voronoi Neighbor Averaging (eVNA) technique (Timin et al, 2010), using the EPA's
9	Model Attainment Test Software (MATS; Abt Associates, 2010b). In this draft, the spatial
10	surfaces are created using EPA's Downscaler software (Berrocal et al, 2012). More details on
11	the ambient measurements, the 2007 CMAQ model simulation, the Downscaler fusion technique,
12	and a technical justification for changing from eVNA to Downscaler can be found in Chapter 4.
13	Three "fused" spatial surfaces were created for: (1) the May-September mean of the 8-hr
14	daily maximum (consistent with the metric used by Smith et al. 2009); (2) the June-August mean
15	of the 8-hr daily mean from 10am to 6pm (consistent with the metric used by Zanobetti and
16	Schwartz 2008); and (3) the April-September mean of the 1-hr daily maximum (consistent with
17	the metric used by Jerrett et al. 2009) O ₃ concentrations across the continental U.S. These fused
18	spatial surfaces each represent one seasonal average across 2006-2008, rather than three separate
19	years of concentrations. Section 4.3.2 presents maps, distributions, and statistical
20	characterizations of these O_3 concentrations metrics across the U.S., including how they compare
21	to 2006-2008 design values.

22

8.1.1.2 Concentration-Response Functions

23 While Chapter 7 assessed both mortality and morbidity risks associated with O₃ 24 concentrations, due to limitations in baseline morbidity incidence rates, the national scale 25 assessment focuses on mortality risks only. To quantify the impact of O₃ concentrations on 26 mortality, we apply risk estimates drawn from two major short-term epidemiological studies and 27 one long-term epidemiological study. These studies are consistent with those used in the analysis 28 of O₃-related risk in selected urban areas (Section 7.2) and those mortality endpoints concluded 29 to have a causal or suggestive causal relationship with O₃ exposure by the 2013 Integrated 30 Science Assessment for O₃ and Related Photochemical Oxidants (U.S. EPA 2013). 31 For short-term mortality, we use city-specific and national average risk estimates drawn 32 from the Smith et al. (2009) study of O₃ and mortality in 98 U.S. urban communities between 33 1987 and 2000 as our main results, and the Zanobetti and Schwartz (2008) study of O₃ and

1 mortality in 48 U.S. cities between 1989 and 2000 as a sensitivity analysis, consistent with the

2 urban case study analysis in Chapter 7. City-specific effect estimates for both studies are

3 provided in Appendix 4-A.

4 Smith et al. (2009) found that the average non-accidental mortality increase across all 98 5 urban areas was $0.32\% \pm 0.08$ (95% posterior interval [PI], 0.41%-0.86%) for a 10 ppb increase 6 in the 8-hr daily maximum O₃ concentration, based on April to October O₃ observations. Since 7 the national-scale analysis requires a single modeling period definition but some monitors only 8 collect data from May to September, the corresponding city-specific effect estimates are applied 9 to each day from May to September in BenMAP using May to September average 8-hr daily 10 maximum O₃ concentration based on 2006-2008 observed concentrations fused with 2007 11 modeled concentrations. The length of the O₃ season can affect the magnitude of mortality effect 12 estimates – a longer season may yield higher effect estimates per unit O_3 concentration since O_3 13 concentrations over the longer season may be lower than the O₃ concentrations over the warmest 14 months only. Conversely, if the longer period captures periods of lower O_3 -related mortality 15 incidence, the effect estimates may be lower than effect estimates for the warmest months only. 16 Our application of the Smith et al. (2009) April to October effect estimates to May to September 17 O₃ concentrations likely introduces some bias in the results, but it is unclear in which direction. 18 Zanobetti and Schwartz (2008) found that the average total mortality increase across all 19 48 cities was 0.53% (95% confidence interval, 0.28%-0.77%) for a 10 ppb increase in June-20 August 8-hr daily mean O_3 concentration from 10 am to 6 pm, using a 0-3 day lag. We apply the 21 city-specific effect estimates that correspond to this national average effect estimate each day 22 from June to August in BenMAP using the June to August, mean 8-hr daily mean O₃ 23 concentration based on 2006-2008 observed concentrations fused with 2007 modeled 24 concentrations. Consistent with Chapter 7, these results are presented as a sensitivity analysis. 25 As in Chapter 7, we use city-specific risk estimates from the short-term epidemiology 26 studies, but apply them here only to the counties that were included in the epidemiology studies 27 rather than to the entire core-based statistical area (CBSA). Chapter 7 estimated risk across entire 28 CBSAs to more completely capture expected O₃ changes across broader areas and avoid bias 29 resulting from including only those areas where O_3 is expected to increase under alternative 30 standards. The inclusion of the entire CBSA in that analysis required the application of a single 31 effect estimate to the entire CBSA. However, the national-scale assessment is a gridded analysis, 32 which allows greater spatial resolution in the application of effect estimates. In addition, eight 33 CBSAs nationwide included multiple cities defined separately by Smith et al. (2009), some of 34 which showed considerable heterogeneity in effect estimates within the same CBSA. 35 Heterogeneity among effect estimates within a single CBSA implies that effect estimates from 36 one county may not be accurate representations of effect estimates in nearby

8-4

1 counties. However, since city-specific effect estimates often have low power due to small 2 population size, we are unable to draw a strong conclusion regarding how well one county's 3 effect estimates represents those in nearby counties. For this national-scale assessment, we 4 apply effect estimates from each city as defined in the epidemiology studies to retain the full set 5 of information available from those studies. In addition, for counties not included by the 6 epidemiology studies, we apply the average effect estimate derived from all the urban areas 7 included in each of the studies ("national average") as it takes advantage of a wider and more 8 diverse population. 9 Since both national average estimates from these studies are based on urban areas only,

10 we have higher confidence in their application to other U.S. urban areas than to rural areas. To 11 demonstrate the magnitude of the results for which we have the highest confidence, we present 12 the percentage of estimated deaths occurring within the urban areas included in the 13 epidemiological studies and within all urban areas across the U.S. Lower confidence in the 14 results for rural areas does not indicate that the mortality risk among populations living in such 15 areas is unaffected by O₃ pollution. Rather, the level of understanding for the O₃-mortality 16 relationship in these areas is simply lower due to a lack of available epidemiological data at these 17 levels. We also examine the effect of varying the effect estimate applied between the cities

18 included by the epidemiology studies in a sensitivity analysis.

19 We quantify long-term O₃-related respiratory mortality in this REA since the Integrated 20 Science Assessment for O₃ and Related Photochemical Oxidants (O₃ ISA) concluded that the 21 evidence supports a likely to be causal relationship between long-term O_3 exposure and respiratory effects, including respiratory morbidity and respiratory-related mortality (U.S. EPA, 22 23 2013). As detailed in Chapter 7, we quantify long-term O_3 -related mortality using the respiratory 24 mortality effect estimates from the Jerrett et al. (2009) two-pollutant model that controlled for 25 PM_{2.5} concentrations, applied to each gridcell across the entire United States. This model found 26 that a 10 ppb increase in the April-September average of the 1-hr daily maximum O₃ 27 concentration was associated with a 4% (95% confidence interval, 1.0%-6.7%) increase in

28 respiratory mortality.

8.1.1.3 Demographic Inputs

This analysis uses the same baseline mortality rates and population estimates as were used in the urban case study area analysis in Chapter 7. We derive baseline incidence rates for mortality by age, cause, and county from the CDC Wonder database (CDC, 2004-2006). As this

32 database only provides baseline incidence rates in 5-year increments, we use data for the year

33 2005, the closest year to the analysis year 2007 used for the population and air quality modeling.

1 We use 2007 population because it matches both the year of the emissions inventory and

2 meteorology used for the air quality modeling.

The starting point for estimating the size and demographics of the potentially exposed population is the 2010 census-block level population, which BenMAP aggregates up to the same grid resolution as the air quality model. BenMAP back-casts this 2010 population to the analysis year of 2007 using county-level growth factors based on economic projections (Woods and Poole Inc., 2012).

8

9 **8.1.2 Results**

Table 8.1 summarizes the estimated O₃-related premature mortality associated with 20062008 average O₃ concentrations under various assumptions for the health impact function.
Applying Smith et al. (2009) effect estimates for May-September, we estimate 15,000 (95% CI,
1400-28 000) premature O₂-related non-accidental deaths annually for 2007. As a sensitivity

13 1,400-28,000) premature O₃-related non-accidental deaths annually for 2007. As a sensitivity

14 analysis, we apply Zanobetti and Schwartz (2008) effect estimates for June-August, finding

15 16,000 (95% CI, 6,000-25,000) premature O_3 -related all-cause deaths annually for 2007. Figure

16 8.2 Figure 8.4 show that estimated O₃-related mortality is most concentrated in highly populated

17 counties or those counties with urban areas found to have high effect estimates by Smith et al.

18 (2009) or Zanobetti and Schwartz (2008). For the application of Jerrett et al. (2009) national

19 average effect estimate for April-September, we estimate 45,000 (95% CI, 17,000-70,000)

21 Because the epidemiological studies included only selected urban areas, we are more 22 confident in the magnitude of the estimated O₃-related deaths occurring within those urban areas. 23 As shown in Table 8.1, approximately 43% of the O₃-related deaths estimated using Smith et al. 24 (2009) effect estimates occur in the 98 urban locations included in that study, and 30% of the O₃-25 related deaths estimated using Zanobetti and Schwartz (2008) effect estimates occur in the 48 26 urban areas included in that study. We are also more confident in extrapolating the national 27 average effect estimates to other urban areas than we are to rural areas, as the national average 28 estimates are based on all urban areas included by the study. To estimate the percentage of total 29 O₃-attributable deaths occurring within all urban areas across the continental U.S., we sum the results for the 12km gridcells that have a total population greater than 12,000 (approximately 30 equal to the 95th percentile of gridcell populations across the continental U.S.). The percentage of 31 32 O₃-attributable deaths occurring within urban areas defined in this way is 65% for results based 33 on Smith et al. (2009) effect estimates and 64% for results based on Zanobetti and Schwartz

34 (2008) effect estimates. While our confidence is lower when the national average effect estimates

²⁰ premature O_3 -related respiratory deaths among adults age 30 and older.

- 1 are extrapolated to rural areas, less certainty in the magnitude of O₃-related deaths in rural areas
- 2 does not imply that O_3 has no effect on health in these areas.
- 3

4Table 8-1Estimated annual O3-related premature mortality in 2007 associated with 2006-52008 average O3 concentrations (95th percentile confidence interval)

Source of risk estimate and modeling period	Exposure duration	Age	City-specific effect estimates ¹	National average effect estimate ²
Smith et al. (2009), May-September 95% confidence interval % occurring within the 98 cities	Short-term	>0	15,000 (1,400-28,000) 43%	16,000 (7,200-22,000)
Zanobetti and Schwartz (2008), June-August 95% confidence interval % occurring within the 48 cities	Short-term	>0	16,000 (6,000-25,000) 30%	15,000 (8,300-22,000)
Jerrett et al. (2009), April-September 95% confidence interval	Long-term	≥30 years	-	45,000 (17,000-70,000)

¹City-specific effect estimates are applied to the gridcells lying within the cities defined in the epidemiological studies. Average effect estimates across all cities included in the epidemiological studies (national average) are applied to all other gridcells. For the application of Smith et al. (2009) effect estimates, city-specific effect
estimates were applied to 2,227 gridcells and the national average to 44,064 gridcells. For the application of Zanobetti and Schwartz (2008) effect estimates, city-specific effect estimates were applied to 925 gridcells and the national average to 45,366 gridcells.

12 13

² National average effect estimates are based on the average of all cities included in the epidemiological studies applied to all 12km gridcells nationally.

Table 8.1 also shows O₃-related deaths estimated by applying the national average risk estimate from the epidemiological studies to all gridcells in the U.S. Compared with applying city-specific effect estimates to the gridcells corresponding to each urban area, using the national average effect estimate for all gridcells yields equivalent central estimates. However, applying the national average also results in tighter confidence intervals since the national average effect estimates had higher statistical power and thus tighter confidence bounds compared with the effect estimates for individual cities.

Table 8.2 shows the mean, median, 2.5 percentile and 97.5 percentile of the estimated percentage of mortality attributable to ambient O₃ across all counties in the U.S. Using Smith et al. (2009) effect estimates, O₃-attributable mortality contributes an average of 1.5% (95% confidence interval, 1.1%-1.8%) to county-level May-September non-accidental mortality (all ages) and 0.6% (0.4%-0.7%) to all year all-cause mortality (all ages). For results using Zanobetti

- 1 and Schwartz (2008) effect estimates, O₃-attributable mortality contributes an average of 2.5%
- 2 (95% confidence interval, 1.7%-3.0%) to county-level June-August all-cause mortality (all ages)
- 3 and 0.6% (0.4%-0.8%) to all year all-cause mortality (all ages). For the results using Jerrett et al.
- 4 (2009) effect estimates, O₃-attributable mortality contributes an average of 18.5% (95%
- 5 confidence interval, 15.2%-21.5%) to county-level April-September adult (age 30+) respiratory
- 6 mortality and 1.9% (1.3%-2.6%) to all year all-cause mortality (all ages). Figure 8.5 through
- 7 Figure 8.7 show that the counties with the highest percentage of mortality attributable to O_3 are
- 8 typically those with the highest O₃ levels.
- 9 Figure 8.8 displays the cumulative distribution of the percent of county-level all-cause,
- 10 all-age, and all-year mortality attributable to ambient O_3 using effect estimates from all three
- 11 epidemiological studies. For the results based on Smith et al. (2009) and Zanobetti and Schwartz
- 12 (2008) effect estimates, 0.8% of all-cause, all-age, and all-year mortality is attributable to O_3 for
- 13 approximately 99% of U.S. counties. For the results based on Jerrett et al. (2009) effect
- 14 estimates, 2.8% of all-cause, all-age, and all-year mortality is attributable to O₃ for
- 15 approximately 99% of U.S. counties.
- 16

Mean, median, 2.5 percentile, and 97.5 percentile of the estimated percentage 1 **Table 8-2** of mortality attributable to ambient O₃ for all 3087 counties in the continental **U.S.**¹ 3

		Percentage of total incidence attributable to O ₃				
Source of risk estimate, modeling period, and mortality endpoint used to generate percentage	Total incidence (2005)	Mean (%)	Median (%)	2.5 Percentile (%)	97.5 Percentile (%)	
Smith et al. (2009), May-September Non-accidental mortality, all ages All-cause mortality, all ages All-cause mortality, all ages, all year	964,837 1,028,334 2,454,896	1.5 1.4 0.6	1.5 1.4 0.6	1.1 1.0 0.4	1.8 1.7 0.7	
Zanobetti and Schwartz (2008), June-August All-cause mortality, all ages All-cause mortality, all ages, all year	618,345 2,454,896	2.5 0.6	2.5 0.6	1.7 0.4	3.0 0.8	
Jerrett et al. (2009), April-September Respiratory mortality, ages 30+ All-cause mortality, all ages All-cause mortality, all ages, all year	236,756 1,229,968 2,454,896	18.5 3.8 1.9	18.7 3.7 1.9	15.2 2.6 1.3	21.5 5.2 2.6	

¹ For the mortality endpoints matching the epidemiology studies as a percentage of incidence of the same endpoint for the same seasonal definition, and as a percentage of all-cause mortality for all age groups (both seasonal and all year).



Figure 8.2Estimated annual non-accidental premature deaths (individuals) in2007 associated with average 2006-2008 May-September average 8-hr dailymaximum O3 levels by county using Smith et al. (2009) effect estimates



Figure 8.3 Estimated annual all-cause premature deaths (individuals) in 2007 associated with average 2006-2008 June-August average 8-hr daily mean (10am-6pm) O₃ levels by county using Zanobetti and Schwartz (2008) effect estimates



Figure 8.4 Estimated annual adult (age 30+) respiratory premature deaths (individuals) in 2007 associated with average 2006-2008 April-September average 1-hr daily max O₃ levels by county using Jerrett et al. (2009) effect estimates



Figure 8.5 Estimated percentage of May-September total non-accidental mortality (all ages) attributable to 2006-2008 average O₃ levels by county using Smith et al. (2009) effect estimates



Figure 8.6 Estimated percentage of June-August total all-cause mortality (all ages) attributable to 2006-2008 average O₃ levels by county using Zanobetti and Schwartz (2008) effect estimates



Figure 8.7 Estimated percentage of April-September respiratory mortality among adults age 30+ attributable to 2006-2008 average O₃ levels by county using Jerrett et al. (2009) effect estimates





 $^{^{2}}$ Estimated O₃-attributable deaths are based on the mortality cause, age group, and season inherent to the epidemiological study upon which it is based (May-September non-accidental mortality for all ages for results based on Smith et al. (2009) effect estimates, June-August all-cause mortality for all ages for results based on Zanobetti and Schwartz (2008) effect estimates, and April-September respiratory mortality for ages 30+ for results based on Jerrett et al. (2009) effect estimates).

1 Figure 8.9 shows the cumulative distribution of the county-level percent of total O₃-2 related deaths by O_3 concentration. The mortality results based on Smith et al. (2009) 3 concentration-response functions are compared with the May-September average of the 8-hr 4 daily maximum O_3 concentration, those based on Zanobetti and Schwartz (2008) concentration-5 response functions are compared with the June-August average of the 8-hr mean O_3 6 concentration from 10am to 6pm, and those based on Jerrett et al. (2009) concentration-response 7 functions are compared with the April-September average of the 1-hr daily maximum O₃ 8 concentration, consistent with the O₃ concentration metrics used in each study. The mortality 9 results based on Zanobetti and Schwartz (2008) effect estimates are shifted to the right of the 10 mortality results based on the Smith et al. (2009) concentration response functions because the 11 seasonal averaging time for the results based on Zanobetti and Schwartz (2008) is limited to the 12 summer months when O₃ tends to be highest. Similarly, the mortality results based on Jerrett et 13 al. (2009) effect estimates are shifted to the right of the mortality results based on Zanobetti and 14 Schwartz (2008) and Smith et al. (2009) because Jerrett et al. (2009) results use the seasonal 15 average of the 1-hr daily maximum, which tends to be higher than the seasonal average of 8-hr 16 daily maximum and seasonal average of 8-hr daily mean metrics (see Figure 4-18). For all three 17 epidemiology studies, we find that 90-95% of O₃-related deaths occur in locations where the 18 May to September average 8-hr daily maximum, June to August average 8-hr daily mean (10am-19 6 pm), or April to September average 1-hr daily maximum O₃ concentrations are greater than 40 20 ppb. A seasonal average concentration of 40 ppb corresponds to 2006-2008 design values 21 ranging from approximately 50 to 90 ppb, depending on the seasonal average concentration 22 metric (see Figure 4-19). 23





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8.1.3 Sensitivity analysis

10 For the results presented above, the national average effect estimate for results based on Smith et al. (2009) and Zanobetti and Schwartz (2008) was applied to all gridcells between the 11 12 cities included in the studies. However, O₃-mortality effect estimates have been shown to exhibit 13 significant regional variability across the U.S. (e.g. Smith et al. 2009). Smith et al. (2009) found 14 that using the national average effect estimate may overestimate risk in cities that have low effect 15 estimates, including Los Angeles and Denver, but may underestimate risk in cities that have high 16 effect estimates, including New York City and Chicago. We conduct two sensitivity analyses 17 aimed at characterizing the sensitivity of estimated O_3 -attributable premature deaths to the use of 18 national average effect estimates between the cities that were included by Smith et al. (2009) and 19 Zanobetti and Schwartz (2008). 20 First, we examine the sensitivity of estimated O_3 -attributable premature deaths to the application of the 5th highest and 5th lowest effect estimates of all the cities included in the Smith 21

et al. (2009) and Zanobetti and Schwartz (2008) studies to the gridcells between the cities

included in these studies (Table 8.3). As in the main results, city-specific effect estimates are 1 applied to the gridcells in which the cities lie. Applying the 5th highest effect estimate from Las 2 Vegas to the gridcells between the cities included by Smith et al. (2009) yields a 36% lower 3 4 estimate of O₃-attributable deaths as compared with the main results. Applying the 5th lowest effect estimate from Dallas/Ft. Worth yields a 42% higher estimate of O₃-attributable deaths as 5 compared with the main results. Applying the 5th lowest effect estimate from Los Angeles to the 6 gridcells between the cities included by Zanobetti and Schwartz (2008) yields a 37% lower 7 estimate of O₃-attributable deaths as compared with the main results. Applying the 5th highest 8 9 effect estimate from Columbus, OH, yields a 30% higher estimate of O₃-attributable deaths as

- 10 compared with the main results.
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12Table - 8.3Sensitivity of estimated O3-attributable premature deaths to the application of13the 5th lowest and 5th highest city-specific risk estimates found by Smith et al.14(2009) and Zanobetti and Schwartz (2008) to the gridcells between the cities15included in those studies.

			Percent
			change
		O ₃ -attributable	from main
Source of risk estimate and sensitivity study	Beta and city	mortality	results
Smith et al. (2009), May-September			
5 th lowest city beta	0.00014	9,600	-36%
	Las Vegas, NV	(-20,000 - 38,000)	
5 th highest city beta	0.000538	21,000	+42%
	Dallas/Ft. Worth, TX	(-1000 – 43,000)	
Zanobetti and Schwartz (2008), June-August			
5 th lowest city beta	0.000274	9,800	-37%
	Los Angeles, CA	(-3,700 – 23,000)	
5 th highest city beta	0.000739	20,000	+30%
	Columbus, OH	(100 - 40,000)	

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18 Second, we examine the sensitivity of estimated O₃-attributable premature deaths to the

19 application of Smith et al. (2009) Bayesian-shrunken city-specific estimates using regional

20 average priors rather than the national average prior (Table 8.4). For gridcells between the cities

21 included by Smith et al. (2009), we apply the regional average effect estimate, rather than the

22 national average effect estimate as in the main results. Regional definitions are shown in Figure

1 8.10. Estimated O_3 -attributable deaths using the regional prior city-specific effect estimates and 2 the regional average effect estimates between the 98 cities included by Smith et al. (2009) are 3 approximately 20% larger than the main results, with 38% of estimated deaths occurring in the 4 98 cities rather than 43%. The 95% confidence interval for the results using the regional prior 5 spans zero, whereas the 95% confidence interval for the results using the national prior does not. 6 Since the regional average effect estimates are all based on fewer data points (in some regions, 7 the regional average is based on only seven cities; see Appendix 8-A) than is the national 8 average, the confidence interval for each regional average effect estimate is large and sometimes 9 spans zero. The large confidence intervals for the regional average effect estimates drive the 10 confidence interval that spans zero for O₃-attributable mortality estimated using regional prior 11 effect estimates. Confidence intervals that span zero do not imply that higher O₃ is associated 12 with decreased mortality, as there is no biologically plausible mechanism for such an effect, and 13 in no case do we see a significant negative central estimate. Rather, confidence intervals 14 spanning zero indicate a lack of statistical power to precisely determine the magnitude of an 15 effect. Figure 8.11 shows estimated O₃-attributable deaths by region using the national average 16 17 prior compared with using the regional average priors from Smith et al. (2009). Results generally 18 follow conclusions made by Smith et al. (2009) based on the magnitude of the regional effect 19 estimates. For example, using the national average effect estimate may substantially 20 underestimate O₃-attributable deaths in the North East and Industrial Midwest where regional 21 effect estimates are large. Using the national average effect estimate may also overestimate O₃-22 attributable deaths in the Upper Midwest, Southern California, and South West, which were 23 found to have small effect estimates. However, these three regions have very large confidence 24 intervals which all span zero, since these regional averages are based on few cities (7, 7, and 9, 25 respectively, compared with 26 in the South East, 19 in Industrial Midwest, 16 in North East, and 26 12 in North West; see Appendix 8-A).

1Table - 8.4Sensitivity of estimated O3-attributable premature deaths to the application of2Smith et al. (2009) regional prior Bayes-shrunken city-specific and regional3average effect estimates, as compared with the national prior Bayes-shrunken4city-specific and national average effect estimates as in the main results.

Risk estimate	O ₃ -attributable premature deaths	Percent O ₃ - attributable deaths in 98 cities
City-specific, national prior with national average	15,000 (1,400 – 28,300)	43%
City-specific, regional prior with regional averages	18,000 (-2,000 – 24,000)	38%



Figure 8.10 Regions used in the sensitivity analysis based on the Smith et al. (2009)
 regional-prior Bayes-shrunken city-specific and regional average effect
 estimates (Source: Samet et al. 2000).



national prior regional prior

Figure 8.11 O₃-attributable premature deaths by region as calculated by applying Smith et al. (2009) regional prior Bayes-shrunken and regional average effect estimates, as compared with the national prior Bayes-shrunken and national average effect estimates as in the main results

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7 8.1.4 Discussion

8 We estimated the total all-cause deaths associated with short-term exposure to recent O_3 9 levels across the continental U.S., using average 2006-2008 observations from the O₃ monitoring 10 network fused with a 2007 CMAO simulation and city-specific O₃-mortality effect estimates 11 from two short-term epidemiology studies. Applying Smith et al. (2009) effect estimates for 12 May-September, we estimate 15,000 (95% CI, 1,400-28,000) premature O₃-related non-13 accidental deaths (all ages) annually for 2007. Using Smith et al. (2009) effect estimates, O₃-14 attributable mortality contributes an average of 1.5% (95% confidence interval, 1.1%-1.8%) to 15 county-level May-September non-accidental mortality (all ages) and 0.6% (0.4%-0.7%) to all 16 year all-cause mortality (all ages). As a sensitivity, we apply Zanobetti and Schwartz (2008) effect estimates for June-August, finding 16,000 (95% CI, 6,000-25,000) premature O₃-related 17 18 all-cause deaths (all ages) annually for 2007. For results using Zanobetti and Schwartz (2008) 19 effect estimates, O₃-attributable mortality contributes an average of 2.5% (95% confidence 20 interval, 1.7%-3.0%) to county-level June-August all-cause mortality (all ages) and 0.6% (0.4%-0.8%) to all year all-cause mortality (all ages). For the application of Jerrett et al. (2009) effect 21 22 estimates for April-September, we estimate 45.000 (95% CI, 17.000-70,000) premature O₃-

related adult (age 30 and older) respiratory deaths. For the results using Jerrett et al. (2009) effect

1 estimates, O₃-attributable mortality contributes an average of 18.5% (95% confidence interval,

2 15.2%-21.5%) to county-level April-September adult (age 30+) respiratory mortality and 1.9%

3 (1.3%-2.6%) to all year all-cause mortality (all ages). For all three epidemiology studies, we find

4 that 90-95% of O_3 -related deaths occur in locations where the May to September average 8-hr

5 daily maximum, June to August average 8-hr daily mean (10am-6pm), or April to September

6 average 1-hr daily maximum O₃ concentrations are greater than 40 ppb. A seasonal average

7 concentration of 40 ppb corresponds to 2006-2008 design values ranging from approximately 50

8 to 90 ppb, depending on the seasonal average concentration metric.

9 A previous analysis estimated that short-term O_3 exposure was associated with 4,700 10 (95% CI, 1,800-7,500) premature deaths nationwide annually, based on 2005 O₃ concentrations 11 and Bell et al. (2004) national average effect estimates (Fann et al., 2012). The results estimated 12 here are higher, resulting mainly from two important differences. First, Fann et al. (2012) 13 estimated risk only above North American background, simulated O₃ concentrations in the 14 absence of North American anthropogenic emissions, which was set to 22 ppb in the east and 30 15 ppb in the west. Fann et al. (2012) also used a national average mortality effect estimate for 8-hr 16 daily maximum O_3 during the warm season only, calculated using ratios of 24-hr mean 17 concentrations to 8-hr daily maximum concentrations (see Abt Associates 2010). The Smith et 18 al. (2009) national average beta used here, 0.000322, is based on April-October O₃ data and is 19 approximately 23% larger than that used by Fann et al. (2012), 0.000261. Since the risk 20 modeling period (and the seasonal definition for the seasonal average 8-hr daily maximum 21 concentration) was May to September for both studies, the higher beta used here yields a larger O₃ mortality estimate. These two differences in methods explain the larger O₃ mortality estimates 22 23 of this analysis compared with the previous estimate by Fann et al. (2012).

Estimated O₃-attributable premature deaths based on Jerrett et al. (2009) effect estimates are approximately three times larger than results based on Smith et al. (2009) and Zanobetti and Schwartz (2008) effect estimates. The mean estimated county-level percent of all-cause, all-year,

and all-age mortality is also three times larger for results based on Jerrett et al. (2009) effect

28 estimates, indicating that the larger estimate does not simply result from a longer modeling

29 period or different population subset (e.g. adult respiratory disease for Jerrett et al. (2009) effect

30 estimates versus all-age non-accidental or all-cause mortality for Smith et al. (2009) and

31 Zanobetti and Schwartz (2008) effect estimates). Recent studies using long-term O₃-mortality

32 relationships found by Jerrett et al. (2009) to quantify the burden of mortality due to

anthropogenic O_3 globally (Anenberg et al. 2010, 2011) and for the U.S. specifically (Fann et al.

34 2012) have also found that using Jerrett et al. (2009) long-term effect estimates yields O₃-related

35 mortality burden estimates that are approximately two to four times larger than estimates based

36 on short-term effect estimates. Since long-term mortality relationships include both acute and

1 chronic exposure effects, the significantly larger mortality estimates calculated using long-term

- 2 concentration-mortality relationships suggest that considering only short-term mortality may
- 3 exclude a substantial portion of O₃-related risk. However, since the short-term mortality
- 4 relationships include a larger population (all ages versus adults ages 30 and older only) and all
- 5 mortality causes, the short-term mortality relationships may capture some O₃ effects that are not
- 6 captured by Jerrett et al. (2009). It is likely that some portion of the estimated premature deaths
- 7 attributable to short-term O₃ exposure is captured by estimated premature deaths attributable to
- 8 long-term O₃ exposure, but the extent of the overlap between these estimates is unknown.
- 9

10

EVALUATING THE REPRESENTATIVENESS OF THE URBAN STUDY AREAS 8.2 11 IN THE NATIONAL CONTEXT

12 To further support interpretation of risk estimates generated in Section 7.2, this section 13 presents three analyses that assess the representativeness of the 12 urban study areas in the 14 national context. First, we assess the degree to which the urban study areas represent the range of 15 air quality levels and key O_3 risk-related attributes that vary spatially across the nation. We have 16 partially addressed this issue by selecting urban study areas in different geographical regions of 17 the country (see Section 7.2). In this section, we evaluate how well the selected urban areas represent the overall U.S. for a set of spatially-distributed O₃ risk related variables (e.g. weather, 18 19 demographics including socioeconomic status, baseline health incidence rates; Section 8.2.1). 20 Section 8.2.2 identifies where our 12 urban study areas fall along the distribution of O₃-21 attributable mortality risk across the U.S. This analysis allows us to assess the degree to which 22 the 12 urban study areas capture locations within the U.S. likely to experience elevated levels of 23 risk related to ambient O_3 . Finally, we give a national context to the estimated O_3 responses to 24 emission changes in the urban study areas by assessing how well these 12 areas and the 3 25 additional exposure areas represent air quality trends and responses to emissions across the entire 26 U.S. (Section 8.2.3). 27 We do not attempt to assess the representativeness of the 15 urban study areas considered 28 in the exposure assessment for O_3 related risk because data limitations preclude us from being 29 able to characterize individual-level exposure across the U.S. However, the urban study areas

- 30 considered in both the exposure and risk assessments shared common selection criteria,
- 31 including consideration of O_3 concentrations, availability of adequate monitoring data,
- 32 demographics, and exposure factors. Therefore, conclusions from this analysis of the
- 33 representativeness of the 12 urban study areas for risk would also apply to those areas for

34 exposure.

8.2.1 Analysis Based on Consideration of National Distributions of Risk-Related Attributes

This section evaluates how well the urban study areas reflect national-level variability in a series of O_3 risk-related variables. For this analysis, we first generate distributions for riskrelated variables across the U.S. and for the specific urban study areas considered in Section 7.2 from generally available data (e.g. from the 2000 Census, Centers for Disease Control (CDC), or other sources). We then plot the specific values of these variables for the selected urban study areas on these distributions, and evaluate how representative the selected study areas are of the national distributions for these individual variables.

10 Estimates of risk (either relative or absolute, e.g. number of cases) within our risk 11 assessment framework are based on four elements: population, baseline incidence rates, air 12 quality, and the coefficient relating air quality and the health outcome (i.e. the O_3 effect 13 estimates). Each of these elements can contribute to heterogeneity in risk across urban locations, 14 and each is variable across locations. In addition, there may be other identifiable factors that 15 contribute to the variability of the four elements across locations. In this assessment, we examine 16 the representativeness of the selected urban area locations for the four main elements, as well as 17 factors that have been identified as influential in determining the magnitude of the C-R function 18 across locations.

19 While personal exposure is not incorporated directly into O₃ epidemiology studies, city-20 specific O₃ effect estimates are affected by differing levels of exposure which in turn are related 21 to variability in exposure determinants. The correlation between monitored O_3 and personal O_3 22 exposure also varies between cities. The O₃ ISA has comprehensively reviewed epidemiological 23 and toxicological studies to identify variables which may affect the O_3 effect estimates used in 24 the city-specific risk analysis in Section 7.2 and the national-scale risk analysis in Section 8.1 25 (U.S. EPA 2013). Determinants of the O₃ effect estimates used in risk assessment can be grouped 26 into four broad areas:

Demographics: education, income, age, unemployment rates, race, body mass index and
 physical conditioning, public transportation use, and time spent outdoors.

- Baseline health conditions: asthma, chronic obstructive pulmonary disease,
 cardiovascular disease (atherosclerosis, congestive heart disease, atrial fibrillation,
 stroke), diabetes, inflammatory diseases, and smoking prevalence.
- Climate and air quality: O₃ levels, co-pollutant levels (annual mean PM_{2.5}), temperatures
 (days above 90 degrees, mean summer temp, 98th percentile temp).
- Exposure determinants: air conditioning prevalence.

Although data limitations preclude our ability to conduct a national-scale exposure assessment as we have done for O_3 -attributable risk in Section 8.1, we assess the representativeness of the

1 urban study areas across the national distribution of climate, air quality, and air conditioning 2 prevalence, factors which influence individual exposure. As discussed in detail in Chapter 5, no 3 available data base is sufficient to assess the national representativeness of time spent outdoors, 4 another important personal exposure determinant, among persons residing in each of the urban 5 case study areas. However, previous analyses suggest that children's time spent outdoors varies 6 little across U.S. regions (section 8.10.2 of U.S. EPA, 2009). In addition, as discussed in Section 7 5.1.1, time spent outdoors and the percent of person-days having at least one minute outdoors 8 (participation rate) does not appear to vary much over the past few decades based on analyses 9 using the CHAD database, nor does there appear to be a temporal trend over the past decade 10 based on analyses using the American Time Use Survey (ATUS). In considering that many of 11 the activity pattern studies in CHAD were from national surveys conducted in metropolitan areas 12 and that the evaluation results indicate little difference in time expenditure over broad 13 geographic areas and survey collection years, it is likely that the distribution of time spent 14 outdoors generated for the simulated persons in the 15 urban study areas (Chapter 5) reasonably 15 reflects the most important elements of a national distribution of time spent outdoors. 16 Based on these identified potential risk determinants, we identify datasets that could be

17 used to generate nationally representative distributions for each parameter. We are not able to 18 identify readily available national datasets for all variables. In these cases, if we are able to 19 identify a broad enough dataset covering a large enough portion of the U.S., we use that dataset 20 to generate the parameter distribution. In addition, we are not able to find exact matches for all of 21 the variables identified through our review of the literature. In cases where an exact match is not 22 available, we identify proxy variables to serve as surrogates. For each parameter, we report the 23 source of the dataset, its degree of coverage, and whether it is a direct measure of the parameter 24 or a proxy measure (Table 8.5). Summary statistics for the most relevant variables are provided 25 in Table 8.6.

26 Figure 8.12 through Figure 8.18 show the cumulative distribution functions (CDF) 27 plotted for the nation for the four critical risk function elements (population, air quality, baseline incidence, and the O_3 effect estimate), as well as where the urban study areas fall on the 28 29 distribution. While the urban-scale analysis in Chapter 7 includes the full core-based statistical 30 area for the selected cities, we consider here only the counties included in each city as defined by 31 the epidemiological studies, since we only have information on O_3 effect estimates for these 32 counties. This approach is consistent with the national-scale assessment of O₃-attributable risk in 33 Section 8.1, from which we draw county-level O₃-attributable risk estimates for the 34 representativeness analysis in Section 8.2.3. These figures focus on critical variables representing 35 each type of risk determinant, e.g. we focus on all-cause and non-accidental mortality rates, but 36 we also have conducted analyses for cardiovascular and respiratory mortality separately. The

8-23

1 vertical black lines in each graph show the values of the variables for the individual urban study

2 areas. The city-specific values that comprise the national CDF for mortality risks found by

3 Zanobetti and Schwartz (2008) are also displayed on the graphs of those attributes, as the number

4 of cities included in that study is smaller (48 cities). The complete set of analyses is provided in

5 Appendix 4-A.

6 These figures show that the selected urban study areas represent the upper percentiles of 7 the distributions of population and do not represent the locations with lower populations (urban

8 study areas are all above the 90th percentile of U.S. county populations). This is consistent with

9 the objectives of our case study selection process, e.g. we are characterizing risk in areas that are

10 likely to be experiencing excess risk due to O_3 levels above alternative standards. The urban

11 study areas span the full range of seasonal average 8-hr daily maximum O₃ concentrations in

12 monitored U.S. counties and the full distribution of O₃ risk coefficients across the cities included

13 by Smith et al. (2009) and Zanobetti and Schwartz (2008). The urban study area analysis

14 includes the two cities with the highest risk coefficients found by Smith et al. (2009) – New York

15 City and Philadelphia – as well as the two highest found by Zanobetti and Schwartz (2008) –

16 New York City and Detroit. In Table 8.6, respiratory and cardiovascular mortality have higher

17 concentration-response relationships than non-accidental and all-cause mortality because they

18 are based on a smaller baseline population and are the diseases most affected by O_3 exposure.

19 The urban study areas do not capture the upper end of the distribution of baseline mortality,

20 including all-cause (Figure 8.15) and non-accidental mortality (Figure 8.16), as well as

21 cardiovascular and respiratory mortality (see Appendix 8-B). The interpretation of this is that the

22 case study risk estimates may not capture the additional risk that may exist in locations that have

23 the highest baseline mortality rates.

				Degree of
Potential risk				national
determinant	Metric	Year	Source	coverage
Demographics				
Age	Percent age 85 years and	2005	County Characteristics, 2000-2007 Inter-	All counties
	older		university Consortium for Political and	
			Social Research	
Age	Percent age 65 years and	2005	County Characteristics, 2000-2007 Inter-	All counties
	older		university Consortium for Political and	
			Social Research	
Age	Percent age 14 years and	2005	County Characteristics, 2000-2007 Inter-	All counties
	younger		university Consortium for Political and	
			Social Research	
Education	Population with less than	2000	USDA/ERS,	All counties
	high school diploma		http://www.ers.usda.gov/Data/Education/	
Unemployment	Percent unemployed	2005	County Characteristics, 2000-2007 Inter-	All counties
			university Consortium for Political and	
			Social Research	
Income	Per capita personal income	2005	County Characteristics, 2000-2007 Inter-	All counties
			university Consortium for Political and	
			Social Research	
Race	Percent nonwhite	2006	County Characteristics, 2000-2007 Inter-	All counties
			university Consortium for Political and	
			Social Research	

Table - 8.5 Data Sources for O3 risk-related Attributes

				Degree of
Potential risk				national
determinant	Metric	Year	Source	coverage
Population	Total population	2008	Cumulative Estimates of Resident	All counties
			Population Change for the United States,	
			States, Counties, Puerto Rico, and Puerto	
			Rico Municipios: April 1, 2000 to July 1,	
			2008, Source: Population Division, U.S.	
			Census Bureau	
Population density	Population/square mile	2008	Cumulative Estimates of Resident	All counties
			Population Change for the United States,	
			States, Counties, Puerto Rico, and Puerto	
			Rico Municipios: April 1, 2000 to July 1,	
			2008, Source: Population Division, U.S.	
			Census Bureau (calculated "as the crow	
			flies")	
Urbanicity	ERS Classification Code	2003	County Characteristics, 2000-2007 Inter-	All counties
			university Consortium for Political and	
			Social Research	
Climate and Air Que	ality			
O ₃ levels	Monitored 4 th high 8-hr	2007	EPA Air Quality System (AQS)	725 Monitored
	daily maximum			counties
O ₃ levels	Seasonal mean 8-hr daily	Avg. 2006-2008	AQS	671 Monitored
	maximum			counties
O ₃ levels	Seasonal mean 1-hr daily	Avg. 2006-2008	AQS	671 Monitored

				Degree of
Potential risk				national
determinant	Metric	Year	Source	coverage
	maximum			counties
O ₃ levels	Seasonal mean	Avg. 2006-2008	AQS	671 Monitored
				counties
PM _{2.5} levels	Monitored annual mean	2007	AQS	617 Monitored
				counties
Temperature	Mean July temp	1941-1970	County Characteristics, 2000-2007 Inter-	All counties
			university Consortium for Political and	
			Social Research	
Relative Humidity	Mean July RH	1941-1970	County Characteristics, 2000-2007 Inter-	All counties
			university Consortium for Political and	
			Social Research	
Exposure Determind	ants			
Ventilation	Percent residences with no	2004	American Housing Survey	76 cities
	air conditioning			
Baseline Health Cor	ıditions			
Baseline mortality	All Cause		CDC Wonder 1999-2005	All counties
Baseline mortality	Non Accidental		CDC Wonder 1999-2006	All counties
Baseline mortality	Cardiovascular		CDC Wonder 1999-2007	All counties
Baseline mortality	Respiratory		CDC Wonder 1999-2008	All counties
Baseline morbidity	Acute myocardial	2007	Behavioral Risk Factor Surveillance System	184 metropolitan
	infarction prevalence		(BRFSS)	statistical areas

				Degree of
Potential risk				national
determinant	Metric	Year	Source	coverage
				(MSA)
Baseline morbidity	Diabetes prevalence	2007	BRFSS	184 MSA
Baseline morbidity	Stroke prevalence	2007	BRFSS	184 MSA
Baseline morbidity	Congestive heart disease prevalence	2007	BRFSS	184 MSA
Obesity	Body Mass Index	2007	BRFSS	184 MSA
Level of exercise	Vigorous activity 20 minutes	2007	BRFSS	184 MSA
Level of exercise	Moderate activity 30 minutes or vigorous activity 20 minutes	2007	BRFSS	184 MSA
Respiratory risk	Current asthma	2007	BRFSS	184 MSA
factors				
Smoking	Ever smoked	2007	BRFSS	184 MSA
C-R Estimates				
Mortality risk	Non Accidental	2009	Smith et al. (2009)	98 cities
Mortality risk	All Cause	2008	Zanobetti and Schwartz (2008)	48 cities
Mortality risk	Cardiovascular	2008	Zanobetti and Schwartz (2008)	48 cities
Mortality risk	Respiratory	2008	Zanobetti and Schwartz (2008)	48 cities

									Sampl	e Size nties or
	Aver	age	Standard Deviation		Maximum		Minimum		cities)	
Risk Attribute	Urban Study Areas	U.S. Dataset	Urban Study Areas	U.S. Dataset	Urban Study Areas	U.S. Dataset	Urban Study Areas	U.S. Dataset	Urban Study Areas	U.S. Dataset
Demographics										
Population per county	1,642,198	97,020	1,972,403	312,348	9,862,049	9,862,049	354,361	42	23	3143
Population density (Pop/sq mile)	10,378	258	16,550	1,757	71,758	71,758	1,313	0	23	3143
Median age (Years)	35.7	38.6	2.3	4.4	40.0	55.3	32.1	20.1	23	3141
% Age 0 to 14 years	20.7	19.0	2.4	2.9	24.6	36.8	14.7	0.0	23	3141
% Age 65+ years	11.3	14.9	2.5	4.1	15.2	34.7	5.8	2.3	23	3141
% Age 85+ years	1.7	2.1	0.6	0.9	2.5	7.7	0.5	0.1	23	3141
Unemployment rate (%)	5.7	5.4	1.2	1.8	8.6	20.9	4.1	1.9	23	3133
% with less than high school diploma	20.9	22.6	7.9	8.8	37.7	65.3	8.7	3.0	23	3141
Income per capita (\$)	40305	27367	14238	6604	93377	93377	23513	5148	23	3086
% Non-white	36.4	13.0	15.3	16.2	86.7	95.3	31.7	0.0	23	3141
% Commute by public transportation*	7.1	1.6	8.1	2.5	30.7	30.7	1.5	0.0	12	366
Health Conditions										
Prevalence of CHD (%) *	3.6	4.3	0.8	1.3	4.6	8.7	2.6	1.8	11	184
Prevalence of asthma (%) *	8.5	8.1	1.3	1.9	11.2	13.2	6.0	3.6	11	184
Prevalence of diabetes (%) *	8.1	8.5	1.2	2.1	10.6	16.5	5.4	2.2	11	184
Prevalence of AMI (%) *	3.6	4.1	0.6	1.3	4.8	10.2	2.8	1.7	11	184
Prevalence of obesity (%) *	24.7	26.0	4.0	4.1	32.7	35.7	18.7	14.0	11	182
Prevalence of stroke (%) *	2.6	2.7	0.7	1.0	3.7	6.5	1.5	0.7	11	184
Prevalence of ever smoked (%)*	18.3	19.6	3.1	4.0	23.1	34.4	14.2	6.5	11	184
Prevalence of exercise (20 minutes,										
%)*	29.5	28.0	2.7	4.8	33.8	44.1	23.7	15.4	11	183
Prevalence of exercise (30	50.2	40.7	2.2	 – – –	55.0	(7.1	477 4	27.2	11	102
Minutes,%)*	50.2	49.7	2.3	5.4	55.3	67.1	47.4	57.3	11	182
100,000 people)	756.2	950.6	204.1	249.6	1139.5	1958.4	361.6	117.7	23	3142

Table - 8.6 Summary Statistics for Selected O₃ Risk-related Attributes

	A wor	19.00	Standard	Deviation	Movi		Mini		Sample (# of cou	e Size nties or
Risk Attribute	Urban Study Areas	U.S. Dataset								
All cause mortality (deaths per 100.000 people)	810.1	1022.3	217.4	258.6	1257.8	2064.2	402.5	176.8	23	3142
Cardiovascular mortality (deaths per 100,000 people) Respiratory mortality (deaths per	310.5	392.1	93.9	121.0	459.6	970.4	122.4	37.5	23	3142
100,000 people)	66.2	97.3	17.0	32.3	90.1	351.0	34.8	13.3	23	3136
O_3 4th high maximum 8-hr average (ppm) O_3 seasonal mean (ppb)	0.087 33.9	0.077 34.5	0.009 5.4	0.010 6.6	0.105 51.0	0.126 64.8	0.072 25.8	0.033 8.6	23 22	725 671
O_3 seasonal mean of maximum 8-hr average (ppb)	50.7	48.6	7.5	7.2	70.2	79.7	40.8	13.3	22	671
O_3 seasonal mean of 1-hr daily maximum (ppb)	58.8	54.7	7.5	8.0	85.1	92.4	46.5	17.6	22	671
$PM_{2.5}$ annual mean (µg/m3)	14.1	11.7	2.6	3.1	16.9	22.5	8.4	3.4	23	617
$PM_{2.5}$ 98th %ile daily average (µg/m3)	35.8	30.7	8.1	9.3	59.0	81.1	21.2	9.1	23	617
Average temperature (°F) July temperature long term average (°F)	57.2 76.0	57.2 75.9	5.0 3.4	7.9 5.4	70.3 83.3	76.2 93.7	50.1 68.5	39.0 55.5	23 23	202 3104
July Relative Humidity long term average (%)	61.5	56.2	10.2	14.6	70.0	80.0	28.0	14.0	23	3104
Exposure Determinants										
% No air conditioning*	15.5	16.6	85.7	79.1	42.9	86.7	0.4	0.0	12	76
C-R Estimates										
Non-accidental mortality O_3 risk*	0.000388	0.000322	0.000217	0.000131	0.000917	0.000917	0.000148	-0.000033	12	98
All Cause mortality O3 risk*	0.000627	0.000527	0.000314	0.000205	0.001092	0.001092	0.000163	0.000096	12	48
Respiratory mortality O ₃ risk*	0.000877	0.000800	0.000282	0.000186	0.001424	0.001424	0.000307	0.000307	12	48
Cardiovascular mortality O3 risk*	0.000898	0.000825	0.000173	0.000124	0.001064	0.001064	0.000418	0.000418	12	48

*Attribute for which only city-specific data were available.



Figure 8.12 Comparison of county-level populations of urban case study area counties to the frequency distribution of population in 3,143 U.S. counties.



Figure 8.13 Comparison of county-level seasonal mean 8-hr daily maximum O₃
 concentrations in urban case study area counties to the frequency distribution
 of seasonal mean 8-hr daily maximum O₃ concentrations in 671 U.S. counties
 with O₃ monitors.


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Figure 8.15 Comparison of county-level all-cause mortality in urban case study area
 counties to the frequency distribution of all-cause mortality in 3,137 U.S.
 counties.



Figure 8.16 Comparison of county-level non-accidental mortality in urban case study area
 counties to the frequency distribution of non-accidental mortality in 3,135
 U.S. counties.



Figure 8.17 Comparison of city-level all-cause mortality risk coefficients from Zanobetti
 and Schwartz (2008) in urban case study areas to the frequency distribution
 of all-cause mortality risk coefficients from Zanobetti and Schwartz (2008) in
 48 U.S. cities.



Figure 8.18 Comparison of city-level national prior Bayes-shrunken non-accidental mortality risk coefficients from Smith et al. (2009) in urban case study areas to the frequency distribution of national prior Bayes-shrunken non-accidental mortality risk coefficients from Smith et al. (2009) in 98 U.S. cities.

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7 Figure 8.19 through Figure 8.24 show national CDFs and the urban study area values for 8 several selected potential risk attributes. These potential risk attributes do not directly enter the 9 risk equations, but have been identified in the literature as potentially affecting the magnitude of 10 the O₃ C-R functions reported in the epidemiological literature. Comparison graphs for other risk 11 attributes are provided in Appendix 4-A. The selected urban study areas do not capture the 12 higher end percentiles of several risk characteristics, including populations 65 years and older, 13 baseline cardiovascular disease prevalence, baseline respiratory disease prevalence, and smoking 14 prevalence. Summarizing the analyses of the other risk attributes, we conclude that the urban 15 study areas provide adequate coverage across population, population density, O₃ levels (seasonal 16 mean, seasonal mean 8-hr daily maximum, and seasonal mean 1-hr daily maximum), PM_{2.5} co-17 pollutant levels, temperature and relative humidity, unemployment rates, percent non-white 18 population, asthma prevalence obesity prevalence, income, and less than high school education.

1 We also conclude that while the urban study areas cover a wide portion of the distributions, they

- 2 do not provide coverage for the upper end of the distributions of percent of population 65 and
- 3 older (below 60th percentile), percent of population 85 years and older (below 75th percentile),
- 4 prevalence of angina/coronary heart disease (below 70th percentile), prevalence of diabetes
- 5 (below 85th percentile), stroke prevalence (below 90th percentile), prevalence of heart attack
- 6 (below 80th percentile), prevalence of smoking (below 85th percentile), all-cause mortality rates
- 7 (below 85th percentile), non-accidental mortality rates (below 80th percentile), cardiovascular
- 8 mortality rates (below 75th percentile) and respiratory mortality rates (below 50th percentile), and
- 9 percent of residences without air conditioning (below 90th percentile). In addition, the urban
- 10 study areas do not capture the highest or lowest ends of the distribution of exercise prevalence
- 11 and do not capture the low end of the distribution of public transportation use (above the 65th
- 12 percentile).



13 14

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Figure 8.19 Comparison of county-level percent of population 0 to 14 years old in urban case study area counties to the frequency distribution of percent of population 0 to 14 years old in 3,141 U.S. counties.



 Figure 8.20
 Figure 8.20
 Comparison of county-level percent of population age 65 years old and older in urban case study area counties to the frequency distribution of percent of population age 65 and older in 3,141 U.S. counties.



Figure 8.21 Comparison of county-level income per capita in urban case study areas t the frequency distribution of income per capita in 3,141 U.S. counties.



Figure 8.22 Comparison of county-level July temperature in urban case study area
 counties to the frequency distribution of July temperature in all U.S. counties.



Figure 8.23 Comparison of city-level asthma prevalence in urban case study areas to the frequency distribution of asthma prevalence in 184 U.S. cities.



Figure 8.24 Comparison of city-level air conditioning prevalence in urban case study
 areas to the frequency distribution of air conditioning prevalence in 76 U.S.
 cities.

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6 Based on the above analyses, we can draw several inferences regarding the 7 representativeness of the urban case studies. First, the case studies represent urban areas that are 8 among the most populated in the U.S. Second, they represent areas with relatively high levels of 9 O₃ (4th high 8-hr daily maximum, seasonal mean 8-hr daily maximum, seasonal mean 1-hr daily 10 maximum, and seasonal mean). Third, they capture well the range of city-specific effect 11 estimates found by Smith et al. (2009) and Zanobetti and Schwartz (2008) studies. These three 12 factors would suggest that the urban study areas should capture well overall risk for the nation, 13 with a potential for better characterization of the high end of the risk distribution. 14 However, there are several other factors that suggest that the urban study areas may not 15 be representing areas that may have a high risk per ppb of O_3 . Several of the factors with 16 underrepresented tails, including age and baseline mortality are spatially correlated (R=0.81), so

- 17 that certain counties which have high proportions of older adults also have high baseline
- 18 mortality and high prevalence of underlying chronic health conditions. Because of this, omission

2 Florida, may lead to underrepresentation of high risk populations. However, with the exception 3 of areas in Florida, most locations with high percentages of older populations have low overall 4 populations, less than 50,000 people in a county. And even in Florida, the counties with the 5 highest O_3 levels do not have a high percent of older populations. This suggests that while the 6 risk per exposed person per ppb of O₃ may be higher in these locations, the overall risk to the 7 population is likely to be within the range of risks represented by the urban case study locations. 8 Due to data limitations, we were only able to assess the representativeness of the urban 9 study areas in terms of one exposure-related attribute, air conditioning prevalence. Assessing the 10 representativeness of the urban study areas in terms of air conditioning prevalence, we found that 11 the urban study areas do not capture the highest end of percent of residences without air 12 conditioning. If the cities with the lowest air conditioning prevalence also have high O_3 levels,

of certain urban areas with higher percentages of older populations, for example, cities in

13 we could be missing a high risk portion of the population that is exposed to O_3 indoors as air 14 infiltrates indoors from outdoors. However, 4th highest 8-hr daily maximum O_3 levels in the 15 cities in the top 10th percentile of percentage of residences without air conditioning (mainly in 16 northern California and Washington) are approximately average (0.08 ppm) or lower than

17 average. Since these concentrations are not the highest found across the U.S., we are likely not

18 excluding a high risk population that has both low air conditioning prevalence and high O_3

19 concentrations, and the overall risk to the population is likely to be within the range of risks

20 represented by the urban case study locations.

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8.2.2 Analysis Based on Consideration of National Distribution of O₃-Related Mortality Risk

24 In this section we discuss the second representativeness analysis which identifies where 25 the 12 urban study areas examined in Chapter 7 fall along the distribution of estimated national-26 scale mortality risk. This assessment reveals whether the baseline O_3 mortality risks in the 12 27 urban case study areas represent more typical or higher end risk relative to the national risk 28 distribution presented in Section 8.1. For consistency, we compare the national O_3 mortality risk 29 distribution to the O₃ mortality risk results for the urban study areas that were generated from the 30 national-scale assessment in Section 8.1, rather than the results from the urban study area 31 analysis in Chapter 7 which uses different methods. To be consistent with the national-scale 32 assessment, we define the urban study areas here as they were defined in the epidemiology 33 studies, rather than including full core-based statistical areas as in Chapter 7. The results of this 34 representativeness analysis are presented graphically in Figure 8.25 through Figure 8.27, which 35 display the cumulative distribution of total mortality attributable to ambient O_3 at the county 36 level developed as part of the national-scale analysis. Values for the 23 counties included in the

1 urban case study areas as defined in the epidemiology studies are then superimposed on top of

2 the cumulative distribution to assess the representativeness of the urban case study areas.

3 For the results based on Smith et al. (2009) effect estimates, New York City and

4 Philadelphia have the highest percentage of May-September non-accidental mortality attributable

5 to ambient O_3 of the 12 urban study areas and are located at the highest end of the distribution of 6 U.S. O_3 -related mortality risk (Figure 8.25). Of the 12 urban study areas, these two cities had the

7 highest effect estimates found by Smith et al. (2009; See Appendix 4-A). Boston and Los

8 Angeles had the lowest O₃-related mortality risk of the 12 urban study areas and are located at

9 the lowest end of the U.S. distribution. Overall, O_3 mortality risk in the 12 urban study areas are

10 representative of the full distribution of U.S. O₃-related mortality risk, with the mean percentage

of May-September non-accidental mortality for all ages of 1.5% (95% confidence interval, 1.11.8%).

13 For the results based on Zanobetti and Schwartz (2008) effect estimates, Detroit and New 14 York City are at the very highest end of the U.S. distribution of county-level risk of June-August 15 all-cause mortality due to ambient O₃ (Figure 8.26). These two cities had the highest effect 16 estimates of the 48 cities included in the study (see Appendix 4-A). The high effect estimates in 17 Detroit and New York City could be due to high rates of public transportation use (for New York 18 City), low air conditioning prevalence, high smoking prevalence (in Detroit), high incidence of 19 mortality and other adverse health outcomes (e.g. diabetes, stroke, acute myocardial infarction, 20 etc.), and high unemployment rates. Houston and Los Angeles had the lowest risk and were 21 located at the very lowest end of the U.S. distribution of county-level risk of mortality due to ambient O₃. These two cities had the lowest effect estimates found by Zanobetti and Schwartz 22 23 (2008), possibly because they cover a large spatial extent and have high rates of time spent 24 driving, which could lead to exposure misclassification in the underlying epidemiologic study. 25 Houston also has a very high rate of air conditioning use (nearly 100% of residences) and Los 26 Angeles has been shown to have high rates of adaptive behavior on high ambient O₃ days (i.e. 27 more time spent indoors as a result of high ambient O₃ concentrations; Neidell 2009, 2010), both

of which would lead to lower personal O_3 exposure relative to other cities. Overall, O_3 mortality

risk in the 12 urban study areas are representative of the full distribution of U.S. O₃-related

30 mortality risk, with the mean percentage of June-August all-cause mortality for all ages of 2.5%

31 (95% confidence interval, 1.7-3.0%).

For the results based on Jerrett et al. (2009) effect estimates, the 12 urban study areas are centered more in the middle of the distribution of U.S. county-level risk of adult (ages 30 and older) respiratory mortality due to ambient O_3 exposure. These results are based on the application of a single national average effect estimate to all gridcells across the U.S., rather than city-specific effect estimates as were applied for the results based on Smith et al. (2009) and

- 1 Zanobetti and Schwartz (2008) effect estimates. Therefore, the location of the urban study areas
- 2 on the distribution of county-level risk is driven mainly by O_3 concentration and not by the effect
- 3 estimate. While Denver, Atlanta, Sacramento, and Los Angeles are at the highest end of the U.S.
- 4 distribution, Figure 8.27 shows that some counties have a higher percentage of mortality
- 5 attributable to O_3 than these four cities. Overall, O_3 mortality risk in the 12 urban study areas are
- 6 representative of the full distribution of U.S. O₃-related mortality risk, with the mean percentage
- 7 of April-September respiratory mortality for adults ages 30 and older of 18.7% (95% confidence
- 8 interval, 15.2-21.5%). However, we are not capturing the very highest end of O₃-related risk
- 9 based on Jerrett et al. (2009) effect estimates in the 12 urban study areas.
- 10



Results based on Smith et al. (2009) effect estimates

- Selected urban study area
- Figure 8.25
 Figure 8.25
 Cumulative distribution of county-level percentage of May-September nonaccidental mortality for all ages attributable to 2006-2008 average O₃ for the U.S. and the locations of the selected urban study areas along the distribution, using Smith et al. (2009) effect estimates.



-----Results based on Zanobetti and Schwartz (2008) effect estimates

Selected urban study areas

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Figure 8.26 Figure 8.26 Cumulative distribution of county-level percentage of June-August all-cause mortality for all ages attributable to 2006-2008 average O₃ for the U.S. and the locations of the selected urban study areas along the distribution, using Zanobetti and Schwartz (2008) effect estimates.



- 17 characterize past phenomena. In section 8.2.3.2, we look at air quality model predictions of
- 18 temporal and spatial patterns of O₃ changes in response to further NOx reductions from 2007

1 levels. This analysis is subject to typical model limitations but has the advantage of isolating the 2 effects of precursor emissions changes and has the ability to simulate how O_3 would change in 3 response to NOx (and VOC) emissions reductions (relative to recent 2007 levels) similar to those

4 used in the HDDM adjustment scenarios for just meeting existing and alternative standards.

- 5 These two complimentary analyses give qualitatively similar results, building confidence that the
- 6 overarching conclusions are robust across the US as a whole.

8.2.3.1 Ambient patterns in trends of measured O₃ concentrations

7 This section describes how annual distributions of O3 measurements collected by EPA's 8 national monitoring network have changed between 1998 and 2011. These years were chosen 9 because large reductions in anthropogenic NOx emissions have occurred over this time period 10 especially in the Eastern half of the United States. From 2000 to 2011 nationwide NOx emissions were cut almost in half (from 22.6 to 12.9 million tons per year)³. However, it should be noted 11 12 that these reductions did not occur uniformly across the country. Improvements in vehicle 13 emissions standards helped reduce NOx emissions in many locations throughout the country. In 14 contrast, EPA rules like the NOx SIP call were focused on controlling emissions from power 15 plants in the Eastern US and consequently there have been relatively larger reductions in NOx emissions in the East. In addition, some urban areas which have traditionally had high O₃ levels, 16 17 like Los Angeles and Houston, have substantially cut local NOx and VOC emissions to improve 18 their air quality. Also, there may be some localized areas in which NOx emissions have 19 increased due to population growth, new sources such as oil and gas development, or increased 20 wildfire activity. Appendix 8-C provides plots of emissions trends by region of the U.S. These 21 plots show that each of nine regions of the U.S. have experienced decreasing NOx emissions 22 ranging from approximately a 20% decrease to a 45% decrease from 2002 to 2011 depending on 23 the region. Conversely, VOC emissions have increased in some regions since 2002 (the South, 24 the Southwest, and the West-North-Central) and decreased in others. Due to non-linear O_3 25 formation chemistry and the potential for changes in local chemical regimes resulting from these 26 emissions reductions, past trends may not reflect the ambient changes which will occur from 27 future emissions reductions. Nonetheless, these ambient data provide information on actual O₃ 28 changes in response to emissions reductions and can give insight into the types of changes in O_3 that have occurred both within and outside the urban study areas. 29 First, we look at national maps which show changes in 50th percentile and 95th percentile 30

31 summer season (April-October)⁴ 8-hour daily maximum O_3 values (Figure 8.28, Figure 8.29).

³ Data were accessed from EPA's emission trend website on August 15, 2013: http://www.epa.gov/ttn/chief/trends/trends06/national_tier1_caps.xlsx

⁴ The April-October time period corresponds to the required monitoring season for most of the 12 urban areas. Therefore, in selecting a consistent time period that could be analyzed for the urban case study areas, we chose to

- 1 These maps reflect the absolute (ppb) difference between O₃ percentiles from two three-year periods $(2001-2003 \text{ and } 2008-2010)^5$. Figure 8.28 shows that increases in median O₃ 2 3 concentrations occurred in many large urban areas including both study area locations in Chapter 4 7^{6} and non-study area locations⁷. Only a few monitors with increasing median O₃ appear outside 5 of cities, most notably in southwestern Colorado and central Kansas. The increases in urban 6 areas are likely explained by O₃ "disbenefits" to NOx reductions which were described in 7 Chapter 4, Appendix 4-C and in the following section of this chapter. Widespread decreases of 8 median O_3 in suburban and rural locations suggest the efficacy of large NOx emissions 9 reductions on reducing O₃ over large regions of the country. Finally, the less frequently observed 10 cases of median O₃ increases in rural areas are likely caused by different phenomena. Cooper et al. (2012) suggested that increasing rural O_3 in the Western US may be due to increasing oil and 11 gas development, wildfires and O₃ transport from Asia. Conversely, Figure 8.29 shows that 95th 12 13 percentile O₃ values for these two sets of years decrease in almost all urban as well as rural areas of the country. Only a few sites in Colorado, Nevada, and California show any increases in 95th 14 15 percentile O₃ between 2001-2003 and 2008-2010. The consistent decreases across most of the 16 United States indicate that the large NOx reductions from power plants and mobile sources have 17 been quite successful in reducing O_3 on the highest O_3 days. These results suggest that many of 18 the urban case study areas may show O_3 responses that are typical of other large urban areas in
- 19 the U.S. However, decreasing O_3 in large non-urban portions of the country may not be fully

20 captured in the urban case studies.

use the April-October time period in Chapter 4 for composite monitor distributions to summarize ozone values relevant to the epidemiology-based risk assessment.

⁵ These two three-year periods were chosen to represent years before and after most NOx emission reductions were in place. In addition the 2001-2003 period was used to designate areas for the 1997 8-hour ozone standard and the 2008-2010 period was used to designate areas for the 2008 8-hour ozone standard. Data from these two time periods have undergone extensive quality checks.

⁶ Los Angeles, Denver, Houston, Atlanta, Chicago, Detroit, Cleveland, New York, Philadelphia, and Washington D.C.

⁷ San Francisco, Reno, Phoenix, New Orleans, Birmingham, Miami, and Cincinnati



Change in April - October Median Daily Maximum 8-hour Ozone Concentration from 2001 - 2003 to 2008 - 2010

Figure 8.28 Change in 50th percentile summer season (April-October) daily 8-hr maximum
 O₃ concentrations between 2001-2003 and 2008-2010.



Change in April - October 95th Percentile Daily Maximum 8-hour Ozone Concentration from 2001 - 2003 to 2008 - 2010

Figure 8.29 Change in 95th percentile summer season (April-October) daily 8-hr maximum O₃ concentrations between 2001-2003 and 2008-2010.

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4 To examine these trends further, we evaluate the 1998-2011 data from the 15 case-study areas. Only monitors within the 15 study areas⁸ were analyzed, and within each study area, 5 6 monitors were put into three groups based on the degree of urbanization. The degrees of 7 urbanization were determined by the population density of the census tract containing the 8 monitor (plotted in Figure 8.30). Population data were obtained from the U.S. Census Bureau⁹, 9 and the classes were determined by breaks in the population density calculated from those data: "high population density" (> 1000 people/km²), "medium population density" (between 400 and 10 11 1000 people/km²), and "low population density" (< 400 people/km²). Data were additionally split 12 out into three different time periods (all months, warm months: May through September, and 13 cool months: October through April). These warm and cool season categorizations were chosen 14 to isolate effects that are observed at different times of year. The April-October time period

⁸ These 15 areas are the 12 urban case study areas in the epidemiological-based risk assessment and the 3 additional exposure urban case study areas.

⁹ Obtained from: http://www2.census.gov/geo/tiger/TIGER2010DP1/Tract_2010Census_DP1.zip

- 1 which was examined in Figure 8.28 and Figure 8.29 include all warm season and two cold
- 2 season months and thus show behavior that has influences from both. Summaries were thus
- 3 calculated for groups of monitors specific to 1) Study Area, 2) Month subset, and 3) Urban class.
- 4



6 Figure 8.30 Population density at each O₃ monitor.

7

8 Figure 8.31, Figure 8.32, and Figure 8.33 display the data described above in ribbon plots 9 for high, medium, and low population density monitor locations in each case study area. The lines bordering the dark and light red ribbons in this plot are (from top to bottom) the 95th, 75th, 10 25th, and 5th percentiles of the annual data indicated by each panel, and the median (i.e. 50th 11 percentile) is shown by the line in the middle of the central lighter ribbon. The colors of the lines 12 13 separating the ribbons depict significant trends (dark blue for decreasing and light blue for 14 increasing) or no significant trend (white). Statistical significance for multi-year O₃ trends was 15 determined using the Spearman rank order correlation coefficient (p-value < 0.05). Plots showing a characterization of the entire O₃ distribution (not just discrete cut points of 5th, 25th, 50th, 75th, 16 17 and 95th percentiles) are provided in Appendix 8-C. These plots show consistent trends over the past 13 years for O_3 , with high O_3 values 18 19 decreasing fairly uniformly across different regions and areas of different degrees of

20 urbanization. Conversely, mean and median trends appear quite different in high, medium, and

1 low population density areas. Mid-range O_3 concentrations at low population density locations 2 within the case study areas (so still relatively close to a major city) have generally decreased over a period of substantial NOx emissions reductions¹⁰. This decrease is most pronounced in the 3 4 summer months and in the Eastern half of the U.S. (low population density monitors in 3 out of the 5 Western¹¹ case-study areas and in only 4 out of the 10 Eastern¹² case study areas do not 5 have significant decreases in summertime median O₃ concentrations). Mid-range O₃ 6 7 concentrations in many, but not all, high population density areas have significantly increased in 8 winter months. Wintertime increases were significant in 11 of the 15 areas (only Atlanta, Boston, 9 Houston and Sacramento did not increase significantly). Thirteen out of 15 summertime high population density area trends in median O_3 were not significant¹³, but combining winter and 10 summer measurements to determine annual trends showed that Denver, Los Angeles, New York 11 and Philadelphia high population density sites had significantly increasing annual median O_3 12 while Boston, Chicago, Dallas and St. Louis had significantly increasing 25th percentile O₃ but 13 14 no significant median trend. These results reflect increasing mid-range O₃ concentrations mainly 15 confined to urban centers during periods of NOx reductions. One important point to note is that the design value monitor (the monitor with the highest average (over three years) of 4th highest 16 17 daily maximum value) in most of the case-study locations is located outside of the high 18 population density area (as defined here). Downward trends in medium and low population 19 density areas are therefore generally representative of the behavior at the highest O_3 monitor in 20 an area, whereas trends in urban centers may be important from an exposure perspective. 21 In summary, any increasing O_3 trends occur more in highly populated areas, during cool 22 months, and at the lower end of the O_3 distribution. Conversely, any decreasing O_3 trends occur 23 more during warm months, in lower population areas, and at the upper end of the O₃ distribution. 24 One result of these two phenomena is a narrowing of the range of O_3 concentrations over this 25 period of decreasing NOx emissions. For instance, there are many cases where the top and 26 bottom of a single distribution exhibit different trends. For example, the low population density 27 monitors of Dallas, Los Angeles, Philadelphia and Saint Louis and the high population density 28 monitors for Baltimore, Dallas, and Philadelphia for all months show a significant increase in the 5th percentile and a simultaneous significant decrease in the 95th percentile. More common is a 29

¹⁰ Denver is an outlier among the case study areas with consistently increasing mid-range ozone trends across seasons and urban classifications. Denver may be subject to increasing emissions from large wildfires and oil and gas development which are not typical of other urban case study areas. In addition, Denver is particularly susceptible to influences from stratospheric intrusions and international transport due to its high altitude

¹¹ Western case study areas for this purpose include: Dallas, Denver, Houston, Los Angeles, and Sacramento

¹² Eastern case study areas for this purpose include: Atlanta, Baltimore, Boston, Chicago, Cleveland, Detroit, New York, Philadelphia, St. Louis, and Washington D.C.

¹³ Only Houston and Dallas had statistically significant trends in median summertime urban ozone

1 significant change in one end of the distribution, but no significant change in the other (e.g., the

- 2 summer months at high population density monitors in all case study areas except Baltimore,
- 3 Chicago, and Detroit). It is important to note that there are also cases where both ends of the
- 4 distribution change in the same manner and there is therefore no narrowing of the range of O_3
- 5 concentrations in these areas.
- 6



Figure 8.31 Distributions of O₃ concentrations for high population density monitors by
 different subsets of months over a 13-year period. From top to bottom in each
 ribbon plot, the blue and white lines indicate the spatial mean of the 95th, 75th,
 50th, 25th, and 5th percentiles for each monitor for every year from 1998-2011.



Figure 8.32 Distributions of O₃ concentrations for medium population density monitors
by different subsets of months over a 13-year period. From top to bottom in
each ribbon plot, the blue and white lines indicate the spatial mean of the 95th,
75th, 50th, 25th, and 5th percentiles for each monitor for every year from 19982011.



Figure 8.33 Distributions of O₃ concentrations for low population density monitors by different subsets of months over a 13-year period. From top to bottom in each ribbon plot, the blue and white lines indicate the spatial mean of the 95th, 75th, 50th, 25th, and 5th percentiles for each monitor for every year from 1998-2011.

6

7 Maps of ambient trends in both New York City and Chicago most clearly show these 8 trends and further illustrate this behavior. Figures 8.34 and 8.35 show trends in daily maximum 8-hour O₃ values these two cities for May-September. Plots for other case-study areas are 9 10 provided in Appendix 8-C. For both cities, the fourth highest 8-hr daily maximum O₃ value 11 either has a downward trend or no trend at all monitors. In New York (Figure 8.34), mean and 12 median O_3 values significantly decrease at downwind locations in New York and Connecticut. Conversely, median O₃ values significantly increase from 1998 to 2011 at two core urban sites 13 14 (one at City College of NY in upper Manhattan and one near Queen's college) and at a nearby 15 site on Long Island. Similarly, in Chicago (Figure 8.35), mean and median trends in O_3 are 16 downward or insignificant in Indiana and in suburban Illinois locations and show increases near 17 the highly populated urban core. 18



Figure 8.34 Map of O_3 trends at specific monitors in the New York area. All upward and downward facing triangles represent statistically significant trends from 1998-2011 (p < 0.05), circles represent locations with no significant trends. Sites used in Smith et al (2009) and the Zanobetti and Schwartz (2008) epidemiology studies are represented by colored dots. Only monitors with at least seven years of data are displayed. The pink star indicates the site with the higher design value in 2011. The MSA border as defined by the U.S. census bureau is delineated by the light blue line. Left panel shows trends in annual 4th highest 8-hr daily maximum O_3 values, center panel shows trends in annual mean 8-hr daily maximum O_3 values, and right panel shows trends in annual median 8-hr daily maximum O_3 values.



Figure 8.35 Map of O_3 trends at specific monitors in the Chicago area. All upward and downward facing triangles represent statistically significant trends from 1998-2011 (p < 0.05), circles represent locations with no significant trends. Sites used in Smith et al (2009) and the Zanobetti and Schwartz (2008) epidemiology studies are represented by colored dots. Only monitors with at least seven years of data are displayed. The pink star indicates the site with the higher design value in 2011. The MSA border as defined by the U.S. census bureau is delineated by the light blue line. Left panel shows trends in annual 4th highest 8-hr daily maximum O_3 values, center panel shows trends in annual mean 8-hr daily maximum O_3 values, and right panel shows trends in annual median 8-hr daily maximum O_3 values.

1 To demonstrate how changes in emissions of NOx and anthropogenic VOCs might be 2 driving these trends, Table 8-7 shows trends of O₃ in high and low population areas and annual National Emissions Inventories (NEI) for 2002, 2005, 2008 and 2011¹⁴ aggregated to the level of 3 the NOAA Climate Regions¹⁵. There is moderate correspondence between the decreases in NOx 4 5 emissions across the regions with the observed decreases in warm season O₃ concentrations in 6 low population areas. VOCs show little correspondence to any of the O₃ trends, which is likely 7 due to complications from 1) the mix of chemicals with a large range of reactivities; 2) complex 8 non-linear chemistry; and 3) the potential impact of the much larger magnitude of biogenic vs. 9 anthropogenic emissions on a regional scale. Details of these calculations can be found in 10 Appendix 8-C.

11

12Table - 8.7Broad Regional Annual Trends of Concurrent O3 Concentrations and13Emissions of NOx and VOCs over the 2000-2011 Time Period

Trend	Central	East North	North East	South	South	South West	West
		Central			East		
High Pop Dens, May2Sep O_3	none	none	none	down	none	none	none
High Pop Dens, Oct2Apr O_3	up	up	up	low %'s up	none	up	up
Low Pop Dens, May2Sep O_3	down	down	down	down	down	low %'s up	top %'s down
Low Pop Dens, Oct2Apr O_3	none	none	top %'s up	none	none	up	none
NO_x Emission	down	down	down	down	down	down	down
VOC Emission	none	none	down	up	down	up	down

12 15

8.2.3.2 Modeled O₃ response to emissions reductions across the United States

16 This section presents an analysis of the CMAQ modeling of O₃ responses to "across-theboard" U.S. anthropogenic precursor emissions reductions described in Appendix 4b. In this 17 18 analysis, we compare the modeled responses of O₃ concentrations in the case study areas to the 19 modeled O_3 responses in the rest of the U.S. For this purpose, we used five CMAQ model 20 simulations: 1) a base simulation which included 2007 emissions of all O_3 precursors, 2) a 50% 21 NOx cut simulation in which U.S. anthropogenic NOx emissions were reduced by 50% from 22 2007 levels, 3) a 90% NOx cut simulation in which U.S. anthropogenic NOx emission were 23 reduced by 90% from 2007 levels, 4) a 50% NOx/VOC cut simulation in which both U.S. 24 anthropogenic NOx and VOC emission were reduced by 50% from 2007 levels, and 5) a 90% 25 NOx/VOC cut simulation. These simulations are analyzed to characterize responses in O_3 to

¹⁴ http://www.epa.gov/ttnchie1/trends/

¹⁵ http://www.ncdc.noaa.gov/monitoring-references/maps/us-climate-regions.php

- 1 "across the board" emissions cuts at four distinct levels and do not represent the exact adjustment
- 2 cases that were used to estimate O₃ concentrations consistent with individual case study areas
- 3 just meeting various potential levels of the NAAQS standard. However, these four cases
- 4 generally span the range of emissions perturbations that were applied in the HDDM adjustment
- 5 methodology described in Chapter 4 and in Appendix 4d.
- 6 In this analysis we focus on seasonal mean O_3 and population as proxies for 7 epidemiology based risk estimates in Chapter 7. Since the epidemiology studies used in Chapter 8 7 show relatively linear response of health outcomes to O_3 concentrations throughout the entire 9 range of measured O_3 values, examining seasonal mean values should provide some 10 understanding of locations where O_3 health effects are expected to increase and decrease as a 11 result of precursor emission reductions. By combining population information with these spatial 12 distributions of seasonal O_3 responses, we can better understand expected O_3 behavior in
- 13 locations where people live. This is not a detailed risk assessment but can provide information on
- 14 the representativeness of the case-study areas to the nation as a whole in terms of expected O_3 -
- 15 related health outcomes.

16 To begin, we examine maps displaying ratios of mean O_3 concentrations in the emissions 17 cut simulations to mean O_3 concentrations in the 2007 base simulation. Figure 8.36 and Figure 18 8.37 show the ratio of seasonal (April-October) mean O₃ in the two NOx emissions reduction 19 simulations to that in the base simulation for the entire model domain. Figure 8.38 and Figure 20 8.39 depict the ratio of January mean O_3 for the two NOx cut simulations. Figures showing the 21 ratios based on the May-September seasonal average and figures for the NOx/VOC emissions 22 reductions scenarios are provided in Appendix 8-C. The maps show widespread decreases (i.e., 23 ratios less than 1) in seasonal mean O_3 across the country. These decreases are especially 24 pronounced in the Eastern U.S. and in California. O_3 increases (i.e., ratios greater than 1) are 25 confined to urban core areas except in January. The spatial extent of these O_3 increases are 26 generally less for the 90% NOx cut simulation than for the 50% NOx cut simulation although the 27 magnitude is greater over very limited areas in Chicago, Seattle, and San Francisco. The O₃ 28 increases are most widespread in the cooler months (January, April, and October). For the April-29 October seasonal average O_3 concentrations, VOC in addition to NOx cuts did not substantially 30 change the ratios of O_3 in the emissions reduction scenarios to O_3 in the base scenario. In the 31 Northeast and Midwest, increases in seasonal mean O₃ concentrations were mainly confined to 32 urban case study areas of New York, Detroit, Chicago, and St. Louis. In the Southeastern U.S., 33 the urban areas which show up as having increased seasonal mean O₃ in the 50% NOx cut 34 simulations include Miami, Orlando, Tampa, and New Orleans (only Miami has O₃ increase in 35 the 90% NOx cut simulation). The only case-study area in the southeast, Atlanta, does not 36 experience increases in seasonal mean O_3 in the model simulation (this is consistent with the

1 changes in risk estimated for Atlanta, which show no increases in total risk as alternative

- 2 standards are simulated). In the central U.S., seasonal mean O₃ in the case study areas of Denver,
- 3 Houston, and Dallas and non-case-study areas of San Antonio, Duluth, and Minneapolis
- 4 increased with 50% NOx reductions. Seasonal mean O₃ increases were seen only in Houston,
- 5 Minneapolis, and Duluth with 90% reductions in simulated NOx emissions. The Northwestern
- 6 U.S. showed some of the most widespread increases in seasonal mean O_3 in the 50% and 90%
- 7 NOx cut simulations covering the Seattle and Portland metro areas as well the San Francisco Bay
- 8 area and in a single model grid cell for Sacramento (50% NOx reduction case only). Sacramento
- 9 is the only city in the Northwest that was included as a case study area. Finally, Los Angeles (a
- 10 case study area), San Diego, Phoenix, and Bakersfield were the areas for which CMAQ predicted
- 11 seasonal mean O₃ increases with the 50% NOx cut simulation. These O₃ increases disappeared
- 12 (or were largely diminished in the case of LA) in the 90% NOx cut case. Based on these maps, it
- 13 appears that in the Northeast and the Central U.S., the case-study area selection likely
- 14 oversampled these O₃ increases on a geographic basis since all locations outside of city centers
- 15 experienced decreasing seasonal mean O₃ with the NOx reduction model simulations. However
- 16 in two regions, the Southeast and the Northwest, the urban case study area did not experience
- 17 increases in seasonal mean O₃ concentrations while other urban areas in the region did. In these
- 18 two regions, the urban case-study selection likely under-sampled the locations which
- 19 experienced increases in seasonal mean O₃.





Figure 8-36 Ratio of April-October seasonal average O₃ concentrations in the brute force 50% NOx emissions reduction CMAQ simulations to April-October seasonal average O₃

3 4 concentrations in the 2007 base CMAQ simulation.



Figure 8.37 Ratio of April-October seasonal average O₃ concentrations in the brute force 90% NOx emissions reduction CMAQ simulations to April-October seasonal average O₃ concentrations in the 2007 base CMAQ simulation.





emissions reduction CMAQ simulations to January monthly average O₃ concentrations in the 2007 base CMAQ simulation.



Figure 8.39 Ratio of January monthly average O₃ concentrations in brute force 90% NOx
 emissions reduction CMAQ simulations to January monthly average O₃
 concentrations in the 2007 base CMAQ simulation.

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In order to characterize the representativeness of case study areas in a more quantitative 6 manner, paired O₃ concentrations and population data¹⁶ were extracted from each model grid cell 7 8 and categorized in various manners. Figure 8.40 and Figure 8.41 depict the percent of U.S. 9 population living in areas with increases or decreases in monthly or seasonal mean O_3 under the 10 emissions reductions scenarios (50% NOx cut and 90% NOx cut respectively) compared to O₃ in 11 the base modeling scenario. The top panels show data for January monthly mean O₃, the center 12 panels show data for seasonal mean O₃ (June-August), and the bottom panels show data for seasonal mean O₃ (April-October). Tabulated results and equivalent plots for the combined 13

¹⁶ Block level population data from the 2010 Census was aggregated to the 12km CMAQ grid cell level. The 2007 population was then calculated using population growth factors developed by Woods and Poole Economics, Inc.

- NOx/VOC cut simulations are provided in Appendix 8-C. Month by month break-outs for each
 case study area are also available in Appendix 8-C.
- 3 The vast majority of the U.S. population lives in areas where the CMAQ simulations 4 predict mean O₃ decreases for the June-August and April-October time periods. The majority of 5 population living in case-study areas also lives in locations with decreasing seasonal mean O_3 6 concentration under NOx reduction scenarios. As discussed previously, more locations have 7 increasing mean O_3 in the cooler months as demonstrated by the fact that almost all of the U.S. 8 population lives in locations where the model predicts increases in mean O_3 in January. The case 9 study areas represent 29% of the total U.S. population. These areas account for 20-30% of the 10 U.S. population that experience decreasing seasonal mean O₃ for April-October in the NOx cut simulations and 50-60% of the U.S. population that experience increasing seasonal O₃ for April-11 12 October. Consequently, the urban-case study areas over-sample populations living in locations 13 with increasing seasonal mean O_3 in response to NOx cuts compared to populations living in 14 locations with decreases in seasonal mean O₃. In all panels displayed in Figures 8.28 and 8.29, 15 most of the population lives in locations where increases or decreases in mean O₃ were greater
- 16 than 1 ppb.
- 17



Figure 8.40 Histograms of U.S. population living in locations with increasing and decreasing mean O₃. Values on the x-axis represent change in mean O₃ (ppb) from the 2007 base CMAQ simulation to the 50% NOx cut CMAQ simulation. The percentages of the U.S. population living in areas that have changes less than -1 ppb, from -1 to +1 ppb, and greater than 1 ppb are shown on the y-axis. Left plots show population numbers in locations not included in one of the cases study areas while right plots show population numbers in locations included in one of the case study areas. Top plots show changes in January monthly mean O₃, middle plots show changes in seasonal mean June-August O₃, and bottom plots show changes in seasonal mean April-October O₃.


Figure 8.41 Histograms of U.S. population living in locations with increasing and decreasing mean O₃. Values on the x-axis represent change in mean O₃ (ppb) from the 2007 base CMAQ simulation to the 90% NOx cut CMAQ simulation. The percentages of the U.S. population living in areas that have changes less than -1 ppb, from -1 to +1 ppb, and greater than 1 ppb are shown on the y-axis. Left plots show population numbers in locations not included in one of the cases study areas while right plots show population numbers in locations included in one of the case study areas. Top plots show changes in January monthly mean O₃, middle plots show changes in seasonal mean June-August O₃, and bottom plots show changes in seasonal mean April-October O₃.

1 The proportion of the population living in locations of increasing seasonal mean O_3 in response to NOx emissions reductions varies considerably between case study areas. Figure 8.42 2 3 and Figure 8.43 show these proportions by city for the 50% and 90% NOx reduction scenarios. 4 For the 50% NOx reduction scenario, the CMAQ results predict that four out of fifteen study 5 ares (Chicago, Detroit, Los Angeles, and New York) have more than 50% of their populations 6 living in locations with increasing mean O₃ for April-October. Most other urban case study areas 7 have between 5% and 30% of their populations living in these areas with increasing mean O₃ 8 levels. For the 90% NOx reduction scenario, the percent of population living in such locations 9 decreases substantially for all cities, leaving four out of fifteen study areas (Detroit, Houston, 10 Los Angeles, and New York) with more than 5% of their populations living in areas with

- 11 increasing mean O_3 levels.
- 12



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Figure 8.42 Population (as % of total case-study area population) living in locations of
 increasing April-October seasonal mean O₃ in the 50% NOx reduction
 CMAQ simulation.



Figure 8.43 Population (as % of total case-study area population) living in locations of increasing April-October seasonal mean O₃ in the 90% NOx reduction CMAQ simulation.

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6 We can further understand these results by looking at them in terms of population density 7 in the case-study areas versus across the U.S. as a whole. As in Section 8.2.3.1, we define census 8 tracts with population density greater than 1000 people/km² as high population density, but the 9 low-mid population density classification used here is a combination of the low and medium 10 classifications in that section. Figure 8.44 and Figure 8.45 split out the April-October results from Figure 8.40 and Figure 8.41 into high and low-mid sub-categories. Appendix 8-C provides 11 12 similar breakouts for the other panels in Figure 8.40 and Figure 8.41. First, based on these 13 definitions, we see that 57% of the population in case-study areas lives in high population 14 density locations while only 27% of the U.S. population does. As discussed above, the high 15 population areas are more likely to experience increases in mean O₃ as a result of NOx emission 16 reductions compared to lower population areas. Therefore, the fact that the case-study areas used 17 in the risk and exposure assessments are more densely populated than the country as a whole 18 means that these analyses may estimate higher risks under emissions reduction scenarios than 19 would be experienced, on average, across the country. Figure 8.44 and Figure 8.45 show 20 generally similar shapes for the high population density histograms in the study-area and non-21 study area locations. In the 50% NOx cut simulation, 69% of the population living in high 22 density case-study areas would experience increases in mean seasonal O₃ compared to 63% of 23 the population the population living in high density areas of the country as a whole. Similarly in the 90% NOx cut simulation, 28% of the population in high density locations both within the 24 25 study areas and across the U.S. as a whole lives in locations of increasing seasonal mean O_3 . This

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- 1 suggests that the selected study areas adequately represent population-weighted changes in mean
- 2 O₃ for people living in high density areas. Similarly, less densely populated locations within the
- 3 case-study areas show O_3 increases equivalent to those seen in less densely populated areas in
- 4 the U.S. as a whole. In the 50% NOx cut simulation, 7% of people in low-mid density study area
- 5 locations live where mean seasonal O₃ is increasing, while 5% of people in all low-mid density
- 6 U.S. locations live where mean seasonal O_3 is increasing. Similarly, in the 90% NOx cut
- 7 simulation, the numbers are 2% for both low-mid density study area populations and for low-mid
- 8 density populations in the U.S. as a whole. Thus the oversampling of populations living in
- 9 locations of increasing mean seasonal O_3 in response to NOx cuts, as shown in Figures 8.28 and
- 10 8.30, appears to be entirely due to the fact that the study areas oversample populations living in
- 11 high density areas compared to the U.S. population as a whole.
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Figure 8.44 Histograms of U.S. population living in locations with increasing and decreasing mean O₃. Values on the x-axis represent the change in seasonal mean (April-October) O₃ from the 2007 base CMAQ simulation to the 50% NOx cut CMAQ simulation. The percentages of the U.S. population living in areas that have changes less than -1 ppb, from -1 to +1 ppb, and greater than 1 ppb are shown on the y-axis. Left plots show population numbers in locations not included in one of the cases study areas while right plots show population numbers in locations included in one of the urban case study areas. Bottom plots show histograms for low-mid population density areas while top plots show histograms for high population density areas.



Figure 8.45 Histograms of U.S. population living in locations with increasing and decreasing mean O₃. Values on the x-axis represent the change in seasonal mean (April-October) O₃ from the 2007 base CMAQ simulation to the 90% NOx cut CMAQ simulation. The percentages of the U.S. population living in areas that have changes less than -1 ppb, from -1 to +1 ppb, and greater than 1 ppb are shown on the y-axis. Left plots show population numbers in locations not included in one of the cases study areas while right plots show population numbers in locations included in one of the urban case study areas. Bottom plots show histograms for low-mid population density areas while top plots show histograms for high population density areas.

- 1 **8.2.5 Discussion**
- 2

We evaluated two different questions, 1) to what degree are the 15 cities evaluated in the exposure and risk analyses representative of the overall U.S. population with regards to total O_3 risk?, and 2) to what degree are they representative of the overall U.S. population with regards to the degree of risk reduction that might be observed in response to just meeting the existing and alternative standards.

8 Regarding the first question, we observe that the 12 urban study areas considered in the 9 urban-scale risk assessment presented in Section 7.2 capture urban areas that are among the most 10 populated in the U.S., have relatively high O₃ levels, and represent the range of city-specific 11 effect estimates found by Smith et al. (2009) and Zanobetti and Schwartz (2008). These three 12 factors suggest that the urban study areas capture overall risk for the nation well, with a potential 13 for better characterization of the high end of the risk distribution. We find that the urban study 14 areas are not capturing areas with the highest baseline mortality rates, those with the oldest 15 populations, and those with the lowest air conditioning prevalence. These areas tend to have 16 relatively low O_3 concentrations and low total population, suggesting that the urban study areas 17 are not missing high risk populations that have high O₃ concentrations in addition to greater 18 susceptibility per unit O₃. We also find that the 12 urban study areas represent the full range of 19 county-level O₃-related risk across the entire U.S. We conclude from these analyses that the 12 20 urban study areas adequately represent O₃-related risk across the U.S. 21 Concerning the second question, we observe that the 15 urban areas considered in the 22 exposure and risk assessment case study areas over-sample populations living in locations with

23 increasing seasonal mean O_3 in response to NOx cuts. This suggests that the selected study areas

24 adequately represent population-weighted changes in mean O_3 concentrations for urban

25 populations, but may be under-representing decreasing median O_3 concentrations in suburban

and rural areas. As a result, the risk estimates for populations in the selected urban study areas

27 may understate the risk reductions that might be achieved across the broader U.S. population.

2 8.3 REFERENCES

3	Abt Associates, Inc. 2010. Model Attainment Test Software (Version 2), prepared for EPA.
4	Research Triangle Park, NC: EPA Office of Air and Radiation, Office of Air Quality
5	Planning and Standards. Research Triangle Park, NC. Available at:
6	<http: modelingapps.mats.htm="" scram001="" www.epa.gov="">.</http:>
7	Abt Associates, Inc. 2010. Environmental Benefits and Mapping Program (Version 4.0),
8	prepared for EPA. Research Triangle Park, NC: EPA Office of Air Quality Planning and
9	Standards. Available at <http: air="" benmap="" www.epa.gov="">.</http:>
10	• Anenberg, S.C.; J.J. West; L.W. Horowitz and D.Q. Tong. 2010. "An Estimate of the
11	Global Burden of Anthropogenic Ozone and Fine Particulate Matter on Premature
12	Human Mortality Using Atmospheric Modeling." Environmental Health Perspective,
13	118:1189-1195.
14	Anenberg, S.C.; J.J. West; L.W. Horowitz and D.Q. Tong. 2011. "The Global Burden of Air
15	Pollution Mortality: Anenberg et al. respond." Environmental Health Perspective,
16	119:A158-A425.
17	Bell, M.L.; A. McDermott; S.L. Zeger; J.M. Samet and F. Dominici. 2004. "Ozone and Short-
18	term Mortality in 95 U.S. Urban Communities, 1987-2000." Journal of the American
19	Medical Association, 292:2372-2378.
20	Byun, D. and K. L. Schere. 2006. "Review of Governing Equations, Computational Algorithms,
21	and Other Components of the Models-3 Community Multi-scale Air Quality (CMAQ)
22	Modeling System." Applied Mechanics Reviews, 59:51-77.
23	Centers for Disease Control: Wide-ranging Online Data for Epidemiological Research (CDC-
24	Wonder) (data from years 2004-2006), Centers for Disease Control and Prevention
25	(CDC), U.S. Department of Health and Human Services. Available on the Internet at
26	< <u>http://wonder.cdc.gov</u> >.
27	Cooper, O.R.; R. Gao; D. Tarasick; T. Leblanc and C. Sweeney. 2012. Long-term Ozone Trends
28	at Rural Ozone Monitoring Sites Across the United States, 1990-2010." Journal of
29	Geophysical Research, 117, D22, doi:10.1029/2012JD018261.

1	Fann N.; A. D. Lamson; S. C. Anenberg; K. Wesson; D. Risley; B. J. Hubbell. 2012.
2	"Estimating the National Public Health Burden Associated with Exposure to Ambient
3	PM _{2.5} and Ozone." <i>Risk Analysis</i> , 32:81-95.
4	George, B.J. and T. McCurdy. 2009. "Investigating the American Time Use Survey from an
5	exposure modeling perspective." Journal of Exposure Science and Environmental
6	Epidemiology, 21:92-105.
7	Graham, S. and T. McCurdy. 2004. "Developing Meaningful Cohorts for Human Exposure
8	Models." Journal of Exposure Analysis and Environmental Epidemiology, 14:23-43.
9	Hollman, F.W.; T. J. Mulder and J. E. Kallan. 2000. "Methodology and Assumptions for the
10	Population Projections of the United States: 1999 to 2100." Population Division Working
11	Paper No. 38, Population Projections Branch, Population Division, U.S. Census Bureau,
12	Department of Commerce.
13	Jerrett, M.; R.T. Burnett; C. A. Pope, III; K. Ito; G. Thurston; D. Krewski; Y. Shi; E. Calle; M.
14	Thun. 2009. "Long-term O3 Exposure and Mortality." New England Journal of Medicine,
15	360:1085-1095.
16	Neidell, M. 2009. Information, avoidance behavior and health. J Human Res. 44:450-478.
17	Neidell, M. 2010. Air quality warnings and outdoor activities: evidence from Southern California
18	using a regression discontinuity approach design. J Epidemiol Community Health.
19	64:921-926.
20	Samet, J. M.; S. L. Zeger; F. Dominici; F. Curriero; I. Coursac; D.W. Dockery; J. Schwartz and
21	A. Zanobetti. (2000). "The National Morbidity, Mortality, and Air Pollution Study Part
22	II: Morbidity and Mortality from Air Pollution in the United States." Health Effects
23	Institute, Boston, MA, Number 94, Part II.
24	Smith R.L.; B. Xu and P. Switzer. 2009. "Reassessing the Relationship Between Ozone and
25	Short-term Mortality in U.S. Urban Communities." Inhalation Toxicology, 21:37-61.
26	Timin, B.; K. Wesson and J. Thurman. "Application of Model and Ambient Data Fusion
27	Techniques to Predict Current and Future Year PM2.5 Concentrations in Unmonitored
28	Areas. (2010)," pp. 175-179 in Steyn DG, Rao St (eds). Air Pollution Modeling and Its
29	Application XX. Netherlands: Springer.

1 2 2	U.S. Environmental Protection Agency. 2013. <i>Integrated Science Assessment for Ozone and</i> <i>Related Photochemical Oxidants</i> . Research Triangle Park, NC: EPA. (EPA document
3	number Er A 600/R-10/0701').
4	U.S. EPA. 2010. Quantitative Health Risk Assessment for Particulate Matter. Research Triangle
5	Park, NC: EPA. (EPA document number EPA-452/R-10-005).
6	U.S. EPA. 2009. Risk and Exposure Assessment to Support the Review of the SO2 Primary
7	National Ambient Air Quality Standards: Final Report. Research Triangle Park, NC:
8	U. S. EPA. (EPA document number EPA-452/R-09-007).
9	U.S. EPA. 2007. Ozone Population Exposure Analysis for Selected Urban Areas. Research
10	Triangle Park, NC: EPA. (EPA document number EPA-452/R-07-010). Available at:
11	http://www.epa.gov/ttn/naaqs/standards/ozone/data/2007_07_ozone_exposure_tsd.pdf >.
12	Woods and Poole Economics Inc. 2012. "Complete Demographic Database." Woods and Poole
13	Economics, Inc. Washington, DC. Available at:
14	< <u>http://www.woodsandpoole.com/index.php</u> .>.
15	
16	Zanobetti, A. and J. Schwartz. 2008. "Mortality Displacement in the Association of Ozone with
17	Mortality: An Analysis of 48 Cities in the United States." American Journal of
18	Respiratory and Critical Care Medicine, 177:184-189.
19	Zanobetti, A. and J. Schwartz. 2011. "Ozone and Survival in Four Cohorts with Potentially
20	Predisposing Diseases." American Journal of Respiratory and Critical Care Medicine,
21	194:836-841.
22	Zhang, L.; D. J. Jacob; N.V. Smith-Downey; D. A. Wood; D. Blewitt; C. C. Carouge; A. van
23	Donkelaar; D. B. A. Jones; L.T. Murray and Y. Wang. 2011. "Improved Estimate of the
24	Policy-relevant Background Ozone in the United States Using the GEOS-Chem Global
25	Model with 1/2°x2/3° Horizontal Resolution Over North America." Atmospheric
26	Environment, 45:6769-6776.
27	

9 SYNTHESIS

2 9.1 INTRODUCTION

3 This assessment estimates exposures to O_3 and resulting mortality and morbidity health 4 risks based on the findings of the O_3 ISA (U.S. EPA, 2013) that short-term O_3 exposures are 5 causally related to respiratory effects, and likely causally related to cardiovascular effects, and 6 that long term O_3 exposures are likely causally related to respiratory effects. The assessment 7 evaluates total exposures and risks associated with the full range of observed O₃ concentrations, 8 as well as the incremental changes in exposures and risks between just meeting the existing 9 standard of 75 ppb and just meeting alternative standard levels of 70, 65, and 60 ppb using the form and averaging time of the existing standard: the annual 4th highest daily maximum 8-hour 10 11 O₃ concentration, averaged over three consecutive years. We evaluated alternative standard 12 levels of 70, 65, and 60 consistent with recommendations from CASAC to consider alternative 13 standard levels between 60 and 70 ppb (Frey and Samet, 2012). 14 Following the conceptual framework described in Chapter 2, the assessment evaluates 15 exposures and lung function risk in 15 urban case study areas, and mortality and morbidity risks 16 based on concentration-response functions derived from epidemiology studies in 12 of these urban case study areas¹. The results from these assessments will help inform consideration of the 17 18 adequacy of the existing primary O₃ standards, and potential risk reductions associated with 19 several alternative levels of the standard (for the current form and averaging time). In addition, to 20 place the urban case study area analyses in a broader context, Chapter 8 of this assessment 21 estimates the national burden of mortality associated with recent O₃ levels, and evaluates the 22 representativeness of the 15 urban case study areas in characterizing O₃ exposures and risks 23 across the U.S. This synthesis focuses on the urban case study area assessments of exposure and 24 risk for the scenarios of just meeting the existing and alternative standards. For this synthesis, we 25 discuss the results of the national-scale assessment as they relate to understanding the breadth of 26 O₃ risks across the U.S. and to the national representativeness of the urban case study area risk 27 results.

To facilitate interpretation of the results of the exposure and risk assessment, this chapter provides a synthesis of the various results, focusing on comparing and contrasting those results to identify common patterns, or important differences. These comparisons will focus on patterns

¹ Three additional urban case study areas were evaluated for the human exposure assessment and lung function risk assessment to provide greater geographic representation. There was insufficient information available to conduct the epidemiology-based risk assessment in these 3 additional areas. Also, we originally planned to include Seattle, WA as a 16th urban case study area, but due to limitations in the available air quality monitoring data, we determined that it would not be appropriate to model exposure and risks for Seattle (see appendix 4-E).

1 across urban case study areas, across years of analysis, and across alternative standards. In

- 2 addition, factors related to each specific type of analysis that may influence comparisons
- 3 between the analyses are identified and discussed. The degree to which the integrated results are
- 4 representative of national patterns of exposure and risk is evaluated. Overall confidence in the
- 5 results, as well as relative confidence between the different analyses is also assessed. The chapter
- 6 concludes with an overall integrated characterization of exposure and risk in the context of key
- 7 policy-relevant questions raised in Chapter 2.

8 9.2 SUMMARY OF KEY RESULTS

9 9.2.1 Air Quality Considerations (Chapter 4)

10 Table 9-1 below gives information on the monitoring network, population, and observed 11 peak O₃ concentrations for the 15 case study areas, for the years included in the exposure, lung function risk, and epidemiology based risk assessments. The number of counties, number of O₃ 12 13 monitors, population, and design values (DV) are based on the area definitions used in the 14 exposure modeling and clinical-based lung function risk assessments, while the 2007 and 2009 annual 4th highest values are based on the Core Based Statistical Areas (CBSAs) used in the 15 epidemiology-based risk assessment. The "N/A" values in the 2007 and 2009 4th high columns 16 are for the three urban areas not included in the epidemiology-based risk assessment. The data 17 show a trend of lower peak O3 concentrations (i.e., the 2008-2010 design values and 2009 4th 18 high values are generally lower than the 2006-2008 design values and 2007 4th high values. 19 20 respectively).

Area Name	# of Counties	# of O ₃ Monitors	Population (2010)	2006-2008 DV (ppb)	2007 4 th high (ppb)	2008-2010 DV (ppb)	2009 4 th high (ppb)
Atlanta	33	13	5,618,431	95	102	80	77
Baltimore	7	7	2,710,489	91	92	89	83
Boston	10	14	5,723,468	83	89	77	75
Chicago	16	26	9,686,021	78	N/A	74	N/A
Cleveland	8	13	2,881,937	82	83	77	72
Dallas	11	20	6,366,542	89	N/A	86	N/A
Denver	13	26	3,390,504	86	97	77	79
Detroit	9	12	5,218,852	81	93	75	73
Houston	10	22	5,946,800	91	90	85	91
Los Angeles	5	54	17,877,006	119	105	112	108
New York	27	31	21,056,173	90	94	84	81
Philadelphia	15	19	7,070,622	92	102	83	74
Sacramento	7	26	2,755,972	102	93	102	96
St. Louis	17	17	2,837,592	85	94	77	74
Washington	26	22	5,838,518	87	N/A	81	N/A

2 Table 9-1 Area and Monitoring Information for the 15 Case Study Areas

4 In this analysis, we employed a photochemical model-based adjustment methodology 5 Simon et al. (2012) using the Higher-Order Decoupled Direct Method (HDDM) capabilities in 6 the Community Multi-scale Air Quality Model (CMAQ) (hereafter referred to as HDDM air 7 quality adjustment). The HDDM air quality adjustment methodology replaced the quadratic 8 rollback technique used in the first draft REA to estimate O₃ concentrations consistent with just 9 meeting existing and alternative O₃ standards. The HDDM air quality adjustment procedure 10 estimates the change in observed hourly O_3 concentrations at a given set of monitoring sites 11 resulting from national across-the-board reductions in U.S. anthropogenic NOx and/or VOC 12 emissions. In this analysis, we adjusted O_3 concentrations to just meet the existing standard of 75

- 13 ppb^2 and potential alternative standards of 70, 65, and 60 ppb at ambient monitoring sites in the
- 14 15 case study areas for the 2006-2008 and 2008-2010 periods. In most locations, only NOx
- 15 reductions were used to adjust the distribution of O_3 concentrations, because of the
- 16 ineffectiveness of VOC reductions in reducing peak O₃ concentrations needed to meet the

² Attainment with the existing standard level of 75ppb is determined by the 4th highest maximum 8-hour O₃ concentration, averaged over 3 years (hereafter referred to as the existing standard).

existing and alternative standard levels. Sensitivity analyses were also conducted in some
 locations to evaluate the impact of decreasing both NOx and VOC emissions.

3 The HDDM air quality adjustment methodology represents a substantial improvement 4 over the quadratic rollback method used to adjust O_3 concentrations in previous reviews. First, 5 quadratic rollback was a purely mathematical technique which attempted to reproduce the 6 distribution of observed O_3 concentrations just meeting various standards, while the new 7 methodology uses photochemical modeling to simulate the response in O₃ concentrations due to 8 changes in precursor emissions based on current understanding of atmospheric chemistry and 9 transport. Second, quadratic rollback used the same mathematical formula to adjust 10 concentrations at all monitors within each case study area for all hours, while HDDM allows the 11 adjustments to vary both spatially across each case study area and temporally across hours of the 12 day and across seasons. Finally, quadratic rollback was designed to only allow decreases in O₃ 13 concentrations, while the HDDM air quality adjustment allows both increases and decreases in 14 O₃ concentrations in response to reductions in NOx or VOC emissions. For example, in response 15 to reductions in NOx emissions, the HDDM methodology is able to capture increases in O₃ 16 concentrations that can occur in urban cores characterized by titration of O_3 by fresh NO 17 emissions and decreases in O₃ concentrations downwind.

18 Following HDDM adjustment of O₃ concentrations, several general trends are evident in 19 the changes in O_3 patterns across the case study areas and across the alternative standard levels. 20 In all 15 case study areas, peak O_3 concentrations tended to decrease while the O_3 concentrations 21 in the lower part of the distribution of O_3 tended to increase as the concentrations were adjusted to meet the existing and alternative standards. In addition, O3 concentrations in the high and mid-22 23 range portions of the O₃ distribution generally decreased in the outer, more rural and suburban 24 portions of the urban case study areas, while the O₃ response to NOx reductions was more varied within the urban cores. In particular, while the peak (annual 4th highest daily maximum 8-hour) 25 26 concentrations upon which the existing and alternative standards are defined generally decreased 27 in the urban core of the case study areas in response to modeled reductions in primarily NOx 28 emissions, the O₃ responses near the center of the O₃ distribution at these locations followed one 29 of three patterns when focusing on the mean of the daily maximum 8-hour O_3 concentrations 30 from May to September, as shown in Table 9-2.

Table 9-2 General Patterns in Seasonal (May-Sept) Mean of Daily Maximum 8-hour O₃ Concentrations after Adjusting to Meet Existing and Alternative Standards*

After Adjusting to Meet Existing Standard	After Further Adjusting to Meet Lower Alternative Standards	Case Study Areas Showing Pattern
Decreased	Continued to decrease	Atlanta Sacramento
Decreased	Continued to decrease	Washington, D.C
		Baltimore
		Cleveland,
		Dallas
Increased	Decreased	Detroit
lifereased	Decreased	Los Angeles
		New York
		Philadelphia
		St. Louis
Increased		Boston
	Continued to increase or remained	Chicago
Increased	constant	Denver
		Houston

3 4 * These patterns refer to O_3 responses in the urban core of each urban case study area based on analysis of the interpolated monitor values used as inputs to the exposure and lung function risk analyses.

5

6 The air quality inputs to the exposure modeling and clinical-based lung function risk 7 assessments were estimated hourly O_3 concentrations at each census tract in the 15 case study 8 areas. These values were interpolated from the observed and HDDM-adjusted monitoring data 9 using the Voronoi Neighbor Averaging (VNA) technique. This technique was shown to be an 10 improvement upon the nearest neighbor technique used in the first draft REA and previous O₃ 11 NAAQS reviews (see Appendix 4-A for details). Consequently, the spatial variability of 12 observed and HDDM-adjusted O_3 is better accounted for in these analyses compared to those in 13 the first draft REA. 14 The air quality inputs to the epidemiology based risk assessment were "composite 15 monitor" values, a time-series of the spatially averaged monitoring data, in 12 of the 15 case study areas. Consequently, in cases of urban case study areas within which O₃ was predicted to 16

17 increase in some locations and decrease in others, the air quality inputs to this analysis represent

- 18 a "net" effect for each case study area. The spatial extent of the case study areas used in the
- 19 composite monitor averages were CBSAs. These CBSA areas are larger than the Zanobetti &
- 20 Schwartz, 2008 (Z & S) study areas (used in the first draft REA) which include only a subset of
- 21 the CBSA focused on urban cores. Figure 9-1 (reproduced from Figure 4-7) shows box plots of
- 22 the composite monitor values for 2006-2008 based on the observed data (black), data adjusted to
- 23 meet the existing standard using quadratic rollback (blue), and the HDDM adjustment procedure

- 1 (red) for the Z & S areas. The two adjustment methods were generally comparable in terms of
- 2 the changes in the upper quartile of the distribution. However, by design, quadratic rollback
- 3 always estimated decreases in the 75th percentile, median, and 25th percentile of the composite
- 4 monitor values, while HDDM estimated decreases in these values in some urban case study
- 5 areas, and increases in other areas consistent with atmospheric chemistry. HDDM-based
- 6 adjustments always produced increases in the lower tail of the distribution, while the lower tail
- 7 values generally remained unchanged with quadratic rollback. The differences between the two
- 8 adjustment procedures were the most pronounced in Los Angeles and New York, where the
- 9 largest reductions in NOx were required in order to meet the existing and alternative standards.
- 10 These large reductions in NOx caused a relatively large increase in lower O₃ concentrations
- 11 because of the reduction in NOx titration of O₃. As was noted in Chapter 4, the HDDM-based O₃
- 12 estimates become more uncertain for larger changes in NOx and VOC emissions, and thus there
- 13 was less overall confidence in those results. Even in these cases, the HDDM approach is still
- 14 preferable because it captures better the overall shift in the distribution of O₃ concentrations.
- 15

Boston: Z & S, April-October, 2006-2008





Baltimore: Z & S, April-October, 2006-2008





Atlanta: Z & S, April-October, 2006-2008

Detroit: Z & S, April-October, 2006-2008





Denver: Z & S, April-October, 2006-2008

Cleveland: Z & S, April-October, 2006-2008



LosAngeles: Z & S, April-October, 2006-2008

Houston: Z & S, April-October, 2006-2008



Philadelphia: Z & S, April-October, 2006-2008

10 8 (ppb) 60 03 (r 2 0 base 75



NewYork: Z & S, April-October, 2006-2008



Sacramento: Z & S, April-October, 2006-2008

SaintLouis: Z & S, April-October, 2006-2008



1 2

3

Figure 9-1 Distributions of composite monitor 8-hour daily maximum O₃ concentrations from ambient measurements (black), quadratic rollback (blue), and the HDDM adjustment methodology (red) for meeting the existing standard.

1 9.2.2 Human Exposure Modeling (Chapter 5)

The population exposure assessment evaluates exposures to O₃ using the Air Pollution Exposure (APEX) model for the general population, all school-aged children (ages 5-18), asthmatic school-aged children (ages 5-18), asthmatic adults (ages > 18), and older persons (ages 65 and older), with a focus on populations engaged in moderate or greater exertion (e.g. children engaged in outdoor recreational activities). The strong emphasis on children, asthmatics, and older adults reflected the findings of the last O₃ NAAQS review (U.S. EPA, 2007) and the ISA (U.S. EPA, 2013, Chapter 8) that these are important at-risk groups.

9 We assessed exposure in 15 urban case study areas – Atlanta, Baltimore, Boston, 10 Chicago, Cleveland, Dallas, Denver, Detroit, Houston, Los Angeles, New York, Philadelphia, 11 Sacramento, St. Louis, and Washington, D.C. – for recent O₃ concentrations (2006-2010) and for 12 O₃ concentrations adjusted to just meet existing and alternative standards for two time periods $(2006-2008 \text{ and } 2008-2010)^3$. The analysis provided estimates of the percent of several 13 14 populations of interest exposed to concentrations above three health-relevant 8-hour average O_3 15 exposure benchmarks: 60, 70, and 80 ppb. The ISA includes studies showing statistically 16 significant effects at each of these benchmark levels (U.S. EPA, 2013). These benchmarks were 17 selected to provide some perspective on the public health impacts from exposures to various 18 concentrations that have been associated with O₃-related health effects (e.g., lung inflammation 19 and increased airway responsiveness) in controlled human exposure and toxicological studies, 20 but cannot currently be evaluated in quantitative risk assessments. In addition, the exposure 21 assessment also identified the specific microenvironments and activities that contribute most to 22 exposure and evaluated at what times and how long individuals were in key microenvironments 23 and were engaged in key activities. This assessment focused on persons experiencing the highest 24 daily maximum 8-hour exposure within each study area. The assessment found that: 25 Childhood is an important lifestage where higher exposures and risks can occur, 26 due to the higher time spent outdoors by children, the higher exposure 27 concentration experienced by children while outdoors (i.e. when they are 28 dismissed from school in the afternoon and during the summer, when they may be 29 at an outdoor camp all day) and engagement in moderate or high exertion level 30 activities.

³ Attainment with the O_3 standard is based on the 4th highest maximum 8-hour O_3 concentration, averaged over 3 years. We evaluated two different 3-year periods in determining how air quality in each of the analytical years would respond to just meeting the existing and alterative levels of the standard. This was done to evaluate the effect of variability in meteorology and emissions on exposures and risks associated with just meeting the existing and alternative standards. For the exposure and lung function risk analyses, which provide estimates for each of the five analytical years, this results in two estimates for 2008, because 2008 is included in each of the 3-year averaging periods and there are separate analytical results for 2008 for the adjusted air quality resulting from simulating attainment in each of the two 3-year periods.

- Persons spending a large portion of their time outdoors during afternoon hours
 experienced the highest 8-hour O₃ exposure concentrations given that O₃
 concentrations in other microenvironments were simulated to be lower than
 ambient concentrations.
- Highly exposed children spend half of their outdoor time (on average) engaged in
 moderate or greater exertion levels, such as in sporting activities. Highly exposed
 adults also spent their outdoor time engaged in moderate or greater exertion levels
 though on average, not as frequently as children.

Across the 15 urban case study areas, we find that children are of greatest concern for O₃ exposures compared to other lifestages due to the greater amount of time they spend outdoors engaged in moderate or higher exertion activities. The exposure analysis estimates that children have the highest percent of exposures of concern of any of the at-risk populations or lifestages. As a result, we focus on the results for children (ages 5-18) in the remainder of this discussion. Figure 9-2 (reproduced from Figure 5-11) shows the results of the exposure assessment for all 15 urban case study areas, showing trends across the analytical years for the percent of children with

16 at least one 8-hour exposure greater than the 60, 70, and 80 ppb benchmarks.

17 The limited availability of longitudinal activity diary data and the general population 18 modeling approach used may underestimate the correlation in activity patterns for certain 19 susceptible populations (e.g., outdoor workers), and underestimate how often there are repeated 20 exposures to O_3 concentrations above the exposure benchmarks. As a result, although we are 21 able to report the percent of the population with at least one exposure greater than the alternative 22 exposure benchmarks, we are less confident in the estimated percent of the population 23 experiencing more than one exposure. Individuals with repeated exposures may be at greater risk 24 of significant health effects (U.S. EPA, 2013, Section 6.2.1.1). In addition, the limited data on 25 responses to air quality alerts (e.g., averting behavior) indicates that a small percentage of the 26 population may engage in averting behavior in response to air pollution, which may overstate 27 actual exposures if individuals reduce their exposure during periods of high O_3 .

28 The benchmark exposures of concern are not equivalent to ambient standard levels, as 29 exposures reflect the full pattern of O_3 concentrations throughout a season, coupled with time 30 spent outdoors and indoors engaged in different activities. Thus, just meeting the existing 31 standard will result in shifts in the entire distribution of O_3 over a three year period, and will 32 change the percent of populations experiencing each of the exposure benchmarks of concern. 33 Figure 9-2 shows that the percent of children above the 60 ppb benchmark declines consistently 34 across the 15 urban case study areas when just meeting potential alternative standards of 70, 65, 35 and 60. For most urban case study areas and years, the percent of children above the 60 ppb

36 benchmark is reduced by over half when O_3 is adjusted to meet the 65 ppb alternative standard

relative to the 75 ppb standard. In many urban case study areas and years, just meeting the 65
 ppb alternative standard results in close to zero percent of children above the 60 ppb benchmark.
 For the 70 and 80 ppb benchmarks, meeting an alternative standard of 70 ppb results in a small
 percentage of children exceeding the benchmarks.

5 Year-to-year variability is relatively pronounced for exceedances of the 60 ppb 6 benchmark. In addition, we observe a geographic pattern to the years with the maximum percent 7 of exceedances of the exposure benchmarks reflecting the regional O₃ patterns across years. In 8 general, northeastern urban case study areas saw the highest percentage of exceedances during 9 2007, while southern and western urban case study areas saw a higher percentage of exceedances 10 during 2006. However, these patterns are somewhat dependent on the 3-year averaging period 11 used to determine whether the standards are met. In general variability in the percent of children 12 exceeding the 60 ppb exposure benchmark across urban case study areas is similar to the 13 variability across years. 14 The percent of children with multiple exposures above the exposure benchmarks is 15 generally much lower compared to the percent of children with single exposures above the

benchmarks. However, as noted above, we have lower confidence in these estimates. Even for
the lowest benchmark level of 60 ppb, most locations and years have less than 10 percent of
children experiencing 2 or more exposures when just meeting the existing standard of 75 ppb,

less than 5 percent when just meeting an alternative standard of 70 ppb, and less than 1 percentwhen just meeting an alternative standard of 65 ppb. For most urban case study areas and years,

21 less than 1 percent of children experience 2 or more exposures above the 70 ppb exposure

22 benchmark when just meeting the existing standard.



Figure 9-2 Effects of just meeting existing (column 1) and alternative (columns 2 through 4) standards on percent of children (ages 5-18) with at least one O₃ exposure at or above 60, 70, and 80 ppb-8hr while at moderate or greater exertion, years 2006-2010.⁴

⁴ We were not able to adjust air quality to just meet the 60 ppb alternative standard in the New York City by reducing U.S. NOx and VOC emissions (see chapter3 and appendix 4-D for details). Detroit was already meeting the existing standard for 2008-2010.

1 Table 5-6 summarized the percent of the population of children (ages 5-18) with at least 2 one daily 8-hour exposure above the 60, 70, and 80 ppb benchmarks, providing both the mean 3 and maximum percentage across the five analytical years for each urban case study area. For O₃ 4 adjusted to just meet the existing standard of 75 ppb, the highest maximum percentage of 5 children exceeding the 60 ppb benchmark across years, 26 percent, occurs in Denver, which also 6 has the highest mean percentage across years. After just meeting the existing standard, Los 7 Angeles has the lowest maximum (10 percent) and mean (9.5 percent) percentage of children 8 exceeding the 60 ppb benchmark across years, likely reflecting the highly skewed nature of O_3 9 concentrations in that urban case study area. For example, just meeting the existing standards in 10 Los Angeles moves the majority of O₃ concentrations (sites and days) well below 60 ppb (See 11 Appendix 4-D). Patterns across urban case study areas are generally similar after just meeting 12 alternative standards of 70, 65, and 60 ppb, with the exception that the lowest maximum and 13 mean percentage of children for the alternative standard level of 65 ppb occurs in the New York 14 City urban case study area, which had very large (greater than 90 percent) reductions in NOx 15 emissions that were used to adjust air quality to just meet the 65 ppb standard level in that urban case study area. This resulted in the distribution of O₃ concentrations covering most days of the 16 17 year and most monitoring sites shifting dramatically downward, with most concentrations well 18 below 60 ppb across the New York City urban case study area. The level of confidence in the 19 results for the New York City and Los Angeles study areas for just meeting the alternative 20 standards is lower than that for some of the other urban case study areas due to the HDDM-based 21 O₃ estimates becoming more uncertain for very large changes in precursor emissions. 22 Figure 9-3 (reproduced from Figure 5-19) shows the results of the exposure assessment

23 for all 15 urban case study areas, showing the effect on the percent of children with one or more 24 exposures above the 60 ppb benchmark of just meeting the existing and alternative standards. 25 For each alternative standard, Figure 9-2 shows the maximum percent of children exceeding the 26 benchmark across the modeled years 2006-2010. Patterns of results are similar for the 70 ppb 27 and 80 ppb benchmarks, however, the maximum percents of children exceeding those higher 28 benchmarks are much smaller for all alternative standards. The percent of children exceeding the 29 80 ppb benchmark is close to zero once the existing standard is met. The percent of children with 30 two or more exposures exceeding the 60 ppb benchmark level is substantially lower when just 31 meeting the existing standard, and is close to zero for the 70 ppb and 80 ppb benchmarks. This 32 percentage drops substantially when meeting the 70 ppb standard, and is close to zero in most 33 urban case study areas when meeting the 65 ppb and 60 ppb alternative standards. Patterns for 34 asthmatic children are very similar to patterns for all children.



2

3

Figure 9-3 Effects of just meeting existing (75 ppb) and alternative standards on percent of children (ages 5-18) exceeding 60 ppb exposure benchmark, highest value across years for each urban case study area, 2006-2010.⁵

4 5

6 9.2.3 Health Risks Based on Controlled Human Exposure Studies (Chapter 6)

Using the estimates of exposure from APEX combined with results from controlled
human exposure studies, we estimated the number and percent of at-risk populations or lifestages
(all children aged 5-18, children with asthma aged 5-18, adults aged 18-35, adults aged 36-55,
and outdoor workers) experiencing selected decrements in lung function. The analysis focuses on
estimates of the percent of each at-risk population or lifestage experiencing a reduction in lung
function (mostly for durations of one to five hours) for three different levels of impact, 10, 15,

⁵ We were not able to adjust air quality to just meet the 60 ppb alternative standard in New York City urban case study area by reducing U.S. NOx and VOC emissions (see chapter3 and appendix 4-D for details). Detroit was already meeting the existing standard for 2008-2010.

1 and 20 percent decrements in FEV1. These levels of impact were selected based on the literature

2 discussing the adversity associated with increasing lung function decrements (US EPA, 2013,

3 Section 6.2.1.1). Consistent with the exposure assessment, we focus this summary on lung

4 function decrements in children as they are the lifestage likely to have the greatest percentage at-

5 risk due to higher levels of exposure and exertion. Within the overall population of children,

6 asthmatic children may have less reserve lung capacity to draw upon when faced with

7 decrements, and therefore a $\geq 10\%$ decrement in lung function may be a more adverse event in an

8 asthmatic child than a healthy child.

9 Lung function risks (based on experiencing an estimated 10, 15, or 20 percent decrement

10 in lung function) were estimated for each of the 15 urban case study areas in which human

11 exposures were modeled. Two models were used to estimate lung function risks: one based on

12 application of a population level exposure-response (E-R) function consistent with the approach

13 used in the previous O₃ NAAQS review, and one based on application of an individual level E-R

14 function (the McDonnell-Stewart-Smith (MSS) model), introduced in this review, which

15 incorporates individual differences in physiology, age, and activity patterns (McDonnell et al.,

16 2012). Because the individual level E-R function approach allows for a more complete estimate

17 of risk (incorporating risk responses at varying activity levels, not just moderate or greater

18 exertion), we focus on the results of that approach for this discussion.

19 The MSS model as implemented in APEX has a term that adjusts the lung function 20 response according to an individual's age. The MSS model was fit using data from subjects who 21 ranged in age from 18 to 35. Thus, the MSS model is not able to account for differences in lung 22 function at different age groups between the ages of 5 and 18. However, age does have a 23 pronounced effect on lung function response in the APEX model. APEX models differences in 24 physiological parameters due to age, and these result in age-dependent predictions of ventilation 25 rates, which are used in the MSS model. Ventilation rates also depend on the activities being 26 performed, which are also age-dependent. As a result of differences in physiology and activities, 27 the lung function responses vary by age (see Appendix 6-E).

Figure 9-4 (reproduced from Figure 6-6) shows the results of the lung function risk assessment for all 15 urban case study areas, showing trends across the analytical years for the percent of children with predicted lung function decrements greater than or equal to 10 percent⁶.

 $^{^{6}}$ We have introduced a new method (relative to the O₃ NAAQS review completed in 2008) for calculating the percent of the at-risk populations (all children and asthmatic children) experiencing lung function decrements, based on modeling of individual level responses to O₃ exposures. This model yields significantly higher estimates of the percent of children experiencing lung function decrements greater than 10, 15, and 20 percent. This may be partly due to the specific data inputs from clinical studies used to derive the function, but is also to be expected because the MSS model can reflect greater sensitivity of children to O₃ exposures because it allows for age variability in the relationship between O₃ and FEV1 decrements, and younger populations are more responsive to O₃ exposures than older populations.

- 1 Specifically, Figure 9-4 shows that the percent of children (age 5-18) with greater than or equal
- 2 to 10 percent lung function decrement declines consistently across the 15 urban case study areas
- 3 when just meeting the existing 75 ppb standard, as well as the alternative standards of 70, 65, and
- 4 60. The percent of children at-risk at the 10 percent decrement level remains at or above 10
- 5 percent in many locations after just meeting the 60 ppb alternative standard. The percentage of
- 6 children with greater than or equal to a 15 or 20 percent lung function decrement is much lower
- 7 for all alternative standards, with close to zero percent of children at-risk when just meeting the
- 8 alternative standard of 60 ppb. In general variability in percent of children at-risk across urban
- 9 case study areas is similar to variability across years.



- 1
- 2 Figure 9-4 Effects of just meeting existing (column 1) and alternative (columns 2-4)
- 3 standards on percent of children (ages 5-18) with FEV₁ decrement > 10, 15, and 20%, years
- 4 **2006-2010.**⁷

⁷ We were not able to adjust air quality to just meet the 60 ppb alternative standard in New York City urban case study area by reducing U.S. NOx and VOC emissions (see chapter3 and appendix 4-D for details). Detroit was already meeting the existing standard for 2008-2010.

Figure 9-5 (reproduced from Figure 6-11) shows the results of the lung function risk
assessment for all 15 urban case study areas, showing the effect on the risk of a 10 percent or
greater lung function decrement in children (ages 5-18) of just meeting the existing and
alternative O₃ standards. For each alternative standard, Figure 9-5 shows the maximum percent
risk over all of the modeled years 2006-2010.

6 There is no consistent pattern in the percent of children with 10 percent or greater lung 7 function decrement across urban case study areas just meeting the existing standard of 75 ppb. 8 The 5-year maximum estimated percent of children at-risk ranges from 17 to 22 percent across 9 urban case study areas. The percent reduction in 5-year maximum risk when just meeting the 70 10 ppb alternative standard is more consistent across urban case study areas, ranging from 8 to 23 11 percent (excluding New York City, which had a reduction of 29 percent). Reductions in risk 12 when just meeting the 65 ppb alternative standard are also generally consistent across urban case 13 study areas, with the exception of New York City. Incremental reductions in risk when just 14 meeting the alternative 65 ppb standard compared with just meeting the 70 ppb alternative 15 standard range from 17 to 31 percent excluding New York City, which has a reduction in risk of 16 more than twice as much as the next largest reduction. Incremental reductions in risk from just 17 meeting the alternative 60 ppb standard compared with just meeting the 65 ppb standard are 18 generally consistent, ranging from 16 to 46 percent, with somewhat larger reductions in risk 19 occurring in Cleveland and Denver. Overall, the 5-year maximum percent of children at-risk for 20 lung function decrements of 10 percent or more exceeds 13 percent, 10 percent, and 5 percent in 21 all urban case study areas except New York after just meeting alternative standards of 70, 65, 22 and 60, respectively. Patterns of risk reductions are also similar for the alternative lung function 23 decrement levels of 15 percent and 20 percent. However, the initial percent of the population 24 experiencing these decrements when just meeting the existing standard are substantially lower. 25 Patterns of risk responses using the population level exposure-response model are similar 26 to the MSS individual risk model. However, the starting values for the percent of the population 27 at risk are lower, reflecting the limits of the model in reflecting individual level responses, and

- the limited coverage of the model for exposures at lower exertion levels. For children, the MSS
 model gives results typically a factor of three higher than the population level E-R model used in
- 30 the previous O_3 NAAQS review.
- 31



Figure 9-5 Impact of just meeting existing (75 ppb) and alternative standards on percent of children (ages 5-18) with FEV₁ decrement > 10%, highest value for each urban case study area, 2006-2010.⁸

4

5 9.2.4 Health Risks Based on Epidemiological Studies (Chapters 7 and 8)

6 The epidemiology-based risk assessment evaluated mortality and morbidity risks from 7 short-term O₃ exposures and mortality risks from long-term exposures to O₃ by applying 8 concentration-response (C-R) functions derived from selected epidemiology studies. The 9 analysis included both a set of urban case study area case studies and a national-scale 10 assessment. The urban case study analyses evaluated mortality and emergency department (ED) 11 visits, hospitalizations, and respiratory symptoms associated with recent O₃ concentrations 12 (2006-2010) and with O₃ concentrations adjusted to just meet the existing and alternative O₃

⁸ We were not able to adjust air quality to just meet the 60 ppb alternative standard in New York City by reducing U.S. NOx and VOC emissions (see chapter3 and appendix 4-D for details). Detroit was already meeting the existing standard for 2008-2010.

1 standards (see section 9.2.1 and Chapter 4). Mortality and hospital admissions (HA) were

- 2 evaluated in 12 urban case study areas, while ED visits and respiratory symptoms were evaluated
- 3 in a subset of areas with supporting epidemiology studies. The 12 urban case study areas were:
- 4 Atlanta, GA; Baltimore, MD; Boston, MA; Cleveland, OH; Denver, CO; Detroit, MI; Houston,
- 5 TX; Los Angeles, CA; New York, NY; Philadelphia, PA; Sacramento, CA; and St. Louis, MO.

6 The urban case study analyses focus on risk estimates for the middle year of each three-year

7 design value period (2006-2008 and 2008-2010) in order to provide estimates of risk for a year

8 with generally higher O_3 concentrations (2007) and a year with generally lower O_3

9 concentrations (2009).

10 Most of the endpoints evaluated in epidemiology studies cover the entire study 11 population including children and adults. Because most mortality and hospitalizations occur in 12 older persons, these epidemiology-based risk estimates are better indicators of effects in adults 13 than in children. This is an important distinction from the human exposure and lung function risk 14 assessments, which focus on children. The only endpoints specific to children are asthma and all 15 respiratory hospital admissions using the New York specific epidemiology study, respiratory ER 16 visits in Atlanta, and respiratory symptoms in asthmatic children in Boston.

17 Both the urban case study area and national-scale assessments provide the absolute 18 incidence and percent of incidence attributable to O₃. In addition, risks are presented in terms of 19 incidence per 100,000 population to control for the differences in the sizes of the populations 20 across urban case study areas, and to allow for comparison of risks using different definitions of 21 urban extent. In previous reviews, O_3 risks have only been estimated for the portion of total O_3 22 attributable to North American anthropogenic sources (above what was referred to in previous 23 reviews as "policy-relevant background O_3 "). In contrast, this assessment estimates risk for O_3 24 concentrations down to zero, reflecting the lack of evidence for a detectable threshold in the C-R 25 functions (U.S. EPA, 2013, Chapter 2), and the understanding that U.S. populations may 26 experience health risks associated with O₃ resulting from emissions from all sources, both natural 27 and anthropogenic, within and outside the U.S. In order to better reflect how O₃ distributions are 28 likely to respond to just meeting existing and potential alternative standard levels, we adjusted 29 O₃ concentrations to just meet existing and potential alternative standard levels using reductions 30 in only U.S. anthropogenic emissions of O₃ precursors. Thus, the estimated changes in risk 31 between just meeting the existing standards and just meeting potential alternative standard levels 32 only reflect reductions in U.S. anthropogenic emissions. 33

However, consistent with the conclusions in the O_3 ISA (U.S. EPA, 2013), we have relatively lower certainty about the shape of the C-R function towards the lower end of the distribution of O_3 concentrations used in fitting the function due to the reduction in the number of O_3 measurements in this portion of the distribution. We discuss this source of uncertainty below. In addition, we provide the distribution of mortality incidence across the range of O₃
 concentrations in Chapter 7 to inform discussions of uncertainty in the results.

3 9.2.4.1 Urban Case Study Results

- 4 Figures 9-6 and 9-7 (reproduced from Figures 7-4 and 7-5) show the results of the
- 5 mortality and adult (ages 65 and older) respiratory hospital admissions risk assessments for all 12
- 6 urban case study areas, showing the effect on the incidence per 100,000 population just meeting
- 7 the existing 75 ppb standard and alternative O_3 standards of 70, 65, and 60 ppb in 2007 and
- 8 2009.

2007 Simulation year



2009 Simulation year



- 1 2
- Figure 9-6 Impacts of just meeting existing (75 ppb) and alternative standard levels on
 mortality risk per 100,000 population for 2007 and 2009.⁹
- 5

⁹ As noted earlier, we were not able to adjust air quality to just meet the 60 ppb alternative standard in New York City by reducing U.S. NOx and VOC emissions (see chapter3 and appendix 4-D for details). Detroit was already meeting the existing standard for 2008-2010.



2007 Simulation year

2009 Simulation year



1 2

3

4

5

Figure 9-7 Impacts of just meeting existing and alternative standard levels on adult (ages 65 and older) respiratory hospital admissions risk per 100,000 population for 2007 and 2009.¹⁰

¹⁰ We were not able to adjust air quality to just meet the 60 ppb alternative standard in New York City urban case study area by reducing U.S. NOx and VOC emissions (see chapter3 and appendix 4-D for details). Detroit was already meeting the existing standard for 2008-2010.

- In some urban case study areas which have large NOx emissions (e.g. from heavy downtown traffic), O_3 levels are artificially low because the NOx emissions remove O_3 through a chemical reaction (see section 9.2.1 and Chapter 4). In these places, when NOx emissions are decreased to reduce peak O_3 concentrations across the entire CBSA, which often includes locations outside of the urban core areas, lower concentrations of O_3 can go up. This can also happen in other areas on the lowest O_3 days. This phenomenon occurs in some locations when meeting lower alternative standards as well.
- meeting lower alternative standards as well. 8 The overall trend across urban case study areas is small decreases in mortality and 9 morbidity risk as O₃ concentrations are adjusted to just meet incrementally lower alternative 10 standard levels. In New York, there are somewhat greater decreases in these risks, reflecting the 11 relatively large emission reductions used to adjust air quality to just meet the 65 ppb alternative 12 standard, and the substantial change in the distribution of O_3 concentrations that resulted. We 13 were not able to adjust O_3 concentrations to just meet the 60 ppb alternative standard in the New 14 York City urban case study area. Risks vary substantially across urban case study areas; 15 however, the general pattern of reductions across the alternative standards is similar between 16 urban case study areas. Because of the generally lower baseline O_3 concentrations in 2009, risks 17 are generally slightly lower in 2009 relative to 2007; however, the patterns of reductions in risk
- 18 are very similar between the two years.

19 Mortality and morbidity risks generally do not show large responses to meeting existing 20 or alternative levels of the standard for several reasons. First, these risks are based on C-R 21 functions that are approximately linear along the full range of concentrations, and therefore 22 reflect the impact of changes in O₃ along the complete range of 8-hour average O₃ concentrations. This includes days with low baseline¹¹ O₃ concentrations that are predicted to 23 24 have increases in O₃ concentrations, as well as days with higher starting O₃ concentrations that 25 are predicted to have decreases in O_3 concentrations as a result of just meeting existing and 26 alternative standards. Second, these risks reflect changes in the urban-area wide monitor average, 27 which will not be as responsive to air quality adjustments as the design value monitor, and which 28 includes monitors with both decreases and increases in 8-hour concentrations. Third, the days 29 and locations with predicted increases in O_3 concentrations (generally those with low to 30 midrange starting O₃ concentrations) resulting from just meeting the existing or alternative 31 standard levels generally are frequent enough to offset days and locations with predicted 32 decreases in O_3 . The heat maps presented in Figures 7-2 and 7-3 demonstrate that just meeting 33 progressively lower alternative standard levels narrows the distribution of risk across the range

¹¹ By low baseline concentrations, we mean area-wide average O₃ concentrations between approximately 10 and 40 ppb prior to adjustments to just meet the existing and alternative standards.

1 of O₃ concentrations. In addition, the distribution of risk tends to be more centered on area-wide 2 average concentrations in the range of 25 to 55 ppb after just meeting an alternative standard of 3 60 ppb. The focus of the epidemiological studies on urban case study area-wide average O_3 4 concentrations, and the lack of thresholds coupled with the linear nature of the C-R functions 5 mean that in this analysis, the impact of a peak-based standard (which seeks to reduce peak 6 concentrations regardless of effects on low or mean concentrations) on estimates of mortality and 7 morbidity risks based on results of those studies is relatively small. For example, for mortality 8 and hospital admissions, we find a less than 10 percent reduction in risk for most urban case 9 study areas when just meeting the 70 ppb and 65 ppb alternative standards compared to just 10 meeting the existing standard, and a less than 25 percent reduction in risk for all urban case study 11 areas when just meeting the 60 ppb standard compared to just meeting the existing standard. The 12 general pattern for other morbidity risks is similar to hospital admissions. However, we are not 13 able to draw strong conclusions about the results across urban case study areas, because of the 14 limited number of urban case study areas represented for most of the endpoints. 15 We have applied city-specific mortality effect estimates to each urban case study area

16 based on the largest multi-city epidemiological study. However, for many of the urban case study 17 areas, the risk estimates have wide confidence intervals that can include zero, due to the lower 18 statistical power of some of the city-specific effect estimates relative to the national combined 19 effect estimate across cities. Furthermore, there is significant variability in these effect estimates 20 across the 12 urban case study areas, with some urban case study areas having effect estimates from 5 to 7 times greater than other cites (see Chapter 7, section 7.4.1).¹² The variability in effect 21 22 estimates, along with differences in O_3 concentrations, is a driver for the overall variability in the 23 risk results across cities. Smith et al (2009) reports an overall significant national mortality effect 24 estimate with confidence intervals that do not include zero, reflecting the much greater statistical 25 power available when pooling information across urban case study areas.

We also evaluated mortality risks in the 12 urban case study areas associated with longterm O_3 exposures (based on the seasonal average (April to September) of the peak daily onehour maximum concentrations). Risks from long-term exposures after just meeting the existing standard are substantially greater than risks from short-term exposures, ranging from 16 to 20 percent of respiratory mortality across urban case study areas. However, the percent reductions in long-term mortality risks are similar to those for mortality from short-term exposures. For example, we find a less than 10 percent reduction in risk relative to just meeting the existing

¹² This substantial heterogeneity in effect estimates can reflect a number of factors including differences in population susceptibility and behavior related to O₃ exposure and risk (e.g., proximity to roadways, use of air conditioning, commuting patterns, time spent outdoors) and differences in the degree to which the O₃ monitoring network used in the epidemiological study reflects patterns of population exposure.

- 1 standard in most areas when just meeting the 70 ppb and 65 ppb alternative standards, and a less
- 2 than 20 percent reduction when just meeting the 60 ppb alternative standard level. Risk
- 3 reductions for the New York City urban case study area are much greater when just meeting the
- 4 65 ppb alternative standard compared to just meeting the existing standard, with a 24 percent
- 5 reduction in risk in 2007.

6 New York and Los Angeles have characteristics that make epidemiological risk estimates 7 particularly uncertain. In the case of New York, the expansion of the urban case study area 8 definition to the CBSA adds uncertainty due to the large and diverse nature of the CBSA. The 9 New York CBSA includes two urban case study areas which have separate effect estimates 10 available from the Smith et al. (2009) study. These separate effect estimates (for Newark, NJ and 11 Jersey City, NJ) are smaller than the effect estimate for New York, however, they are also based 12 on much smaller populations, and have relatively wider confidence bounds, reflecting low 13 statistical power. For consistency with other urban case study areas and to allow for comparison 14 between the CBSA-based risk estimates and the smaller study area based estimates (see the 15 sensitivity analyses in Chapter 7), we elected to apply the New York city effect estimate, which 16 is based on a very large population and has high precision, to all of the counties in the New York 17 CBSA. While this adds substantial uncertainty to the absolute incidence of mortality for the New 18 York CBSA, it does not affect the pattern of risk reductions when just meeting alternative 19 standards. In addition, as noted earlier, the O_3 adjustments to meet existing and alternative 20 standards in New York and Los Angeles also have additional uncertainties relative to the other 21 10 urban case study areas. 22 We conducted a number of sensitivity analyses based on a population normalized

23 mortality risk metric, e.g. mortality risk per 100,000 population. Maintaining the general linear, 24 no-threshold functional form, mortality risks per 100,000 population are generally robust to 25 alternative specifications of the C-R functions, although in several urban case study areas, using 26 effect estimates from Smith et al. (2009) which were derived using regional priors rather than national priors results in higher risk estimates¹³. Using the effect estimates from Zanobetti and 27 28 Schwartz (2008) has no consistent effect on risk results across the urban case study areas. Using 29 effect estimates based on a copollutant model with PM₁₀, mortality risks are higher in some 30 locations and lower in others. However, in all locations the confidence intervals are substantially

¹³ In Bayesian modeling, effect estimates are "updated" from an assumed prior value using observational data. In the Smith et al (2009) approach, the prior values are either a regional or national mean of the individual effect estimates obtained for each individual city. The Bayesian adjusted city specific effect estimates are then calculated by updating the selected prior value based on the relative precision of each city-specific estimate and the variation observed across all city-specific individual effect estimates. City-specific estimates are pulled towards the prior value if they have low precision and/or there is low overall variation across estimates. City-specific estimates are given less adjustment if they are precisely estimated and/or there is greater overall variation across estimates.

1 wider using the copollutant model with PM_{10} (due to fewer days with both pollutants measured), 2 which makes it difficult to determine whether the increases and decreases in estimates relative to 3 the core estimates are real or the result of statistical error.

- 4 We selected the CBSA as the spatial definition for the urban case study areas. We made 5 this selection to address a downward bias that we identified resulting from a mismatch between 6 the smaller urban core areas used in the epidemiology studies and the larger areas where O_3 7 concentrations are expected to change as a result of meeting the existing and alternative standard 8 levels (see Chapter 7). We included a sensitivity analysis evaluating the result of using a smaller 9 geographic area including only the counties used in the epidemiology study. As expected, using a 10 smaller geographic extent for the urban case study areas results in smaller, and in some cases 11 negative risk reductions when compared to using the CBSA definitions. This reflects the fact that the controlling¹⁴ monitor in many of the 12 urban case study areas is located outside of the small 12 set of counties included in the Smith et al. (2009) urban case study area definitions, and some of 13 14 the monitors that are within that more limited spatial extent are more prone to O_3 titration due to 15 local NOx emission sources. As a result, those monitors are more likely to see increases in O₃ 16 which will, if other monitors with higher concentrations in the broader regions are not included, 17 lead to estimated increases in risk due to the application of a linear, no threshold C-R function. 18 This bias can be substantial, especially in St. Louis and several urban case study areas in the 19 Northeast, including Boston, New York, and Philadelphia, where the highest concentration 20 monitors are outside the Smith et al. (2009) urban case study area definitions. 21 Sensitivity analyses were conducted for scenarios of just meeting existing and alternative standards using combinations of NOx and VOC emissions reductions (as compared to NOx 22 23 reductions alone). The addition of VOC emissions reductions had little impact with the exception
- of New York and Los Angeles, where risk was decreased relative to the NOx-only reductionscenario.

26 9.2.4.2 National-scale Assessment Results

The national-scale assessment evaluated only mortality associated with recent O_3 concentrations across the entire U.S for 2006-2008. The national-scale assessment is a complement to the urban scale analysis, providing both a broader geographic assessment of O_3 related health risks across the U.S., as well as an evaluation of how well the 12 urban study areas represented the full distribution of O_3 -related health risks in the U.S. The national-scale assessment demonstrates that there are O_3 risks across the U.S, not just in urban case study areas, even though the O_3 concentrations in many areas were lower than the existing standard level.

¹⁴ The controlling monitor is the monitor with the highest design value within a defined non-attainment area.
1 While we did not assess the changes in risk at a national level associated with just meeting 2 existing and alternative standards, just meeting existing and alternative standards would likely 3 reduce O_3 concentrations both in areas that are not meeting those standards and in locations 4 surrounding those areas, leading to risk reductions that are not included in the urban-scale 5 analysis.

6 7 8

9.3

COMPARISON OF RESULTS ACROSS EXPOSURE, LUNG FUNCTION RISK, AND EPIDEMIOLOGY-BASED MORTALITY AND MORBIDITY RISK ANALYSES

9 In considering the overall results across the human exposure, lung function risk, and 10 epidemiology-based risk assessments, we focus on the key policy-relevant metrics and levels for 11 each type of assessment. For the human exposure assessment, we selected exposures above the 12 60 ppb exposure benchmark for all children (ages 5-18). We select this exposure metric because 13 children represent a key at-risk lifestage, and the 60 ppb exposure benchmark is the lowest 14 exposure level associated with significant findings in controlled human exposure studies. For the 15 lung function risk assessment, we selected the results for lung function decrements greater than 16 or equal to 10 percent for all children (ages 5-18). We select this lung function risk metric 17 because children represent a key at-risk lifestage, and a 10 percent lung function decrement 18 represents a potentially more adverse event in asthmatic children. For the epidemiology-based 19 risk assessment we selected the core short-term exposure mortality results and the respiratory 20 hospital admission results, because these endpoints were estimated for all of the 12 urban case 21 study areas. Generally speaking, these metrics provide the most differentiation between the 22 alternative standards, helping to inform policy-relevant questions regarding adequacy of the 23 existing standard, and public health impacts of meeting alternative standards. The other metrics 24 analyzed in this REA (e.g. other exposure benchmarks and other lung function decrements) show 25 less response to just meeting the existing standard or potential alternative standard levels. 26 As discussed in Chapter 2, we designed the exposure and risk assessment to help inform 27 two fundamental questions related to the adequacy of the existing standard in protecting public 28 health and the degree of exposure and risk reductions associated with alternative standards

compared with the existing standard. The following discussion evaluates the three types of
analyses we conducted in terms of the consistency of the information provided to inform these
questions.

32 9.3.1 Evaluation of Exposures and Risks After Just Meeting the Existing Standard

To compare the results of the three assessments in urban case study areas, we plot the key metrics from each analysis across urban case study areas for the two common years of analysis (i.e., 2007 and 2009). For three urban case study areas (i.e., Chicago, Dallas, and Washington

9-27

1 D.C.) we have only the exposure and lung function risk assessments, as these urban case study 2 areas did not have sufficient information to estimate epidemiology-based risks. The 3 epidemiology-based metrics are the percent of baseline short-term exposure mortality, based on 4 the core estimates using the C-R functions from Smith et al. (2009), and respiratory hospital 5 admissions based on the core estimates using the C-R functions from Medina-Ramon (2006), 6 attributable to O_3 . Figure 9-8 presents the exposures and risks after just meeting the existing 7 standard of 75 ppb. Each row represents one of the key analytical results; each column gives the 8 results for 2007 and 2009 for each urban case study area. The scale of each analytical metric for 9 each analysis differs, and thus the comparisons across analyses should focus on overall patterns 10 rather than on direct comparisons of numeric estimates.

11 All of the metrics show substantial variability among urban case study areas, although 12 there appears to be less variability in lung function risk and hospital admission risk compared 13 with the exposure metric and mortality risk. The differences between estimates for 2007 and 14 2009 are much higher for some urban case study areas (e.g. Baltimore and Philadelphia) for the 15 exposure metric than any of the risk metrics. This may reflect the explicit threshold nature of the 16 exposure metric, which focused on exposures above a benchmark level of 60 ppb. Differences 17 between years in exposures above the 60 ppb benchmark after just meeting the existing standard 18 are dependent on the number of days during each year with decreases in higher O_3 19 concentrations, as well as the magnitude of the decreases in O_3 on those higher O_3 concentration 20 days. These in turn are sensitive to the shape of the O₃ distribution in the analytical year prior to 21 just meeting the existing standard (which determines the starting number of days above 60 ppb) 22 and the response to emissions reductions applied in meeting the existing standard for 2007 or 23 2009. There is some consistency between metrics in the urban case study areas with highest 24 values for the exposure and lung function risk metrics. However, there were still differences, 25 especially for Los Angeles, which had one of the higher values for lung function risk in 2009, 26 but had one of the lower percentages of children exposed above the 60 ppb benchmark. This 27 again points to the importance of the threshold nature of the exposure metric, combined with the 28 tendency for more substantial decreases in peak O₃ concentrations relative to mid-range and low 29 concentrations when just meeting the existing standard.

There is little consistency within urban case study areas between the epidemiology risk
 metrics and the exposure and lung function risk metrics, and there is also little consistency

9-28

- 1 between the mortality and hospital admission risks. Houston has the lowest metric values in 2007
- 2 (except for mortality risk), but in 2009 has some of the higher risk metrics (except for hospital
- 3 admission risk). New York has the highest mortality risk in 2007 and 2009 but has among the
- 4 lowest hospital admission risks in both years.



Figure 9-8 Comparison of Exposure (Row 1) Lung Function Risk (Row 2) and Epidemiology-Based Risk (Rows 3 and 4) Metrics after Just Meeting the Existing 75 ppb Standard.

1 9.3.2 Reductions in Exposure and Risk Metrics after Just Meeting Alternative Standards

2 To compare the results of the three assessments for urban case study areas after just 3 meeting alternative standards relative to the existing standard, we express each result as a percent 4 of the metric value when just meeting the existing standard. Figure 9-9 presents the percent 5 reduction in exposures and risks after just meeting alternative standards relative to just meeting 6 the existing standard of 75 ppb. In this plot, each row represents one of the key analytical results 7 and each column gives the results for 2007 and 2009 for each urban case study area. The scales 8 are the same between analyses, and as such, it is informative to examine both the overall patterns 9 of change between alternative standards, and also the absolute value of the percent reductions in 10 risk metrics between analyses. In interpreting this chart, higher values mean greater reductions in 11 risk or exposure relative to just meeting the existing standards. Because these are percent 12 reductions, the maximum value is one hundred percent, which if reached would indicate that 13 risks or benchmark exposures are completely eliminated when the alternative standard is met in 14 the urban case study area as was seen for the 60 ppb exposure benchmark.

15 Many of the differences in results across the metrics are driven by how each metric is 16 affected by the O_3 data input to the analysis. In general, the impact of the HDDM adjustments to 17 O_3 vary based on three main considerations: 1) the degree to which the exposure or risk metric is 18 sensitive to changes across the various ranges of O₃ concentrations (e.g. high, mid-range, low); 19 2) whether the exposure or risk metric uses individual census tract concentrations or area-wide 20 average concentrations; and 3) changes in the distribution of O_3 concentrations in the year of 21 analysis between recent O_3 concentrations and adjusted (meeting the existing or alternative 22 standards) O_3 scenarios. With respect to 1), the exposure benchmark metric, which focuses only 23 on exposures above 60 ppb, will not be sensitive at all to changes in O₃ concentrations in the 24 range below 60 ppb. The lung function risk metric, which depends on the dose rate and 25 individuals' characteristics, does not have a concentration threshold. However, because of the 26 logistic form of the response function, it is less sensitive to lower O_3 concentrations and has very 27 few FEV1 responses greater than 10 percent when exposure concentrations are below 20 ppb and 28 very few FEV1 responses greater than 15 percent when exposure concentrations are below 40 29 ppb. On the other hand, the mortality and hospital admission risk metrics are based on non-30 threshold, approximately linear C-R functions, and therefore will be sensitive to changes in O_3 31 along the full range of O_3 . As discussed in Chapter 4, because O_3 at lower concentrations may 32 increase following HDDM adjustment in some locations and on some days to just meet alternative standards¹⁵, this can lead to increases in risk on some days, which can lead to a net 33

¹⁵ The frequency and magnitude of increases in spatially averaged mean concentrations in an urban case study area occur during a season when adjusting air quality to just meet a standard vary considerably between the existing

- 1 increase or decrease in risk over the entire year, depending on whether the days with increased
- 2 risk exceed days with decreased risk (generally due to a preponderance of days with lower O₃
- 3 concentrations). With respect to 2), the exposure and lung-function risk metrics are based on
- 4 concentrations at individual census tracts since they depend on O_3 exposure modeled by moving
- 5 each individual through their environment. Because of this, the exposure and lung-function risk
- 6 metrics are most affected by the spatial and temporal variability of O_3 concentrations across the
- 7 urban case study area. The mortality and hospital admission risk metrics are calculated applying
- 8 C-R functions to area-wide, daily maximum 8-hr average O₃ concentrations. As a result, the
- 9 spatial variability in O_3 concentrations between the monitors will only influence the
- 10 epidemiology-based risk estimates in how they influence the area-wide average. With respect to
- 11 3), all three metrics are influenced by how the distribution of O_3 concentrations changes between
- 12 recent O₃ conditions and after adjustment to just meet existing and alternative O₃ standard levels.

and alternative standards. The highest frequency of occurrence of days with increasing O_3 happens when adjusting air quality to just meet the existing standards, and decreases as air quality is further adjusted to just meet lower alternative standard levels.



Figure 9-9 Comparison of Percent Reduction in Key Risk Metrics for Alternative Standard Levels Relative to Just Meeting the Existing 75 ppb Standard.

1 The exposure and lung function risk metrics are most affected by the reductions in the 2 individual monitors' peak O₃ concentrations, including the magnitude of these reductions and the 3 number of days that experience these reductions. In contrast, the mortality and hospital 4 admission risk metrics are affected by changes in the mean of the seasonal, area-wide average O_3 5 concentrations, where the mean is determined by the frequency and magnitude of increases versus decreases in area-wide, maximum daily 8-hr O_3 concentrations¹⁶. In addition to O_3 6 concentrations, there are other factors that affect the variability across urban case study areas for 7 8 these three metrics, such as activity data and exposure factors for the exposure and lung function 9 risk metrics and the study-specific C-R functions for the mortality and hospital admission risk 10 metrics.

11 One clear observation is that the percent reductions in risk from meeting alternative 12 standard levels relative to meeting the existing standard for the two epidemiology-based 13 endpoints are much smaller than for the exposure benchmark and lung function risk endpoints. 14 The maximum percent reduction in the mortality and hospital admissions risk relative to just 15 meeting the existing standard across years, locations, and alternative standards is less than 25 16 percent, and for many years/locations, the reductions in these risks when just meeting the lowest 17 alternative standard, 60 ppb, are less than 10 percent. The exposure benchmark results show the 18 most reductions when comparing just meeting the existing standard to just meeting alternative 19 standards. Just meeting the 65 ppb standard results in reductions in the percent of children 20 exceeding the 60 ppb exposure benchmark by over 50 percent in all urban case study areas, and 21 by over 75 percent in 12 of the 15 urban case study areas evaluated. For most locations and 22 years, just meeting the 60 ppb alternative standard reduced the percent of children exceeding the 23 60 ppb exposure benchmark by over 90 percent compared to just meeting the existing standard. 24 Reductions in lung function risk were also much higher than reductions in mortality and hospital 25 admissions risk. Just meeting the 65 ppb standard results in reductions in lung function risk by 26 over 25 percent in most locations and years, and just meeting the 60 ppb standard results in 27 reductions by over 40 percent in most locations and years.

There is general consistency in the city-to-city patterns of reductions in the exposure and lung function risk metrics, although the decreases in lung function risk are less than half as large as the reductions in the percent of children exceeding the 60 ppb exposure benchmark (with the clear exception of New York city, which we will discuss further below). The patterns of reductions in mortality and hospital admission risk are generally consistent with the patterns for

¹⁶ As noted previously, changes in the spatial extent of the urban case study areas over which monitors are averaged can change the magnitude and sign of the change in the spatial average O_3 concentration for an urban case study area. For example, we found that we bias the risk estimates low when using urban case study area definitions that include only urban core counties and not the counties with monitors experiencing the most reductions in O_3 .

1 exposure and lung function risk for 2007, with the exception of Houston and Philadelphia.

- 2 However, for 2009, the patterns for mortality and hospital admission risk are quite different, both
- 3 from the 2007 results, and from the exposure and lung function risk results. This is due to the
- 4 generally lower O_3 concentrations in 2009, which results in a greater number of days with
- 5 predicted increases in O_3 concentrations at low concentrations, fewer days with very high
- 6 concentrations where predicted reductions in O_3 occur, and a smaller predicted decrease in O_3
- 7 concentrations on those high days. This affects the mortality and hospital admissions risk more
- 8 than the exposure and lung function risk metrics because those metrics incorporate thresholds,

9 and therefore are not responsive to changes in O_3 concentrations below those thresholds.

- Additional considerations are important in interpreting the reduction in exposure and risk
 between the existing standard and alternative standards. The REA analyses focus on reducing
- 12 peak O₃ concentrations, in particular the 4th high O₃ concentration averaged over 3-years so as to
- 13 simulate meeting the existing standard or various alternative standards. In addition, the air
- 14 quality adjustments are based on applying reductions in U.S. anthropogenic emissions. In this
- 15 way, the adjusted air quality reflects day-to-day O₃ concentrations that could occur when
- 16 focusing on reducing high O_3 concentrations rather than on reducing mean O_3 concentrations. In
- 17 addition, because the analyses do not include reductions of O₃ precursor emissions from sources
- 18 other than U.S. anthropogenic emissions (e.g. international emissions, biogenics, etc), the O₃
- 19 concentrations in the adjusted air quality account for O₃ created from natural and international
- sources, even if 100 percent emissions reductions are applied to U.S. anthropogenic sources in
 adjusting air quality scenarios.
- 22 Finally, with respect to the epidemiology based analyses, we note that 2007, which had 23 generally higher O₃ concentrations than 2009, had more days where O₃ concentrations decreased 24 as a result of adjusting peak O_3 concentrations to just meet alternative standards. Thus just 25 meeting alternative standards resulted in net decreases in risk in all locations, with the exception 26 of Houston for just meeting the 70 ppb alternative standard. In contrast, 2009, which had 27 generally lower concentrations than 2007, had more days in the range where O₃ concentrations 28 were increased as a result of adjusting peak O₃ concentrations to just meet alternative standards, 29 and thus the patterns reflect some locations where mortality and hospital admissions risk 30 increases. However, for 2009, in all locations, when just meeting the lowest alternative standard 31 of 60 ppb, mortality and hospital admission risks are decreased relative to just meeting the 32 existing standard.
- 33

19.4OVERALL ASSESSMENT OF REPRESENTATIVENESS OF EXPOSURE AND2RISK RESULTS

9.4.1 Representativeness of Selected Urban Case Study Areas in Reflecting Areas Across the Nation with Elevated Risk

5 We selected urban case study areas for the exposure and risk analyses based on several 6 criteria (e.g. recent elevated O₃ concentrations and presence of at-risk populations and lifestages) 7 we identified as likely indicators of areas and populations likely to experience high O_3 exposures 8 and risks (see Section 7.3.1). We then conducted several analyses to determine the extent to 9 which our selected urban case study areas actually represent the highest mortality and morbidity risk areas. We compared the distributions of risk characteristics¹⁷ and mortality risk (based on 10 11 recent O₃ concentrations) for the 12 urban case study areas used in the epidemiology-based risk 12 assessment with the corresponding national distributions. We also evaluated the degree to which 13 our selected urban case study areas represent the patterns of O₃ concentration changes 14 experienced by the overall U.S. population.

15 Based on the comparisons of distributions of risk characteristics, the selected urban case 16 study areas represent urban case study areas that are among the most populated in the U.S., have 17 relatively high peak O₃ concentrations, and capture well the range of city-specific mortality risk 18 effect estimates. These three factors alone would suggest that the case study urban case study 19 areas should capture well the overall risk for other heavily populated urban case study areas in 20 the nation, with a potential for better characterization of the high end of the risk distribution. The 21 selected urban case study areas do not include those with the highest numbers of some at-risk 22 populations or lifestages, specifically older people with high baseline mortality rates. However, 23 most locations in the U.S. (except Florida) with high percentages of older people have low 24 overall populations, less than 50,000 people in a county, or low O_3 concentrations. This suggests 25 that while the risk per exposed person per ppb of O_3 may be higher in these locations, the overall 26 risk to the population is likely to be within the range of risks represented by the urban case study 27 locations. 28 Based on the comparisons of distributions of short-term O_3 exposure mortality risk (using

the percent of mortality metric) for recent O₃ concentrations, the 12 selected urban case study areas are representative of the full distribution of U.S. O₃-related mortality risk in urban case study areas. Two of the selected areas, New York and Philadelphia are representative of the highest end of the distribution of short-term O₃ mortality risk. Overall, O₃ mortality risk for

33 short-term O₃ exposures in the 12 urban study areas are representative of the full distribution of

¹⁷ In this context, risk characteristics are the elements of populations, air quality, and inputs to the C-R functions that are expected to be correlated with estimated mortality risks (see Chapter 8).

1 U.S. urban O_3 -related mortality, representing both high end and low end risk counties. For the 2 long-term O_3 exposure mortality risk metric (again using the percent of mortality), the 12 urban 3 study areas are representative of the central portion of the distribution of risks across all U.S. 4 counties; however, the selected 12 urban case study areas do not capture the very highest (greater 5 than 98th percentile) or lowest (less than 25th percentile) ends of the national distribution of long-6 term exposure-related O_3 -related risk.

7 8

9.4.2 Representativeness of Selected Urban Case Study Areas in Reflecting Responsiveness of Risk to Just Meeting Existing and Alternative O₃ Standards

9 While we selected urban case study areas to represent those populations likely to 10 experience elevated risks from O_3 exposure, we did not include among the selection criteria the 11 responsiveness of O_3 in the urban case study area to decreases in O_3 precursor emissions that 12 would be needed to just meet existing or alternative standards.

13 In our preliminary evaluations of risk modeling results, we observed a consistent 14 presence of days with low to midrange starting O_3 concentrations for which O_3 concentrations 15 (using the 8-hour maximum metric) increased after adjustments to just meet the existing and 16 alternative standards across the selected urban case study areas. As noted above, this led to 17 estimates of increased risk on those days, and in some cases, estimates of increased risk over the 18 course of the O_3 season, reflecting both the magnitude and frequency of the predicted increases 19 relative to the predicted decreases in O_3 concentrations. As explained above, this pattern was 20 more pronounced when using a more spatially limited definition of the urban case study areas, 21 but even when using the CBSA definitions, there were still days when the area-wide average O_3 22 increased, primarily due to predicted increases in O_3 in the core counties of the urban case study 23 areas.

24 In order to better understand how prevalent this type of air quality response was across

25 the U.S., we conducted several additional analyses of O_3 concentrations. These included

26 evaluations of trends at O₃ monitors during a period of time with significant O₃ precursor

emission reductions, and evaluations of temporal and spatial patterns of O₃ changes across the

28 U.S., based on air quality modeling results, to simulate how O_3 would change across the U.S. in

29 response to NOx (and VOC) emissions reductions (relative to recent 2007 levels) similar to those

30 used in the HDDM adjustments (see section 9.2.1 above). The latter analysis includes an

31 assessment of the association of different types of O_3 responses with population counts to help

32 characterize the degree to which populations in the U.S. experience O_3 conditions like those in

the selected 15 urban case study areas (see Chapter 8).

Overall, both types of analyses showed that decreases in O₃ precursor emissions lead to decreases in O₃ concentrations in areas with higher starting O₃ concentrations, which tend to be rural or suburban case study areas, and on days with higher O₃ concentrations. The analyses also

- 1 indicate that in urban core areas (those with high levels of fresh NOx emissions), decreases in
- 2 NOx emissions can lead to increases in O₃, primarily for days when initial O₃ concentrations are
- 3 suppressed due to NOx titration. The observed widespread decreases of median O_3 in suburban
- 4 and rural locations when NOx emissions are decreased suggest the efficacy of large NOx
- 5 emissions reductions on reducing O_3 over large regions of the country.
- 6 These results suggest that many of the urban case study areas may show O_3 responses 7 that are typical of other large urban case study areas in the U.S., but may not represent the 8 response of O_3 in other populated areas of the U.S., including suburban case study areas, smaller 9 urban case study areas, and rural areas. These smaller urban case study areas would be more 10 likely than our urban case study areas to experience area-wide average decreases in mean O_3 11 concentrations as O_3 standards are met. Even though large urban case study areas throughout the 12 U.S. have high population density, 73 percent of the U.S. population lives outside of these high population density areas¹⁸, and thus, a large proportion of the population is likely to experience 13 14 greater mortality and morbidity risk reductions in response to reductions in 8-hour O₃ 15 concentrations than are predicted by our modeling in the selected urban case study areas. The 16 analyses presented in Section 8.2.3.2 show that populations in the case study areas we selected 17 are approximately twice as likely to experience increasing mean O_3 concentrations as 18 populations in the U.S. as a whole. Because our selection strategy for risk modeling was focused 19 on identifying areas with high risk, we tended to select large urban population centers. As 20 discussed in the previous section, this strategy was largely successful in including those urban 21 case study areas in the upper end of the O_3 risk distribution. However, this also has led to an 22 overrepresentation of the populations living in locations where we estimate increasing mean 23 seasonal O_3 in response to adjusting air quality to just meet the existing and alternative standards 24 using NOx emissions reductions. The implication of this is that our estimates of mortality and 25 morbidity risk reductions for the selected urban case study areas are likely to understate the 26 average risk reduction that would be experienced across the population and should not be seen as 27 representative of potential risk reductions for most of the U.S. population.

28 9.5 OVERALL ASSESSMENT OF CONFIDENCE IN EXPOSURE AND RISK 29 RESULTS

30 As with any complex analysis using estimated parameters and inputs from numerous data 31 sources and models, there are many sources of uncertainty that may affect our exposure and risk

- 32 estimates. These sources of uncertainty are discussed in each of the chapters related to air
- 33 quality, exposure, lung function risk, and epidemiology based mortality and morbidity risk. The

¹⁸ High population density areas are defined here as locations with population densities greater than 1000 people/km²

overall effect of the combined set of uncertainties on confidence in the interpretation of the
 results of the analyses is difficult to quantify. However, we provide our judgment of our overall
 confidence here, with an understanding that alternative judgments may also be supported.

The degree to which each analysis was able to incorporate quantitative assessments of uncertainty differed, due to differences in available information on uncertain parameters and complexities in propagating uncertainties through the models. In general, we followed the World Health Organization tiered approach to uncertainty characterization (WHO, 2008), which includes both quantitative and qualitative assessments. Each chapter includes a table identifying and characterizing the potential impact of key uncertainties on risk estimates, including the degree to which we were able to quantitatively address those uncertainties.

In considering our overall confidence in the results, there are several key considerations
 discussed below related to sources of uncertainty which we were not able to fully quantify, but
 which may have a large impact on both overall confidence and confidence in individual analyses.

14 9.5.1 Uncertainties in Modeling O₃ Responses to Meeting Standards

15 There is inherent uncertainty in all deterministic air quality models, such as CMAQ, the 16 photochemical grid model used to develop the model-based O_3 adjustment methodology. 17 Evaluations of air quality models against observed pollutant concentrations build confidence that 18 the model performs with reasonable accuracy despite both structural and parametric 19 uncertainties. A comprehensive model performance evaluation provided in Appendix 4-B shows 20 generally acceptable model performance that is equivalent to or better than typical state-of-the 21 science regional modeling simulations described in Simon et al. (2012). Two additional sources 22 of uncertainties in the HDDM adjustment methodology are the applicability of HDDM 23 sensitivities over large emissions perturbations and the variability in data used to create 24 regressions which allowed the application of these sensitivities to ambient data. Both sources of 25 uncertainty are shown to be reasonably small in chapter 4 with the first having a mean error of 26 less than 1ppb for 50% NOx cuts and less than 4 ppb for 90% NOx cuts. The uncertainty 27 introduced from the application of regressions to determine sensitivities were quantified by 28 propagating uncertainties in the sensitivities through to uncertainties in the final predicted O₃ 29 concentrations which had standard errors less than 1.4 ppb for all adjustment scenarios. New 30 York and Los Angeles had the largest uncertainties in these two areas due to the fact that they 31 required the largest reductions in NOx emissions. Uncertainties stemming from the application 32 of 8-months of model data to 5-years of ambient data and the across-the-board emissions cut 33 assumptions are further discussed in chapter 4 but are not expected to substantially degrade 34 confidence in the air quality results.

9.5.2 Uncertainties in Modeling Exposure and Lung-function Risk

2 With regard to the exposure and lung-function risk estimates, the modeling explicitly 3 incorporates population variability in many of the modeling inputs. We did not attempt to 4 probabilistically incorporate the many sources of uncertainty in model parameters or input data 5 due to limitations in the ability to specify distributions characterizing our confidence in those 6 variables. To explore the impacts of some of the more important sources of uncertainty, we 7 conducted a limited set of sensitivity analyses. For the exposure assessment, the estimate of 8 repeated exposures above exposure benchmarks is based on the limited set of diaries of activity 9 data available in the Consolidated Human Activity Database (CHAD) database (see Chapter 5). 10 The method for constructing activity patterns over the course of an O_3 season may not fully 11 capture the behavior of children who have systematically high outdoor activity levels. As a 12 result, while we are able to report the percent of children with two or more exposures, modeling 13 of the distribution of multiple exposures is limited, and the ability to identify the percent of the 14 population with unusually high numbers of multiple exposures is not possible. 15 For the lung function risk assessment, sensitivity analyses indicate that the MSS model 16 parameter related to the impact of the ventilation rate was most influential in determining the 17 estimated number of children with FEV₁ decrements greater than 10 percent. Estimates of lung 18 function decrements are also influenced by how much variability in individual response is 19 assumed in the MSS model. Sensitivity analyses indicate that when a greater amount of 20 variability is allowed in the MSS model, the percent of children ages 5-18 with FEV_1 decrements 21 greater than 10 percent can increase substantially. In addition, we performed analyses to 22 understand the age-related factors in APEX that could influence the estimated FEV₁ decrements. 23 It was found that the four most influential factors influencing the relationship between the 24 predicted FEV_1 decrement and age are the decreasing level of exertion, the decreasing equivalent 25 ventilation rate (with increasing age), the higher time spent outdoors by children, and the higher 26 exposure concentration experienced by children while outdoors. These all lead to children having 27 higher FEV₁ decrements than adults, and are more influential than the MSS model age term.

28 9.5.3 Uncertainties in Modeling Epidemiological-based Risk

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A major issue in using the results of the epidemiology studies in estimating risk is the narrow geographic definition used for urban case study areas in the epidemiology studies. In

- 31 many of the urban case study areas, we observe two distinct patterns of O_3 response to the
- 32 reductions in precursor emissions we evaluated to just meet the existing and alternative standard
- 33 levels. The first pattern generally occurs in areas outside the urban core (e.g. suburban and rural
- 34 areas), and on days when O_3 concentrations are on the higher end of the distribution of O_3
- 35 concentrations, and is characterized by predicted decreases in 8-hour O₃ concentrations. These

1 tend to be the locations where the highest 8-hour design values occur. The second pattern 2 generally occurs in the urban core, and on days when O_3 concentrations are on the lower end of 3 the distribution of O_3 concentrations, and is characterized by predicted increases in 8-hour O_3 4 concentrations. The narrow definitions of urban case study areas used in the epidemiological 5 studies generally included the urban core areas, but did not include all of the suburban or rural 6 areas. The narrow geographic definitions led to a clear downward bias in the estimates of risk 7 changes that would be associated with just meeting the standards in the urban case study areas, 8 because the risk changes would reflect the locations with a tendency towards increases in 8-hour 9 O_3 , but would not include locations outside the urban core with decreases in O_3 . In many cases, 10 the narrowly defined geographic definitions used in the epidemiology studies did not even 11 include the location with the monitor that was violating the standard. We addressed this bias by 12 expanding the urban case study area to the CBSA. However, this adds additional uncertainty to 13 the risk estimates, and reduces our confidence that we have a good match between the basis of 14 the C-R function (just urban core locations) and the risk analysis context (including both urban 15 core counties and other counties in the CBSA). A clear implication of this decision is that the 16 absolute incidence estimates will be larger than if the analysis was limited to a smaller number of 17 counties. For this reason, we have placed more emphasis on risk metrics that have been 18 normalized for population size (e.g. risks per 100,000 population and percent risk), so as to 19 facilitate comparisons between cities of different population sizes and to reduce the influence of 20 population size on the risk metrics.

21 The epidemiology studies used as the source for C-R functions for short-term exposure 22 mortality and morbidity endpoints all use time-series approaches to estimate the effect of daily 23 variations in O₃ concentrations on daily mortality or morbidity incidence. The effect estimates 24 developed in these epidemiology studies were based on air quality and health information 25 observed over periods of time in the past (1987-2000). These effect estimates were based on day-26 to-day variations in area-wide O₃ concentrations estimated from observed concentrations at 27 monitors that reflect a specific set of emissions and atmospheric conditions. In our REA 28 analyses, we apply these effect estimates to adjusted air quality scenarios that are reflective of 29 substantial changes in O_3 concentrations across an area due to, in some cases, large decreases in 30 NOx and VOC emissions reductions. The resulting spatial and temporal patterns of O₃ may not 31 be the same as the spatial and temporal patterns of O_3 that existed at the time of the 32 epidemiology study. The potential for different spatial and temporal patterns in O_3 33 concentrations between the adjusted air quality scenarios and the air quality observed during the 34 epidemiology study period potentially adds uncertainty to the estimates of risk, as it is not clear 35 the degree to which the exposure surrogate used in the epidemiology study correlates with the 36 exposure surrogate used in the risk analysis. The degree of this potential uncertainty increases

9-41

1 with the amount of emissions reductions applied in the adjusted air quality scenario. This is

- 2 because as the amount of emissions reductions applied increases, the spatial and temporal
- 3 patterns of O₃ concentrations become increasingly different from those patterns observed for
- 4 recent O₃ concentrations (2006-2010) that are more similar, although not identical (due to
- 5 reductions in NOx between 2000 and 2006), to the patterns for the time period covered by the
- 6 epidemiology studies (1987-2000). We are not able to quantify the effect or magnitude of this
- 7 uncertainty, because we do not know the relationship between O₃ variability and the C-R
- 8 functions. However, to the extent that the uncertainty is shown to be important, it seems
- 9 reasonable to conclude that the larger the adjustment to the O_3 distributions, the more likely there
- 10 could be a mismatch in the exposure surrogates.

Overall, these sources of uncertainty cause us to have reduced confidence in estimates of short-term risk based on modeling the larger (CBSA-based) study areas using the multi-city time series-based effect estimates. This reduces the utility of the risk assessment in directly informing the decision regarding the level of the standard since we have lower confidence in estimates of absolute risk associated with a given standard level. However, the risk assessment can still be useful in providing estimates of the general magnitude and direction of changes in risk associated with an alternative standard level.

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9.6 OVERALL INTEGRATED CHARACTERIZATION OF RISK IN THE CONTEXT OF KEY POLICY RELEVANT QUESTIONS

20 Our analyses set out to inform two questions: 1) what are the magnitudes of exposures of 21 concern and risks for O_3 -related health effects that are estimated to occur with O_3 concentrations 22 that just meet the existing O₃ standard?; and 2) to what extent do alternative standards reduce 23 estimated exposures and risks of concern attributable to O₃, focusing on at-risk populations and 24 lifestages? In evaluating risk, we did not limit the assessment to just the absolute risk that is 25 attributable to U.S. or North American emissions, as this is not relevant to answer the two 26 questions. Instead, we estimated total risk from all O₃ concentrations and the distribution of risk 27 over the range of O₃ concentrations. Our estimates of changes in risk from meeting alternative O₃ 28 standard levels relative to meeting the existing standard reflect only the impact of reductions in 29 U.S. precursor emissions on O₃ distributions, recognizing that these emissions are most likely to 30 be affected by implementation of the standards.

- To inform these questions, we conducted air quality, exposure, and risk analyses for selected urban case study areas. We evaluated changes in the distribution of O₃ concentrations along the full range of O₃ concentrations down to zero. We have utilized a new method (compared to the O₃ NAAQS review completed in 2008) for estimating O₃ concentrations
- 35 consistent with attaining existing and alternative standards, based on modeling the response of

1 O_3 concentrations to reductions in U.S. anthropogenic NOx and VOC emissions, using the

2 HDDM capabilities in CMAQ. This modeling incorporates all known emissions, including

3 emissions from non-anthropogenic sources and anthropogenic emissions from sources in and

4 outside of the U.S. As a result, background O₃ concentrations are directly modeled and, therefore,

5 do not need to be separately specified. Application of this approach also addresses the

6 recommendation by the National Research Council of the National Academies (NRC, 2008) to

7 explore how emissions reductions might affect temporal and spatial variations in O₃

8 concentrations, and to include information on how NOx versus VOC control strategies might

9 affect exposure to O_3 and potential risks.

We estimated exposures and risks using several different metrics. Consistent with the available evidence, we estimated the percentages of different study populations and lifestages with exposures exceeding several health-based exposure benchmarks. We estimated lung function risks based on a model of individual risk of lung function decrements that incorporates a dose-equivalent threshold and individual exposures, activity levels, and physiology. We estimated mortality and morbidity risks based on non-threshold C-R functions derived from

epidemiology studies. These three different analyses result in differing sensitivities of results to changes in the O_3 concentration distribution. Because the three metrics are affected differently in

the analyses by changes in O_3 at low concentration levels, it is important to understand these

19 changes in O_3 at low concentrations in interpreting differences in the results across metrics.

20 We also evaluated the degree to which exposures of concern and lung function risk were 21 reduced in the portions of urban case study areas (urban core areas) that were more likely to 22 experience an increase in low concentrations of O₃, and in some cases an overall net increase in 23 epidemiology based mortality and morbidity risk (results for this assessment are presented in 24 Appendix 9A). We compared these estimates of changes in exposures and lung function risk to 25 estimates of changes in exposures and lung function risk in the areas outside of the urban core 26 areas to judge whether for exposures of concern and lung function risk we see the same pattern 27 of risk reduction between those areas.

28 Both exposures of concern and lung-function risk estimates in the core urban case study 29 areas showed similar patterns compared with the areas outside the urban cores when just meeting 30 the existing and potential alternative standards. Thus, we observe that in urban core areas which 31 in some cases showed overall increases in epidemiology based mortality and morbidity risk 32 when looking across these same air quality scenarios (see section 9.5.3), we generally see 33 reductions in exposures of concern and lung function risk. These findings illustrate that 34 populations within core urban case study areas are likely to experience risk reductions for health 35 endpoints reflected in the exposure and lung-function analyses.

1 The mortality and morbidity risk assessment is the analysis that is most sensitive to the 2 increases in O_3 in the lower part of the distribution of initial O_3 concentrations at some monitors 3 and on some days after meeting the existing and alternative standards in some urban case study 4 areas. As demonstrated in the heat maps (Figures 7-2 and 7-3), the increases in O_3 (and resulting 5 estimated increases in risk) occur largely on days with initial O₃ concentrations in the range of 10 6 to 40 ppb. In addition, mean O_3 concentrations for the urban case study areas change little 7 between air quality scenarios for meeting the existing and alternative standards, because mean 8 concentrations reflect both the increases in O_3 at lower concentrations and the decreases in O_3 9 occurring on days with high O₃ concentrations. This leads to small net changes in mortality and 10 morbidity risk estimates for many of the urban case study areas. For New York, we find there is 11 a larger decrease relative to other urban case study areas (nearly five times as large as the next 12 largest result for Los Angeles), in mortality and respiratory hospital admissions when just 13 meeting the 65 ppb alternative standard compared to just meeting the existing standard, 14 reflecting the large degree of air quality adjustment needed to meet the standard at all monitors in 15 New York. Both the net change in risk and the distribution of risk across the range of O_3 16 concentrations in the urban case study areas may be relevant in considering the degree of 17 additional protection provided by just meeting existing and alternative standards. 18 The dampened response of short-term mortality risk can be contrasted with lung function 19 risk estimates based on application of results from controlled human exposure studies. The lung 20 function risk estimates primarily reflect changes in the upper end of the O₃ distribution and 21 reflect counts of exceedances of lung function decrement benchmarks, rather than summing risks

across all days in the season. In addition, lung function risks are based on detailed microenvironmental exposure modeling which uses individual monitor values instead of composite monitor values, thereby resulting in less dampening of spatial variability in O₃ within a given urban study area.

26 The exposure benchmark analysis is the least sensitive to changes in O_3 in the lower part 27 of the distribution of initial O_3 concentrations, because the lowest of the exposure benchmarks is at 60 ppb, well above the portion of the distribution of initial O₃ concentrations that increased. 28 29 Since the modeled exposures will always be less than or equal to the monitor concentrations, a 30 benchmark of exposure at 60 ppb is above the range of O₃ concentrations where the HDDM 31 approach estimates increases in concentrations. Thus, this metric is most reflective of the 32 decreases in O₃ at high concentrations that are expected to result from just meeting the existing 33 and alternative standards.

The lung function risk analysis is less sensitive than the mortality and morbidity risk assessments to increases at very low concentrations of O_3 , because the risk function is logistic and shows little response at lower O_3 dose rates that tend to occur when ambient concentrations are lower (generally less than 20 ppb for the 10 percent FEV₁ decrement and generally less than
 40 ppb for the 15 percent FEV₁ decrement). However, because there are still some increases in
 O₃ concentrations that occur in the 50 to 60 ppb range where the estimated risk is more
 responsive, there may be some reduction in the magnitude of the risk decrease (this is evident
 when comparing the lung function risk metric with the exposure benchmark metric in figure 9-

6 8).

7 The exposure-based lung function risk assessment is based on controlled human exposure 8 studies which studied responses in healthy adults. Although the lung function model based on 9 this population shows less responsiveness at lower ambient concentrations, the applicability of 10 this model to the responses of more sensitive populations and lifestages, including children and 11 asthmatics, is uncertain. In addition, although the most complete information for generating an 12 exposure-response function is available for FEV1 as a measure of lung function, there are other, 13 potentially more public health relevant effects, such as lung inflammation, which have also been 14 shown to respond to O_3 . As such, the lung function risk analysis should be seen as providing 15 useful but not complete information on risks of health responses to O_3 .

16 Exposures above health benchmarks and risks remain after adjusting O_3 to just meet the 17 existing standard. The percentage of children with at least one 8-hour O_3 exposure exceeding 60 18 ppb is greater than 10 percent in at least one of the five analytical years for all of the 15 urban 19 case study areas. The percent of children with a predicted decrement in lung function greater 20 than or equal to 10 percent is greater than 16 percent in at least one of the five analytical years 21 for all of the 15 urban case study areas, and for a 15 percent decrement is less than 7 percent for 22 all years and areas. O₃-attributable mortality is slightly less than one percent up to four percent of 23 total mortality across the 12 urban case study areas, with little variation between 2007 and 2009. 24 O₃-attributable respiratory hospital admissions are between 2 and 3 percent across the 12 urban 25 case study areas, with little variation between 2007 and 2009. The percent attributable risk for 26 other morbidity endpoints is somewhat higher than for respiratory hospital admissions, but we 27 only estimated these endpoints for a more limited set of urban case study areas due to data 28 limitations.

29 The degree of reduction in exposures and risks when adjusting O_3 from just meeting the 30 existing standard to just meeting lower alternative standard levels varies considerably between 31 metrics. The greatest degree of reduction occurs in exposures above the 60 ppb exposure 32 benchmark, followed by reductions in lung function decrements greater than or equal to 10 33 percent, with the smallest changes in mortality and respiratory hospital admissions. Although the 34 magnitude of reduction differs between the different exposure and risk metrics, there are 35 generally the same patterns of reductions for the exposure benchmark and lung function risk 36 metrics, showing consistent reductions across all 15 urban case study areas. Risk reductions also

1 occur in most of the urban case study areas for mortality and respiratory hospital admissions. 2 However, these reductions are small, and reflect net changes in risk that include days with risk 3 increases as well as risk decreases. For most urban case study areas, the greatest incremental 4 reductions in exposures above the 60 ppb benchmark occurred when just meeting 70 ppb 5 compared to just meeting the existing standard. Just meeting lower standards of 65 ppb and 60 6 ppb had incrementally smaller reductions in the percent of children exposed above 60 ppb. 7 Incremental lung function risk reductions are more even between alternative standards, with 8 similar or greater incremental reductions for the 65 ppb and 60 ppb alternatives compared with 9 the incremental reductions for just meeting 70 ppb. Incremental reductions in mortality and 10 respiratory hospital admissions risk are small between alternative standards, but more urban case 11 study areas have somewhat larger risk reductions when comparing just meeting the 60 ppb 12 alternative to just meeting the 65 ppb standard, than when comparing 65 ppb to 70 ppb or 70 ppb 13 to 75 ppb. Long-term exposure mortality risk results show larger absolute estimates of mortality 14 risk and more consistent reductions across urban case study areas. However, percent changes in 15 long-term exposure mortality are similar to those for short-term exposure mortality.

16 In conclusion, we have estimated that exposures and risks remain after just meeting the 17 existing standards and that in many cases, just meeting alternative standard levels results in 18 reductions in those exposures and risks. Meeting alternative standards has larger impacts on 19 metrics that are not sensitive to changes in lower O_3 concentrations. When meeting the 70, 65, 20 and 60 ppb alternative standards, the percent of children experiencing exposures above the 60 21 ppb health benchmark falls to less than 20 percent, less than 10 percent, and less than 3 percent 22 in the worst O₃ year for all 15 case study urban case study areas, respectively. Lung function risk 23 also drops considerably as lower standards are met. When meeting the 70, 65, and 60 ppb 24 alternative standards, the percent of children with lung function decrements greater than or equal 25 to 10 percent falls to less than 21 percent, less than 18 percent, and less than 14 percent in the 26 worst O₃ year for all 15 case study urban case study areas, respectively. Mortality from short-27 and long-term O_3 exposures and respiratory hospitalization risk is not greatly affected by 28 meeting lower standards, reflecting the impact of increasing O_3 on low concentration days, and 29 the non-threshold nature of the C-R function.

1 9.7 REFERENCES

- Frey, C. and J. Samet. 2012. CASAC Review of the EPA's Policy Assessment for the Review of
 the O₃ National Ambient Air Quality Standards (First External Review Draft August
 2012). U.S. Environmental Protection Agency Science Advisory Board, (Document
 number EPA-CASAC-13-003).
- Medina-Ramon, M.; A. Zanobetti and J. Schwartz. 2006. "The Effect of O₃ and PM₁₀ on Hospital
 Admissions for Pneumonia and Chronic Obstructive Pulmonary Disease: A National
 Multicity Study."*American Journal of Epidemiology*, 163(6), 579-588.
- McDonnell, W.F.; P. W. Stewart; M. V. Smith; C. S. Kim C.S. and E. S. Schelegle. 2012.
 "Prediction of lung function response for populations exposed to a wide range of O₃ conditions." *Inhalation Toxicology*, 24:619-633.
- Simon, H.; K. R. Baker; F. Akhtar; S. L. Napelenok; N. Possiel; B. Wells and B. Timin. 2013.
 "A Direct Sensitivity Approach to Predict Hourly O₃ Resulting from Compliance with the National Ambient Air Quality Standard." *Environmental Science and Technology*, 47, 2304-2313.
- Smith, RL; B. Xu and P. Switzer. 2009. "Reassessing the Relationship Between O₃ and Short term Mortality in U.S. Urban Communities." *Inhalation Toxicology*, 21: 37-61.
- U.S. Environmental Protection Agency. 2007. O₃ *Health Risk Assessment for Selected Urban case study areas*. Research Triangle Park, NC: Office of Air Quality Planning and
 Standards. (EPA document number EPA 452/R-07-009). Available at:
 .
- U.S. EPA. 2013. Integrated Science Assessment for O₃ and Related Photochemical Oxidants:
 Final. Research Triangle Park, NC: U.S. Environmental Protection Agency. (EPA document number EPA/600/R-10/076F).
- World Health Organization. 2008. Part 1: *Guidance Document on Characterizing and Communicating Uncertainty in Exposure Assessment, Harmonization Project Document No. 6.* Published under joint sponsorship of the World Health Organization, the
 International Labour Organization and the United Nations Environment Programme.
 WHO Press, World Health Organization, 20 Avenue Appia, 1211 Geneva 27,
 Switzerland.
- Zanobetti, A. and J. Schwartz. 2008. "Mortality Displacement in the Association of O₃ with
 Mortality: An analysis of 48 Cities in the United States." *American Journal of Respiratory and Critical Care Medicine*, 177:184-189.
- 34

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