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SAI AIRSHED MODEL OPERATIONS MANUALS

Volume I -- User's Manual

ATMOSPHERIC SCIENCES RESEARCH LABORATORY  
OFFICE OF RESEARCH AND DEVELOPMENT  
U.S. ENVIRONMENTAL PROTECTION AGENCY  
RESEARCH TRIANGLE PARK, NORTH CAROLINA 27711

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Volume I -- User's Manual

by

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## NOTICE

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## ABSTRACT

This report presents a general view of the Systems Applications, Inc. (SAI) Airshed Model as well as detailed operating instructions for the user. Included are discussions of all the files needed to run the model, the data preparation programs that produce each file, the input formats and data preparation methods, samples of input and output, and information flow diagrams that illustrate the job stream control on any computer.

Chapter II presents a technical summary of the SAI Airshed Model; Chapter III contains an overview of the model as a system of computer programs, outlining the five main functions that must be performed by users and programs together. Chapter IV discusses some of the concepts and terms basic to an understanding of the system, followed in Chapter V by descriptions of the job control notations that will be used throughout this manual. Chapters VI through XI examine in detail the five main functions of the system, providing input and output definitions and examples, and job control diagrams for each program that must be exercised.

The Systems Manual, the companion to the User's Manual, describes the system from a programmer's point of view. Included are descriptions of all subroutines and how they fit together, run-time core allocation techniques, internal methods of segment handling using secondary storage, and detailed structure definitions of all files in the system. The Systems Manual also discusses procedures for implementing the programs on different computers, and the addition, removal, or replacement of computing algorithms and data preparation methods.

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## ABBREVIATIONS

act	activation
add	additive
b	boundary
bd	bound
beg	beginning
B.T.	Boundary Table
chem	chemistry
coeff	coefficient
col	column
conc	concentration
const	constant
crit	criterion
dark	darkness
diam	diameter
fact	factor
ht	height
i	initial
iter	iteration
LBL	Lawrence Berkeley Laboratory
loc	location
low	lower
L.T.	Local Table
max	maximum
M.E.B.	meteorology, emissions, and initial and boundary conditions
meth	method
min	minimum
molec	molecular
mult	multiplicative
num	numerical
param	parameter

pr	print
prof	profile
pt	point
srce	source
reac	reaction
ref	reference
rel	relative
resis	resistance
spec	species
S.S.T.	Segment Scratch Table
S.T.	Segment Table
stn	station
st-st	steady-state
subreg	subregion
surf	surface
temp	temperature
tol	tolerance
u-def	user-defined
up	upper
UTM	Universal Transverse Mercator
val	value
var	variable
vartn	variation
vel	velocity
vert	vertical
wt	weight

## I INTRODUCTION

This report presents a general view of the Systems Applications, Inc. (SAI) Airshed Model as well as detailed operating instructions for the user. Included are discussions of all the files needed to run the model, the data preparation programs that produce each file, the input formats and data preparation methods, samples of input and output, and information flow diagrams that illustrate the job stream control on any computer.

Chapter II presents a technical summary of the SAI Airshed Model; Chapter III contains an overview of the model as a system of computer programs, outlining the five main functions that must be performed by users and programs together. Chapter IV discusses some of the concepts and terms basic to an understanding of the system, followed in Chapter V by descriptions of the job control notations that will be used throughout this manual. Chapters VI through XI examine in detail the five main functions of the system, providing input and output definitions and examples, and job control diagrams for each program that must be exercised.

The Systems Manual, the companion to the User's Manual, describes the system from a programmer's point of view. Included are descriptions of all subroutines and how they fit together, run-time core allocation techniques, internal methods of segment handling using secondary storage, and detailed structure definitions of all files in the system. The Systems Manual also discusses procedures for implementing the programs on different computers, and the addition, removal, or replacement of computing algorithms and data preparation methods.

## II THE SAI AIRSHED MODEL

Systems Applications, Incorporated (SAI) designed its Airshed Model to calculate the concentrations of both reactive and inert air pollutants through simulation of the numerous physical and chemical processes that take place in the atmosphere. Much of the current interest in the model arises from the need to calculate ozone concentrations as a part of various environmental assessment studies (such as federally required State Implementation Plans and New Source Reviews). The results of recent evaluation and applications studies indicate that the SAI Airshed Model provides a sound, practical, and effective means for analyzing a variety of regional air quality problems.

Interactions among organic compounds (including pure and oxygenated hydrocarbons and other hydrocarbon derivatives) and nitrogen oxides ( $\text{NO}_x$ ) are usually responsible for the relatively high ozone concentrations observed in and downwind of urban areas. Among the main factors affecting photochemical air quality in urban areas are:

- > The spatial and temporal distribution of emissions of  $\text{NO}_x$  and organic species.
- > The composition of the emitted organic species.
- > The spatial and temporal variations of the wind field.
- > The stability of the atmosphere and the associated dynamics of the mixing layer.
- > The chemical reactions involving organic species and  $\text{NO}_x$ .
- > The diurnal variations of ultraviolet radiation.
- > The loss of ozone and its precursors by surface uptake processes.
- > The ambient background concentrations of ozone, organic species, and  $\text{NO}_x$  immediately upwind of the urban area and within an elevated inversion layer.

Thus, development of an accurate ozone prediction relationship requires investigation of the relevant physical and chemical atmospheric phenomena that influence the magnitude and distribution of both ozone and its precursors.

To limit ozone concentrations, one must control all or part of the ozone formation process. Practically, man has the most influence on anthropogenic emissions. If ozone concentrations were proportional to emissions of organic compounds or  $\text{NO}_x$ , it would be simple to calculate the reduction in anthropogenic emissions necessary to achieve a desired reduction in ozone concentrations. But the relationship between ozone air quality and emissions of  $\text{NO}_x$  or organic species is complicated and nonlinear. Furthermore, the relationship changes with location and even with time at a given location because of variations in emissions, meteorology, and other factors.

To make available an appropriate quantitative treatment of this relationship, the U.S. Environmental Protection Agency (EPA) has, through a multiyear research effort, supported the development of the SAI Airshed Model. Initial model development efforts and applications (to the Los Angeles area) are fully described in a 15-volume series of reports by Roth et al. (1971) and Reynolds et al. (1973a). These reports are summarized in the three papers by Reynolds et al. (1973b, 1974) and Roth et al. (1974). Evaluation of the model's predictions and its components at that time showed that improved treatments of some physical and chemical processes were necessary. As a result, another series of research efforts was carried out to improve the model. These efforts are described in a 7-volume series of reports by Jerskey and Seinfeld (1976), Jerskey et al. (1976), Killus et al. (1977), Lamb (1976), Lamb et al. (1977), Liu et al. (1976a), and Reynolds et al. (1976), and in the papers by Lamb et al. (1975), Liu and Seinfeld (1975), Liu et al. (1976b), and Reynolds (1977). A summary of the technical features included in this model is given in the report by Reynolds et al. (1982).

## A. THE TECHNICAL FEATURES OF THE SAI AIRSHED MODEL

The SAI Airshed Model can simulate the dynamic behavior of up to 20 pollutants. When photochemical simulations are carried out, 11 species must be included: nitric oxide (NO), nitrogen dioxide ( $\text{NO}_2$ ), ozone ( $\text{O}_3$ ), single-bonded carbon atoms (PAR), double-bonded carbon atoms except ethylenes (OLE), aromatic-bonded carbon atoms (ARO), carbonyl-bonded carbon atoms, ethylene, benzaldehyde (BZA), peroxyacetyl nitrate (PAN), and carbon monoxide (CO). In addition, the model can provide predictions of  $\text{SO}_2$  and aerosol.\* The user can exercise any of the following options with respect to the pollutants to be simulated:

- > Up to 20 unreactive pollutants.
- > Photochemical species, CO,<sup>†</sup> and up to 9 other unreactive pollutants.
- > Photochemical species, CO,  $\text{SO}_2$ , and up to 8 other unreactive pollutants.
- > Photochemical species, CO,  $\text{SO}_2$ , aerosol,<sup>§</sup> and up to 7 other unreactive pollutants.
- > Photochemical species, CO, aerosol,\*\* and up to 8 other unreactive pollutants.

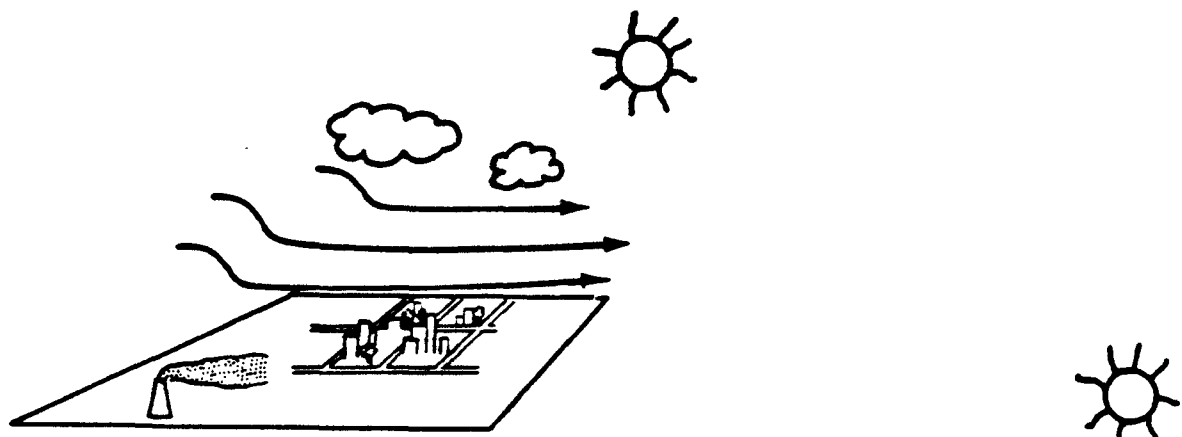
As illustrated in Figure II-1, simulations are performed on a three-dimensional grid selected by the user to cover the region of interest. Figure II-1 also shows the various atmospheric processes treated in each grid cell. The base of an elevated inversion layer is often selected as the top of the grid, but the model includes provisions to facilitate treatment of the inversion layer itself. The model's computer codes can handle any number of cells in the horizontal and vertical directions.

\* The aerosol component includes emitted particulates and organic, nitrate, and sulfate aerosols formed in the atmosphere.

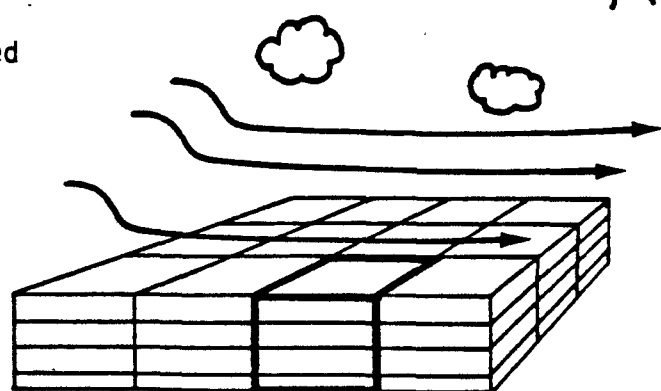
† Photochemical species include the first 10 pollutants cited above.

§ This aerosol consists of organic, nitrate, and sulfate components.

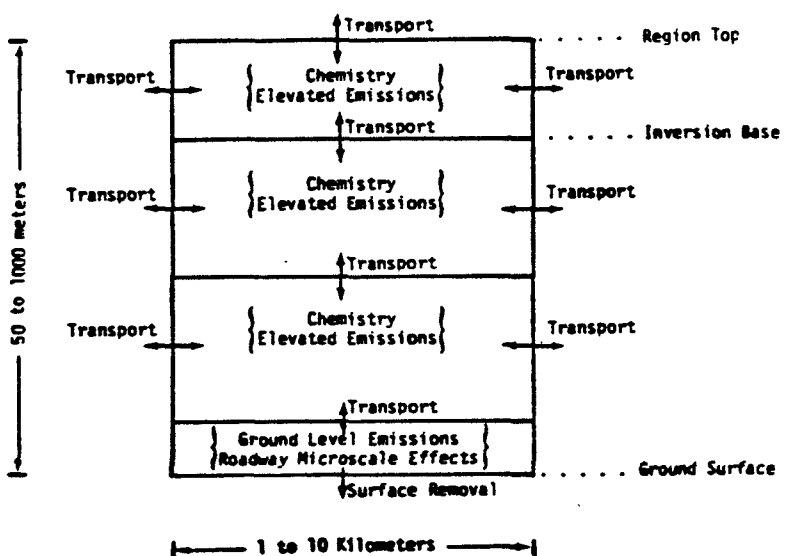
\*\* This aerosol consists of only the organic and nitrate components.



(a) The Area To be Modeled



(b) Specification of the Grid



(c) Atmospheric Processes Treated in a Column of Grid Cells

Figure II-1. Schematic Illustration of the Grid Used and Treatment of Atmospheric Processes in the SAI Airshed Model

## 1. Treatment of Atmospheric Chemistry

Ozone is not emitted from sources; rather, it is formed by chemical reactions in the atmosphere. Consequently, to calculate ozone concentrations, a physicochemical model must contain a kinetic mechanism, that is, a group of chemical reactions and rate constants intended to represent the ozone formation process. Because of computing time limitations, it is not possible to include explicitly each chemical species of interest; thus, most of these kinetic mechanisms treat organic compounds in groups, often on the basis of the reactive functional groups they contain. For example, propylene, 1-butene, and 1-pentene, which each contain a carbon-carbon double bond, can be aggregated together as a single "lumped" species. Such a mechanism was used in the airshed model until 1976, when the Carbon-Bond Mechanism was developed.

Because of the association of reactions and reactivities with carbon bonds, the range of reactions and rate constants to be treated in a kinetic mechanism can be narrowed considerably when each carbon atom is treated according to its bond type. This concept is the basis for the Carbon-Bond II Mechanism, a 65-reaction mechanism developed at SAI and employed in the SAI Airshed Model (Whitten and Hogo, 1977; Whitten, Killus and Hogo, 1980). In this mechanism, the carbon atoms of each organic compound are assigned to one of the following groups: single-bonded carbon atoms, double-bonded carbon atoms except ethylene, ethylene, aromatic-bonded carbon atoms, carbonyl-bonded carbon atoms, and benzaldehyde. Further discussion of the definition and treatment of these groups is given in the final report on the application of the model to the Tulsa urban area (Reynolds et al., 1982). The user can add to the mechanism four reactions describing  $\text{SO}_2$  oxidation and reaction expressions accounting for the formation of sulfate, organic, and nitrate aerosol products.\* Simulations of smog chamber experiments indicate that the Carbon-Bond Mechanism performs significantly better than those previously employed in the airshed model (Whitten, Hogo, and Killus, 1980).

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\* The 69-reaction mechanism is shown in Chapter IX and is discussed in more detail in Reynolds et al. (1982).

## 2. Treatment of Advective Pollutant Transport

Pollutants are transported primarily by advection, that is, by the mean or bulk motion of the wind. A major difficulty in treating advection is wind shear, the variation of the wind with altitude. The wind speed near the ground typically increases with height, but the wind several hundred meters aloft may be faster or slower--and frequently from a different direction. In the original airshed model, wind shear was neglected, but later sensitivity studies indicated that inclusion of wind shear effects in the model could significantly alter NO<sub>2</sub> and ozone predictions (Reynolds et al., 1976). Consequently, the capabilities of the model were expanded to treat wind shear phenomena. Theoretical wind shear relationships were derived by Lamb (1976) using the predictions of a planetary boundary layer model developed by Deardorff (1972). These relationships are useful when wind measurements aloft are not available. When wind measurements are available both at the surface and aloft, one or more objective techniques are used to prepare appropriate three-dimensional wind inputs to the model. Two algorithms in the model are described by Liu et al. (1973) and Killus et al. (1977).

## 3. Treatment of Turbulent Diffusion

Pollutants are transported and dispersed largely by the action of the wind rather than by molecular diffusion. The treatment of advection discussed above does not represent pollutant transport completely because it ignores the influence of small-scale features of the wind, called eddies. Describing these microscale features of the wind deterministically is difficult because available data are insufficient to characterize winds on such a fine scale. Consequently, it is necessary to parameterize the microscale turbulent diffusion processes. In the SAI Airshed Model, and most other grid models as well, transport of a pollutant by turbulent diffusion is assumed to be proportional to the rate of change of concentration with position (i.e., concentration gradient). The

proportionality factor is termed the eddy diffusivity coefficient. These coefficients are sometimes treated as constants, but they can more accurately be treated as functions of atmospheric stability, surface roughness, and height above the ground.

Because of the empirical nature of the eddy diffusivity concept, it has heretofore been difficult to obtain precise measured or theoretical estimates for the diffusivity coefficients. At SAI, considerable effort has been devoted to the study of turbulent transport processes. As a result of these efforts, we have developed a novel procedure for estimating diffusivity coefficients. In this approach, control theory techniques are employed in conjunction with the results of a sophisticated planetary boundary layer model to generate "optimal" diffusivity values. For further details, the reader is referred to the reports by Lamb (1976) and Lamb et al. (1977).

#### 4. Treatment of Other Microscale Phenomena

Several phenomena besides turbulent diffusion have characteristic spatial or temporal scales too small for explicit, deterministic resolution in urban and regional grid models. Two important examples are the rise of a hot, buoyant emissions plume from a large point source and the scavenging of ozone near roadways by NO emitted from vehicles. Studies of these phenomena and others at SAI resulted in the incorporation of an improved treatment of plume rise from point sources. In addition, a preliminary version of a module to treat the microscale effects of the emissions from a network of line sources (e.g., highways) in the lowest layer of grid cells has also been developed. In this layer, taken to be about 20 meters deep, NO emissions from vehicles on a roadway are allowed to react only with the ozone in the immediate vicinity of the roadway. More detailed descriptions of these algorithms and other microscale phenomena are given by Lamb (1976) and Lamb et al. (1977). Since the performance of the microscale module has not been thoroughly evaluated, we do not recommend that this optional model feature be exercised at this time.

## 5. Treatment of Surface Removal Processes

Many types of pollutants, including NO<sub>2</sub>, ozone, and SO<sub>2</sub> in particular, can be removed from the atmosphere by chemical reaction, adsorption, or absorption at various surfaces found in urban areas. A methodology for calculating the removal of gaseous pollutants by surface sinks has been incorporated in the model. Surface removal is assumed to take place in two steps: transport to the surface followed by uptake by the surface. Parameterization of this two-step process is accomplished by defining a resistance to mass transport and a resistance to surface removal. The transport resistance is estimated from theoretical considerations of turbulent transfer in the atmospheric boundary layer, and the surface resistance is obtained from experimental data on the uptake of pollutants by various types of surfaces (Killus et al., 1977).

## 6. Other Technical Features

The numerical procedure used to solve the conservation-of-mass equations for each pollutant is an important component of a grid model. To minimize the propagation of numerical errors, techniques for treating horizontal (advective) pollutant transport--an element of the model particularly difficult to handle numerically--were examined. The results of these studies indicated that, of the techniques examined, the SHASTA integration scheme provided the best balance between accuracy of prediction and computational speed (Reynolds et al., 1976; Killus et al., 1977).

The SAI Airshed Model contains features designed to reduce cost by minimizing labor and computing expenses. For example, computerized data preparation programs are available for automating the conversion of various types of data into the proper input format. These features have no effect on the model's treatment of atmospheric processes, but they do aid the user in preparing the inputs required by the Airshed Model.

The model output consists of the predicted pollutant concentrations for every grid cell. These predictions are generally averaged over a period of one hour and are saved on a computer file for subsequent display and analysis. An example of the printed output produced by the display program is given in Figure II-2. If other computer graphics capabilities are available, then the results can be plotted as shown in Figures II-3 and II-4. Note, however, that the computer programs described in this manual produce only printed displays of the predicted concentrations. Gridded census data can be used in conjunction with the model predictions to estimate the dosages and exposures that are experienced by the human population within the modeling region. The results of an exposure analysis carried out for Denver are shown in Figure II-5. Of course, other displays and analyses can be prepared depending on the needs of the user.

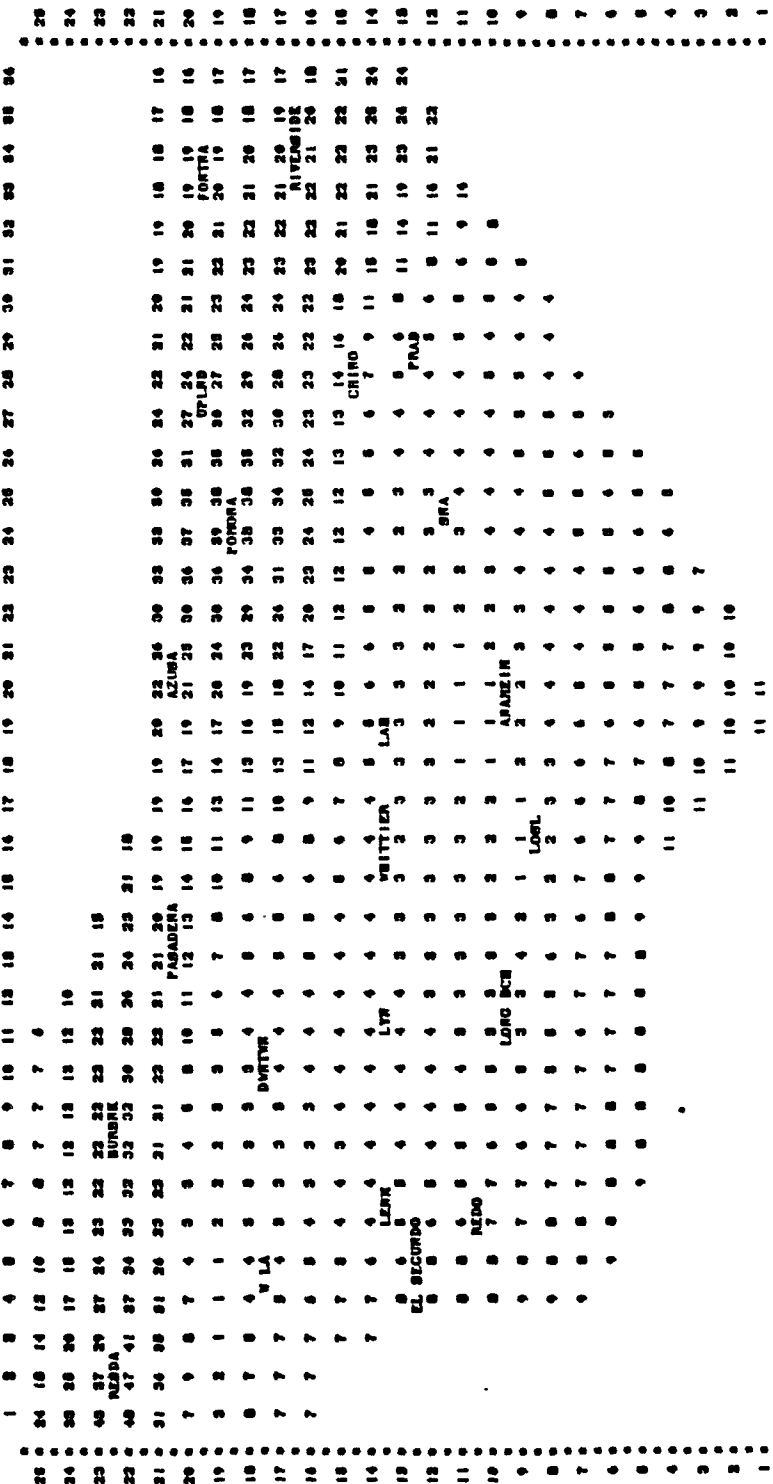
## B. MODEL USAGE

In general, studies that employ the SAI Airshed Model can be segmented into three phases: adaptation, evaluation of validity and performance, and application. The specifics of each phase depend on the region to be modeled and the intended application, but certain characteristics apply to each case.

The primary objective of Phase I is to adapt the model for application to a particular region. Efforts involved in this undertaking include:

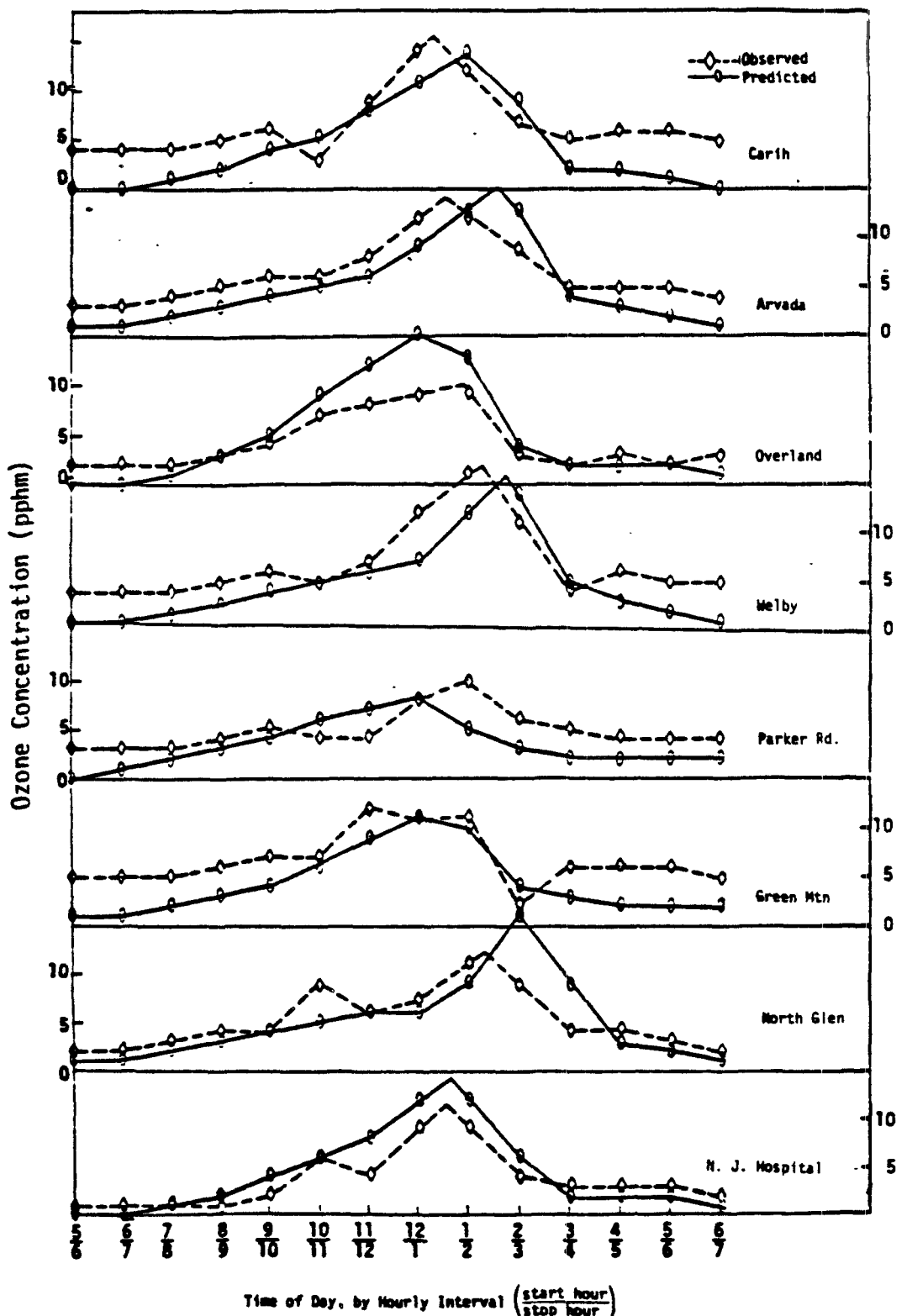
- > Review of pertinent air quality studies of the region.
- > Examination and assessment of available emissions and aerometric data.
- > Specification of supplemental data (if any) needed for verification and application of the model.
- > Collection of supplemental data, if necessary, through intensive field monitoring studies.
- > Adaptation of existing, or development of new, input data preparation programs.

Upon completion of these tasks, the user is in a position to carry out photochemical simulations with the model.



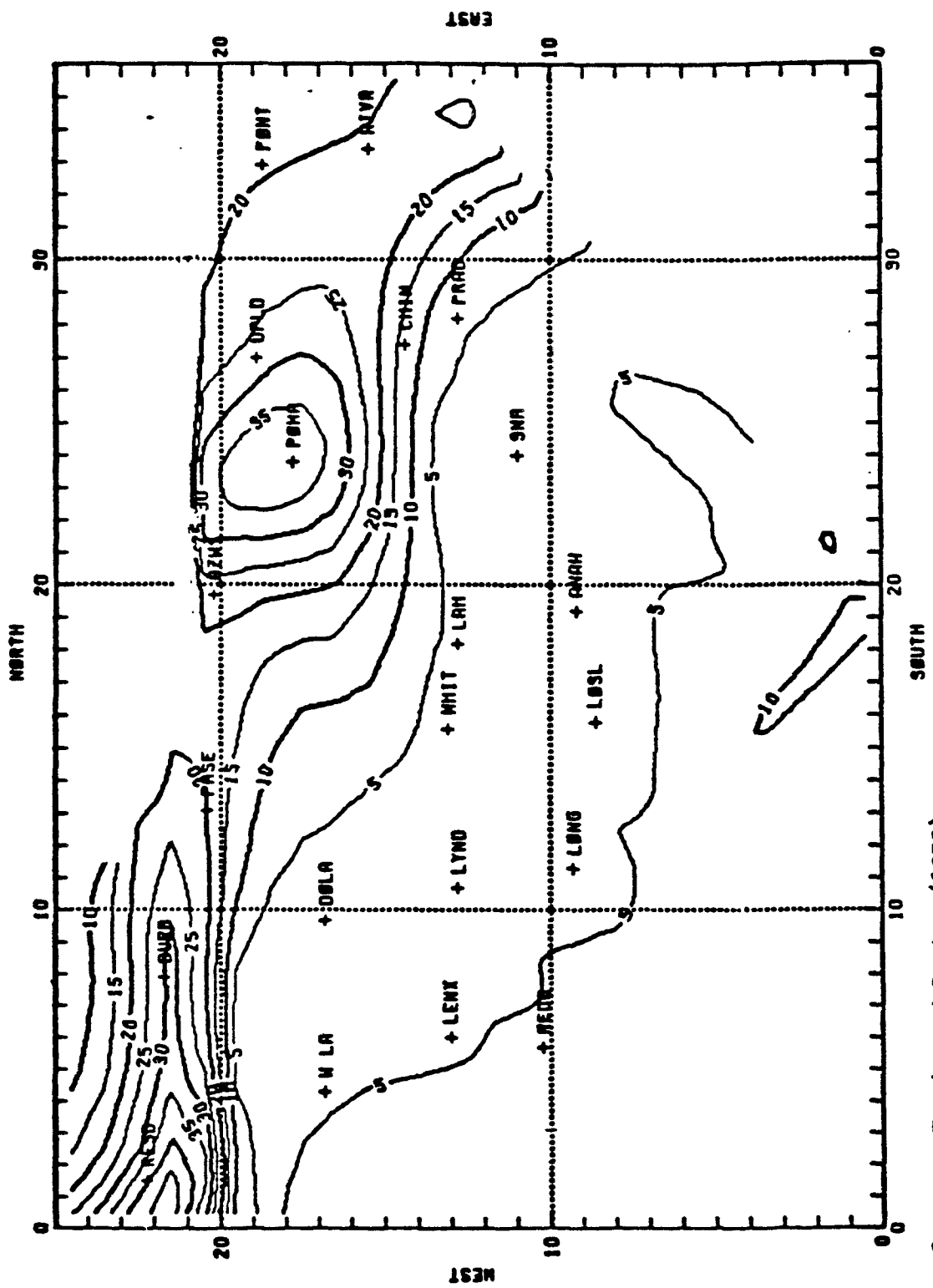
Source: Tesche and Burton (1978).

Figure II-2. Example of a Printed Map of Ground-Level Concentration Predictions. Average ground-level O<sub>3</sub> concentrations (pphm) are given for Los Angeles on 4 August 1975 between 1:00 and 2:00 p.m. PST.



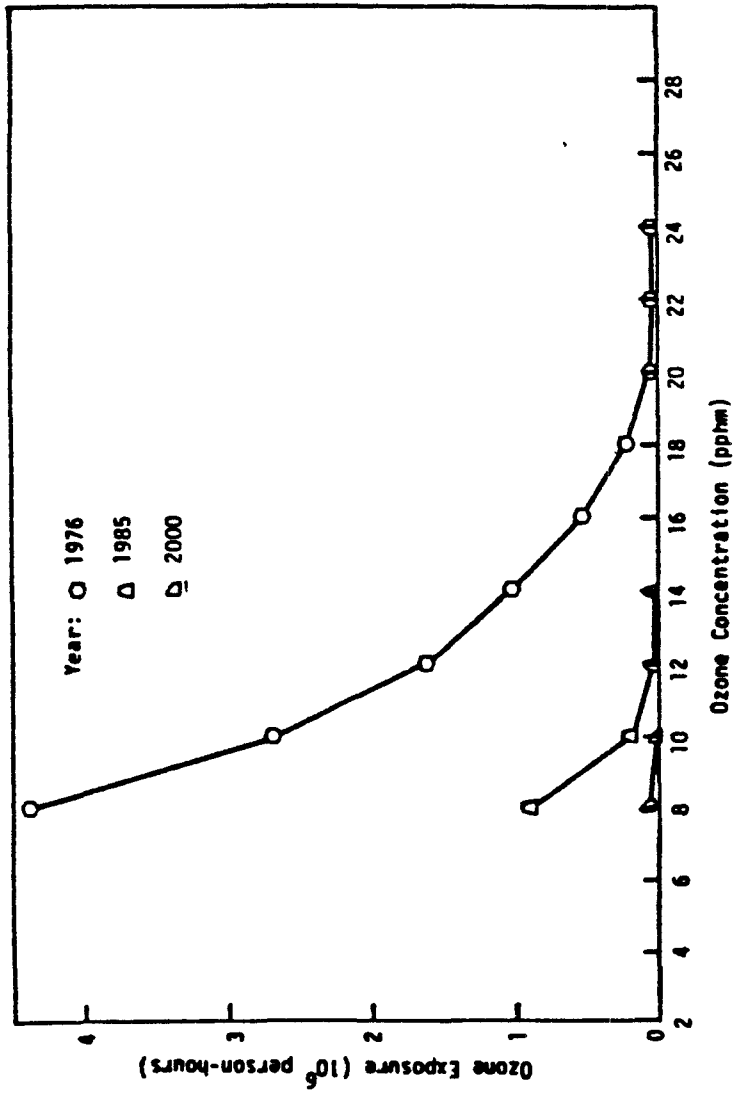
Source: Anderson et al. (1977).

Figure II-3. Example of Plots Illustrating Model Predictions and Actual Measured Concentrations. Ozone concentrations measured at various stations in the Denver area on 28 July 1976 are compared with the corresponding predictions of the SAI Airshed Model as modified for Denver.



Source: Tesche and Burton (1978).

Figure II-4. Example of a Plotted Contour Map of Ground-Level Concentration Predictions. Average ground-level  $\text{O}_3$  concentrations (pphm) are given for Los Angeles on 4 August 1975 between 1:00 and 2:00 p.m. PST.



Source: Anderson et al. (1977)

Figure II-5. Example of the Results of a Population Exposure Analysis:  
Denver

Phase II is devoted to assessing the performance of the model. First, the model is used to simulate the formation of photochemical smog on one or more days in the past chosen to represent typical or episodic meteorological conditions. Model predictions are then examined in light of actual observations in an effort to characterize model performance (see Figure II-3). Tasks in this phase include:

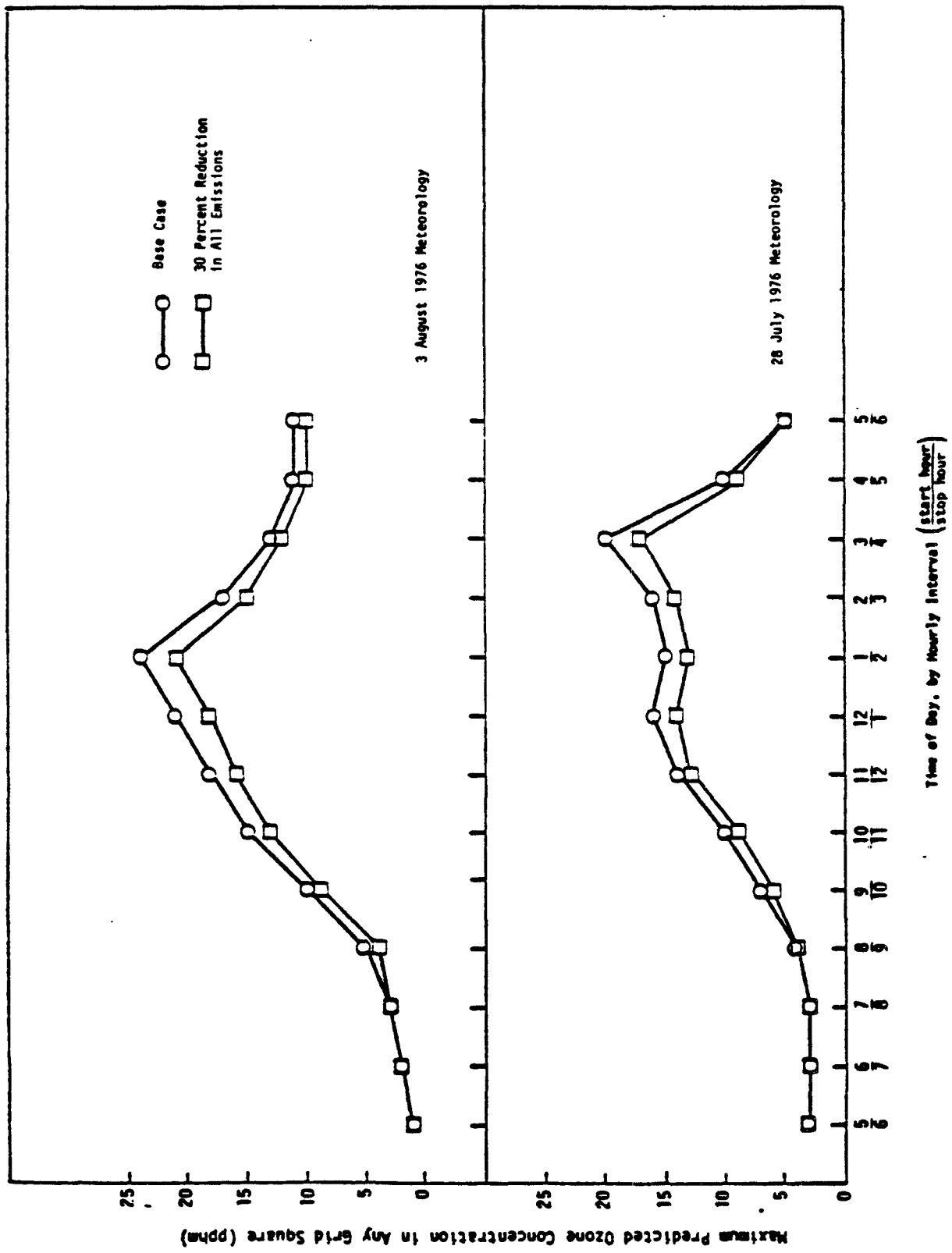
- > Selection of days for testing.
- > Preparation of model inputs for each day.
- > Performance of model simulations and analysis of results.
- > Collection of additional data, modification of the model or input data preparation programs, or both, as needed, until satisfactory model performance is obtained.

After the model has been demonstrated to perform adequately, further applications studies can be carried out with reasonable confidence.

The intent of the air quality modeling study is not fully realized until Phase III, when control issues are addressed. Applications of the model generally involve the following tasks:

- > Definition of the scenarios to be considered.
- > Specification of input conditions for each scenario (for example, adverse meteorological conditions, weekday emissions, and so on).
- > Performance of model simulations and analysis of results.

In evaluating the effects of some change in emissions, one compares the predictions for that case with those for a base case (see Figure II-6). The base case might, for example, represent conditions without the imposition of emissions controls. Several measures can be examined in assessing the effects of emissions alterations, including maximum one-hour-average concentrations, population dosages, and area-wide concentration changes. Of course, the model predictions can also be compared



Source: Anderson et al. (1977).

Figure 11-6. Sample Results of Phase III--Simulation of Emission Control Scenarios. Ozone concentrations predicted by the SAI Airshed Model for the Denver area for 1976 emissions and a 30 percent emissions reduction are shown; note that reducing emissions of both  $\text{NO}_x$  and organic species by 30 percent reduced the maximum predicted ozone concentration by about 15 percent.

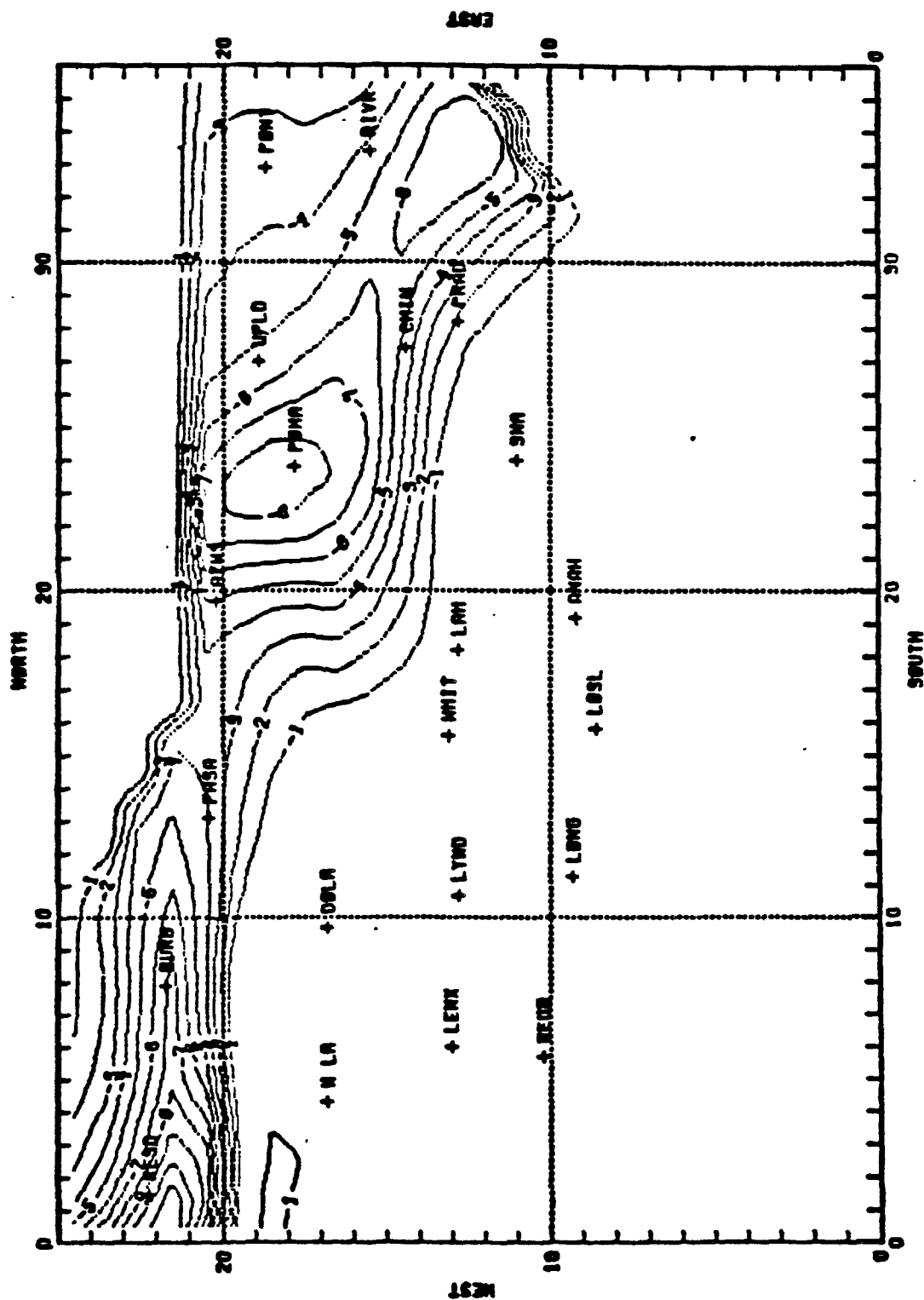
with air quality standards. As shown in Figures II-5 through II-7, various graphical techniques can be employed to display the model results.

### C. MODEL APPLICATIONS

Many seemingly straightforward but inherently difficult issues arise in planning for the abatement and control of air pollution in an urban area. Some of these issues can best be addressed through the use of an airshed model; others can be addressed only in this way. This section discusses potential model applications and summarizes studies performed through June 1978 that employed the SAI Airshed Model.

Analyses that can be carried out with the SAI Airshed Model include:

- > Development of an emissions control policy. For example, the model can be applied to cities to ascertain the influence of various potential motor vehicle emissions standards on ambient CO, ozone, and NO<sub>2</sub> concentrations.
- > Evaluation of proposed emissions control strategies. The model would be particularly useful in helping to establish estimates of the relative emissions reductions of organic species and NO<sub>x</sub> required to achieve the ozone and NO<sub>2</sub> air quality standards.
- > Selection of sites for future sources of air contaminants. Of particular interest here is the examination of alternative future source development scenarios that would minimize the possibility of subsequent violations of air quality standards.
- > Planning for the control of air pollution episodes. The model can be used to ascertain either the extent of intermittent controls required to relieve episodic conditions or the effectiveness of alternative control measures, such as emissions reductions in various portions of a region.



Source: Tesche and Burton (1978).

Figure II-7. Contour Plot Illustrating the Predicted Effect of a Control Strategy. This figure indicates the predicted effect on O<sub>3</sub> levels in the Los Angeles area of a 50 percent reduction in motor vehicle emissions of NO<sub>x</sub> and organic species. The O<sub>3</sub> isopleths (pphm) represent the difference between the control strategy and base case predictions for the hours 1:00 to 2:00 p.m. PST.

- > Assessment of the contributions responsible for existing levels of air pollution. The model can be exercised both with and without particular sources included in the emissions inputs to examine their contributions to existing air quality.
- > Estimation of air quality in areas for which pollutant concentration data are unavailable. Model predictions can be used to locate regions experiencing high pollutant concentrations and thus aid in selecting additional monitoring sites.

In general, applications such as those cited above are carried out by altering any of four inputs to the model: the spatial distribution of emissions, the temporal distribution of emissions, the composition of organic compounds emitted from sources, and the initial and boundary pollutant concentrations. Control strategies that attempt to alter the reactivity of organic species emitted into the air, such as solvent substitution measures, can be handled in the model by altering the magnitude and composition of the emissions from the appropriate sources. In addition, emissions from one city may be responsible for increased background pollutant concentrations in a city downwind. The effects of alterations in background concentrations can be examined through changes in the initial and boundary pollutant concentrations used as input to the model.

Besides these applications, the SAI Airshed Model can be used in other photochemical air pollution analyses. Sensitivity studies, for example, can help assess the relative influences of various physical and chemical processes on pollutant concentrations in a particular region. Such information might be used as a basis for simplifying the treatments of various processes in the model or for reducing the amount of input data to be collected. Moreover, the model can serve as an educational tool by enhancing the existing understanding of the causes of photochemical air pollution in the region. The SAI Airshed Model can also be employed to test new theories, algorithms, and parameterizations of atmospheric processes. Thus, the model is useful in applications ranging from the evaluation of

emissions control strategies to the advancement of knowledge in atmospheric science.

Initially, most applications of the SAI Airshed Model were in research projects aimed at improving the model's components or in evaluative studies to ascertain its performance. However, several studies have been carried out that are illustrative of the applications cited earlier. We list below several of these investigations carried out through 1978 that the user can refer to for information on applying the model to suit specific needs.

- > An evaluation of four motor vehicle emissions control strategies for Los Angeles. The scenarios included (1) a 30 percent reduction in vehicle mileage, (2) a 100 percent reduction in vehicular emissions, (3) 79 percent, 73 percent, and 62 percent reductions in the emissions of organic species, NO<sub>x</sub> and CO, respectively (strategy proposed by the Environmental Quality Laboratory at Caltech), and (4) a 90 percent reduction in vehicle mileage in the downtown Los Angeles area (Reynolds et al., 1973a).
- > An examination of a proposed State Implementation Plan for Los Angeles (June 1973). Three scenarios were considered in this study: (1) baseline 1969 conditions, (2) baseline 1977 conditions (with existing legislation in force), and (3) 1977 conditions with additional emission reductions resulting from gasoline rationing, carpooling, implementation of special bus lanes and parking restrictions, and limited control of stationary sources (Reynolds and Seinfeld, 1975).
- > An analysis of air quality for the Denver metropolitan region for the years 1976, 1985, and 2000. This study examined the impact on air quality of the urban growth associated with the construction of new wastewater treatment facilities. Model runs were also made to ascertain the sensitivity of the predictions to various land use scenarios that involved alterations to the spatial

distribution of emissions (Anderson et al., 1977). Samples of the results are given in Figures II-3, II-5, and II-6.

- > A demonstration of the model's utility in selecting sites for aerometric monitoring. As part of a study sponsored by the EPA Environmental Monitoring and Support Laboratory in Las Vegas, the SAI model was applied to Las Vegas and the predictions were employed to estimate the best locations for CO monitors (Liu et al., 1977).
- > An examination of the influence of highway location, design, and operation on photochemical smog formation. This study, sponsored by the Federal Highway Administration, is evaluating various transportation control strategies and will provide guidance to highway personnel in the use of photochemical models (Reynolds et al., 1979).
- > An analysis of model sensitivity to ascertain the relationships between the quantity of data collected for use in estimating model inputs and the performance of the SAI Airshed Model. This information will be integrated into a larger study for the EPA Office of Air Quality Planning and Standards (OAQPS) aimed at examining means for employing the SAI model in evaluating the performance of simple oxidant relationships (Tesche et al., 1981).
- > An examination of the capability of the SAI model to resolve the impact of emissions from major elevated point sources on regional air quality. Sponsored by the Southern California Edison Company, this study focused on the influence of NO<sub>x</sub> and SO<sub>2</sub> emissions reductions on the expected incremental changes in NO<sub>2</sub>, ozone, and sulfate levels aloft and at the ground in the Los Angeles area (Tesche and Burton, 1978).
- > An evaluation of alternative population growth scenarios for the Los Angeles metropolitan area. The purpose of this study is to help regional planners at the Southern California Association of Governments determine the extent to which future air quality levels in 1990 will be sensitive to the

manner in which the area is allowed to develop over the next 12 years.

Other ongoing studies that are not direct applications of the model but that do have some bearing on model usage include:

- > An investigation to establish guidelines for setting model performance measures and standards. This study for the OAQPS will yield information that will aid in evaluating the suitability of a model for use in a particular application (Hayes, 1978).
- > A study to establish guidelines for validation of air quality models. This effort will provide guidance to users in the examination of model performance. This study is also being carried out for OAQPS (Hillyer, Reynolds, and Roth, 1979).

In addition to these studies, versions of the SAI Airshed Model have been implemented for use in several cities, both in this country and in Europe. Table II-1 summarizes these applications.

TABLE II-1. USE OF THE SYSTEMS APPLICATIONS (SAI) AIRSHED MODEL  
IN STUDIES OF AIR QUALITY IN URBAN AREAS.

User	Model Version	Study Site	Sponsor and Reference
Systems Appl., Inc.	1973	Los Angeles	EPA, Contracts CPA 70-148 and 68-02-0339 (Reynolds, 1973)
Systems Appl., Inc./ CAL TRANS/CARB	1973	Los Angeles	California Department of Transportation, Contract K-7319; California Air Resources Board, Contract ARB 4-250
Systems Appl., Inc.	1973	Los Angeles	Texaco
Systems Appl., Inc.	1973	Los Angeles	Western Oil and Gas Association
Systems Appl., Inc.	1977	Los Angeles	DOT, Contract DOT-FH-11-8529 (Reynolds et al., 1979)
Systems Appl., Inc.	1977	Los Angeles	Southern California Edison Company (Tesche and Burton, 1978)
Systems Appl., Inc.	1977	Los Angeles	Southern California Association of Governments (Reynolds et al., 1978)
Systems Appl., Inc.	1978	Los Angeles	Southern California Association of Governments and South Coast Air Quality Management District
Systems Appl., Inc.	1978	Los Angeles	NCAQ Contract (Souten, Tesche, and Oliver, 1981)
Systems Appl., Inc.	1982	Los Angeles	Western Oil and Gas Association (Roth et al., 1984)
Systems Appl., Inc.	1978	Tulsa	EPA Contract 68-02-3370 (Reynolds et al., 1982)
Systems Appl., Inc./ TNO	1978	The Netherlands	TNO (Builtjes et al., 1982)
Systems Appl., Inc./ Freie Universität Berlin/TNO	1978	Bonn/Cologne	Stern and Scherer (1982)
Systems Appl., Inc.	1978	California Central Valley	PG&E
Systems Appl., Inc.	1978	Los Angeles	EPA, Contract 68-02-2870 (Tesche et al., 1981)

Continued

TABLE II-1 (Concluded)

User	Model Version	Study Site	Sponsor and Reference
Systems Appl., Inc.	1978	Los Angeles	EPA, Contract RP-1375-1-2 (Seigneur et al., 1983)
Systems Appl., Inc./ State of Colorado	1973	Denver	Colorado Department of Highways, Contract 75-109
Systems Appl., Inc.	1976	Denver	EPA Region VIII, Contract 68-01-4341 (Anderson et al., 1977)
Systems Appl., Inc./ State of Colorado	1976	Denver	Colorado State Health Department and Department of Highways
Systems Appl., Inc.	1977	Denver	DOT, Contract DOT-FH-11-8529 (Reynolds et al., 1979)
Systems Appl., Inc.	1977	Denver	ERT/Colorado Interstate Pipeline Company
Systems Appl., Inc.	1973	Las Vegas	EPA, Contract 68-03-2174 (Liu et al., 1977)
Systems Appl., Inc./ British Leyland	1973	Turin, Italy	British Leyland
Systems Appl., Inc.	1977	Phoenix	EPA, Contract 68-03-2446
Systems Appl., Inc.	1977	Tampa	EPA, Contract 68-03-2446
Systems Appl., Inc.	1978	St. Louis	EPA, Contract 68-02-2429
Systems Appl., Inc.	1978	St. Louis	EPRI, Contract RF-1375-1-2 (Seigneur et al., 1983)
Systems Appl., Inc.	1978	Sacramento	DOT, Contract DOT-FH-11-8529 (Reynolds et al., 1979)
Systems Appl., Inc.	1982	Las Vegas	Southern California Edison Company (Tesche, Oliver, and Haney, 1982)
SRI, International	1982	Philadelphia	EPA, Contract 68-02-3538
Systems Appl., Inc.	1983	Philadelphia	EPA, Contract 68-02-3582 (Haney, Tesche, and Killus, 1983)

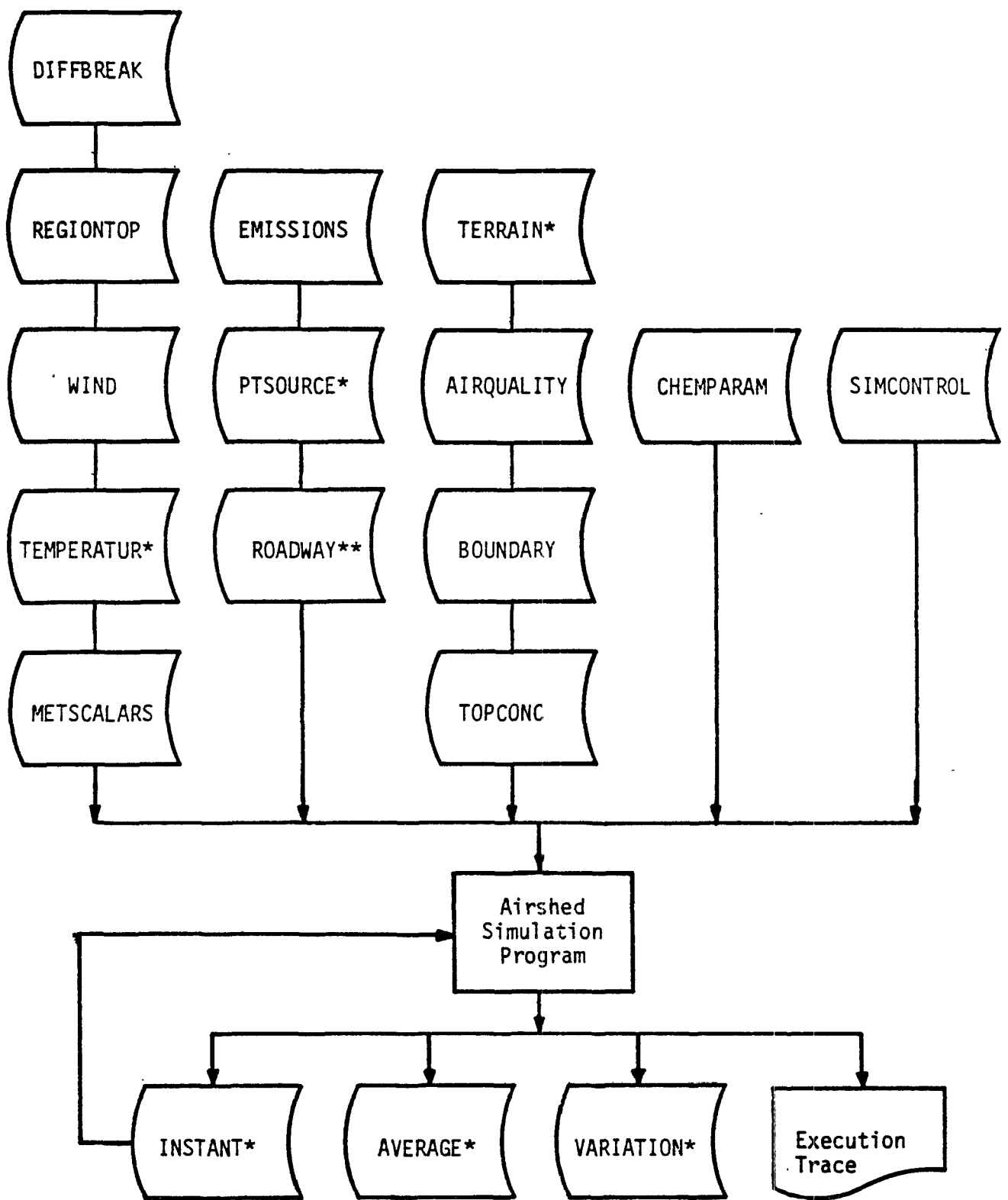
### III OVERVIEW OF THE SYSTEM

The SAI Airshed Model System contains at its core the Airshed Simulation Program, the input data that consist of 10 to 14 files, depending on the program options chosen. Each file is created by a separate data preparation program. The output from the Airshed Simulation Program consists of three data results files and an execution trace report. The output data files can be further processed by the display and analysis programs and can also be used for restarting the simulation. Figure III-1 shows the Airshed Simulation Program and identifies its input and output files.

The SAI Airshed Model System can be described according to the five major tasks or functions that the user and the programs must perform:

- > M.E.B. (meteorology, emissions, and initial and boundary conditions) file preparation.
- > M.E.B. file segmentation.
- > Control data preparation.
- > Airshed simulation.
- > Display and analysis.

This classification of programs and tasks is used as the organizational basis for both this User's Manual and the companion Systems Manual. Each of the five major functions is represented by a rectangular box in Figure III-2, which illustrates the flow of information through the system and shows the position of each major function within the system. The following sections of this chapter briefly discuss each function in terms of the activities required of the user, and the remaining chapters of this report present the detailed information necessary for the completion of each task.



\* Optional files.

\*\* Optional file not recommended for usage at this time.

Figure III-1. Airshed Simulation Program with Input and Output Files

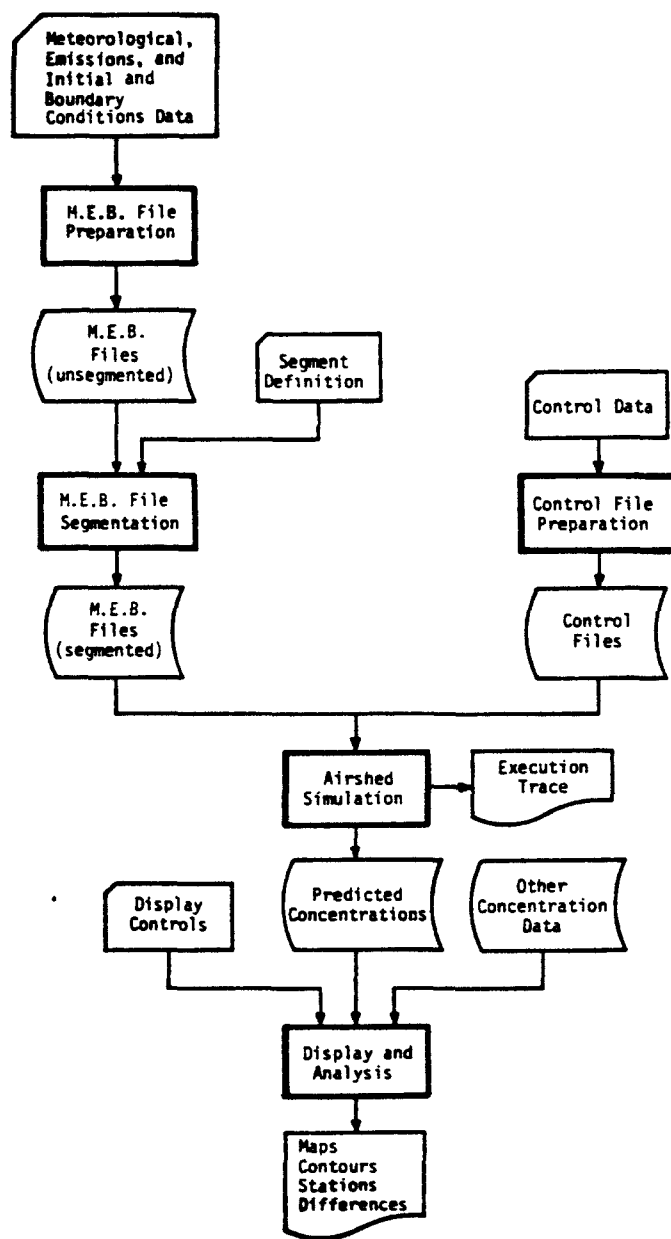


Figure III-2. Information Flow in the SAI Airshed Model System

## A. M.E.B. DATA FILE PREPARATION

Of the 14 files input to the Airshed Simulation Program, two are classified as "control files" and are discussed separately. The preparation of the remaining 12 files, which contain meteorological, emissions, and initial and boundary conditions data (referred to in this report as the "M.E.B. files"), is discussed here. The Airshed Simulation Program requires appropriate meteorological, emissions, and initial conditions data for each cell of the region's rectangular grid. In addition, the concentration of each pollutant must be specified at each point on the boundary where the wind is flowing into the region. These data enter the program through the M.E.B. files. Table III-1 lists the 12 files, grouped by type; the 10-character code name as well as the descriptive name is given for each file. Also, the data items contained in each are listed and classified according to the nature of the variability (spatial, temporal, or pollutant) exhibited by each item.

For each file, there is a data preparation program that takes either observations or estimates at specific locations in the region, creates a fully gridded data field, and writes the data to the file in the appropriate format. The 12 M.E.B. data preparation programs are designed to be used together as a unified, user-oriented package. The input formats to all the programs are standardized and reasonably self-documenting. The interpolation methods are intended to accommodate input data of any complexity, from variables for which little or no data are available to those for which there are many observations and well-known interactions. Also, the units of measure are standardized, and a flexible unit conversion scheme is built into the programs.

The M.E.B. data preparation function consists of the following tasks for the user:

Table III-1. M.E.B. Data Files

<u>File Description</u>	<u>Nature of Variability</u>			
	<u>Axes</u>	<u>Vertical Direction</u>	<u>Species</u>	<u>Time</u>
<b>Meteorological Files</b>				
WIND--Wind				
$ u _{\max}^+$ , $ v _{\max}^+$				t
Average wind speed along each boundary				t
u-velocity component	x	y	z	t
v-velocity component	x	y	z	t
DIFFBREAK--Diffusion Break				
Diffusion break	x	y		t
REGIONTOP--Top of the Region				
Top of the modeling region	x	y		t
*TEMPERATUR--Temperature				
Ground-level temperature	x	y		t
METSCALARS--Meteorological Scalars				
Temperature gradient below diffusion break				t
Temperature gradient above diffusion break				t
Exposure index				t
Diurnal photolysis rate constant				t
Concentration of H <sub>2</sub> O				t
Atmospheric pressure				t
<b>Emissions Files</b>				
EMISSIONS--Ground-Level Emissions				
Ground-level emissions (lumped mobile and nonpoint stationary sources)	x	y	sp	t
*PTSOURCE--Point Source Emissions				
Location of source	(x,y)			
Effective plume height	(x,y)			t
Vertical cell index	(x,y)			t
Emissions rate	(x,y)	(z)	sp	t

Table III-1 (Concluded)

File Description	Nature of Variability			
	Axes	Vertical Direction	Species	Time
<b>*ROADWAY--Roadway Microscale Parameters **</b>				
Emissions from motor vehicles	x y		sp	t
Roadway microscale parameter	x y			t
<b>Boundary Files</b>				
<b>*TERRAIN--Terrain Surface Parameters</b>				
Surface roughness	x y			
Vegetation factor	x y			
<b>BOUNDARY--Boundary Definition and Concentrations</b>				
Boundary definition	(x,y)			
Boundary concentrations	(x,y)	z	sp	t
<b>AIRQUALITY--Air Quality</b>				
Initial concentrations	x y	z	sp	(t)s
<b>TOPCONC--Concentrations at the Top of the Region</b>				
Boundary concentrations at the top of the modeling region	x y		sp	t

x = east-west axis of grid.

y = north-south axis of grid.

z = vertical axis of grid.

sp = chemical species.

t = time.

( ) = data that vary spatially but are not present over the entire grid.

\* = files that need not be present for a simulation run.

<sup>†</sup> u and v refer to the x and y components of the wind velocity, respectively.

<sup>§</sup> Concentrations must be present at least for the beginning time of the simulation.

\*\* = The roadway microscale procedure is not recommended for usage at this time.

- > Determination of the appropriate files for a given simulation.
- > Detailed examination of the data requirements and optional methods for each file.\*
- > Translation of available data into a form acceptable to the model.
- > Selection of the proper interpolation methods.\*
- > Establishment of the size and location of the region.
- > Establishment of the vertical distribution of cells within the region.
- > Exercise of the data preparation programs, examination of the results, and modifications of the inputs or algorithms until a reasonable set of data is produced.

The preparation of the M.E.B. data files is certainly the most difficult and time-consuming task associated with using the SAI Airshed Model System. It is also the most important task, and careful, intelligent creation of the input is required to yield the best simulation results.

#### B. M.E.B. DATA FILE SEGMENTATION

The SAI Airshed Model System has been designed to enable simulation on a rectangular grid of any size and shape; that is, the dimensions of the region are not fixed within the programs, but are specified at run time. If a specified region does not fit within the core limitations of the host computer, that region can be divided into any number of contiguous subrectangles called "segments"; in this case, the size of each segment, rather than the size of the entire region, is limited by the amount of core available. The Airshed Simulation Program processes each segment in turn and information at the interfaces between segments is maintained.

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\* In some situations, the algorithms embedded in the data preparation programs may not be adequate for the intended model application; in these cases, the user will have to develop and interface the appropriate capability with the SAI Airshed Model System.

To enable simulation processing by segment, the user must create segmented data files from unsegmented files. The M.E.B. file segmentation function consists of the following tasks for the user.

- > Estimation of the size of the largest segment that will fit in core. This size will be determined by the number of species to be simulated, the number of vertical levels, the number of point sources, the program options selected, and the amount of core available on the host computer.
- > Definition of the size and arrangement of segments in the simulation region, and preparation of a segmentation control deck describing that layout.
- > Running the segmentation program on each of the spatially varying M.E.B. files.

### C. CONTROL DATA PREPARATION

The Airshed Simulation Program requires two control files: the Chemistry Parameters file (CHEMPARAM), and the Simulation Controls file (SIMCONTROL), each created by its own data preparation program. Table III-2 summarizes the contents of the control files.

For the Chemistry Parameters file, all chemical species to be simulated are named and their properties specified. This list of species, rather than that appearing on any of the other data files, determines which species will be simulated. If there are any reactive species, the reaction rate constants and their other properties (such as the activation energy and reference temperature) are also specified. The user's tasks in making the Chemistry Parameters file are:

- > Preparation of a card deck describing the species and the chemical mechanism to be used.
- > Running the program that creates the Chemistry Parameters file.

TABLE III-2. Airshed Simulation Program Control Files

CHEMPARAM (Chemistry Parameters)

For each species

Name

Reactivity flag

Steady-state flags

Lower and upper concentration bounds

For each reaction

Rate constant

Photolysis and temperature-dependence flags

Activation energy and reference temperature

For each coefficient

Name

Value

SIMCONTROL (Simulation Controls)

Run identifier

Beginning and ending times

File options

Simulation options

Time slice controls

Minimum chemistry time step

Convergence criteria

Averaging interval

Instantaneous concentration output interval

Print options

---

Note: These files contain no spatially or temporally variable parameters.

The Simulation Controls file includes values for the simulation options, time span, integration controls, and output options, which are the data likely to be changed most frequently. The user's tasks in generating this file are:

- > Preparation of the card deck containing all the simulation control parameters.
- > Running the program that creates the Simulation Controls file.

Although the creation of the Simulation Controls file is described as a separate task among a number of tasks preliminary to running the simulation program, it can be performed just prior to a simulation program run within the same computer job.

#### D. AIRSHED SIMULATION

The Airshed Simulation Program reads the data files that were created in the three previously described steps and performs the simulation as specified by the control files. Its output consists of files containing instantaneous and time-averaged concentration predictions in segmented form, and a printed trace of the progress of the simulation. Concentration predictions are not printed by the simulation program except in a compact form for debugging purposes only. The selective printing of the simulation results is performed by the display and analysis function. The user's tasks in running the airshed simulation are:

- > Proper specification of all input files to be used.
- > Arranging to save the output files that the program creates.

#### E. DISPLAY AND ANALYSIS

The final function of the SAI Airshed Model System is the display and analysis of the simulation results. This function has been removed from the Airshed Simulation Program because often, no matter which output

options are selected for a simulation run, different options are needed at a later time. By writing the concentration predictions of all species in all dimensions to output files, one can select any available output mode at any later time.

The current display program capabilities include printout of either instantaneous or average concentration grid maps for selected species at selected vertical levels at selected times. Instantaneous or average vertical concentration profiles can also be printed at user-specified stations. The user's tasks in running the display program are:

- > Proper specification of all input files required (including the desired output file from the Airshed Simulation Program).
- > Preparation of an input deck containing parameters for the selection of times, species, and vertical levels to be printed, and station locations where vertical concentration profiles will be printed.

In addition, the design of the display program provides for the inclusion of additional analysis capabilities without extensive modification of the program. Specifically, the structure permits the comparison of two concentration data files. The two data files could be the output from two simulation runs (comparing different control strategies, for example), or one file could be simulation predictions and the other a gridded representation of station observation data.

## IV DEFINITION OF SYSTEM CONCEPTS

This chapter discusses several important aspects of the Airshed Model System that must be understood before the model can be used. The concepts explained here include the subdivision of the modeling region into an array of three-dimensional grid cells, the pollutants treated by the model, and the handling of the time variability of various parameters in the model. Also, the segmentation of the entire modeling region into one or more sub-regions for performing the numerical integration process is examined. Finally, the treatment of time increments used in the Airshed Simulation Program is detailed.

### A. THE MODELING REGION (GRID)

The modeling region is a three-dimensional volume subdivided into an array of three-dimensional grid cells. This section first describes the projection of this array onto the x-y, or horizontal, plane and then the method of determining the cell thicknesses in the z, or vertical, direction. Complete instructions for defining the location, size, shape, and resolution of the modeling region are given in Section VI.B.2, which discusses the REGION packet.

#### 1. Horizontal Distribution of Grid Cells

Three pairs of values (which are further discussed below) define the horizontal modeling region:

- > The x- and y-coordinates, in meters, of its origin (the lower left-hand corner point) with respect to a reference origin.

- > The grid spacing (cell size), in meters, along each axis.
- > The number of grid cells along each axis.

In the horizontal plane, the region consists of a rectangular grid area with the x-axis parallel to the equator (east-west) and the y-axis perpendicular to the equator (north-south).\* This area is divided into rectangular grid cells of equal size. Although the spacing of cells may be different for one axis than for the other, yielding nonsquare cells, it must be uniform along each axis.

The region specified must include one cell at each boundary of the simulated area. Thus, if the area to be simulated is a 5 x 10 grid, the region specified must be at least 7 x 12, as shown in Figure IV-1. Also, portions of the inner rectangular area can be omitted from the simulation by defining the boundaries appropriately. Figure IV-2 shows a possible boundary for the Los Angeles region for which the prediction of concentrations in the areas over the Pacific Ocean and the San Gabriel Mountains would not be of interest. Boundaries of the simulated area are input to the simulation through the BOUNDARY file (see Section VI.A.6, which describes the method for defining the horizontal boundaries of the region).

The method for describing locations within the specified region has been standardized in the current version of the SAI Airshed Model. First, a "reference origin"--a point on the map defined by its Universal Transverse Mercator (UTM) coordinates--is designated. All other points in the region, including measuring stations, point sources, and the origin of the modeling region itself, are then described in terms of their distances in the x- and y-directions from the reference origin. The reference origin may coincide with the origin of the region, it may lie within the grid, or it may be completely outside the grid. Thus, the x- and y-coordinates of

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\* This orientation of the horizontal grid is used for reference throughout the system. However, some other more useful or convenient grid orientation can be used, provided that point locations and wind vector components are carefully defined.

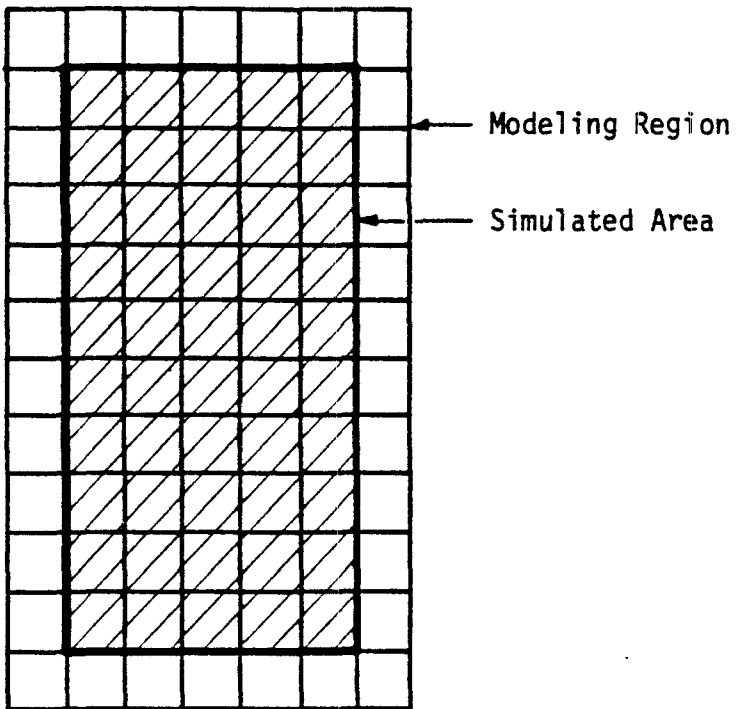


Figure IV-1. Example of a Modeling Region That Includes the Required Boundary Cells

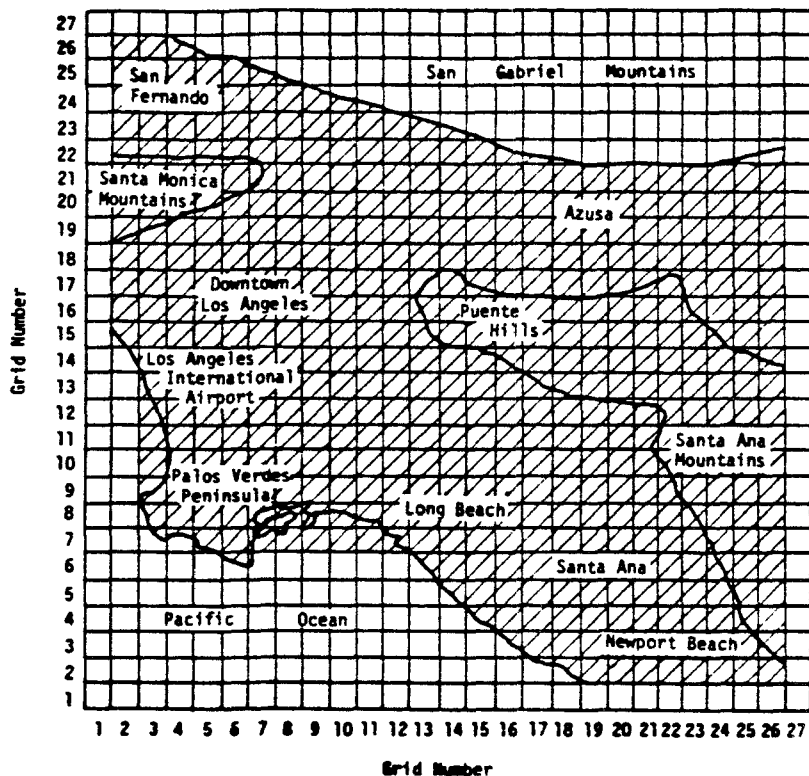


Figure IV-2. 54 Mile by 54 Mile Modeling Region of Los Angeles Basin. Each grid square is 2 miles by 2 miles.

any point with respect to the reference origin may be positive or negative. In Figure IV-3, the reference origin is outside the modeling region. The x-coordinate of the region origin is positive and the y-coordinate is negative. The point sources and measuring stations appear both within the region and outside it. Using this method for locating points, the user can change the location, size, shape, or resolution of the modeling region without changing the coordinates of any of the other input data.

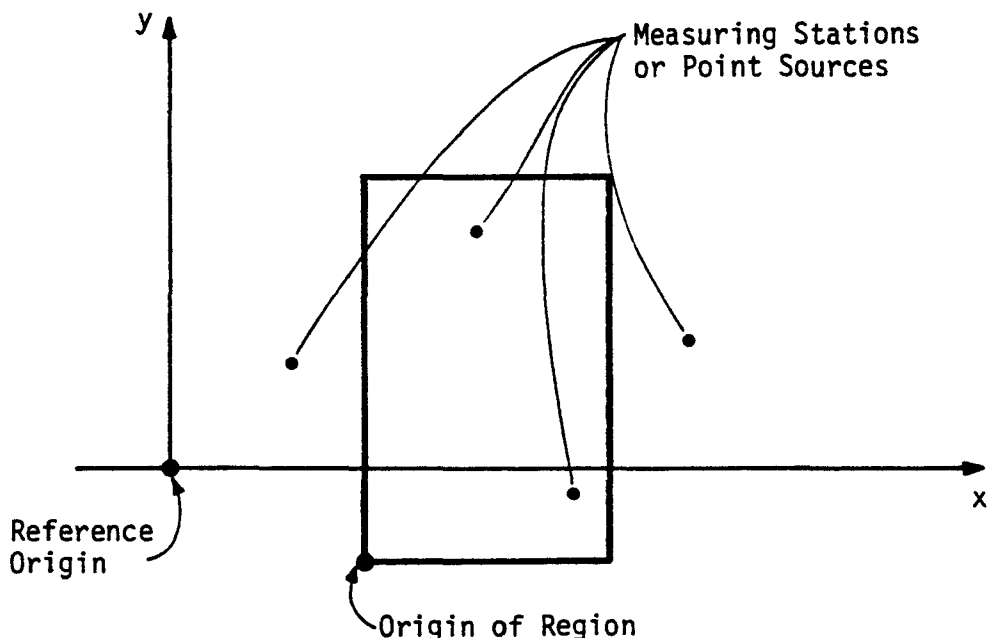


Figure IV-3. Example of the Location of a Modeling Region with Respect to a Reference Origin

It is sometimes necessary to describe the region in terms of its grid cells, generally when dealing with a whole cell (containing spatially averaged values) rather than a single point. Grid cell references instead of point locations are used to designate subregions and regional segments for the data preparation programs and to label printouts of grid values. Grid cells are numbered by their indices in the x- and y-directions, with Cell (1,1) referring to the cell in the lower left (southwest) corner of the grid, as illustrated in Figure IV-4.

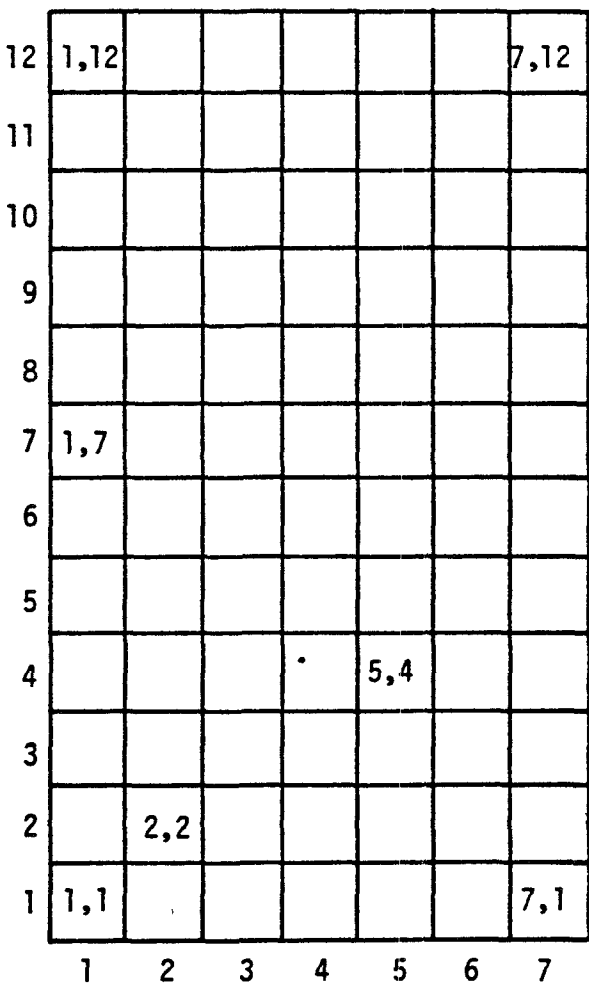


Figure IV-4. Conventional Method for Numbering Grid Cells

## 2. Vertical Distribution of Grid Cells

Although the modeling region in the horizontal plane consists of rectangular cells of equal size, cells in the vertical direction are distributed differently. The vertical distribution of cells is described using the following definitions:

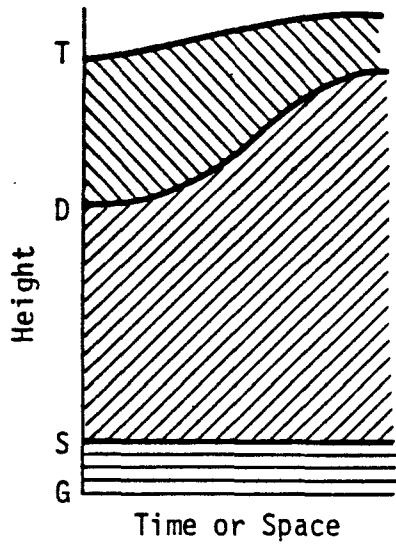
- > The diffusion break is the height above ground at which the diffusion characteristics (or eddy diffusivity) undergo a significant change. For example, when a mixed layer is capped by a stable inversion layer, the diffusion break occurs at the base of the inversion. If a ground-based stable layer is present, the diffusion break then occurs at the top of the inversion layer. Gridded values of the diffusion break are specified as a function of x, y, and time, and are input via the file DIFFBREAK.
- > The top of the region is the height above ground of the top of the simulation region. Gridded values are specified as a function of x, y, and time, and are contained in the file REGIONTOP. The top of the region may be located below, above, or at the same height as the diffusion break.
- > The space between the ground and the top of the region is divided into three layers:
  - The surface layer is one cell deep, and all surface layer cells maintain a constant height over space and time. Microscale calculations are performed in this layer using concentration predictions from the corresponding cell immediately above each surface layer cell. Ground-level emissions, surface deposition, and the reaction products of NO emissions from roadways and ambient ozone are combined to produce a flux of each pollutant into the cell above the surface layer cell. If the surface layer height is set to

zero, there is no surface layer and the microscale calculations are bypassed. Since the microscale prediction capabilities have not been fully tested, it is recommended that the surface layer height be set to zero.

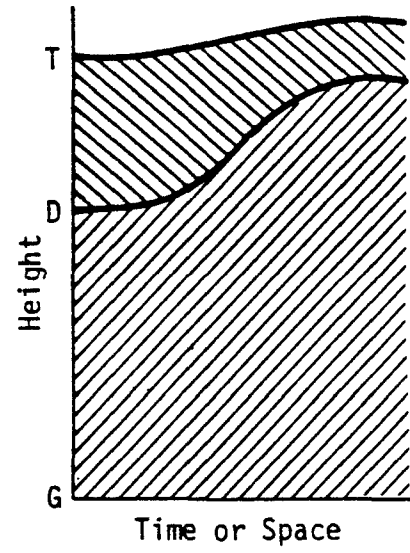
- The lower layer, which must be at least one cell deep, includes the cells between the top of the surface layer (or those at ground level, if there is no surface layer) and the diffusion break.
- The upper layer includes the cells between the diffusion break and the top of the region. Calculations in this layer can be omitted by setting its number of cells to zero.

Figure IV-5 illustrates various "layer" configurations that can be employed in the model. Figure IV-5(a) shows surface, lower, and upper layers. Figure IV-5(b) omits the surface layer. Figure IV-5(c) contains surface and lower layers, but no upper layer; note that, in this case, the top of the region actually modeled is the diffusion break, not the region top parameter input. The user must specify the number of cells and the minimum cell height in the lower and upper layers. Figure IV-5(d) shows the region described in Figure IV-5(a) with two cells in the lower layer and two cells in the upper layer. The number of levels in the region, i.e., the number of cells in the vertical dimension of the concentration matrix, is equal to the number of cells in the lower layer plus the number of cells in the upper layer plus one, if there is a surface layer. The total number of levels is used to calculate the segment size limitation. For example, five levels are described in Figure IV-5(d).

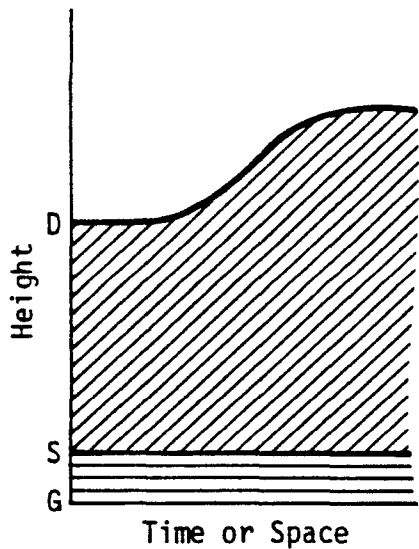
If either the lower or upper layer cells (as defined above) are smaller than the minimum cell height specified, enough space is borrowed from the adjacent layer to make up that height. Such a situation occurs in cases in which the diffusion break is very low or it is higher than the top of the region. Figure IV-5(e) illustrates all possible vertical cell distributions for a region that embodies a surface layer, two cells in the lower layer, and two cells in the upper layer. The top of the region is assumed to be high enough to include the total number of cells



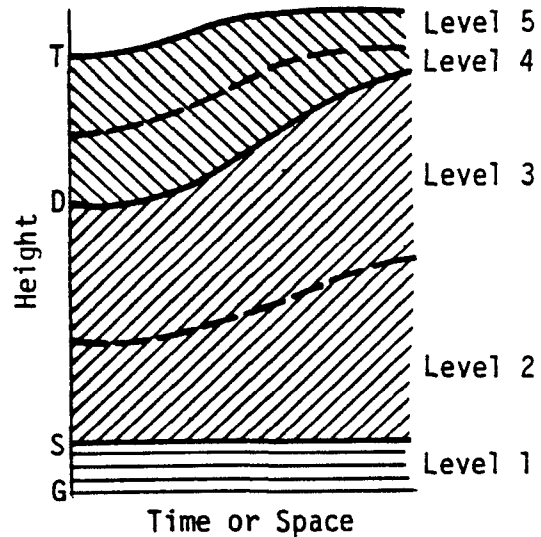
(a) Configuration of a Region into Surface, Lower, and Upper Layers



(b) Configuration of a Region into Lower and Upper Layers

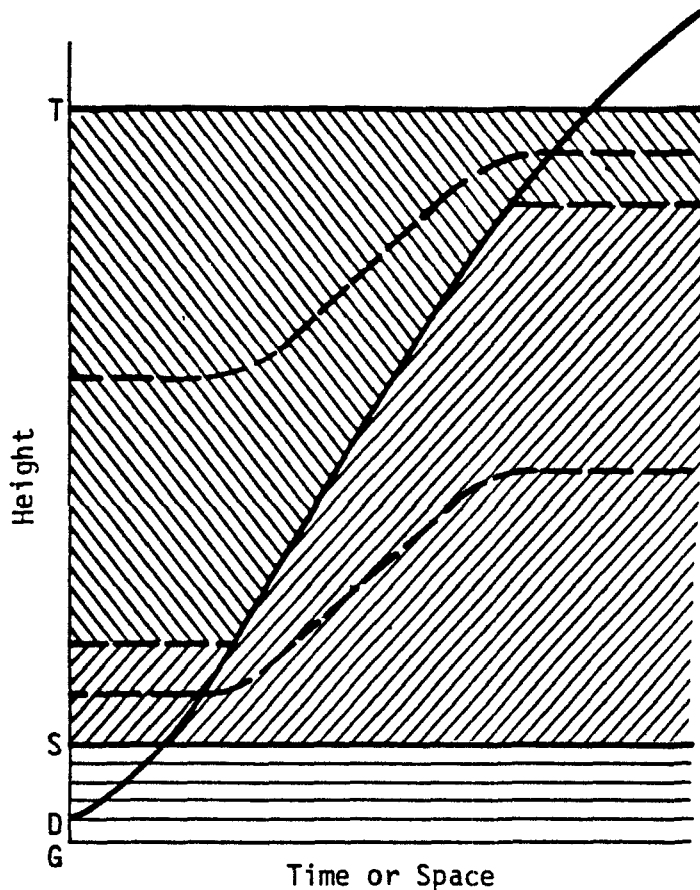


(c) Configuration of a Region into Surface and Lower Layers



(d) The Region Described in Figure IV-5(a) Sectioned into Five Levels

Figure IV-5. Various Vertical Cell Distributions That Can Be Employed in the SAI Airshed Model



- (e) All Possibilities for Vertical Cell Distribution  
of a Region That Contains a Surface Layer and Two  
Cells Each in the Upper and Lower Layers

**G** = Ground.

**D** = Diffusion break.

**S** = Height of the surface layer.

**T** = Top of the region.

===== Surface layer.

/ / / / Lower layer.

\ \ \ \ Upper layer.

Figure IV-5 (Concluded)

at the minimum height. For cases in which the minimum cell height is used to create a pseudo-lower layer or a pseudo-upper layer, the simulation program still uses the actual height of a cell with respect to the diffusion break rather than the cell number to calculate the vertical diffusivity. Instructions for specifying the vertical distribution of cells in the region are given in Section VI.B.2, which discusses the REGION packet.

#### B. CHEMICAL SPECIES

Each cell in the three-dimensional region previously described will carry a concentration value for each chemical species simulated. The Chemistry Parameters (CHEMPARAM) file indicates the species to be considered and their properties (Section IX.A). Initial concentrations of each pollutant are provided by the Air Quality (AIRQUALITY) file. The vertical column of concentrations at each boundary cell of the horizontal plane are provided by the Boundary (BOUNDARY) file. Boundary concentrations for each species at the top of the region are on the Top Concentrations (TOPCONC) file. Instructions for creating these three files are given in Chapter VII.

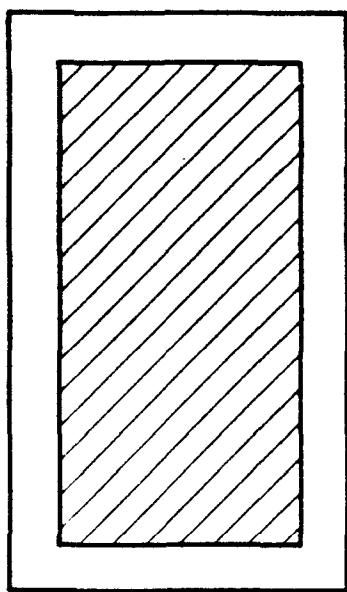
#### C. TIME

Concentrations of each species in each cell are moved ahead from a beginning time to an ending time in increments called "time slices". The beginning time, ending time, and the size of time slice are specified on the Simulation Controls (SIMCONTROL) file (Section IX.B). The time intervals for concentration averaging and for the output of the four-dimensional concentration fields are also specified on this file. A further discussion of the treatment of time is given in Section E of this chapter.

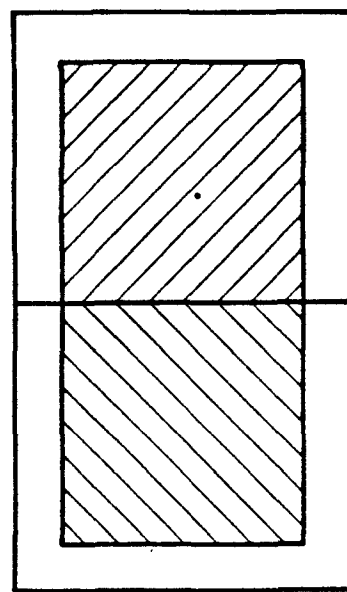
#### D. SEGMENTATION

The segmentation process involves dividing the user-defined modeling region into one or more rectangular segments in the horizontal plane, thereby allowing the Airshed Simulation Program to retain in core the data for the entire area it is processing, and providing flexibility as to the choice of the number and size of grid cells. The maximum horizontal dimensions for each segment are determined by the amount of core storage available on a particular computer, the number of vertical levels and chemical species to be simulated, and the simulation options to be used. Although the data arrays for all segments are maintained in secondary storage, all of the data arrays for a single segment reside in core during the period that the segment is being processed.

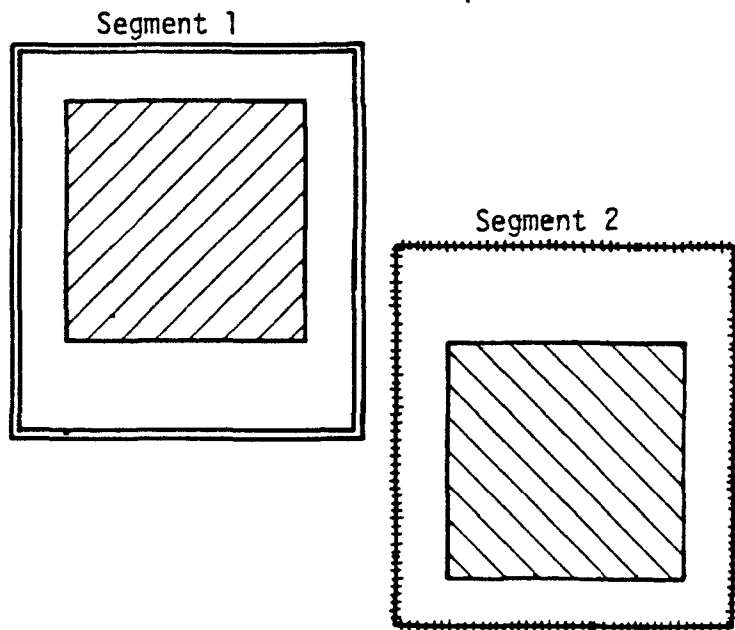
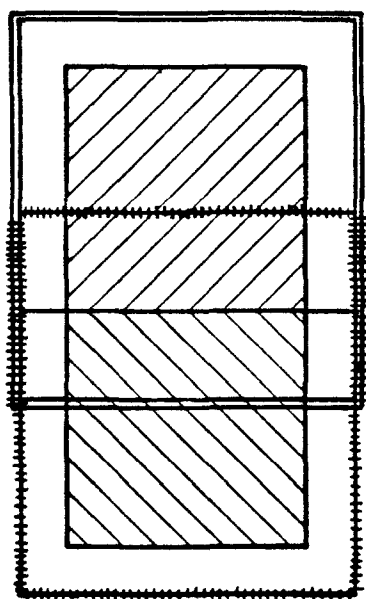
Each segment must be constructed as if it were an independent region being simulated, with its own set of input data and boundary conditions. To provide adequate information at the segment interfaces, adjacent segments must be constructed with a four cell overlap, as illustrated in Figures IV-6(a) through IV-6(c). Figure IV-6(a) shows a 7 x 12 cell region, consisting of a 5 x 10 area to be simulated plus a one-cell boundary on all edges. If the data covering this area do not fit in core, the area can be divided into two segments as shown in Figure IV-6(b). However, since each segment must include two boundary cells from the adjacent segment, the length of each segment is increased by two cells at the interface boundary. Figure IV-6(c) shows the layout of the original region, segmented into two 7 x 8 regions, each with a 5 x 5 simulation area. The two southernmost simulated rows of Segment 1 are the north boundary cells for Segment 2, and the two northernmost simulated rows of Segment 2 are the south boundary cells of Segment 1. Instructions for calculating maximum segment size and defining the regional segment layout are given in Chapter VIII.



(a) A  $5 \times 10$  Cell Region  
with a One-Cell Boundary  
on All Edges



(b) Division of the  $5 \times 10$   
Cell Region into Two  
 $5 \times 5$  Cell Segments



(c) Layout of a Region Showing the Interface of Boundary Cells  
for Two Segments

Figure IV-6. Example of the Segmentation of a Region

## E. TIME SLICES AND INTEGRATION TIME STEPS

The "time slice" is the amount of simulation time during which the integration for any segment will proceed without updating the boundary conditions or any other time-varying data. At the beginning of a time slice, new data from the files are read if necessary, and the concentrations at the segment interfaces are updated (Chapter VII of the Systems Manual details the method of maintaining segment interface concentrations). Then, each segment in turn is entirely processed for one time slice. Because all data for a given segment are in core at once, this scheme reduces the number of I/O transfers.

To describe how the time slice is determined, another time increment--the integration time step--must be defined. The integration time step is the increment over which the numerical integration of the governing equations in a segment actually takes place. The numerical integration procedure consists of four sequential steps:

- > Horizontal advection and diffusion (Steps 1 and 2).
- > Vertical advection and diffusion, including the treatment of emissions, surface deposition, and the surface layer microscale effects (Step 3).
- > Chemistry (Step 4).

The maximum time step allowable for the horizontal transport calculations in a segment is determined by the simulation program on the basis of the maximum wind velocity component over the segment and by the horizontal grid cell size:

$$\Delta t_{\max} = \min\left(\frac{.5\Delta x}{|u|_{\max}}, \frac{.5\Delta y}{|v|_{\max}}\right),$$

where

$\Delta t_{\max}$  = maximum allowable time step for the segment,  
 $\Delta x$  = size of the grid cell in the x-direction,  
 $\Delta y$  = size of the grid cell in the y-direction,  
 $|u|_{\max}$  = maximum absolute value of the wind velocity component in the x-direction over the entire segment for all vertical levels,  
 $|v|_{\max}$  = maximum absolute value of the wind velocity component in the y-direction over the entire segment for all vertical levels.

The size of the time slice is determined in the following way:

- > First, the time slice is set to the maximum size input by the user on the SIMCONTROL file.
- > If necessary, the time slice is reduced to avoid exceeding the time interval for which the data on any input file apply.
- > Also, if necessary, it is further reduced to avoid exceeding the next output time (instantaneous or average).
- > Finally, it is reduced if necessary on the basis of the smallest maximum time step that is allowed in any segment.\*

---

\* The user sets the maximum number of time steps that can be taken in any slice. Thus, the time slice used can be no larger than

$$n(\Delta t_{\max})_{\min} \quad ,$$

where  $n$  = maximum number of steps in a slice (input on the SIMCONTROL file), and  $(\Delta t_{\max})_{\min}$  = the smallest maximum allowable time step calculated for any segment. In general, if  $n$  is set large, the time slice used will be limited by the maximum time slice input. If  $n$  is small, the time slice used will be limited by the smallest allowable time step for any segment.

After the size of the time slice has been determined, the overall integration time step to be used for each segment is calculated. If the maximum time step for the segment is shorter than the calculated time slice, the slice is divided into the smallest number of equal time steps such that each is less than or equal to the maximum time step. Thus, if the time slice is 0.25 hour, and the maximum time step is 0.10 hour, the time slice is divided into three equal time steps of 0.08 1/3 hours each. If the maximum time step is greater than the time slice, the overall time step used will be equal to the time slice.

The overall time step described above (limited by the maximum wind speed and time slice size) may still be too large to allow completion of the chemistry step (Step 4). If the time step used for advection is too large to allow convergence in the chemistry calculations for any given cell, the time step is cut in half only for the chemistry step for that cell. This "chemistry time step" is successively halved until the chemistry calculations converge, and a sufficient number of these steps are taken for that cell to fill out the overall time step. If the chemistry time step for any cell must be decreased to less than the minimum chemistry time step specified by the user (on SIMCONTROL), the simulation will stop and values of all arrays will be printed in the neighborhood of the cell.

## V JOB CONTROL SPECIFICATIONS

This manual does not present specific job control examples because they vary greatly depending not only on the computer system used but also on the intentions and the style of the individual user. Instead, Chapters VII through XI contain diagrams illustrating the flow of information (card decks, binary files, and printed output) among the various computer programs. This chapter discusses a typical information flow diagram, suggests a worksheet for keeping track of programs, and provides a summary of file characteristics to aid in the preparation of job control statements.

### A. INFORMATION FLOW DIAGRAMS

Each information flow diagram provides the following details, which are specified for each program being described:

- > The names of the files associated with the program
- > The FORTRAN unit number assigned to each file
- > The program name
- > Any necessary subroutine libraries
- > Any optional flow patterns.

Figure V-1 is an example of such a diagram. The input is a card deck read on Unit 5 (the standard read unit), and a binary file named BINARY, read on Unit 3. An optional input file--BIN2--if present, is read on Unit 2. The program to be executed--PROGM--requires subroutines that are stored in SUBLIB. The output from the program is another binary file called BINOUT, written on Unit 4, and a printout of results on Unit 6, the standard print unit.

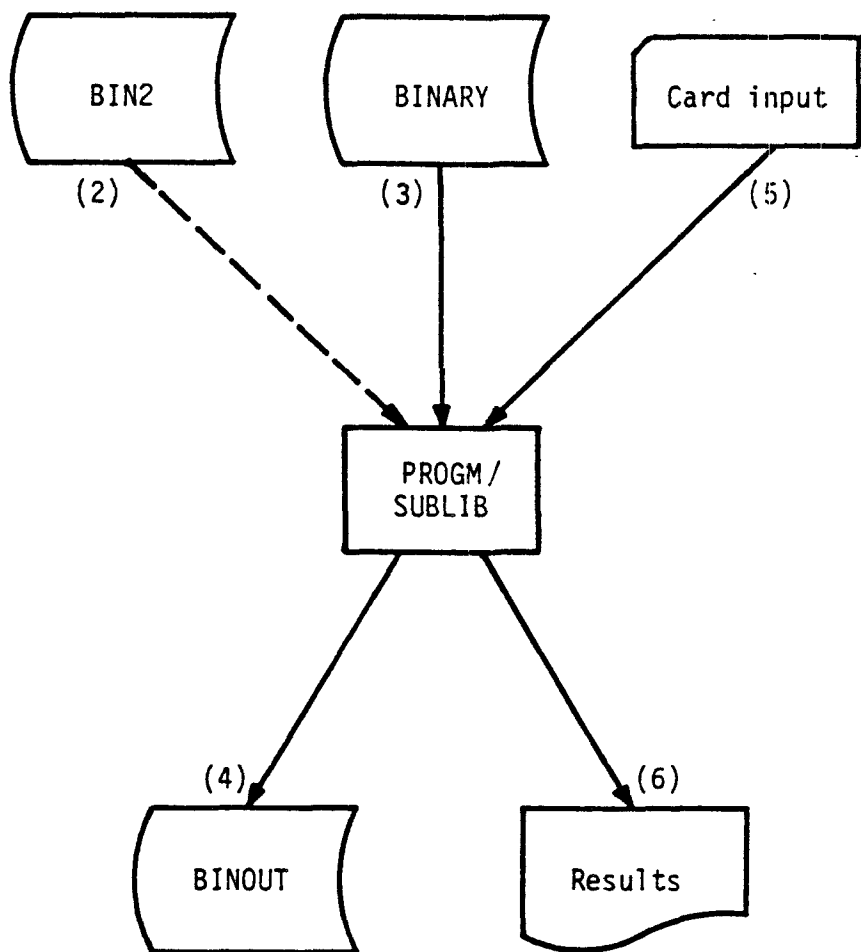


Figure V-1. Example of an Information Flow Diagram  
for Use in Job Control

## B. PROGRAM WORKSHEET

This manual assumes that the programs have already been compiled and that they exist in an executable form. Exhibit V-1(a) is a worksheet that can be used to locate and identify each program in the system; Exhibit V-1(b) shows that worksheet filled out for use on the Lawrence Berkeley Laboratory (LBL) system. To use a different system, users who implement the programs must provide the information necessary to run them, including link/loading information, storage requirements, and the location of predefined procedures.

## C. FILE CHARACTERISTICS

The 17 data files associated with the SAI Airshed Model are described in detail in Chapter IX of the Systems Manual. This section simply tabulates the essential features of each file to aid in the preparation of job control statements.

With the exception of the card-image input files and the output print files, all files in the SAI Airshed Model are binary (unformatted) files of variable record lengths. Each file contains one to four standard header records followed by data records specific to the particular file. The following terms are used to express the number of records on a file and the maximum record length:

sg = number of segments,  
sp = number of species,  
 $x_{max}$  = largest number of cells in the x or y dimension  
for any segment,  
 $xy_{max}$  = largest number of grid cells in the horizontal  
plane for any segment,  
z = number of vertical levels,  
 $pt_{max}$  = largest number of point sources in any segment.

Program	Source	Object	Libraries	Scratch Vector	Size	Other Information
AIRQUL						
BNDARY						
DIFFBK						
EMISSN						
METSCL						
PTSRC						
REGINTP						
ROADWY						
TMPTR						
TERAIN						
TOPCON						
WNDSET						
UTILITY						
FILUTIL						
SEGDEF						
CPREP						
SPREP						
AIRSIM						
DISPLAY						

(a) Format

Exhibit V-1. Sample SAI Airshed Model Program Worksheet for Use in Job Control

Program	Source	Object	Libraries	Scratch Vector	Size	Other Information
AIRQL	AIRQLU	AIRQLAL	i, F	14,500	170,000 <sub>s</sub>	
BNDARY	BNDARYU	BOUNDARY	(	17,000		
DIFFBK	DIFFBKU	DIFFBK		26,500		
EMISSN	EMISSNU	EMISSION		20,000		
METSCL	METSCLU	METSCAL		40,000		
PTSRCE	PTSRCEU	PSOURCE		23,500		
REGNTPL	REGNTPU	REGNTPO		22,500		
ROADWY	ROADWYU	ROADWAY		20,000		
TMPPTR	TMPPTRU	TMPPTR		26,500		
TERAIN	TERAINU	TERRAIN		27,000		
TOPCON	TOPCONU	TOPCON		20,000		
WNDSET	WNDSETU	WNDSSET		11,900		
UTILITY	UTLTYU	UTILITY		-	-	
FILUTL	FILUTLU	FILUTL		-	-	
SEGDEF	SEGDEFU	SEGDEF		1		
CPRP	JCPREP	JCPREP		32,000 <sub>s</sub>		
SPREP	JSPREP	JSPREP		27,000 <sub>s</sub>		
AIRSIM	JTSIMU	JTSIM		30,000	170,000 <sub>s</sub>	Overlay form -- requires AUTODAY control deck
DISPLAY	DISPLAY	DISPLAY		-	134000 <sub>s</sub>	

(b) Worksheet Filled Out for Use on the LBL System

Exhibit V-1 (Concluded)

Table V-1 shows the number of words in each of the four header records. Table V-2 lists the 17 files, indicates the number of records appearing on each one, and gives the maximum length of a nonheader record. If this length is smaller for the largest segment than that of the longest header record, then the longest header determines the maximum record length for the whole file. In addition to the 17 data files shown in Table V-2, some temporary scratch files might be used by a particular program, depending on the implementation and the size of the region to be modeled. Chapter IX of the Systems Manual explicitly defines these files.

Table V-1. Header Record Lengths for the  
SAI Airshed Model Data Files

<u>Record Name</u>	<u>Length (decimal words)</u>
File description header record	76
Region description header record	15
Segment description header record	4 sg
Species description header record	10 sp

Table V-2. Characteristics of the SAI Airshed Model Data Files

Name of File	Header Records				Number of Time-Invariant Records	Number of Records per time Interval	Maximum Data Record Length (decimal words)
	File	Region	Segment	Species			
AIRQUALITY	✓	✓	✓	✓	--	1 + sg*sp*z	11 + xy <sub>max</sub>
BOUNDARY	✓	✓	✓	✓	4sg	1 + 4*sg*sp	12 + x <sub>max</sub> *z
DIFFBREAK	✓	✓	✓		--	1 + sg	11 + xy <sub>max</sub>
EMISSIONS	✓	✓	✓	✓	--	1 + sg*sp	11 + xy <sub>max</sub>
METSCALARS	✓	✓			--	2	66
PTSOURCE	✓	✓	✓	✓	2sg	1 + sg*(2 + sp)	6*pt 11 + pt
REGIONTOP	✓	✓	✓		--	1 + sg	11 + xy <sub>max</sub>
ROADWAY						1 + sg*(1 + sp)	11 + xy <sub>max</sub>
TEMPERATUR	✓	✓	✓		--	1 + sg	11 + xy <sub>max</sub>
TERRAIN	✓	✓	✓		2sg	--	11 + xy <sub>max</sub>
TOPCONC	✓	✓	✓	✓	--	1 + sg*sp	11 + xy <sub>max</sub>
WIND	✓	✓	✓		--	1 + 2sg*z	11 + xy <sub>max</sub>
CHEMPARAM	✓			✓	2-4	--	62 18 sp 5*reactions 11*coefficients
SIMCONTROL	✓				1	--	36
AVERAGE	✓	✓	✓	✓	--	1 + sg*sp*z	11 + xy <sub>max</sub>
INSTANT	✓	✓	✓	✓	--	1 + sg*sp*z	11 + xy <sub>max</sub>
VARIATION	✓	✓	✓	✓	--	1 + sg*sp	11 + xy <sub>max</sub>

## VI PACKETS USED IN CREATING THE M.E.B. FILES

As many as 12 different data preparation programs are required for constructing the M.E.B. files--one program for each file. This chapter discusses the data modules--packets--that are used by the data preparation programs. All input data are divided into packets; each packet contains a specific type of information necessary to some aspect of data preparation. The following sections first present general rules that apply to the preparation of all the packets, and then explicit rules and formats for each one. Chapter VII contains rules for the preparation of the data files using these packets, and examples of the output from the programs.

### A. ELEMENTS OF PACKET CONSTRUCTION

The ordering and internal structure of the data packets have been designed for consistency of formats, flexibility of use, and ease of visual analysis. Each packet begins with a header card identifying the packet and ends with a termination/trailer card that reads END or ENDTIME. The packet names, their classifications, and their proper ordering are shown in Table VI-1.

Of the time-invariant data, the CONTROL and REGION packets are mandatory, and they are entered first and second, respectively, in the input deck. The CONTROL packet defines input and output options and maximum variable counters used by the program to set internal array dimensions. The REGION packet defines the location, size, and resolution of the modeling region. If present, the UNITS packet follows the REGION packet. The UNITS packet names user-defined variables and specifies unit conversions.

**Table VI-1. Packets Used by the SAI Airshed Model  
Data Preparation Programs**

Time-Invariant Packets

CONTROL  
REGION  
UNITS  
STATIONS  
POINT SOURCES  
BOUNDARIES

Time-Varying Packets

TIME INTERVAL  
SUBREGIONS  
METHOD  
VERTICAL METHOD  
CONSTANTS  
GRID VALUES  
STATION READINGS  
EMISSIONS VALUES  
EMISSIONS FACTORS  
BOUNDARY READINGS  
SCALARS  
VERTICAL PROFILES

The remaining time-invariant packets, STATIONS, BOUNDARIES, and POINT SOURCES, which define fixed locations in the region, are optional depending on the file being created and the method used.

The TIME INTERVAL packets appear next. Each TIME INTERVAL packet contains the other time-varying packets to be used during the interval specified and ends with an ENDTIME card. The time intervals used must cover the time span specified in CONTROL with no gaps or overlaps. The time-varying packets included within each TIME INTERVAL packet define the data preparation methods to be used and supply the time-varying data.

Different interpolation methods can be used for different variables in different areas of the region. The SUBREGION packet defines the areas, and the METHOD packet defines the method to be used for each variable in each subregion. The VERTICAL METHOD packet describes the method to be used for calculating values for variables that vary vertically. Although any of these packets can be changed in subsequent time intervals, for most runs the subregions and methods supplied in the first time interval will be used for the entire duration of the run. The other time-varying packets define the values for the input variables. The particular packets used for any given run depend on the file being created and the methods selected. After the first TIME INTERVAL packet has defined the information to be written on the file, this information will persist into succeeding time intervals as described in Section A.3 of this chapter.

## 1. Rules for Card Format

Each data card is divided into two sections: Columns 1 to 60 contain input data; Columns 61 to 80 are reserved for any desired user identification information. All input fields are 10 columns wide except packet headers and file identifiers, which can occupy the entire width available for input, i.e., Columns 1 to 60.

Integers are input in Format I10 and must be right justified. Floating point variables are input as F10.0. Alphanumeric information can occupy any of the columns available for input of that data as long as it is correctly ordered and no extraneous or erroneous symbols are included. Thus, for example, \_C\_ON\_TROL\_\_ would be recognized as "CONTROL"; CNOTROL and CONTROLZ would not. The 60-column file identifier is not subject to validation and, therefore, can contain any information the user desires.

## 2. The Reserved Word ALL

The word ALL is a special designation that is reserved to enable the user to specify information globally; therefore, it should not be used to indicate a specific alphanumeric entity, such as a variable, subregion, station, point source, or boundary name. When used in the input field usually associated with an alphanumeric name, ALL means that the subsequent information applies to all the names that could occupy that input field. ALL can generally be used wherever it makes sense. The following examples illustrate both the proper and improper usages of this command:

- > Proper usages of ALL.
  - A card in the METHOD packet:

```
|ALL -----|_WX_____|GRIDVALUE| .
```

This card specifies that in all subregions the values of the variable WX will be input by the GRID VALUE method. If ALL also occurred in place of WX, e.g.,

```
|ALL-----|ALL-----|GRIDVALUE| .
```

the values of all input variables in all subregions would also be input by the GRID VALUE method.

- A card in the STATION READINGS packet:

```
|WEST_____|_ALL_____|-9.0_____| .
```

This information indicates that the values of all variables reported by Station WEST are to be set to -9.0 (a value that implies data are missing).

- > Improper usage of ALL. A card in the UNITS packet:

```
|ALL_____|KG/D_____|a           |b           |c           | ,
```

where a, b, and c are numbers. This card says that all input variables will have values input in kg/day, to which the conversion parameters a and b and the molecular weight c are to be applied. It is improper because different species have different molecular weights.

### 3. Persistence of Data

As previously mentioned, the information provided in any TIME INTERVAL packet remains in effect until it is replaced. This persistence rule applies to all time-varying data, including subregion and method definitions and input values. For example, METHODS for computing each variable in each subregion need be specified only in the first TIME INTERVAL. They can be changed in a later TIME INTERVAL if desired, but in most cases only one definition is required.

If no data are available for a given station over a certain time interval, the user can omit that station in that time interval, and the data previously input will be used. However, if the data are actually to be treated as missing, the station must be included and must be assigned values that are interpreted by the program as "missing data." In general,

missing integer and real values are represented by -9 and -9.0, respectively. Although the use of this device can save considerable duplication of effort, it places on the user the burden of ensuring that all changes in status are specified.

#### 4. Units of Measure

The Airshed Simulation Program assumes a standard and consistent set of units for all of its computations, and the files input to the program must contain data expressed in these "internal" units. The data input to the file preparation programs, however, may be in other than internal units. To accommodate such data, the programs contain a set of standard alternative unit designations for use in directing automatic conversion of input units to internal units. Additionally, the user can specify the use of nonstandard conversion factors. Section B.3 of this chapter discusses the internal units for each variable, alternative units and their associated conversion factors, and the method for specifying non-standard unit conversions.

#### 5. Variables

Each program has a set of output variables. For files that vary by species, the output variables are the species names listed in the CONTROL packet. For files that do not vary by species, the output variables are names built into the corresponding data preparation program; the implicit output variable names for each file are presented in the appropriate section of Chapter VII. Values input to the program may be for the output variables themselves, or they may be for variables that will be acted on in some way to produce the output variables. For example, concentrations of total hydrocarbon may be input to the AIRQUALITY program, whereas its output consists of concentrations in each of several carbon-bond classes.

Whenever the input variables are different from the output species or implicit output variables, they are referred to as "user-defined variables," and must be named in the UNITS packet. Designation of the user-defined variables in this way allows the program to allocate space for them internally.

## 6. Methods

This section explains the methods for calculating the ground-level values of each variable within each subregion. (Section B.8 of this chapter gives instructions on dividing the modeling region into subregions.) These methods are designated in the METHOD packet, which is discussed in Section B.9 of this chapter. The first eight methods (a through h) can be used by any of several data preparation programs; the remaining methods (i through r) are each specific to a single program.

### a. CONSTANT

A single value is to be used for this variable in every ground-level cell in this subregion. The value must be defined in a CONSTANTS packet and the program will convert it to internal units if necessary. The CONSTANT method requires no other parameters.

### b. GRID VALUE

A value for this variable will be input for each ground-level grid cell in the subregion. The values will be defined in a GRID VALUES packet and the program will convert them to internal units if necessary. The GRID VALUE method requires no other parameters.

c. STATINTERP (Station Interpolation)

The value for this variable at each ground-level grid cell in the subregion is calculated by the program as the weighted average of values at selected measuring stations. Station locations will be defined in a STATIONS packet; values for the variable at each station will be input in the STATION READINGS packet and the program will convert them to internal units if necessary. To calculate the value at a given grid cell, the program weights each station value by the inverse of the distance of the station from the center of the cell. The following four parameters must be specified to control the selection of the measuring stations to be included in the average:

- > EXTENT--This number determines the acceptability of a station on the basis of the subregion it is in. If EXTENT = 0.0, a station within the radius of influence will be included in the average regardless of the subregion it occupies. If EXTENT ≠ 0.0, only stations within the same subregion will be accepted. If this number is omitted, it will be assumed to be zero.
- > INITRADIUS--initial radius of influence. All stations within this distance of the cell for which values are being calculated will be used. This number is assumed to be in grid units. If this number is set very large, all stations will be included. Omission of this number constitutes an error.
- > RADIUSINCR--If no measuring stations with values are encountered within the initial radius of influence, the radius will be incremented by this amount until at least one station is included. This number is assumed to be

in grid units. When this number is small, the values generated will be distributed more smoothly over the region, but the cost in computing time could be great. Conversely, when this number is large, computing time might be reduced, but at the expense of irregularities in the computed grid values. If this number is omitted, only the initial radius will be used, and the failure to find a valid station constitutes an error.

- > MAXRADIUS--maximum radius of influence. This number is assumed to be in grid units. Failure to find valid station data within this radius constitutes an error. If this number is omitted, the entire modeling region may be used.

d. POISSON

The value for this variable at each ground level grid cell in the sub-region is calculated by the program using the Poisson smoothing method [see Killus et al. (1977), Chapter IV, pp. 135-142]. Values for the variable at selected measuring stations must be input in the STATION READINGS packet. The POISSON method requires three parameters:

- > MAXITER--the maximum number of iterations; the suggested number is ≤ 200.
- > ERRORTOL--error tolerance. This parameter is expressed in internal units of the variable; the suggested value =  $0.01 * \text{expected value}$ .
- > OMEGA--weighting factor to aid convergence; the suggested value = 1.4.

e. SPLIT/COMB (Split or Combine)

Any variable to be output can be formed by splitting or combining input variables (typically, these are species):

$$\text{var}_{\text{out}} = \sum_{i=1}^N \text{var}_i * \text{factor}_i$$

This method requires N parameter cards; on Card i the parameter name is the name of variable<sub>i</sub>, and the value is factor<sub>i</sub>. If the parameter (variable) name is left blank, the corresponding factor is treated as a constant (i.e., var<sub>i</sub> = 1). All values for input variables will already have been converted to internal units when this computation is done, so the factors specified should not include unit conversions.

f. E-W INTERP (East-West Interpolation)

For each row of grid cells within a subregion, a linear interpolation will be carried out between values in the bordering cells in the east and west edges of the row. This subregion must not lie on an edge (i.e., it must be bounded on east and west by other subregions) and values for the bordering subregions must be calculated by a noninterpolative method. The E-W INTERP method requires no other parameters.

g. N-S INTERP (North-South Interpolation)

For each column of grid cells within a subregion, a linear interpolation will be carried out between values in the bordering cells on the north and south edges of the column. This subregion must not lie on an edge (i.e., it must be bounded on north and south by other subregions), and values for the bordering subregions must be calculated by a noninterpolative method. The N-S INTERP method requires no other parameters.

h. USER (User-Supplied Algorithm)

Each data preparation program is designed to call a user-supplied subroutine for any variable for which the method USER has been selected. All available data are passed to the subroutine as arguments. At present, all user subroutines are dummies; as new methods are developed, they can be inserted in user subroutines, and parameter values can be read and passed as for any of the standard methods.

i. BOUNDVALUE

This method is used only for creation of the BOUNDARY file. It specifies that concentration values will be input for each boundary line segment through the BOUNDARY READINGS packet. BOUNDVALUE requires no parameters.

The following two methods are used only for creating the PTSOURCE file.

j. EMVALUES

This method name indicates that point source emissions will be entered for this species for this point source type (the "subregion" field) by means of the EMISSIONS VALUES packet. EMVALUES requires no parameters.

k. EMFACTORS

This method name indicates that for this species and point source type, the emissions values and flow rate previously entered will be modified by factors supplied by means of an EMISSIONS FACTORS packet. This method requires no parameters.

The following two methods are used only for the REGIONTOP file. Both require that the DIFFBREAK file be input and that the vertical definition of the region be included in the REGION packet. Both use NZUPPR (the number of cells in the upper layer) as defined in the REGION packet.

l. FIXDHEIGHT

The REGIONTOP is defined such that there will be NZUPPR cells of a fixed height above the diffusion break, subject to the maximum height indicated on the method card. REGIONTOP can be defined as being equal to the diffusion break, subject to the maximum height indicated on the method card, by specifying NZUPPR = 0. This method requires one parameter: UPCELLHT-- the cell height in the upper layer (above the diffusion break). If NZUPPR > 0, this number must be specified and must be greater than the minimum height of upper layer cells.

m. SAMEHEIGHT

The REGIONTOP is defined such that there will be NZUPPR cells above the diffusion break of the same height as the NZLWR cells between the top of the surface layer and the diffusion break, subject to the maximum height indicated on the method card. No other parameters are required.

The following two methods are used only for the TOPCONC file. Both require that vertical concentration profiles be input by means of a VERTICAL PROFILES packet; i.e., they require that for each species in each subregion, a number of height-concentration pairs be provided. Both methods require the REGIONTOP file. The method RELTOPCONC requires the DIFFBREAK file as well.

n. ABSTOPCONC

The concentration at the top of the region will be calculated from the profile based on the height of the top of the region. The vertical profile

input describes the vertical distribution of concentration values. The height of the top of the region above ground is compared with the height of each profile point to determine the value to be used. The profile is input as a set of pairs  $(H_i, F_i)$ , where  $H$  is height and  $F$  is some profile value. Since it is assumed that the pair  $(H_1, F_1)$  corresponds to ground level, the following transformation is applied to all heights:

$$H'_i = H_i - H_1 \quad .$$

The profile is thus considered to be the set of pairs  $(H'_i, F_i)$  ordered by increasing  $H'$ .

The profile is used for a particular grid cell in the following way: For a cell at a given  $(x,y)$  location on the grid, find the height  $T$  of the top of the region. Then, the concentration,  $C$ , at the top is defined as follows:

$$C = \begin{cases} F_n, & \text{if } T > H'_n ; \\ \cdot & \\ F_{i-1} + \frac{T - H'_{i-1}}{H'_i - H'_{i-1}}(F_i - F_{i-1}), & \text{if } H'_{i-1} \leq T \leq H'_i \quad . \end{cases}$$

#### o. RELTOPCONC

The concentration at the top of the region will be calculated from the profile based on the height of the top of the region relative to the height of the diffusion break. The vertical profile input describes the vertical distribution of concentration values. The height of the top of the region relative to the height of the diffusion break is compared to the height of each profile point relative to its diffusion break to determine the value to be used. The profile is input as a set of pairs  $(H_i, F_i)$ , where  $H$  is height and  $F$  is some profile value. Since it is assumed that the pair  $(H_1, F_1)$  corresponds to ground level, the following transformation is applied to all pairs: For all  $i$ ,

$$H'_i = (H_i - H_1)/DB_p \quad ,$$

where  $DB_p$  = diffusion break at the profile location. The profile is thus considered to be the set of pairs  $(H'_i, F_i)$  ordered by increasing  $H'$ .

The profile is used for a particular grid cell in the following way. For a cell at a given  $(x,y)$  location on the grid, find the height of the top of the region. Then, convert the absolute height  $T$  to the height of the top relative to the diffusion break at that location:

$$T' = \frac{T}{DB} \quad .$$

Then, the concentration,  $C$ , at the top is defined as follows:

$$C = \begin{cases} F_n, & \text{if } T' > H'_i \\ F_{i-1} + \frac{T' - H'_{i-1}}{H'_i - H'_{i-1}} (F_i - F_{i-1}), & \text{if } H'_{i-1} \leq T' \leq H'_i \end{cases} \quad .$$

The following three methods are used only in the WIND program. In this program there are four implicit variables: SPEED and DIRECTION, the variables ordinarily input to the program; and WINDX and WINDY, the  $x$ - and  $y$ -components of the wind that are written to the WIND file. Further discussion of the specification of these variables appears in Chapter VII.

#### p. DIVERGENCE

The DIVERGENCE method uses the algorithms developed by Anderson and Durran and reported in Killus et al. (1977) to calculate a three-dimensional wind field. The method is unique in that it calculates the values for the two output variables (WINDX and WINDY) simultaneously and computes the values for all cells at all levels. DIVERGENCE is, therefore, also a vertical method and must be explicitly named in the VERTICAL METHOD packet

as well. Use of this method requires that station readings and vertical profiles for SPEED and DIRECTION be specified and that the ground-level temperature field (TEMPERATUR data file) be available. Since this method internally performs a Poisson interpolation, it requires the same three parameters as the POISSON method (previously discussed). These parameters should be specified for the first variable (WINDX or WINDY) that appears using the DIVERGENCE method. The other variable (WINDY or WINDX) should also use the DIVERGENCE method, but no parameter need be included. The vertical method for both of these variables should also be DIVERGENCE. This method cannot be specified for SPEED or DIRECTION. The DIVERGENCE method requires that the files DIFFBREAK, REGIONTOP, and TEMPERATUR be attached to the run.

The methods VECTORIZE and POLARIZE are used only to calculate the three-dimensional arrays. Necessary conversion of station values and profile values are handled automatically.

q. POLARIZE

This method can only be specified for the variables SPEED and DIRECTION. The values of SPEED and DIRECTION will be calculated from the vector components WINDX and WINDY. This method should be used if printouts of gridded SPEED and DIRECTION are desired. Some method other than VECTORIZE (discussed next) should be specified for WINDX and WINDY if POLARIZE is used. No parameters are required.

r. VECTORIZE

This method is used to convert SPEED and DIRECTION to WINDX and WINDY. This method should be used to calculate the output variables WINDX and WINDY if gridded SPEED and DIRECTION values are given or calculated by the program. POLARIZE should not be the method used for SPEED and DIRECTION if VECTORIZE is used for WINDX and WINDY.

## 7. Vertical Methods

The vertical methods are used to define values of the output variables at each vertical cell, given the values at ground level. Each method assumes that ground-level values already exist and are in internal units. These methods are designated in the VERTICAL METHOD packet, which is discussed in Section B.10 of this chapter. The first eight vertical methods (a through h) can be used by any of several data preparation programs; the remaining methods (i through k) are each specific to a single program.

### a. CONSTANT

Values in each vertical cell are equal to the ground-level value. This method requires no additional parameters.

There are four vertical methods that require the input of a vertical profile to describe the shape of the vertical distribution of values. These vertical methods require that the DIFFBREAK and REGIONTOP files be attached. For two of these methods, ABSPROFILE and RELPROFILE, the ground-level value only is used in calculating vertical distribution. For the other two, ABSPROFRAT and RELPROFRAT, the value for the variable at the top of the region is required in addition to the ground-level value, and the profile defines the shape of the interpolation between them. These two ratio methods also require the TOPCONC file.

Each of these general vertical profile methods can be used in either the "absolute" or "relative" mode. In the absolute mode, the heights provided with each profile point are directly used to calculate the variable value at a cell of a given height. That is, the absolute height of the cell above ground is used to determine the profile value. In the relative mode, the heights provided with each profile point are used as heights relative to the diffusion break or top of region at the specified location of the profile.

b. ABSPROFILE

The vertical profile input describes the shape of the vertical distribution of values using the ground-level value. The height of each cell above ground is compared with the height of each profile point to determine the scaling factor to be used. The profile is input as a set of pairs  $(H_i, F_i)$ , where  $H$  is height and  $F$  is some profile value. Since it is assumed that the pair  $(H_1, F_1)$  corresponds to ground level, the following transformation is applied to all pairs:

$$H'_i = H_i - H_1 \quad ,$$

$$F'_i = F_i / F_1 \quad .$$

The profile is thus considered to be the set of pairs  $(H'_i, F'_i)$  ordered by increasing  $H'$ .

The profile is used for a particular grid cell in the following way: For a cell at a given  $(x,y)$  location on the grid, find the average height  $(A_k)$  of each vertical cell ( $k$ ) as a function of the diffusion break and top of the region. If the ground-level value at that location is  $V_1$ , then

$$V_k = \text{factor}_k * V_1 \quad ,$$

where

$$\text{factor}_k = \begin{cases} F'_n, & \text{if } A_k > H'_n ; \\ F'_{i-1} + \frac{A_k - H'_{i-1}}{H'_i - H'_{i-1}} (F'_i - F'_{i-1}), & \text{if } H'_{i-1} \leq A_k \leq H'_i \end{cases} .$$

c. REPROFILE

The vertical profile input describes the shape of the vertical distribution of values using the ground-level value. The height of each cell relative to the height of the diffusion break is compared to the height of each profile point relative to its diffusion break to determine the scaling factor to be used. The profile is input as a set of pairs  $(H_i, F_i)$ , where  $H$  is height and  $F$  is some profile value. Since it is assumed that the pair  $(H_1, F_1)$  corresponds to ground level, the following transformation is applied to all pairs: For all  $i$ ,

$$H'_i = (H_i - H_1)/DB_p \quad ,$$

$$F'_i = F_i/F_1 \quad ,$$

where  $DB_p$  = diffusion break at profile location. The profile is thus considered to be the set of pairs  $(H'_i, F'_i)$  ordered by increasing  $H'$ .

The profile is used for a particular grid cell in the following way. For a cell at a given  $(x,y)$  location on the grid, find the average height  $A_k$  of each vertical cell  $k$  as a function of the diffusion break and top of the region. Then convert the absolute height  $A_k$  to the height of the cell relative to the diffusion break at that location:

$$A'_k = \frac{A_k}{DB} \quad .$$

If the ground-level value at that location is  $V_1$ , then

$$V_k = \text{factor}_k * V_1 \quad ,$$

where

$$\text{factor}_k = \begin{cases} F'_n, & \text{if } A'_k > H'_n \quad ; \\ F'_{i-1} + \frac{A'_k - H'_{i-1}}{H'_i - H'_{i-1}} (F'_i - F'_{i-1}), & \text{if } H'_{i-1} \leq A'_k \leq H'_i \quad . \end{cases}$$

d. ABSPROFRAT

The vertical profile input describes the shape of the vertical distribution of values expressed as the relative contribution (weighting) of the ground-level value and another value (which may be the value at the top of the region). The height of each cell above ground is compared with the height of each profile point to determine the contribution factor to be used. The profile is input as a set of pairs  $(H_i, F_i)$ , where  $H$  is height and  $F$  is a profile value  $0 \leq F \leq 1$ .  $F = 0$  means that the ground-level value only should be used.  $F = 1$  means that the other (top) value only should be used. (The exact formula is shown below.) Since it is assumed that the pair  $(H_1, F_1)$  corresponds to ground level, the following transformation is applied to all heights: For all  $i$ ,

$$H'_i = H_i - H_1$$

The profile is thus considered to be the set of pairs  $(H'_i, F_i)$  ordered by increasing  $H'$ .

The profile is used for a particular grid cell in the following way. For a cell at a given  $(x,y)$  location on the grid, find the average height  $A'_k$  of each vertical cell  $k$  as a function of the diffusion break and top of the region. If the ground-level value at that location is  $V_g$  and the other (top) value is  $V_t$ , then

$$V_k = V_g + \text{factor}_k (V_t - V_g) ,$$

where

$$\text{factor}_k = \begin{cases} 1.0 , & \text{if } A'_k > H'_n ; \\ F'_{i-1} + \frac{A'_k - H'_{i-1}}{H'_i - H'_{i-1}} (F'_i - F'_{i-1}) , & \text{if } H'_{i-1} \leq A'_k \leq H'_i . \end{cases}$$

e. RELPROFRAT

The vertical profile input describes the shape of the vertical distribution of values expressed as the relative contribution (weighting) of the ground-level value and another value (which could be the value at the top of the region). The height of each cell relative to the diffusion break is compared to the height of each profile point relative to its diffusion break in order to determine the contribution factor to be used. The profile is input as a set of pairs  $(H_i, F_i)$ , where  $H$  is height and  $F$  is a profile value,  $0 \leq F \leq 1$ .  $F = 0$  means that the ground-level value only should be used.  $F = 1$  means that the other (top) value only should be used. (The exact formula is presented below.) Since it is assumed that the pair  $(H_1, F_1)$  corresponds to ground level, the following transformation is applied to all heights: For all  $i$ ,

$$H'_i = (H_i - H_1)/DB_p ,$$

where  $DB_p$  = diffusion break at profile location. The profile is now considered to be the set of pairs  $(H'_i, F_i)$  ordered by increasing  $H'$ .

The profile is used for a particular grid cell in the following way. For a cell at a given  $(x,y)$  location on the grid, find the average height  $A_k$  of each vertical cell  $k$  as a function of the diffusion break and top of the region. Then convert the absolute height  $A_k'$  to the height of the cell relative to the diffusion break at that location:

$$A'_k = \frac{A_k}{DB} .$$

If the ground-level value at that location is  $V_g$  and the other (top) value is  $V_t$ , then

$$V_k = V_g + \text{factor}_k (V_t - V_g) ,$$

where

$$\text{factor}_k = \begin{cases} 1.0, & \text{if } A'_k > H'_n ; \\ F'_{i-1} + \frac{A'_k - H'_{i-1}}{H'_i - H'_{i-1}} (F'_i - F'_{i-1}), & \text{if } H'_{i-1} \leq A'_k \leq H'_i . \end{cases}$$

f. E-W INTERP (East-West Interpolation)

For each row a linear interpolation will be carried out between values in the bordering cells in the east and west edges of the row. This subregion must not lie on an edge (i.e., it must be bounded on east and west by other subregions), and values for the bordering subregions must be calculated by a noninterpolative method. The E-W INTERP method requires no other parameters.

g. N-S INTERP (North-South Interpolation)

For each column a linear interpolation will be carried out between values in the bordering cells on the north and south edges of the column. This subregion must not lie on an edge (i.e., it must be bounded on north and south by other subregions), and values for the bordering subregions must be calculated by a noninterpolative method. The N-S INTERP method requires no other parameters.

h. VERTUSER (User-Supplied Algorithm)

For each file containing vertically varying data, the file preparation program is designed to call a user-supplied subroutine for any variable for which the vertical method VERTUSER has been selected. All available data are passed to the subroutine as arguments. At present, all vertical user subroutines are dummies; as new methods are developed, they can be inserted in user subroutines, and parameter values can be read and passed as for any of the standard vertical methods.

To create the PTSOURCE file, a vertical method must be chosen for determining the height at which the emissions from each point source enter the modeling region. Two vertical methods have been provided: STACKHGT and PLUMERISE, described below. Each of these vertical methods requires that the DIFFBREAK and REGIONTOP files be attached to the run.

i. STACKHGT

The emissions enter the region in the vertical cell that contains the top of the stack. No additional parameters are required.

j. PLUMERISE

Plume rise is calculated by the Briggs formulae [see Briggs (1975)]. For this vertical method the TEMPERATUR, METSCALARS, and WIND files must also be attached to the run. PLUMERISE requires no other parameters.

k. DIVERGENCE

The vertical method DIVERGENCE is used only for the WIND file. If DIVERGENCE has been specified in the METHOD packet for WINDX and WINDY, it must also be specified in the VERTICAL METHOD packet for those variables. No other parameters are required.

## B. PACKET RULES AND FORMATS

This section presents the following information for each packet:

- > Instructions as to where the packet is used.
- > Special information about the contents.
- > A table outlining the definition of each field on the cards.
- > An exhibit showing the appearance (format) of the packet as a whole.

### 1. CONTROL Packet Rules

The CONTROL packet must always be the first packet input. The first three cards are the packet header, and the name and identifier of the file to be created. The next five cards contain counters for dimensioning arrays, and input and output option specifications; the standard entries are listed in Table VI-2. These control parameters are used in different combinations for different programs; each file preparation description in Chapter VII includes a table of the specific control parameters required for each file. For some files, some control parameters do not apply; these parameters are listed as "Spare" and any values are ignored. Following the control cards are those containing species names, if any, and the time span of the file. The contents of each card in the CONTROL packet are described in Table VI-3, and the format is shown in Exhibit VI-1.

**Table VI-2. Standard Entries for Cards 4 Through 8  
of the CONTROL Packet for M.E.B. Files**

<u>Card Number</u>	<u>Entries</u>
4	Number of species Number of user-defined variables Number of stations Number of subregions Number of parameters Spare
5	Output file number Print input cards Print output grid Spare Spare Spare
6	Print units table Print station locations table Print regional grid Print methods table Print station values table Spare
7	Number of vertical parameters Number of heights in profile Print vertical methods table Print vertical profile tables Spare Spare
8	DIFFBREAK file number REGIONTOP file number TOPCONC file number TEMPERATUR file number METSCALARS file number WIND file number

Table VI-3. CONTROL Packet for M.E.B. Files

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<b>1</b> <b>Packet Header</b>	Header      1-10	60A1	--	Must be CONTROL, left adjusted
<b>2</b> <b>File Name</b>	File name      1-10	10A1	--	Must be the name (type) of the file to be created; misspellings will not be tolerated
<b>3</b> <b>File Identifier</b>	Identifier      1-10	60A1	--	Can be any alphanumeric sequence; it will not be validated, but will be written on the file and used in printed page headings exactly as given
				These cards contain integer-format control information; all five cards must be included for all programs even though a given program does not require all the control information
<b>4</b> <b>Control Counters</b>	Number of Species 1-10	I10	--	For files that vary by species, this is the number of species output to the file; for files that do not vary by species, this number should be zero; in either case, this parameter controls the number of species names read from Cards 9+ in this packet; for files that do not vary by species, the species names are ignored

Table VI-3 (Continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
Number of user-defined variables 11-20	I10	--		If species are input that do not appear in the output list, their number must be provided here; they must also be named in the UNITS packet; for files that do not vary by species, there is no need for user-defined variables by any of the current methods; if a user-defined method requires other than the standard variables, however, the additional variables can be enumerated here and named in the UNITS packet, and values can be supplied in the time-varying data packets
Number of stations (or boundary lines or point sources) 21-30	I10	--		Defines the maximum number of fixed locations in the region that are to be specified in time-invariant packets
Number of sub-regions 31-40	I10	--		Maximum number of different subregions that will be named in all SUBREGION packets; must be at least one
Number of parameters 41-50	I10	--		Maximum number of parameters that are defined or implied for all methods in all subregion-variable pairs; to obtain this value, multiply the number of parameters required for each method by the number of subregion-variable pairs that use the method (ALL counts not as 1, but as the total number); each time a parameter is redefined, it must be counted as a separate entry; add 3 to the maximum number of entries to ensure that enough space is allocated

Table VI-3 (Continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<b>5</b> <b>Output Controls</b>				
	Output file unit number 1-10	I10	--	Depending on the implementation, a specific number may be required for each output file
	Input print option 11-20	I10	--	= 0--don't print cards; = 1--do print input cards
	Output print option 21-30	I10	--	= 0--don't print all output data; = 1--do print all output data
<b>6</b> <b>Print Options</b>				For all options specified on this card, the value 0 means "don't print it" and the value 1 means "do print it"
	Print units table 1-10	I10	--	
	Print station locations (or boundaries or point sources) 11-20	I10	--	
	Print region 21-30	I10	--	
	Print methods table 31-40	I10	--	
	Print station values (or boundary values or point source emissions) 41-50	I10	--	

Table VI-3 (continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<b>7 Vertical Controls</b>				
Number of vertical parameters 1-10	I10	--		Meaningful only for files that vary vertically; for other files, this card should be blank
				Maximum number of parameters that are defined or implied for all vertical methods for all subregion-variable pairs; this value is computed in the same manner as the number of parameters (defined on Card 4)
Number of profile heights 11-20	I10	--		Maximum number of height-value pairs that will be input for any single profile
Print vertical methods table 21-30	I10	--		= 0--don't print it; = 1--do print it
Print vertical profiles table 31-40	I10	--		= 0--don't print it; = 1--do print it
<b>8 File Unit Assignment</b>				
		--		Contains FORTRAN unit numbers for files that might be required as input to a particular data preparation program; if any of these files is not to be read in a run, its field should be left blank or set to zero; for consistency, the same unit number should be assigned to a given file for every data preparation program using that file

Table IV-3 (Continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
DIFFBREAK	unit number 1-10	I10	--	Suggested value = 11, if required
REGIONTOP	unit number 11-20	I10	--	Suggested value = 12, if required
TOPCONC	unit number 21-30	I10	--	Suggested value = 13, if required
TEMPERATUR	unit number 31-40	I10	--	Suggested value = 14, if required
METSCALARS	unit number 41-50	I10	--	Suggested value = 15, if required
WIND	unit number 51-60	I10	--	Suggested value = 16, if required
<sup>9+</sup> <b>Species Names</b>	Species 1-10	10A1	--	One species name per card; the number of species cards to be included here is defined on Card 4

Table VI-3 (Concluded)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
10 Time Span				Defines the time span for which the file contains data; this card must be present
Beginning date 1-10	I10	Julian date (yyddd)		yy is the last two digits of the year and ddd is the three-digit Julian date
Beginning time 11-20	I10	Military time (hhmm)		hh is hours and mm is minutes--this time will be internally converted to hours and fraction
End date 21-30	I10	Julian date (yyddd)		Same format as that for beginning date
End time 31-40	I10	Military time (hhmm)		Same format as that for beginning time
11 Packet Terminator	10A1	--		Must be END, left adjusted
	Terminator 1-3			

<u>Card Number</u>	<u>Card Identifier</u>
1	Packet Header
2	File Name
3	File Identifier
4	Control Counters
5	Output Controls
6	Print Options
7	Vertical Controls
8	File Unit Assignments
9+	Species Names
10	Time Span
11	Packet Terminator

<u>Column Number</u>	<u>FORTRAN Format</u>				
1-10	11-20	21-30	31-40	41-50	51-60
CONTROL					
File name					
# species	# u-def var	# stn	# subreg	# param	spare
output file	pr input	pr output	spare	spare	spare
pr units	pr stn loc	pr region	pr meth	pr stn val	spare
#vert param	# heights	pr vert meth	pr vert prof	spare	spare
DIFFBREAK	REGIONTOP	TOPCONC	TEMPERATUR	METSCALAR	WIND unit #
unit #	unit #	unit #	unit #	unit #	unit #
spec name					
beg date	beg time	end date	end time		
END					

Exhibit VI-1. CONTROL Packet Format

## 2. REGION Packet Rules

The REGION packet must follow the CONTROL packet. It contains a complete definition of the location, size, and resolution of the modeling region. The same REGION packet values should be used to prepare all M.E.B. files for a given region. The contents of each card in the REGION packet are described in Table VI-4 and the format is shown in Exhibit VI-2.

Table VI-4. REGION Packet for M.E.B. Files

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be REGION, left adjusted
2 Reference Origin (UTM)				The origin (in UTM coordinates) used to locate the modeling region and all other fixed points (stations, point sources, boundary lines); these numbers are not used for computation but are compared to the reference origins on other files for consistency
	x-location 1-10	F10.0	m	x-coordinate
	y-location 11-20	F10.0	m	y-coordinate
	Zone 21-30	I10	--	UTM zone
3 Origin of Grid				The location of the lower left-hand corner of the grid with respect to the reference origin; this point is used to place the grid on the map; the grid is assumed to be rectangular and is oriented with the x-direction increasing to the east and the y-direction increasing to the north; the grid origin may be the same as the reference origin, in which case the coordinates are 0.0, 0.0
	x-location 1-10	F10.0	m	x-coordinate
	y-location 11-20	F10.0	m	y-coordinate

Table VI-4 (Continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<b>4</b> <b>Grid Cell Size</b>				The grid consists of a matrix of rectangular cells of the same size and shape; this card defines the length of the cell along each axis
	x-size 1-10	F10.0	m	x-length
	y-size 11-20	F10.0	m	y-length
<b>5</b> <b>Grid Size</b>				Defines the size of the grid in terms of the number of cells in each horizontal direction and the number of cells in the vertical direction
	Number of cells in x-direction 1-10	I10	--	Must be > 0
	Number of cells in y-direction 11-20	I10	--	Must be > 0
	Number of cells in z-direction 21-30	I10	--	Should be equal to the number of cells in the lower layer, plus the number of cells in the upper layer, plus one if there is a surface layer of nonzero height (see Card 6); for files that do not vary vertically, this number can be set to 0
<b>6</b> <b>Vertical Definition</b>				Contains the parameters required to define the vertical distribution of cells within the region; (see Section IV.A.2 for a detailed discussion of vertical distribution parameters); if the number of z cells = 0, this card must be present, but the values will be ignored

Table VI-4 (Concluded)

Card Number and Name	Item and Columns	Format	Units	Comments
Number of cells in lower layer 1-10	I10	--		Must be $\geq 1$
Number of cells in upper layer 11-20	I10	--		Must be $\geq 0$
Surface layer 21-30	F10.0	m		Height of surface layer; must be 0.0 if no surface layer is to be simulated (since microscale features have not been fully tested, set to 0.0)
Lower layer 31-40	F10.0	m		Minimum height of cells in lower layer
Upper layer 41-50	F10.0	m		Minimum height of cells in upper layer
7 Packet Terminator	10A1 Terminator 1-3	--		Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	<u>Column Number</u>					<u>FORTRAN Format</u>
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header						60A1
2	Reference Origin (UTM)	x-loc	y-loc	zone	.	.	2F10.0, 1I0
3	Origin of Grid	x-loc	y-loc				2F10.0
4	Grid Cell Size	x-size	y-size	# z	# y	# z	2F10.0
5	Grid Size	# x	# y	# z up	# z low	# z layer	3I10
6	Vertical Definition	# z low	# z up	ht surf	min cell ht low	min cell ht up	2I10, 3F10.0
7	Packet Terminator	END					10A1

Exhibit VI-2. REGION Packet Format

### 3. UNITS Packet Rules

The UNITS packet must be used whenever values for any species or implicit variable are input in other than internal units. Additionally, all user-defined variables must be named in this packet, even if they are already in internal units. When it appears, this packet should follow the REGION packet and precede all the others, since it can modify coordinates specified in other time-invariant packets.

Table VI-5 shows the internal units for all variables used by the SAI Airshed Model. Table VI-6 shows other standard units that can be used and the associated conversion factors for each variable. Table VI-7 shows the standard molecular weights for certain species. In addition to internal units or built-in standard units, the user can specify his own input units, conversion factors, and molecular weights. The following specific rules apply to this packet:

- > If a species or an implicit variable is not defined in this packet, it is assumed to be already in the appropriate internal units.
- > If the unit name for a variable is left blank, the program will provide space for that variable and no unit conversions will be performed.
- > If the variable is in one of the standard alternate units shown in Table VI-6, the units should be named and the factor fields left blank; the built-in conversion factors will then be used.
- > If the variable will be input in nonstandard units, the unit name should be nonblank and scaling factors provided; conversion will be made using the following formula:

$$\text{var}_{\text{internal}} = a * \text{var}_{\text{input}} + b .$$

Table VI-5. Implicit Variable Names and Internal Units  
Used in the SAI Airshed Model

Variable Name	Internal Units	Code
Time	hr	*
HEIGHT	m	M
COORD	m	M
DIFFBREAK	m	M
REGIONTOP	m	M
ROUGHNESS	m	M
DIAMETER	m	M
SPEED	m/hr	M/HR
WINDX	m/hr	M/HR
WINDY	m/hr	M/HR
CARM	m/hr	M/HR
STACKVEL	m/hr	M/HR
FLOWRATE	m <sup>3</sup> /hr	M3/HR
DIRECTION	radians (from N = 0)	RADN=0
TEMPERATUR	°K	DEGK
STACKTEMP	°K	DEGK
TGRADBELOW	°K/m	DEGK/M
TGRADABOVE	°K/m	DEGK/M
ATMOSPRESS	atm	ATMOSPHERE
EXPCLASS	unitless	*
VEGFACTOR	unitless	*
RADFACTOR	min <sup>-1</sup>	*
Concentrations	ppm ( $\mu\text{g}/\text{m}^3$ for AEROSOLS)	PPM MICROG/M3
CONCWATER	ppm	PPM
Point source emissions	g-mol/hr (g/hr for AEROSOLS)	GM/HR G/HR

\* The units for these variables can not be changed by the UNITS packet.

Table VI-6. Standard Unit Conversions

Variable	Standard Units	Standard Unit Code	a*	b
TEMPERATUR	$^{\circ}\text{K}^+$	DEGK	-9.0	-9.0
	$^{\circ}\text{C}$	DEGC	-9.0	273.15
	$^{\circ}\text{F}$	DEGF	0.5555556	255.38
	$^{\circ}\text{R}$	DEGR	0.5555556	-9.0
HEIGHT	m <sup>†</sup>	M	-9.0	-9.0
COORD	cm	CM	0.01	-9.0
ROUGHNESS		KM	1000.0	-9.0
DIFFBREAK		IN	0.0254	-9.0
REGIONTOP		FT	0.3048	-9.0
DIAMETER	mi	MI	1609.344	-9.0
SPEED	m/hr <sup>†</sup>	M/HR	-9.0	-9.0
WINDX	km/hr	KM/HR	1000.0	-9.0
WINDY		MI/HR	1609.344	-9.0
CARM		M/MIN	60.0	-9.0
STACKVEL		FT/MIN	18.288	-9.0
	m/sec	M/S	3600.0	-9.0
	ft/sec	FT/S	1097.28	-9.0
	knots	KNOTS	1852.0	-9.0
DIRECTION	radians (from N = 0) <sup>†</sup>	RADN=0	-9.0	-9.0
	radians (from S = 0)	RADS=0	-9.0	3.141593
	degrees (from N = 0)	DEGN=0	0.01745329	-9.0
	degrees (from S = 0)	DEGS=0	0.01745329	3.141593
	16-point (from S = 1)	16PTS=1	0.392699	2.748893
	16-point (from S = 16)	16PTS=16	0.392699	3.141593
	36-point (from S = 1)	36PTS=1	0.1745329	2.967059
	36-point (from S = 36)	36PTS=36	0.1745329	3.141593

Table VI-6 (Continued)

<u>Variable</u>	<u>Standard Units</u>	<u>Standard Unit Code</u>	<u>a*</u>	<u>b</u>
TGRADUPPER	°K/m <sup>†</sup>	DEGK/M	-9.0	-9.0
TGRADLOWER	°K/km	DEGK/KM	0.001	-9.0
	°C/m	DEGC/M	-9.0	-9.0
	°C/km	DEGC/KM	0.001	-9.0
	°F/ft	DEGF/FT	1.822689	-9.0
	°F/mi	DEGF/MI	0.0003452	-9.0
	°R/ft	DEGR/FT	1.822689	-9.0
	°R/mi	DEGR/MI	0.0003452	-9.0
ATMPRESS	atm <sup>†</sup>	ATMOSPHERE	-9.0	-9.0
	Bars	BARS	0.986923	-9.0
	mm of Hg	MMHG	0.0013158	-9.0
	ft of H <sub>2</sub> O	FTH2O	0.029499	-9.0
	kg/cm <sup>2</sup>	KG/CM2	0.967841	-9.0
	in of Hg	INHG	0.0334211	-9.0
	lb/in <sup>2</sup>	LB/IN2	0.068046	-9.0
	psi	PSI	0.068046	-9.0
	millibars	MILLIBARS	0.0009869	-9.0
	in of H <sub>2</sub> O	INH2O	0.0024582	-9.0
Concentrations	ppm <sup>†</sup>	PPM	-9.0	-9.0
CONCWATER	pptm <sup>†</sup>	PPTM	0.1	-9.0
	pphm	PPHM	0.01	-9.0
	ppb	PPB	0.001	-9.0
	g mol/m <sup>3</sup>	GM/M3	.24400	-9.0
	lb mol/ft <sup>3</sup>	LBM/FT3	3.876x10 <sup>"</sup>	-9.0
	µg/m <sup>3</sup>	MICROG/M3	-0.0244	-9.0
	g/m <sup>3</sup>	G/M3	-24400	-9.0
	lb/ft <sup>3</sup>	LB/FT3	-3.876x10 <sup>"</sup>	-9.0

Table VI-6 (Concluded)

<u>Variable</u>	<u>Standard Units</u>	<u>Standard Unit Code</u>	<u>a*</u>	<u>b</u>
Point source emissions	g-mol/hr <sup>†</sup>	GM/HR	-9.0	-9.0
	g-mol/day	GM/D	0.0416667	-9.0
	lb-mol/hr	LBM/HR	453.592	-9.0
	lb-mol/day	LBM/D	18.89967	-9.0
	kg/hr	KG/HR	-1000.0	-9.0
	kg/day	KG/D	-41.6667	-9.0
	lb/hr	LB/HR	-453.592	-9.0
	lb/day	LB/D	-18.89967	-9.0
	ton/hr	TON/HR	-907184.0	-9.0
	ton/day	TON/D	-37799.3	-9.0
	ton/year	TON/YR	-103.6	-9.0

\* When the multiplicative factor (a) is negative, but not -9., the molecular weight is used in the denominator, and the absolute value of a is used as the multiplier.

† Internal units.

Table VI-7. Standard Species Names and Molecular Weights Used for Unit Conversion

<u>Species</u>	<u>Molecular Weight</u>
NO	30.
NO <sub>2</sub>	46.
CO	28.
O <sub>3</sub>	48.
H <sub>2</sub> O <sub>2</sub>	34.
SO <sub>2</sub>	64.
HN <sub>0</sub> 2	47.
H <sub>2</sub> O	18.
O <sub>2</sub>	32.
CO <sub>2</sub>	44.
SO <sub>4</sub>	96.
PAN	121.
PAR	16.
OLE	32.
ETH	32.
ARO	96.
AEROSOLS	1.
CH <sub>4</sub>	16.
TOTAL HC	16.
SOX	64.
NOX	46.
OXIDANT	48.

- > If the variable is a species and the input values will be in mass units rather than mole units, the following formula will be used:

$$\text{var}_{\text{internal}} = (\text{a} * \text{var}_{\text{input}} + \text{b}) / \text{mol wt}$$

- > If both a standard unit name and scaling factors are specified, a warning message will be printed and the user's scaling factors will be used.
- > A designation of -9.0 indicates that the conversion factor is not to be applied.
- > If the factor fields are not blank or zero, a molecular weight must be entered on the card. If the variable does not have a molecular weight, e.g., as in TEMPERATUR, either a -9.0 or a +1.0 should be entered in the molecular weight field.

The contents of each card in the UNITS packet are shown in Table VI-8 and the format is given in Exhibit VI-3.

Table VI-8. UNITS Packet for M.E.B. Files

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<b>1</b> <b>Packet Header</b>	<b>Header</b> 1-10	<b>60A1</b>	--	Must be UNITS, left adjusted
<b>2</b> <b>Unit Definition</b>	<b>Variable name</b> 1-10	<b>10A1</b>	--	<p>Can be a species name, implicit variable, user-defined variable, or either of two special names:</p> <ul style="list-style-type: none"> <li>&gt; COORD--for station, point source, boundary, and vertical profile coordinates;</li> <li>&gt; HEIGHT--for station height, point source, stack height, and vertical profile heights (note that this name does not apply to the height of the diffusion break or top of the region);</li> </ul> <p>conversions specified apply to values input in all subsequent packets</p>
				If field is left blank, no unit conversion will be performed; if this is one of the standard unit names listed in Table VI-6, the corresponding conversion factors will be used; if it is any other unit name, conversion factors must be input
	<b>Unit name</b> 11-20	<b>10A1</b>	--	

Table VI-8 (Concluded)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
Multiplicative factor (a) 21-30		F10.0	--	Can be left blank if standard conversion factors are to be used
Additive factor (b) 31-40		F10.0	--	Can be left blank if standard conversions are to be used
Molecular weight (c) 41-50		F10.0	--	Can be left blank if standard molecular weights are to be used; if the unit name does not belong to the standard set, this number should be set to 1., even for nonspecies variables
3				
Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	<u>Column Number</u>					<u>FORTRAN Format</u>
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	UNITS					60A1
2+	Unit Definition	var name	unit name	mult fact	add fact		2(10A1), 3F10.0
3	Packet Terminator	END					10A1

Exhibit VI-3. UNITS Packet Format

#### 4. STATIONS Packet Rules

The STATIONS packet must be included if the methods STATINTERP, POISSON, or DIVERGENCE are used for any variable. The STATIONS packet, if present, must appear before the first time interval; it must include all stations for which values may appear at any time within the span of the file. The number of station cards must not exceed the maximum number of stations specified in the CONTROL packet. Duplication of station names is considered an error resulting in termination of the run. The contents of each card in the STATIONS packet are shown in Table VI-9 and the format is given in Exhibit VI-4.

Table VI-9. STATIONS Packet for M.E.B. Files

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be STATIONS, left adjusted
2+ Station Name and Location				
Station name 1-10		10A1	--	
x-location 11-20		F10.0	m, unless COORD was specified in UNITS packet	Can be any name except the same as that for another station, or blank, or ALL x-coordinate of the station; this number must be specified
y-location 21-30		F10.0	m, unless COORD was specified in UNITS packet	y-coordinate of the station; this number must be specified
z- location 31-40		F10.0	m, unless HEIGHT was specified in UNITS packet	Height of the station above ground; if omitted, this number is assumed to be 0; for files that do not vary vertically, this number is ignored
3 Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	Column Number				<u>FORTRAN Format</u>	
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	STATIONS					60A1
2+	Station Name and Location	stn name	x-loc	y-loc	z-loc		10A1, 3F10.0
3	Packet Terminator	END					10A1

Exhibit VI-4. STATIONS Packet Format

## 5. POINT SOURCES Packet Rules

The POINT SOURCES packet is used for preparing the PTSOURCE file; it must appear before the first time interval and include all point sources for which values may appear at any time within the span of the file. There are two cards (a pair) required for each point source, and the number of pairs must equal the number of point sources specified on Card 4 of the CONTROL packet. Duplication of point source names is considered an error, resulting in termination of the run. The contents of each card in the POINT SOURCES packet are shown in Table VI-10 and the format is given in Exhibit VI-5.

Table VI-10. POINT SOURCES Packet for the PTSOURCE File

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be POINT SOURCES, left adjusted
2+ Point Source Name and Location				
	Point source name 1-10	10A1	--	Can be any alphanumeric sequence except the same as that of another point source or type of source, or blank, or ALL.
	Type 11-20	10A1	--	Can be any alphanumeric sequence except a point source name, or blank, or ALL; point sources of the same type are treated as a group when specifying the method for calculating emissions and when specifying the vertical method; they can also be treated as a group when specifying emissions or emission factors
x-location		F10.1	m, unless COORD was specified in UNITS packet	x-coordinate of the point source with respect to the reference origin; this number must be specified
y-location		F10.1	m, unless COORD was specified in UNITS packet	y-coordinate of the point source with respect to the reference origin; this number must be specified

Table VI-10 (Concluded)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
3+ Point Source properties				
	Stack height 1-10	F10.0	m, unless HEIGHT was specified in UNITS packet	Height of the top of the stack above the ground; this number must be greater than zero
	Stack exit diameter* 11-20	F10.0	m, unless DIAMETER was speci- fied in UNITS packet	Must be specified if it is to be used to calculate the flow rate for the PLUMERISE method
	Stack exit temperature 21-30	F10.0	°K, unless STACKTEMP was spe- cified in UNITS packet	Used to calculate heat flux for the PLUMERISE method
	Stack exit velocity* 31-40	F10.0	m/hr, unless STACKVEL was speci- fied in UNITS packet	Used with stack exit diameter to calcu- late flow rate for the PLUMERISE method
4 Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

\* If the vertical method PLUMERISE is to be used to calculate the height at which the emissions enter the modeling region, either both the stack exit diameter and stack exit velocity or the stack flow rate alone must be specified. Flow rate is specified in the EMISSIONS VALUES packet.

<u>Card Number</u>	<u>Card Identifier</u>	Column Number					<u>FORTRAN Format</u>
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	POINT SOURCES					60A1
2+	Point Source Name and Location	pt srce nam	type	x-loc	y-loc		2(10A1), 2F10.0
3+	Point Source Properties	stack ht	stack diam	stack temp	stack vel		4F10.0
4	Packet Terminator	END					10A1

Exhibit VI-5. POINT SOURCES Packet Format

## 6. BOUNDARIES Packet Rules

The BOUNDARIES packet, used for preparing the BOUNDARY file, must appear before the first time interval. It contains a set of line segments that define the boundary of the simulated area; each line segment is defined by two coordinate pairs that represent its end points. The line segments can appear in any order and, within each one, the end points can appear in any order, but the complete set must represent a closed figure (i.e., every end point must be identical to one, and only one, end point of another line segment). In addition, the figure represented must be nonconcave along each axis; that is, within a single row or column, any two cells within the simulated area must not have any nonsimulated cells between them.

The boundary line segments outline the edge cells within the inner area to be simulated. Figure VI-1 shows a 17 x 22 cell region in which the inner 15 x 20 area is to be simulated. In the figure, the boundary line segments are indicated by dotted lines. To ensure that the program succeeds in matching the end points of touching line segments, each end point should be defined to be in the center of the cell in which it lies. The number of boundary line segments input must not exceed the maximum number specified on Card 4 of the CONTROL packet. Duplication of line segment names is considered an error resulting in termination of the run.

The contents of each card in the BOUNDARIES packet are shown in Table VI-11 and the format is given in Exhibit VI-6.

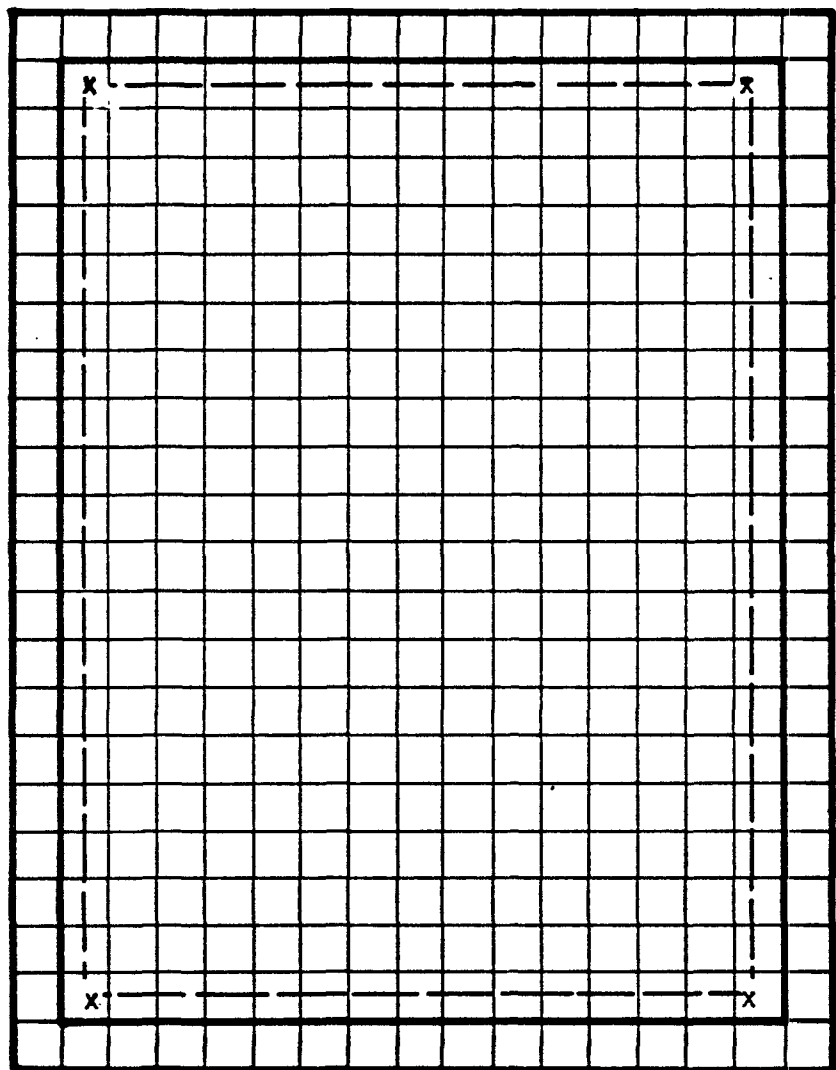


Figure VI-1. Definition of Boundary Line Segments

**Table VI-11. BOUNDARIES Packet for the BOUNDARY File**

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<b>1</b> <b>Packet Header</b>	<b>Header</b> 1-10	<b>60A1</b>	--	Must be BOUNDARIES, left adjusted
<b>2+</b> <b>Boundary Definition</b>				
Line segment name 1-10		10A1	--	Any alphanumeric sequence except the same as that for another line segment name, blank, or ALL
x-beginning 11-20		F10.0	m, unless COORD was specified in UNITS packet	x-coordinate of the first end point with respect to the reference origin; the end point should lie in the center of a cell
y-beginning 21-30		F10.0	m, unless COORD was specified in UNITS packet	y-coordinate of the first end point with respect to the reference origin
x-end 31-40		F10.0	m, unless COORD was specified in UNITS packet	x-coordinate of the second end point with respect to the reference origin
y-end 41-50		F10.0	m, unless COORD was specified in UNITS packet	y-coordinate of the second end point with respect to the reference origin
<b>3</b> <b>Packet Terminator</b>	<b>Terminator</b> 1-3	<b>10A1</b>	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	<u>Column Number</u>	<u>FORTRAN Format</u>				
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	BOUNDARIES					60A1
2+	Boundary Definition	Line name	x-beg	y-beg	x-end	y-end	10A1, 4F10.0
3	Packet Terminator	END					10A1

Exhibit VI-6. BOUNDARIES Packet Format

## 7. TIME INTERVAL Packet Rules

A series of TIME INTERVAL packets follows the time-invariant data. The time intervals must be contiguous, must go forward, and must cover the entire time span of the file as specified in the CONTROL packet. If any time interval extends beyond the time span specified in the CONTROL packet, that time interval will be reset by the program to lie exactly within the time span of the file.

TIME INTERVAL packets contain other time-varying packets that define regional divisions, calculation methods, and time-varying data values. Each TIME INTERVAL packet consists of a header card, a time interval card, other packets as desired, and a terminator card. The contents of each card in the TIME INTERVAL packet is shown in Table VI-12 and the format is given in Exhibit VI-7. The other time-varying packets that can be included within the TIME INTERVAL packet for a given file are shown in the appropriate section of Chapter VII; their formats and rules are described in the remainder of this chapter.

Table VI-12. TIME INTERVAL Packet for M.E.B. Files

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be TIME INTERVAL, left adjusted
2 Time Interval	Beginning date 1-10 Beginning time 11-20 End date 21-30 End time 31-40	I10 I10 I10 I10	Julian date (yyddd) Military time (hhmm) Julian date (yyddd) Military time (hhmm)	yy is the last two digits of the year and ddd is the three-digit Julian date hh is hours and mm is minutes; this time will be internally converted to hours and fraction Same format as beginning date Same format as beginning time
3 + Other Packets				Chapter VII lists the packets allowed here for each data file
4 Packet Terminator	Terminator 1-3	10A1	--	Must be ENDTIME, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	<u>Column Number</u>				<u>FORTRAN Format</u>	
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	TIME INTERVAL					60A1
2	Time Interval		beg date	end date	end time		4I10
3+	Other Packets						
4	Packet Terminator		ENDTIME				10A1

Exhibit VI-7. TIME INTERVAL Packet Format

## 8. SUBREGION Packet Rules

The entire modeling region can be divided into subregions for data preparation purposes. Different data preparation methods or other parameters can be associated with each subregion. (These subregions should not be confused with "segments," which are divisions of the region for simulation purposes.) The subregion definitions may likely be different for the preparation of different types of data files; in fact, a subregion definition can be changed within the time span of a given file depending on the availability of or assumptions about the data.

A subregion name must be assigned to each grid square in the modeling region. The maximum number of subregion names that may be assigned is specified in the CONTROL packet. If this number = 1, the entire region is considered to be one subregion; nevertheless, the SUBREGION packet must be included and must appear at the beginning of the first TIME INTERVAL packet. In subsequent time intervals, the SUBREGION packet can be used to change subregion assignments. Only those portions of the region to be changed need be specified. New subregion names can be added provided that the maximum number allowed is not exceeded.

Subregion names are assigned by row. A typical subregion definition card contains a subregion name and a row in which it appears. For that row, a column number and cell count, n, are specified. Beginning at that column, the subregion name will be repeated for n columns along the row. The following conditions constitute errors resulting in termination of the program:

- > The cell count extends the column count beyond the edge of the region.
- > Any grid cell is unassigned.
- > The number of subregion names exceeds the maximum number specified in the CONTROL packet.

If the entire region is to be treated as one subregion, the SUBREGION packet should contain the following information:

SUBREGION

A-----|-----1|-----1|-----1|

END ,

where A (or some other simple name) is the subregion name. This name (or ALL) must then be used in the subregion field of all subsequent input cards.

The contents of each card in the SUBREGION packet are shown in Table VI-13 and the format is given in Exhibit VI-8.

Table VI-13. SUBREGION Packet for M.E.B. Files

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<b>1</b> Packet Header	Header <b>1-10</b>	60A1	--	Must be SUBREGION, left adjusted
<b>2+</b> Subregion Definition				
Subregion name <b>1-10</b>		10A1	--	Any alphanumeric sequence except blank or ALL
Row number <b>11-20</b>		I10	--	Must be $\geq 1$ , and $\leq$ number of rows (cells in y-direction) in region
Beginning column number <b>21-30</b>		I10	--	Must be $\geq 1$ , and $\leq$ number of columns (cells in x-direction) in region
Cell count <b>31-40</b>		I10	--	Must not exceed [(number of columns) - (column number) + 1]; if omitted, this value is assumed to be = rest of row; if negative, this value is assumed to be = rest of modeling region
<b>3</b> Packet Terminator	Terminator <b>1-3</b>	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	<u>Column Number</u>				<u>FORTRAN Format</u>	
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	SUBREGION					60A1
2+	Subregion Definition	subreg name	row #	beg col #	# cells		10A1, 3I10
3	Packet Terminator	END					10A1

Exhibit VI-8. SUBREGION Packet Format

## 9. METHOD Packet Rules

The METHOD packet is used to define the calculation method to be used for ground-level values of each variable in each subregion; it is required for all of the M.E.B. files except METSCALARS. The METHOD packet must appear in the first time interval, and it must directly follow the SUBREGION packet. In subsequent time intervals, the methods previously defined will persist unless they are respecified. If new subregions are defined in later time intervals, methods must also be defined for them. The contents of each card in the METHOD packet are shown in Table VI-14 and the format is given in Exhibit VI-9. Within the METHOD packet, each Method Definition card must be followed by its Parameter cards, if any. All of the calculation methods and the parameters required by each one are defined in Section VI.A.6. Chapter VII lists the particular methods that can be used for each file.

Table VI-14. METHOD Packet for M.E.B. Files

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be METHOD, left adjusted
2+ Method Definition				
	Subregion name 1-10	10A1	--	Must match a name in the subregion packet; if the name = ALL, this method will apply to this variable in all subregions
	Variable name 11-20	10A1	--	Must be an output species, implicit variable name, or user-defined variable name from the UNITS packet; if the variable name = ALL, this method will be used for all variables in this subregion
	Method 21-30	10A1	--	See Section VI.A.6 for standard method names
	Minimum value 31-40	F10.0	Internal units assumed, unless otherwise specified in UNITS packet	Lower bound for variable; if the variable name = ALL, this value will be converted according to unit specifications for the first variable; care should be taken in setting limits for ALL variables, especially when species (requiring different molecular weights for conversion) are involved; if the value calculated for the variable is less than this number, that value will be set equal to this number; if the field is left blank, it is assumed to be zero; if it is set to -9.0, no lower limit test will be performed

Table VI-14 (concluded)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
Maximum value 41-50		F10.0	Internal units assumed, unless otherwise specified in UNITS packet	Upper bound for variable; if the variable name = ALL, this value will be converted as described above; if the value calculated for this variable is greater than the number, that value will be set equal to this number; if the field is left blank, it is assumed to be zero; if it is set to -9.0, no upper limit test will be performed
Number of parameters 51-60		I10	--	Defines the number of parameter cards that follow (Card 3 below); must be $\geq 0$ ; Section VI.A.6 gives the number of parameters associated with each method
<b>3+ Parameters</b>				
Parameter name 1-10		10A1	--	Standard parameter names and values are defined in Section VI.A.6; the number of parameter cards for a given method must equal the number specified on Card 2
Parameter value 11-20		F10.0	--	
<b>4 Packet Terminator</b>				
Terminator 1-3		10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>
1	Packet Header
2+	Method Definition
3+	Parameters
4	Packet Terminator

<u>Column Number</u>	<u>FORTRAN Format</u>			
1-10	11-20	21-30	31-40	41-50
METHOD				60A1
	subreg name	var name	method	3(10A1), 2F10.0, I10
	param name	value		10A1,F10.0
		END		10A1

Exhibit V1-9. METHOD Packet Format

## **10. VERTICAL METHOD Packet Rules**

In general, where it is necessary to determine the values of variables in grid cells above ground level, a VERTICAL METHOD packet is required. The VERTICAL METHOD packet is used to define the calculation method to be used for obtaining values of each variable above ground level in each sub-region. The VERTICAL METHOD packet must appear in the first time interval; it must follow the METHOD packet and precede the VERTICAL PROFILES packet (if any). In subsequent time intervals, the vertical methods previously defined will persist unless they are respecified. If new subregions are defined in later time intervals, vertical methods must also be defined for them. Within the VERTICAL METHOD packet, each vertical method definition card must be followed by its parameter cards, if any. All of the vertical calculation methods and the parameters required by each one are defined in Section VI.A.7. Chapter VII lists the particular vertical methods that can be used for each file. The contents of each card in the VERTICAL METHOD packet is shown in Table VI-15, and the format is given in Exhibit VI-10.

Table VI-15. VERTICAL METHOD Packet for M.E.B. Files

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be VERTICAL METHOD, left adjusted
2+ Vertical Method Definition				
	Subregion name 1-10	10A1	--	Must match a name in the subregion packet; if the name = ALL, this vertical method will apply to this variable in all subregions
	Variable name 11-20	10A1	--	Must be an output species, implicit variable name, or user-defined variable name from the UNITS packet; if the variable name = ALL, this vertical method will be used for all variables in this subregion
	Method 21-30	10A1	--	See Section VI.A.7 for standard vertical method names
	Minimum value 31-40	F10.0	Internal units, unless otherwise specified in UNITS packet	Lower bound for variable; as in the METHOD packet, if the variable name = ALL, this value will be converted according to unit specifications for the first variable; if the value calculated for the variable is less than this number, that value will be set equal to this number; if the field is left blank, it is assumed to be zero; if it is set to -9.0, no lower limit test will be performed

Table VI-15 (Concluded)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
	<b>Maximum value 41-50</b>	F10.0	Internal units, unless otherwise specified in UNITS packet	Upper bound for variable; if the variable name = ALL, this value will be converted as described above; if the value calculated for this variable is greater than this number, that value will be set equal to this number; if the field is left blank, it is assumed to be zero; if it is set to -9.0, no upper limit test will be performed
	<b>Number of parameters 51-60</b>	I10	--	Defines the number of parameter cards that follow (Card 3 below); must be > 0; Section VI.A.7 gives the number of parameters associated with each vertical method
<b>3+ Parameters</b>				
	<b>Parameter name 1-10</b>	10A1	--	
	<b>Parameter value 11-21</b>	F10.0	--	
<b>4 Packet Terminator Terminator 1-3</b>				
		10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	Column Number							<u>FORTRAN Format</u>
		1-10	11-20	21-30	31-40	41-50	51-60		
1	Packet Header	VERTICAL METHOD							60A1
2+	Vertical Method Definition	subreg name	var name	method	min value	max value	# param		3(10A1), 2F10.0 1I0
3+	Parameters	param name	value						10A1, F10.0
4	Packet Terminator	END							10A1

Exhibit VI-10. VERTICAL METHOD Packet Format

## **11. CONSTANTS Packet Rules**

The CONSTANTS packet is required in the first time interval if the method for any variable in any subregion has been specified as CONSTANT. All constant values can be included in one packet or, if desired, they can be grouped in several packets. In either case, all constant values must be specified in the initial time interval and will persist in subsequent time intervals unless they are explicitly changed. The contents of each card in the CONSTANTS packet are shown in Table VI-16 and the format is given in Exhibit VI-11.

Table VI-16. CONSTANTS Packet for M.E.B. Files

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be CONSTANTS, left adjusted
2+ Constants				
	Subregion name 1-10	10A1	--	Must be identical to a previously named subregion, or ALL; if ALL is specified, all subregions that use the CONSTANT method for this variable will be assigned this value
	Variable name 11-20	10A1	--	Must be a variable named on a method card that specifies CONSTANT; if ALL is specified, all variables that require CONSTANT for this subregion will be assigned this value
	Value 21-30	F10.0	Internal units, unless otherwise specified in UNITS packet	If the variable name = ALL, this value will be converted according to unit specifications for the first variable; care should be taken in specifying values for ALL variables, especially when species (requiring different molecular weights for conversion) are involved
3 Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	Column Number				<u>FORTRAN Format</u>	
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	CONSTANTS					60A1
2+	Constants	subreg name	var name	value			2(10A1), F10.0
3	Packet Terminator	END					10A1

Exhibit VI-11. CONSTANTS Packet Format

## 12. GRID VALUES Packet Rules

The GRID VALUES packet must be included in the first time interval if the method for any variable in any subregion has been specified as GRID VALUE. All grid values can be included in one packet or, if desired, they can be grouped in several packets. In either case, all grid values must be specified in the initial time interval and will persist in subsequent time intervals unless they are explicitly changed. The contents of each card in the GRID VALUES packet are shown in Table VI-17 and the format is shown in Exhibit VI-12.

Table VI-17. GRID VALUES Packet for M.E.B. Files

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be GRID VALUES, left adjusted
2+ Grid Values				
	Subregion name 1-10	10A1	--	Must be identical to a previously named subregion; GRID VALUES must be specified as the method for this variable
	Variable name 11-20	10A1	--	Must be a variable named or implied (by means of ALL) on a method card for this subregion
	Column number 21-30	I10	--	y-index on grid, counting from bottom (SOUTH) row = 1
	Row number 31-40	I10	--	x-index on grid, counting from left (WEST) column = 1
	Value 41-50	F10.0	Internal units, unless otherwise specified in UNITS packet	If the variable name = ALL, this value will be converted according to unit specifications for the first variables; care should be taken in specifying values for ALL variables especially when species (requiring different molecular weights for conversion) are involved.
3 Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	<u>Column Number</u>				<u>FORTRAN Format</u>	
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	GRID VALUES					
2+	Grid Values	subreg name	var name	col #	row #	value	
3	Packet Terminator	END					

Exhibit VI-12. GRID VALUES Packet Format

### **13. STATION READINGS Packet Rules**

The STATION READINGS packet must appear in the first time interval when the method for any variable in any subregion has been specified as STATINTERP, POISSON, or DIVERGENCE. A value that does not appear in the first time interval for a station-variable pair is recognized as "missing", and the station will not be included in the calculation. In subsequent time intervals, the STATION READINGS packet can be used to modify existing data in the following ways:

- > An existing reading can be changed by substituting a new value.
- > An existing reading can be nullified (i.e., changed to "missing") by supplying a value of -9.0.
- > A "missing" reading can be changed to "nonmissing" by supplying a value.

For any station-variable pair not supplied, the previous values, "missing" or not, will persist. The contents of each card in the STATION READINGS packet is shown in Table VI-18 and the format is given in Exhibit VI-13.

Table VI-18. STATION READINGS Packet for M.E.B. Files

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be STATION READINGS, left adjusted
2+ Station Readings				
Station name 1-10		10A1	--	Must be identical to a name specified in the STATIONS packet
Variable name 11-20		10A1	--	Must be a species name, or an implicit or user-defined variable
Value 21-30		F10.0	Internal units, unless otherwise speci- fied	A blank is treated as zero; -9.0 is considered "missing"
3 Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	Column Number				<u>FORTRAN Format</u>	
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	STATION READINGS					60A1
2	Station Readings	stn name	var name	value			2(10A1), F10.0
3	Packet Terminator	END					10A1

Exhibit VI-13. STATION READINGS Packet Format

#### 14. EMISSIONS VALUES Packet Rules

The EMISSIONS VALUES packet is used to create the PTSOURCE file. It provides emissions values for those variables for which the method EMVALUES was specified. The EMISSIONS VALUES packet must be included in the first time interval (unless all emissions are zero); any value not appearing in the first time interval for a point source-species pair will be assumed to be zero. In subsequent time intervals, existing data will persist unless explicitly changed. If all point sources of a given type have the same emissions values, the type can be specified instead of individual point source names.

If the vertical method PLUMERISE has been specified for a point source type, the variable FLOWRATE must be either input in the EMISSIONS VALUES packet or calculated. If it is not input, it will be calculated from the stack diameter and exit velocity input in the POINT SOURCES packet.

The contents of each card in the EMISSIONS VALUES packet are shown in Table VI-19 and the format is given in Exhibit VI-14.

**Table VI-19. EMISSIONS VALUES Packet for the PTSOURCE File**

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be EMISSIONS VALUES, left adjusted
2+ Point Source Emissions	Point source or type 1-10	10A1	--	Must be one of the following: > a name appearing in the POINT SOURCES packet, > ALL--meaning all point sources, > a type appearing in the POINT SOURCES packet, meaning all point sources of a given type
	Variable name 11-20	10A1	--	Must be an output species or a user- defined input variable, or FLOWRATE
	Value 21-30	F10.0	gm-mol/hr (gm/hr for AEROSOLS), unless otherwise spec- ified in UNITS packet	Emissions of the variable from the point source; if the variable is FLOWRATE, the units are m <sup>3</sup> /hr, unless otherwise speci- fied in the UNITS packet
3 Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	Column Number					<u>FORTRAN Format</u>
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	EMISSIONS	VALUES				60A1
2+	Point Source Emissions	pt srce or type	var name	value			2(10A1), F10.0
3	Packet Terminator	END					10A1

Exhibit VI-14. EMISSIONS VALUES Packet Format

## 15. EMISSIONS FACTORS Packet Rules

The EMISSIONS FACTORS packet can be input for a given time interval to scale the emissions values previously input. The factors are used only if the method EMFACTORS was specified for the variable and point source type. The following formula will be used:

$$\text{output emissions (source, species)} = \text{factor (source, species)} \\ \times \text{emissions (source, species)} .$$

The factor is always applied to the last emissions value input in an EMISSIONS VALUES packet, not the last one calculated by a previous emissions factor. If factors are not input, they are assumed to be 1.0 (or the previous value specified). Emissions factors are applied to numbers in internal units--they are thus to be used to describe temporal variation rather than unit conversion. The contents of each card in the EMISSIONS FACTORS packet are shown in Table VI-20 and the format is given in Exhibit VI-15.

Table VI-20. EMISSIONS FACTORS Packet for the PTSOURCE File

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be EMISSIONS FACTORS, left adjusted
2+ Emissions Factors	Point source or type 1-10	10A1	--	<p>Must be one of the following:</p> <ul style="list-style-type: none"> <li>&gt; a name appearing in the POINT SOURCES packet,</li> <li>&gt; ALL--meaning all point sources,</li> <li>&gt; a type appearing in the POINT SOURCES packet, meaning all point sources of a given type</li> </ul>
				<p>Must be an output species or a user-defined input variable; if the name = ALL, this factor will apply to all species and to FLOWRATE</p> <p>Emission factors apply to FLOWRATE as well as to species; when the variable = ALL, these factors are automatically applied; when factors are being provided for individual species, the factor for the flow rate, if it changes, must also be specified</p>
3 Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	<u>Column Number</u>				<u>FORTRAN Format</u>	
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	EMISSIONS	FACTORS				60A1
2+	Emissions Factors	pt srce or type	var name	factor			2(10A1), F10.0
3	Packet Terminator	END					10A1

Exhibit VI-15. EMISSIONS FACTORS Packet Format

## 16. BOUNDARY READINGS Packet Rules

The BOUNDARY READINGS packet is used to create the BOUNDARY file; it provides values for those variables for which the method BOUNDVALUE was specified. The packet must be input for the first time interval. Concentration values at ground level are specified for each species for each boundary line segment; missing values are considered an error. In subsequent time intervals, only the data to be changed need be specified. The contents of each card in the BOUNDARY READINGS packet is shown in Table VI-21, and the format is given in Exhibit VI-16.

**Table VI-21. BOUNDARY READINGS Packet for the BOUNDARY File**

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be BOUNDARY READINGS, left adjusted
2+ Boundary Values				
	Line segment name 1-10	10A1	--	Must be a name specified in the BOUNDARY packet, or ALL
	Variable name 11-20	10A1	--	Must be an output species, a user-defined variable, or ALL
	Value 21-30	F10.0	ppm ( $\mu\text{g}/\text{m}^3$ for AEROSOLS), unless otherwise specified in UNITS packet	Ground-level concentration of the variable along the line segment
3 Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	Column Number				<u>FORTRAN Format</u>	
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	BOUNDARY READINGS					60A1
2+	Boundary Values	line name	var name	value			2(10A1), F10.0
3	Packet Terminator	END					10A1

Exhibit VI-16. BOUNDARY READINGS Packet Format

## 17. SCALARS Packet Rules

The SCALARS packet must be input for the first time interval for the METSCALARS file. For this time interval, all six meteorological scalars (TGRADBELOW, TGRADABOVE, EXPCLASS, RADFACTOR, CONCWATER, and ATMOSPRESS) must be assigned values. Missing values are considered errors. In subsequent time intervals, only the data to be changed need be specified. The contents of each card in the SCALARS packet is shown in Table VI-22 and the format is given in Exhibit VI-17.

Table VI-22. SCALARS Packet for the METSCALARS File

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be SCALARS, left adjusted
2+ Meteorological Scalars				Must match the standard spelling of the scalars (see Chapter VII for instruc- tions on preparing the METSCALARS file)
	Scalar name 1-10	10A1	--	
	Value 11-20	F10.0	Internal units, unless otherwise speci- fied in UNITS packet	
3 Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>
1	Packet Header
2+	Scalars
3	Packet Terminator

<u>FORTRAN Format</u>	Column Number				<u>51-60</u>
	<u>1-10</u>	<u>11-20</u>	<u>21-30</u>	<u>31-40</u>	<u>41-50</u>
SCALARS					60A1
scalar name					10A1,
value					F10.0
END					10A1

Exhibit VI-17. SCALARS Packet Format

## 18. VERTICAL PROFILES Packet Rules

For some vertical methods, a profile is required (see Section VI.A.7 for details). The VERTICAL PROFILES packet must contain a profile for each subregion-variable pair requiring one. Each profile must contain at least two points, which must be entered in order of increasing height. The first point always represents ground level. There can be only one vertical profile specified for a given subregion and variable. Each profile consists of a description card followed by a set of height/value pairs. The contents of each card in the VERTICAL PROFILES packet is shown in Table VI-23 and the format is given in Exhibit VI-18.

Table VI-23. VERTICAL PROFILES Packet for M.E.B. Files

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<b>1</b>	<b>Header</b> <b>1-10</b>	<b>60A1</b>	--	Must be VERTICAL PROFILES, left adjusted
<b>2+</b> <b>Profile Description</b>				
	<b>Subregion name</b> <b>11-10</b>	<b>10A1</b>	--	Must be identical to a previously named subregion, or boundary line segment, or ALL; if ALL is specified, all subregions that require a vertical profile for this variable will use these values
	<b>Variable name</b> <b>11-20</b>	<b>10A1</b>	--	Must be a variable that requires a vertical profile for this subregion; if name = ALL, this profile will be used for all variables in this subregion
	<b>x-location</b> <b>21-30</b>	<b>F10.0</b>	m, unless COORD was specified in UNITS packet	x-coordinate of the profile with respect to the reference origin; this number will be ignored if a "relative" method is not being used
	<b>y-location</b> <b>31-40</b>	<b>F10.0</b>	m, unless COORD was specified in UNITS packet	y-coordinate of the profile with respect to the reference origin; this number will be ignored if a "relative" method is not being used
	<b>Number of points</b> <b>41-50</b>	<b>I10</b>	--	Number of profile points that follow; must be <u>&gt; 2</u> (or = -9 to clear all profile points)

Table VI-23 (Concluded)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<sup>3+</sup> <b>Profile Points</b>				
	Height 1-10	F10.0	m, unless HEIGHT was spe- cified in UNITS packet	Must be in increasing order; the first is assumed to be ground level and will be subtracted from all others to obtain each height relative to the ground  See the appropriate vertical method description in Section VI.A.7
	Value 11-20	F10.0	--	
<sup>4</sup> <b>Packet Terminator</b>		10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	<u>Column Number</u>				<u>FORTRAN Format</u>	
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	VERTICAL PROFILES					60A1
2+	Profile Description	subreg name	var name	x-loc	y-loc	# pts	2(10A1), 2F10.0, I10
3+	Profile	height	value				2F10.0
4	Packet Terminator	END					10A1

Exhibit VI-18. VERTICAL PROFILES Packet Rules

## VII M.E.B. FILE PREPARATION RULES AND EXAMPLES

Chapter VI presented the data preparation packets--the building blocks used to generate the M.E.B. data files. This chapter presents detailed instructions for the creation of each file, using the packets described above.

The preparation of some data files may require that other files be input, depending on the method used. In general, vertically varying files may require information from DIFFBREAK and REGIONTOP to define the vertical distribution of grid cells. The following is a suggested order of file preparation, assuming methods requiring the greatest file dependency. This order may be varied, depending on the requirements of the actual methods used.

- > DIFFBREAK.
- > EMISSIONS.
- > METSCALARS.
- > TEMPERATUR.
- > TERRAIN.
- > REGIONTOP (requiring DIFFBREAK).
- > TOPCONC (requiring DIFFBREAK and REGIONTOP).
- > AIRQUALITY (requiring DIFFBREAK, REGIONTOP, and TOPCONC).
- > BOUNDARY (requiring DIFFBREAK, REGIONTOP, and TOPCONC).
- > WIND (requiring DIFFBREAK, REGIONTOP, and TEMPERATUR).
- > PTSOURCE (requiring DIFFBREAK, REGIONTOP, METSCALARS, TEMPERATUR, and WIND).

The output from each program can include a printout of the input a title page, tables of methods, vertical methods, vertical profiles, a subregion grid, and grids of output variable values. In this manual,

a sample of each type of output is shown only for the AIRQUALITY file (Exhibit VII-2). For the other M.E.B. files, only the title page, methods tables, and an output grid sample are shown.

The data file preparation instructions are presented alphabetically by file name; each section includes:

- > A short description of the file.
- > The variables required and their units.
- > The computation methods that can be used.
- > The time variation represented.
- > The arrangement of packets in the input deck and special instructions for each one.
- > Additional input files required.
- > Sample input and output from a file generation run.

## A. AIRQUALITY

The AIRQUALITY file contains a three-dimensional matrix of concentrations for any number of chemical species. Its primary purpose is to provide the set of initial conditions for the beginning of the modeling period; however, it can also be used to prepare a three-dimensional distribution of measured data in any time interval for use in comparison studies. The program AIRQUL is used to create the AIRQUALITY file. AIRQUL requires subroutines from the libraries UTILITY and FILUTIL. Figure VII-1 is an information flow diagram illustrating the job control to be used. The following sections discuss the input to the AIRQUL program.

### 1. Variables

The output variables for the AIRQUALITY file are the species named in the CONTROL packet. Additional user-defined input variables (e.g., "reactive hydrocarbons") can be specified in the UNITS packet. The internal units for the concentrations of all species except AEROSOLS are parts per million (ppm); for AEROSOLS, the units are micrograms per cubic meter ( $\mu\text{g}/\text{m}^3$ ). The standard names for reactive species currently recognized by the Airshed Simulation Program are listed in Section IX.A. If any of these species does not appear on the AIRQUALITY file, the initial concentrations will default to a value defined in the CHEMPARAM file. If any additional species appear on AIRQUALITY, they will be ignored.

### 2. Methods

The methods that can be used to generate the AIRQUALITY file are:

- > CONSTANT
- > GRID VALUE
- > STATINTERP
- > POISSON

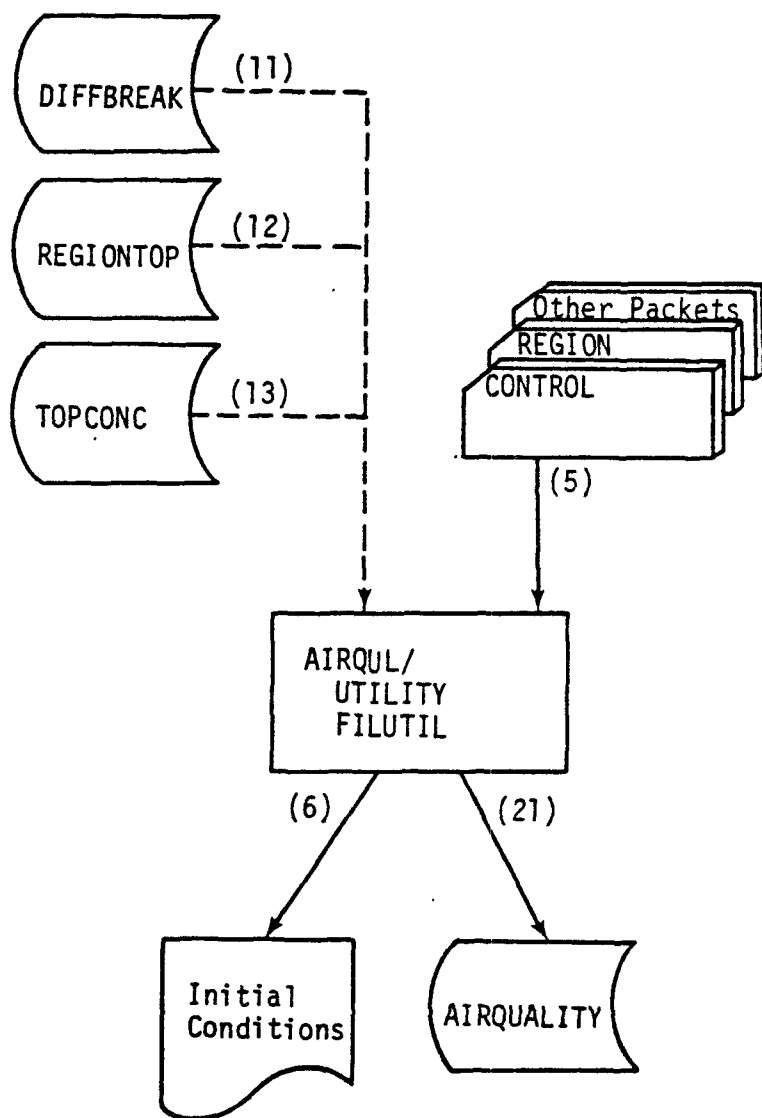


Figure VII-1. Information Flow Diagram for the AIRQUALITY File

- > SPLIT/COMB
- > E-WINTERP
- > N-SINTERP
- > USER.

These methods are discussed in detail in Section VI.A.6.

Since the AIRQUALITY file varies in the vertical direction, a vertical method must also be specified for each output variable in each sub-region. The vertical methods that can be used are:

- > CONSTANT
- > ABSPROFILE
- > RELPROFILE
- > ABSPROFRAT
- > RELPROFRAT
- > E-WINTERP
- > N-SINTERP
- > VERTUSER.

These vertical methods are discussed in detail in Section VI.A.7.

### 3. Time Variation

The concentrations on the AIRQUALITY file are used as initial conditions for the Airshed Simulation Program. The time span of the file must therefore include the beginning simulation time. Furthermore, the time interval containing the data to be used must begin at or before the simulation starting time and must end after the simulation starting time. In the example in Exhibit VII-1 in this chapter, the time interval for the initial concentrations is 0500 to 0501.

#### 4. Packets

Figure VII-2 shows the input deck structure in terms of the packets to be used. Each of these packets is described in detail in Section VI.B. Following are special input packet considerations for the file:

- > CONTROL
  - The file name on Card 2 must be AIRQUALITY.
  - The control variables to be specified on Cards 4 to 8 for AIRQUALITY are shown in Table VII-1.
  - The number of species must be greater than zero.
  - If there are input variables that do not appear as output variables, their number must appear as the number of user-defined variables. All such variables must also be named.
  - If data from measuring stations are to be used (methods STATINTERP or POISSON), the maximum number of such stations must be given.
  - The number of subregions must be at least one.
  - The maximum number of parameters must be sufficient to include all specifications of all parameters.
  - The vertical controls card (No. 7) must include maximum vertical parameter and profile entries as applicable.
  - The file unit assignment card (No. 8) must provide entries for the DIFFBREAK, REGIONTOP, and TOPCONC files if, and only if, these files are required by the vertical methods selected.
  - The beginning and ending dates and times should reflect the time variation considerations discussed in Section 3 above.
  - A set of output species names is required; their number must be the same as the entry in the first control parameter on Card 4. If either the ABSPROFRAT or RELPROFRAT vertical method is selected for any variable, the output species names specified here must match the species names on the TOPCONC file.

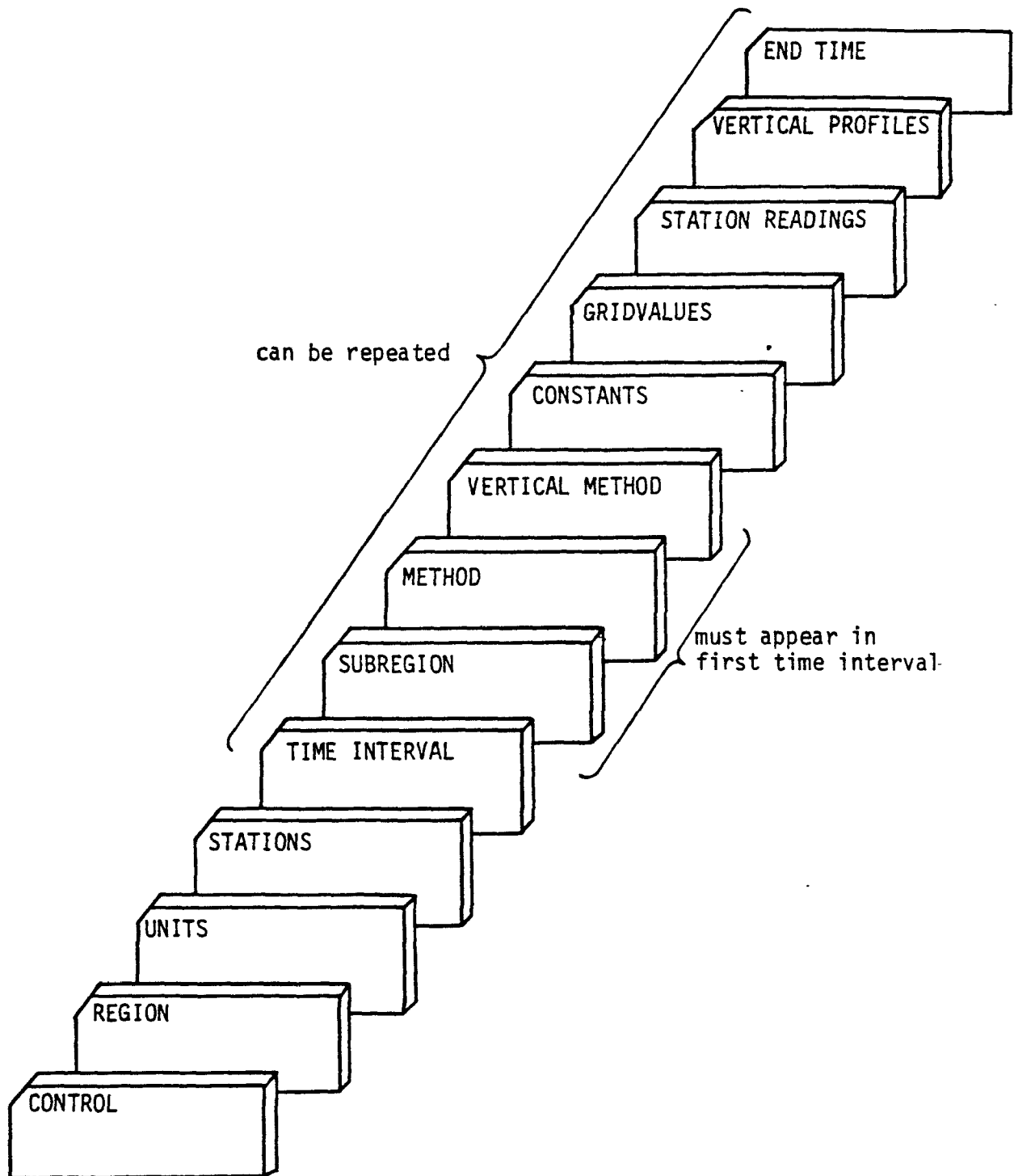


Figure VII-2. Input Card Deck Structure for Preparing the AIRQUALITY File

Table VII-1. CONTROL Packet Entries for  
the AIRQUALITY File

<u>Card Number</u>	<u>Entry</u>
4	Number of species Number of user-defined variables Number of stations Number of subregions Number of parameters Spare
5	Output file number Print input cards Print output grid Spare Spare Spare
6	Print units table Print station locations table Print regional grid Print methods table Print station values table Spare
7	Number of vertical parameters Number of heights in profile Print vertical methods table Print vertical profile tables Spare Spare
8	DIFFBREAK file number REGIONTOP file number TOPCONC file number Spare Spare Spare

- > REGION. This packet must follow the CONTROL packet. The vertical parameters must be provided for the AIRQUALITY file.
- > UNITS. This packet, if present, must follow the REGION packet. The UNITS packet must be provided if:
  - Any input variable will be input in other than internal units.
  - Any user-defined variables are specified.
  - COORD or HEIGHT unit conversions are to be used.The number of user-defined variables must not exceed the maximum specified in the CONTROL packet.
- > STATIONS. This packet is required if either of the methods STATINTERP or POISSON is specified. The number of stations listed must not exceed the maximum specified in the CONTROL packet.
- > TIME INTERVAL. One or more TIME INTERVAL packets must be present. The first time interval must begin at or before the beginning of the time span specified on Card 10 of the CONTROL packet. All time intervals must be contiguous and of nonzero length. Each TIME INTERVAL packet contains one or more of the following packets and ends with an ENDTIME card. Following the first time interval, only those data that are to be changed need be specified.
- > SUBREGION. The first time interval must contain a SUBREGION packet; the inclusion of this packet in other time intervals is optional. The number of subregions must not exceed the maximum specified in the CONTROL packet.
- > METHOD. A method must be provided for every variable--including user-defined variables--in every subregion in the first time interval. Methods can be changed in subsequent TIME INTERVAL packets if desired. Note that each parameter entry contributes to the overall parameter count; the total number of parameters must not exceed the maximum specified in the CONTROL packet.

- > VERTICAL METHOD. A vertical method must be provided for every variable--including user-defined variables--in every subregion in the first time interval. Vertical methods can be changed in subsequent TIME INTERVAL packets if desired. Note that each vertical parameter entry contributes to the overall vertical parameter count; the total must not exceed the maximum specified in the CONTROL packet.
- > CONSTANTS. If the method CONSTANT is assigned to any variable in the METHOD packet, the first time interval must contain a CONSTANTS packet. More than one CONSTANTS packet can appear in any time interval.
- > GRID VALUES. If the method GRID VALUE is assigned to any variable in the METHOD packet, the first time interval must contain a GRID VALUES packet. More than one GRID VALUES packet can appear in any time interval.
- > STATION READINGS. If either the POISSON or STATINTERP method is assigned to any species in the METHOD packet, the first time interval must contain a STATION READINGS packet. More than one STATION READINGS packet can appear in any time interval.
- > VERTICAL PROFILES. If any of the profile methods are assigned to any species in the VERTICAL METHOD packet, the first time interval must contain a VERTICAL PROFILES packet. There must be a vertical profile defined (or implied using ALL) for every variable in every subregion for which a profile method was specified. The number of height-value pairs in any single profile must not exceed the maximum specified in the CONTROL packet. More than one VERTICAL PROFILES packet can appear in any time interval.

## 5. Additional Input Files

If any vertical method besides CONSTANT was selected, the DIFFBREAK and REGIONTOP files must be input to AIRQUL. In addition, if either of

the vertical methods ABSPROFRAT or RELPROFRAT was selected, the TOPCONC file must also be input.

## 6. Examples

Exhibit VII-1 shows a sample input deck for AIRQUALITY file generation. The output from an AIRQUL run using that input deck is shown in Exhibit VII-2.

CONTROL		AIR QUALITY		INITIAL POLLUTANT CONCENTRATIONS		ST LOUIS		500		76195	
		11	1		25	1	30		30		5
		21	1		1						
				5		1		1			
				11	5	1	0				
NO					12	13					
NO2											
O3											
ETH											
OLE											
PAR											
CARB											
ARO											
BZA											
PAN											
CO											
		76195		500		76195		501			
END											
REGION											
		725000.	4252000.		15						
		-19000.	-16000.								
		4000.	4000.								
		17	22		4						
		2	2		0.	50.0		200.			
END											

**Exhibit VII-1.** Sample Input Deck for the AIRQUAL Program

STATIONS	101	19183.	27862.	10.
	102	17518.	34045.	10.
	103	22588.	30467.	10.
	104	22312.	25304.	10.
	105	18706.	24453.	10.
	106	13660.	25566.	10.
	107	15179.	30610.	10.
	108	23407.	39102.	10.
	109	30802.	27886.	10.
	110	22209.	20826.	10.
	111	13812.	20479.	10.
	112	6938.	28913.	10.
	113	12738.	37820.	10.
	114	19320.	45456.	10.
	115	32111.	45799.	10.
	116	37777.	38083.	10.
	117	35560.	20818.	10.
	118	18065.	11256.	10.
	119	4759.	18547.	10.
	120	-1921.	33909.	10.
	121	7414.	50376.	10.
	122	16631.	77223.	10.
	123	52320.	34378.	10.
	124	24275.	-15463.	10.

END	125	-27555.	30240.	10.
TIME INTERVAL	76195	500	76195	501
SUBREGION	A	1	1	-1
END				

Exhibit VII-1 (Continued)

METHOD	ND	POISSON	.00100	1.00	3
A	MAXITER	100			
ERRORTOL	.001000				
OMEGA	1.4				
A	H02	POISSON	.00100	1.00	3
A	MAXITER	100			
ERRORTOL	.001000				
OMEGA	1.4				
A	03	POISSON	.00100	1.00	3
A	MAXITER	100			
ERRORTOL	.001000				
OMEGA	1.4				
A	RHC	POISSON	.05000	4.00	3
A	MAXITER	100			
ERRORTOL	.050000				
OMEGA	1.4				
A	ETH	SPLIT/COMB	.00100	4.00	1
RHC	OLE	.0217	SPLIT/COMB	.00100	4.00
A	RHC	.0440	SPLIT/COMB	.00100	4.00
A	PAR	.6665	SPLIT/COMB	.01000	4.00
RHC	CARB	SPLIT/COMB	.00100	4.00	
A	RHC	.0810	SPLIT/COMB	.00100	4.00
A	ARO	.0248	SPLIT/COMB	.00010	4.00
RHC	A	BZA	SPLIT/COMB	.00000	4.00
RHC	PAN	.0001	CONSTANT	.00001	10
A	PAN	CO	POISSON	.10000	20.00
A	MAXITER	100			
ERRORTOL	.100000				
OMEGA	1.4				
END					
VERTICAL METHOD	ALL	ALL	RELPROFRAT	.0000	100.0
END					
CONSTANTS	PAN	.000010			
A	END				

Exhibit VII-1 (Continued)

STATION	READINGS	INITIAL COND	500
101	03	.024	
102	03	-9.000	
103	03	.027	
104	03	-9.000	
105	03	.002	
106	03	.022	
107	03	.030	
108	03	.027	
109	03	.036	
110	03	.035	
111	03	-9.000	
112	03	.026	
113	03	.034	
114	03	.036	
115	03	-9.000	
116	03	.035	
117	03	.036	
118	03	-9.000	
119	03	.029	
120	03	.019	
121	03	.002	
122	03	.043	
123	03	-9.000	
124	03	.024	
125	03	.003	
101	CO	.254	
102	CO	.050	
103	CO	.174	
104	CO	-9.000	
105	CO	.050	
106	CO	.185	
107	CO	.226	
108	CO	.055	
109	CO	.050	
110	CO	-9.000	
111	CO	.059	
112	CO	.267	
113	CO	.209	
114	CO	.109	
115	CO	.050	
116	CO	-9.000	
117	CO	.148	
118	CO	.050	
119	CO	.320	
120	CO	.239	
121	CO	.108	
122	CO	.050	

Exhibit VII-1 (Continued)

123	CO	.175
124	CO	.256
125	CO	.077
101	RHC	-9.000
102	RHC	.131
103	RHC	.086
104	RHC	.178
105	RHC	-9.000
106	RHC	.196
107	RHC	.146
108	RHC	.060
109	RHC	-9.000
110	RHC	-9.000
111	RHC	.082
112	RHC	.117
113	RHC	.050
114	RHC	-9.000
115	RHC	-9.000
116	RHC	-9.000
117	RHC	-9.000
118	RHC	-9.000
119	RHC	.050
120	RHC	.474
121	RHC	-9.000
122	RHC	-9.000
123	RHC	-9.000
124	RHC	-9.000
125	RHC	-9.000
101	NO	.002
102	NO	.002
103	NO	.003
104	NO	.002
105	NO	.002
106	NO	.002
107	NO	.003
108	NO	.005
109	NO	.002
110	NO	.002
111	NO	.002
112	NO	.002
113	NO	.003
114	NO	.002
115	NO	.002
116	NO	.002
117	NO	.002

Exhibit VII-1 (Concluded)

```
.002  
NO NO NO .003  
119 NO NO .002  
120 NO NO .002  
121 NO NO .002  
122 NO NO .002  
123 NO NO .002  
124 NO NO .002  
125 NO NO .002  
101 NO NO .020  
102 NO NO .004  
103 NO NO .010  
104 NO NO -9.000  
105 NO NO -9.000  
106 NO NO .002  
107 NO NO .011  
108 NO NO .010  
109 NO NO .002  
110 NO NO .005  
111 NO NO .018  
112 NO NO .016  
113 NO NO .010  
114 NO NO .008  
115 NO NO .002  
116 NO NO .002  
117 NO NO .002  
118 NO NO .003  
119 NO NO .011  
120 NO NO .022  
121 NO NO .005  
122 NO NO .002  
123 NO NO .002  
124 NO NO .003  
125 NO NO .002  
END  
VERTICAL PROFILE  
A ALL 35296. 5030.  
0. 0.  
100. 0.  
120. 1.  
500. 1.  
END  
ENDTIME
```

**PROCESSING OF INPUT DATA FOR THE PREPARATION OF FILE AIRQUALITY**

**INITIAL POLLUTANT CONCENTRATIONS ST LOUIS 500 76195**

NUMBER OF USER-DEFINED VARIABLES = 1  
NUMBER OF STATIONS = 25  
NUMBER OF SUBREGIONS = 1  
NUMBER OF METHOD PARAMETERS = 30  
NUMBER OF VERTICAL METHOD PARAMETERS = 5  
NUMBER OF VERTICAL PROFILE HEIGHTS = 5  
PROGRAM OUTPUT WILL BE PLACED ON FILE NUMBER 21  
INPUT DATA FOR DIFFUSION BREAK WILL BE OBTAINED FROM FILE NUMBER 11  
INPUT DATA FOR REGION TOP WILL BE OBTAINED FROM FILE NUMBER 12  
INPUT DATA FOR TOP CONCENTRATIONS WILL BE OBTAINED FROM FILE NUMBER 13  
INPUT CARDS WILL BE PRINTED  
OUTPUT GRID WILL BE PRINTED  
UNIT TABLE WILL NOT BE PRINTED  
STATION LOCATIONS WILL NOT BE PRINTED  
SUBREGION GRID WILL NOT BE PRINTED  
METHOD TABLE WILL BE PRINTED  
STATION VALUES WILL NOT BE PRINTED  
VERTICAL METHOD TABLE WILL BE PRINTED  
VERTICAL PROFILE VALUES WILL NOT BE PRINTED  
BEGINNING DATE OF FILE 76195 BEGINNING TIME OF FILE 5.00  
ENDING DATE OF FILE 76195 ENDING TIME OF FILE 5.02

**Exhibit VII-2. Sample Output From the AIRQUL Program**

A TOTAL OF

11 SPECIES NAMES WERE READ, THE SPECIES NAMES ARE --

NO	NO2	
03	ETH	
OLE		
PAR		
CARB		
ARO		
BZA		
PAs		
CO		
REGION		
72:00.	425200.	15
-19000.	-16000.	
4000.00	4000.00	
:7	22	4
2	2	.00
		50.00
		200.00
END		

Exhibit VII-2 (Continued)

**REGIONAL DEFINITION**

UTM ORIGIN X-COORDINATE	725000.
UTM ORIGIN Y-COORDINATE	4252000.
UTM ORIGIN ZONE NUMBER	15
REGION ORIGIN X-COORDINATE	-19000.
REGION ORIGIN Y-COORDINATE	-16000.
CELL LENGTH IN X-DIRECTION	4000.00
CELL LENGTH IN Y-DIRECTION	4000.00
NUMBER OF CELLS IN X-DIRECTION	17
NUMBER OF CELLS IN Y-DIRECTION	22
NUMBER OF CELLS IN Z-DIRECTION	4
NUMBER OF CELLS IN LOWER VERTICAL LAYER .	2
NUMBER OF CELLS IN UPPER VERTICAL LAYER	2
HEIGHT OF SURFACE LAYER	.00
MINIMUM CELL HEIGHT IN LOWER LAYER	50.00
MINIMUM CELL HEIGHT IN UPPER LAYER	200.00
UNITS	
RHC	PPH
END	.0000
	.0000
	.0000

**Exhibit VII-2 (Continued)**

STATIONS						
101	102	17518.	34045.	10.00		
103	22586.	30467.	10.00			
104	22312.	25304.	10.00			
105	18706.	24453.	10.00			
106	13660.	25566.	10.00			
107	15179.	30610.	10.00			
108	23407.	39102.	10.00			
109	30802.	27886.	10.00			
110	22209.	20626.	10.00			
111	13812.	20479.	10.00			
112	8938.	28913.	10.00			
113	12738.	37820.	10.00			
114	19320.	45456.	10.00			
115	32111.	45799.	10.00			
116	37777.	38083.	10.00			
117	35560.	20818.	10.00			
118	18065.	11256.	10.00			
119	4759.	18547.	10.00			
120	-1921.	33909.	10.00			
121	7414.	50376.	10.00			
122	16631.	77223.	10.00			
123	52320.	34378.	10.00			
124	24275.	-15463.	10.00			
125	-27555.	30240.	10.00			
	0.	0.	.00			
				END		

Exhibit VII-2 (Continued)

TIME INTERVAL	500	76195	501	
SUBREGION				
A	1	1	-1	
END	0	0	0	
METHOD	NO	POISSON	.0010	
MAXITER	1000.0000		1.0000	
ERRORTOL	.0010			
OMEGA	1.4000			
A	NO2	POISSON	.0010	
MAXITER	1000.0000		1.0000	
ERRC2TOL	.0010			
OMEGA	1.4000			
A	03	POISSON	.0010	
MAXITER	1000.0000		1.0000	
ERRORTOL	.0010			
OMEGA	1.4000			
A	RHC	POISSON	.0500	
MAXITER	1000.0000		4.0000	
ERRORTOL	.0500			
OMEGA	1.4000			
A	E7H	SPLIT/COMB	.0010	
RHC	OLE	SPLIT/COMB	.0010	
A	RHC	PAR	SPLIT/COMB	.0100
A	RHC	CARB	SPLIT/COMB	.0010
RHC	A	ARO	SPLIT/COMB	.0001
A	RHC	BZA	SPLIT/COMB	.0000
A	RHC	PAN	CONSTANT	.0000
A	A	CO	POISSON	.1000
MAXITER	1000.0000		20.0000	
ERRORTOL	.1000			
OMEGA	1.4000			
END				

METHODS TABLE								
SUBREGION	VARIABLE	METHOD	MIN. VALUE	MAX. VALUE	PARAMETER	PAR.	VALUE	
A	N0	POISSON	.0010	1.0000	MAXITER ERRORTOL OMEGA	1000.0000 .0010 1.4000		
A	N02	POISSON	.0010	1.0000	MAXITER ERRORTOL OMEGA	1000.0000 .0010 1.4000		
A	03	POISSON	.0010	1.0000	MAXITER ERRORTOL OMEGA	1000.0000 .0010 1.4000		
A	ETH	SPLIT/COMB	.0010	4.0000	MAXITER ERRORTOL OMEGA	1000.0000 .0010 1.4000		
A	OLE	SPLIT/COMB	.0010	4.0000	RHC	.0217		
A	PAR	SPLIT/COMB	.0100	4.0000	RHC	.0440		
A	CARB	SPLIT/COMB	.0010	4.0000	RHC	.6665		
A	ARO	SPLIT/COMB	.0001	4.0000	RHC	.0810		
A	BZA	SPLIT/COMB	.0000	4.0000	RHC	.0248		
A	PAN CO	CONSTANT POISSON	.0000 .1000	.1000 20.0000	MAXITER ERRORTOL OMEGA	1000.0000 .1000 1.4000		
A	RHC	POISSON	.0500	4.0000	MAXITER ERRORTOL OMEGA	1000.0000 .0500 1.4000		
VERTICAL METHOD		RELPROFRAT	.0000 .0000	100.0000 .0000	0	0		
ALL END								

Exhibit VII-2 (Continued)

		METHODS TABLE			
SUBREGION	VARIABLE	METHOD	MIN. VALUE	MAX. VALUE	PARAMETER
					PAR. VALUE
A A A A A A A A A A A A A A A A	NO NO2 O3 ETH OLE PAR CARB ARO BZA PAN CO RHC	RELPROFRAT	.0000 .0000 .0000 .0000 .0000 .0000 .0000 .0000 .0000 .0000 .0000 .0000 .0000 .0000 .0000 .0000	100.0000 100.0000 100.0000 100.0000 100.0000 100.0000 100.0000 100.0000 100.0000 100.0000 100.0000 100.0000 100.0000 100.0000 100.0000	
A	CONSTANTS PAN		.0600		
END			.0000		

Exhibit VII-2 (Continued)

Exhibit VII-2 (Continued)

STATION	READINGS	INITIAL COND	500
101	03	.0240	
102	03	-9.0300	
103	03	.0270	
104	03	-9.0000	
105	03	.0020	
106	03	.0220	
107	03	.0300	
108	03	.0270	
109	03	.0380	
110	03	.0350	
111	03	-9.0000	
112	03	.0280	
113	03	.0340	
114	03	.0360	
115	03	-9.0000	
116	03	.0150	
117	03	.0160	
118	03	-9.0000	
119	03	.0290	
120	03	.0190	
121	03	.0020	
122	03	.0930	
123	03	-9.0000	
124	03	.0240	
125	03	.0010	
101	CO	.5540	
102	CO	.0500	
103	CO	.1740	
104	CO	-9.0000	
105	CO	.0500	
106	CO	.1650	
107	CO	.2260	
108	CO	.0550	
109	CO	.0500	
110	CO	-9.0000	
111	CO	.0590	
112	CO	.2670	
113	CO	.2090	
114	CO	.1090	
115	CO	.0500	
116	CO	-9.0000	
117	CO	.1480	
118	CO	.0500	
119	CO	.1200	
120	CO	.2390	
121	CO	.1060	
122	CO	.0500	
123	CO	.1750	
124	CO	.2560	
125	CO	.0770	

<b>RHC</b>	<b>-9.0000</b>
101 RHC	.1310
102 RHC	.0860
103 RHC	.1780
104 RHC	-9.0000
105 RHC	.1960
106 RHC	.1460
107 RHC	.0600
108 RHC	-9.0000
109 RHC	-9.0000
110 RHC	.0820
111 RHC	.1170
112 RHC	.0500
113 RHC	-9.0000
114 RHC	.0500
115 RHC	-9.0000
116 RHC	.0500
117 RHC	-9.0000
118 RHC	-9.0000
119 RHC	.4740
120 RHC	.0500
121 RHC	-9.0000
122 RHC	-9.0000
123 RHC	-9.0000
124 RHC	-9.0000
125 RHC	-9.0000
101 NO	.0020
102 NO	.0020
103 NO	.0030
104 NO	.0020
105 NO	-9.0000
106 NO	.0020
107 NO	.0030
108 NO	.0050
109 NO	.0020
110 NO	.0020
111 NO	.0020
112 NO	.0020
113 NO	.0030
114 NO	.0020
115 NO	.0020
116 NO	.0020
117 NO	.0020
118 NO	.0020

Exhibit VII-2 (Continued)

119 NO .0020  
120 NO .0030  
121 NO .0020  
122 NO .0020  
123 NO .0020  
124 NO .0020  
125 NO .0020  
101 NO2 -.0200  
102 NO2 -.0040  
103 NO2 .0100  
104 NO2 -.9.0000  
105 NO2 -.9.0000  
106 NO2 .0020  
107 NO2 .0110  
108 NO2 .0100  
109 NO2 .0020  
110 NO2 .0050  
111 NO2 .0180  
112 NO2 .0160  
113 NO2 .0100  
114 NO2 .0080  
115 NO2 .0020  
116 NO2 .0020  
117 NO2 .0020  
118 NO2 .0030  
119 NO2 .0110  
120 NO2 .0220  
121 NO2 .0050  
122 NO2 .0020  
123 NO2 .0020  
124 NO2 .0030  
125 NO2 .0020  
ENU .0000  
END .0000  
ENDTIME 0

A VERTICAL PROFILE  
A ALL 35296.0000 5038.0000 4  
0. .0000  
100. .0000  
120. 1.0000  
500. 1.0000

GRID CELL	FILE NAME -- AIRQUALITY	INITIAL POLLUTANT CONCENTRATIONS										VERF.	LEVEL 1	START DATE/TIME	76195 5.00	STOP DATE/TIME	76195 5.02	PAGE 1
		1	2	3	4	5	6	7	8	9	10							
22	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020
21	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020
20	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0021	.0021	.0021	.0021
19	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0021	.0021	.0021	.0021	.0021
18	.0020	.0020	.0020	.0020	.0020	.0021	.0021	.0021	.0021	.0021	.0021	.0021	.0021	.0021	.0021	.0021	.0021	.0021
17	.0020	.0020	.0020	.0020	.0021	.0021	.0021	.0021	.0021	.0021	.0021	.0021	.0021	.0022	.0022	.0022	.0022	.0022
16	.0020	.0020	.0020	.0021	.0021	.0022	.0022	.0022	.0022	.0022	.0023	.0023	.0023	.0023	.0023	.0023	.0025	.0024
15	.0020	.0020	.0020	.0021	.0022	.0023	.0023	.0024	.0024	.0024	.0026	.0026	.0026	.0026	.0026	.0026	.0026	.0029
14	.0020	.0020	.0022	.0022	.0023	.0025	.0025	.0025	.0025	.0025	.0030	.0030	.0030	.0030	.0032	.0032	.0050	.0034
13	.0020	.0020	.0022	.0022	.0024	.0030	.0030	.0026	.0024	.0024	.0026	.0026	.0026	.0026	.0026	.0026	.0033	.0030
12	.0020	.0020	.0020	.0021	.0023	.0024	.0024	.0023	.0023	.0020	.0025	.0025	.0025	.0025	.0025	.0025	.0030	.0026
11	.0020	.0020	.0020	.0020	.0021	.0022	.0021	.0021	.0021	.0021	.0022	.0022	.0022	.0022	.0020	.0020	.0020	.0021
10	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0021	.0021	.0020	.0020	.0020	.0020	.0020	.0020
9	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020
8	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020
7	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020
6	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020
5	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020
4	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020
3	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020
2	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020
1	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020

FILE NAME -- AIRQUALITY INITIAL POLLUTANT CONCENTRATIONS ST LOUIS 500 76195  
 AIR QUALITY VALUES (IN PARTS PER MILLION) SPECIES NO VERT. LEVEL 1 START DATE/TIME 76195 5.00 STOP DATE/TIME 76195 5.02

GRID CELL	13	14	15	16	17
22	.0020	.0020	.0020	.0020	.0020
21	.0020	.0020	.0020	.0020	.0020
20	.0021	.0021	.0020	.0020	.0020
19	.0021	.0021	.0021	.0020	.0020
18	.0021	.0021	.0021	.0020	.0020
17	.0021	.0021	.0021	.0020	.0020
16	.0020	.0021	.0021	.0020	.0020
15	.0025	.0022	.0021	.0020	.0020
14	.0027	.0023	.0020	.0020	.0020
13	.0026	.0023	.0021	.0020	.0020
12	.0023	.0022	.0021	.0020	.0020
11	.0020	.0020	.0020	.0020	.0020
10	.0026	.0020	.0020	.0020	.0020
9	.0020	.0020	.0020	.0020	.0020
8	.0020	.0020	.0020	.0020	.0020
7	.0020	.0020	.0020	.0020	.0020
6	.0020	.0020	.0020	.0020	.0020
5	.0020	.0020	.0020	.0020	.0020
4	.0020	.0020	.0020	.0020	.0020
3	.0020	.0020	.0020	.0020	.0020
2	.0020	.0020	.0020	.0020	.0020
1	.0020	.0020	.0020	.0020	.0020

Exhibit VII-2 (Concluded)

## B. BOUNDARY

The BOUNDARY file contains a time-invariant definition of the boundaries of the region to be modeled and time-varying matrices of pollutant concentrations in each external boundary cell. These boundary values are used by the Airshed Simulation Program to represent pollutant concentrations passing over the boundary into the modeling region. The program BNDARY is used to create the BOUNDARY file. BNDARY requires subroutines from the libraries UTILITY and FILUTIL. Figure VII-3 is an information flow diagram illustrating the job control to be used. In the following sections we discuss the input to the BNDARY program.

### 1. Variables

The output variables for the BOUNDARY file are the species named in the CONTROL packet. Additional user-defined input variables (e.g., "reactive hydrocarbons") can be specified in the UNITS packet. The internal units for the concentrations of all species except AEROSOLS are parts per million (ppm); for AEROSOLS, the units are micrograms per cubic meter ( $\mu\text{g}/\text{m}^3$ ). The standard names for reactive species recognized by the Airshed Simulation Program are listed in Section IX.A. If any of these species does not appear on the BOUNDARY file, the boundary concentrations will default to a value defined in the CHEMPARAM file. If any additional species appear on BOUNDARY, they will be ignored.

### 2. Methods

The methods that can be used to generate the BOUNDARY file are:

- > CONSTANT
- > BOUNDVALUE
- > SPLIT/COMB
- > USER.

The methods CONSTANT and BOUNDVALUE are equivalent. All these methods are discussed in detail in Section VI.A.6.

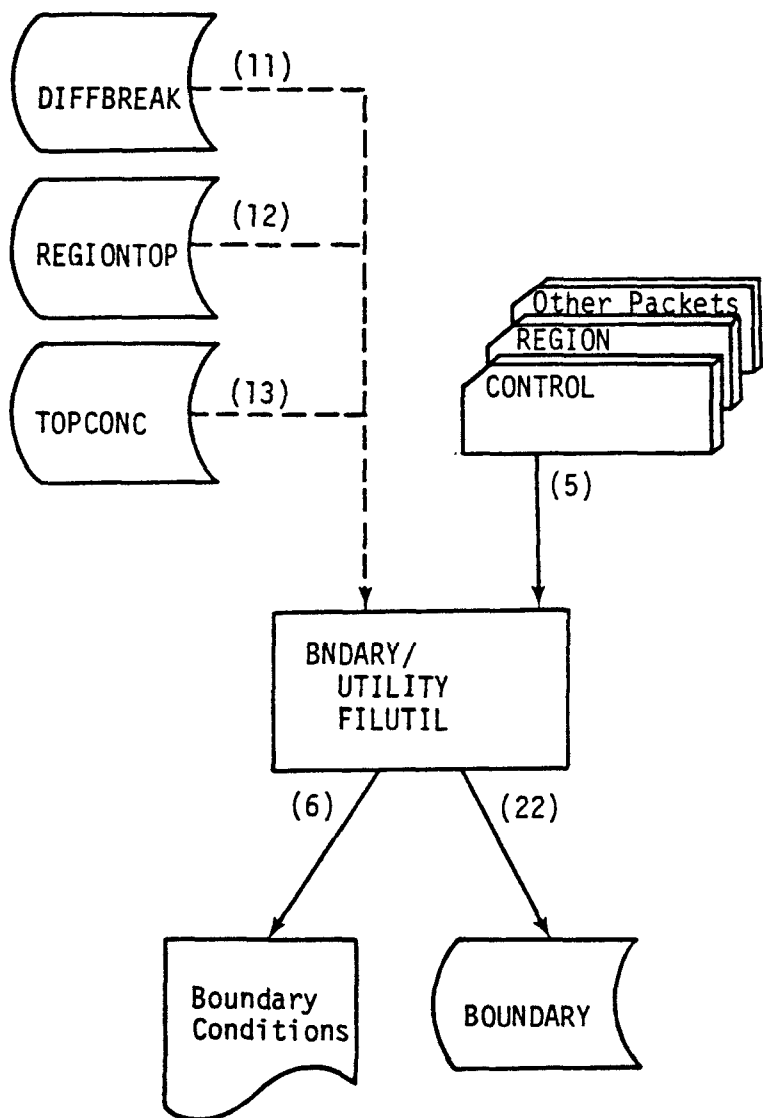


Figure VII-3. Information Flow Diagram for the BOUNDARY File

Since the BOUNDARY file varies in the vertical direction, a vertical method must also be specified for each output variable in each subregion. The vertical methods that can be used are:

- > CONSTANT
- > ABSPROFILE
- > RELPROFILE
- > ABSPROFRAT
- > RELPROFRAT
- > VERTUSER.

These vertical methods are discussed in detail in Section VI.A.7.

### 3. Time Variation

The time span of the BOUNDARY file must include the entire time span of the simulation runs for which it is to be used. Boundary concentrations are considered to be constant during each time interval.

### 4. Packets

Figure VII-4 shows the input deck structure in terms of the packets to be used. Each of these packets is described in detail in Section VI.B. Following are special input packet considerations for the BOUNDARY file:

- > CONTROL
  - The file name on Card 2 must be BOUNDARY.
  - The control variables to be specified on Cards 4 to 8 for BOUNDARY are shown in Table VII-2.
  - The number of species must be greater than zero.
  - If there are input variables that do not appear as output variables, their number must appear as the number of user-defined variables. All such variables must also be named in the UNITS packet.

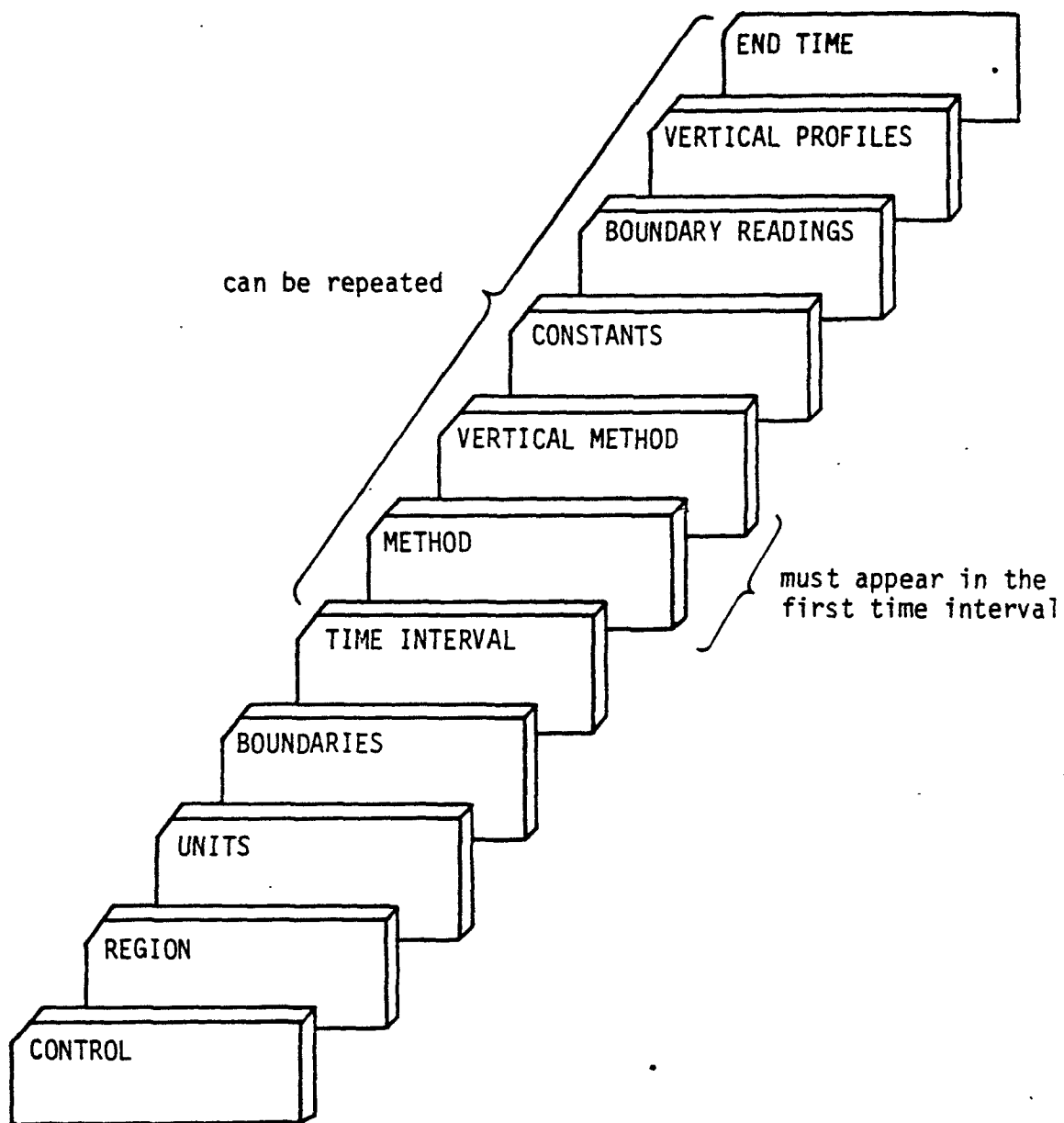


Figure VII-4. Input Card Deck Structure for Preparing the BOUNDARY File

**Table VII-2. CONTROL Packet Entries for the BOUNDARY File**

<u>Card Number</u>	<u>Entry</u>
4	Number of species Number of user-defined variables Number of boundary line segments Spare Number of parameters Spare
5	Output file number Print input cards Print output boundary values Spare Spare Spare
6	Print units table Print boundary line segment locations table Print regional grid Print methods table Print boundary values table Spare
7	Number of vertical parameters Number of heights in profile Print vertical methods table Print vertical profile tables Spare Spare
8	DIFFBREAK file number REGIONTOP file number TOPCONC file number Spare Spare Spare

- The number of boundary line segments must be at least three.
  - The maximum number of parameters must be sufficient to include all specifications of all parameters.
  - The vertical controls card (No. 7) must include maximum vertical parameter and profile entries as applicable.
  - The file unit assignment card (No. 8) must provide entries for the DIFFBREAK, REGIONTOP, and TOPCONC files if, and only if, these files are required by the vertical methods selected.
  - The beginning and ending dates and times should reflect the time variation considerations discussed in Section 3 above.
  - A set of output species names is required; their number must be the same as the entry in the first control parameter on Card 4. If either the ABSPROFRAT or RELPROFRAT vertical method is selected for any variable, the output species names specified here must match the species names on the TOPCONC file.
- > REGION. This packet must follow the CONTROL packet. The vertical parameters must be provided for the BOUNDARY file.
- > UNITS. This packet, if present, must follow the REGION packet. The UNITS packet must be provided if:
- Any input variable will be input in other than internal units.
  - Any user-defined variables are specified.
  - COORD or HEIGHT unit conversions are to be used.
- The number of user-defined variables must not exceed the maximum specified in the CONTROL packet.
- > BOUNDARIES. This packet is required; it names the line segments that define the boundaries of the region. The number of line segments specified must equal the number specified in the CONTROL packet.

- > TIME INTERVAL. One or more TIME INTERVAL packets must be present. The first time interval must begin at or before the beginning of the time span specified on Card 10 of the CONTROL packet. All time intervals must be contiguous and of nonzero length. Each TIME INTERVAL packet contains one or more of the following packets and ends with an ENDTIME card. Following the first time interval, only those data that are to be changed need be specified.
- > METHOD. A method must be provided for every variable--including user-defined variables--for every boundary line segment in the first time interval. In the METHOD packet, the boundary line segment name is entered in the "subregion" field. Methods can be changed in subsequent TIME INTERVAL packets if desired. Note that each parameter entry contributes to the overall parameter count; the total number of parameters must not exceed the maximum specified in the CONTROL packet.
- > VERTICAL METHOD. A vertical method must be provided for every variable--including user-defined variables--for every boundary line segment in the first time interval. In the VERTICAL METHOD packet, the boundary line segment name is entered in the "subregion" field. Vertical methods can be changed in subsequent TIME INTERVAL packets if desired. Note that each vertical parameter entry contributes to the overall vertical parameter count; the total must not exceed the maximum specified in the CONTROL packet.
- > CONSTANTS. If the method CONSTANT is assigned to any variable in the METHOD packet, the first time interval must contain a CONSTANTS packet. More than one CONSTANTS packet can appear in any time interval.
- > BOUNDARY READINGS. If the method BOUNDVALUE is assigned to any variable in the METHOD packet, the first time interval must contain a BOUNDARY READINGS packet. More than one BOUNDARY READINGS packet can appear in any time interval.

> VERTICAL PROFILES. If any of the profile methods are assigned to any species in the VERTICAL METHOD packet, the first time interval must contain a VERTICAL PROFILES packet. There must be a vertical profile defined (or implied using ALL) for every variable for every boundary line segment for which a profile method was specified. In the VERTICAL PROFILES packet, the boundary line segment name is entered in the "subregion" field. The number of height-value pairs in any single profile must not exceed the maximum specified in the CONTROL packet. More than one VERTICAL PROFILES packet can appear in any time interval.

## 5. Additional Input Files

If any vertical method besides CONSTANT was selected, the DIFFBREAK and REGIONTOP files must be input to BNDARY. In addition, if the vertical method ABSPROFRAT or RELPROFRAT was selected, the TOPCONC file must also be input.

## 6. Examples

Exhibit VII-3 shows a sample input deck for BOUNDARY file generation. The output from a BNDARY run using that input deck is shown in Exhibit VII-4.

```

CONTROL
BOUNDARY DEFINITION AND CONCENTRATIONS      10
    11   1   4   1
    22   1   1   1
    1   1   1   1
    5   5   0
    11  12  13
NO
NO2
O3
ETH
OLE
PAR
CARB
ARO
BZA
PAN
CO    76195   500   76195   2400
END
REGION
725000. 4252000.   15
-19000. -16000.
4000.   4000.
17     22     4
2     2     0.
50.   200.
END
UNITS
PPM
END
1.
BOUNDRIES
TOP    -13000.   66000.   43000.   66000.
BOTTOM -13000.  -10000.   43000.  -10000.
LEFT   -13000.  -10000.  -13000.  66000.
RIGHT  43000.  -10000.  43000.  66000.
END

```

TIME	INTERVAL	500	76195	600
METHOD				
ALL	NO	BOUNDFVALUE	0.0	90.
ALL	NO2	BOUNDFVALUE	0.0	90.
ALL	O3	BOUNDFVALUE	0.0	90.
ALL	RHC	BOUNDFVALUE	0.0	90.
ALL	PAN	BOUNDFVALUE	0.0	90.
ALL	CO	BOUNDFVALUE	0.0	90.
ALL	ETH	SPLIT/COMB	0.0	10.
RHC	OLE	SPLIT/COMB	0.0	10.
ALL	RHC	SPLIT/COMB	0.001	10.
RHC	PAR	SPLIT/COMB	0.001	10.
RHC	CARB	SPLIT/COMB	0.0	10.
ALL	RHC	SPLIT/COMB	0.0	10.
ALL	ARO	SPLIT/COMB	0.0	10.
RHC	BZA	SPLIT/COMB	0.0	10.
END				
VERTICAL METHOD				
ALL	ALL	RELPROFRAT	.00000	100.0
END				

Exhibit VII-3 (Continued)

**Exhibit VII-3 (Concluded)**

<b>BOUNDARYREADINGS</b>		<b>PAN</b>		<b>.00001</b>	
ALL	PAN	03	.023	026	.023
TOP	RIGHT	03	.026	026	.026
BOTTOM	LEFT	03	.023	023	.023
TOP	TOP	CO	.121	.121	.121
RIGHT	RIGHT	CO	.192	.192	.192
BOTTOM	LEFT	CO	.121	.121	.121
TOP	TOP	RHC	.079	.079	.079
RIGHT	RIGHT	RHC	.050	.050	.050
BOTTOM	LEFT	RHC	.050	.079	.079
TOP	RIGHT	NO	.003	.002	.002
RIGHT	BOTTOM	NO	.002	.002	.002
BOTTOM	LEFT	NO	.003	.003	.003
TOP	TOP	NO2	.007	.007	.007
RIGHT	RIGHT	NO2	.002	.002	.002
BOTTOM	BOTTOM	NO2	.002	.002	.002
LEFT	LEFT	NO2	.007	.007	.007
EIO					
<b>VERTICAL PROFILE</b>					
ALL	ALL	0.	0.	0.	0.
		102.	0.	122.	1.
		5000.	1.		
END	ENDTIME				

PROCESSING OF INPUT DATA FOR THE PREPARATION OF FILE BOUNDARY

BOUNDARY DEFINITION AND CONCENTRATIONS

NUMBER OF USER-DEFINED VARIABLES = 1  
NUMBER OF BOUNDARY LINES = 4  
NUMBER OF METHOD PARAMETERS = 10  
NUMBER OF VERTICAL METHOD PARAMETERS = 5  
NUMBER OF VERTICAL PROFILE HEIGHTS = 5  
PROGRAM OUTPUT WILL BE PLACED ON FILE NUMBER 22  
INPUT DATA FOR DIFFUSION BREAK WILL BE OBTAINED FROM FILE NUMBER 11  
INPUT DATA FOR REGION TOP WILL BE OBTAINED FROM FILE NUMBER 12  
INPUT DATA FOR TOP CONCENTRATIONS WILL BE OBTAINED FROM FILE NUMBER 13  
INPUT CARDS WILL BE PRINTED  
OUTPUT GRID WILL BE PRINTED  
UNIT TABLE WILL BE PRINTED  
BOUNDARY LINE TABLE WILL BE PRINTED  
BOUNDARY LINE MAP WILL BE PRINTED  
INPUT VALUES METHOD TABLE WILL NOT BE PRINTED  
BOUNDARY VALUES WILL NOT BE PRINTED  
VERTICAL METHOD TABLE WILL NOT BE PRINTED  
VERTICAL PROFILE VALUES WILL NOT BE PRINTED  
BEGINNING DATE OF FILE 76195 BEGINNING TIME OF FILE 5.00  
ENDING DATE OF FILE 76195 ENDING TIME OF FILE 24.00

Exhibit VII-4. Sample Output from BNDARY Program

A TOTAL OF 11 SPECIES NAMES WERE READ, THE SPECIES NAMES ARE --

NO	
ND2	
03	
ETH	
OLE	
PAR	
CARB	
ARO	
BZA	
PAN	
CO	
REGION	
725000.	4252000.
-19000.	-16000.
4000.00	4000.00
17	22
	4
	.00
2	200.00
END	

Exhibit VII-4 (Continued)

UNIT CONVERSION TABLE

VARIABLE NAME	UNIT NAME	MULTIPLICATIVE FACTOR	ADITIVE FACTOR
NO		-9.00000000	-9.00000000
NO2		-9.00000000	-9.00000000
O3		-9.00000000	-9.00000000
ETH		-9.00000000	-9.00000000
OLE		-9.00000000	-9.00000000
PAR		-9.00000000	-9.00000000
CARB		-9.00000000	-9.00000000
ARO		-9.00000000	-9.00000000
BZA		-9.00000000	-9.00000000
PAN		-9.00000000	-9.00000000
CO		-9.00000000	-9.00000000
RHC	PPM	1.00000000	-9.00000000
COORD		-9.00000000	-9.00000000
HEIGHT		-9.00000000	-9.00000000
BOUNDARIES			
TOP	-13000.	66000.	43000.
BOTTOM	-13000.	-10000.	43000.
LEFT	-13000.	-10000.	-13000.
RIGHT	43000.	-10000.	43000.
END	0.	0.	0.

Exhibit VII-4 (Continued)

BOUNDARY LINE LOCATION TABLE

B. LINE NAME	X-BEGINNING(GRID SQ)	X-ENDING (GRID SQ)	Y-BEGINNING (GRID SQ)	Y-ENDING(GRID SQ)
TOP	1.50	15.50	20.50	20.50
BOTTOM	1.50	15.50	1.50	1.50
LEFT	1.50	1.50	1.50	20.50
RIGHT	15.50	15.50	1.50	20.50
***** NOTE ***** BOUNDARY LINES LEFT AND BOTTOM		INTERSECT IN CELL (		
THIS CELL WILL TAKE ITS VALUE FROM THE LATTER LINE		2,	2).	
***** NOTE ***** BOUNDARY LINES RIGHT AND BOTTOM		INTERSECT IN CELL (	16,	2).
THIS CELL WILL TAKE ITS VALUE FROM THE LATTER LINE				
***** NOTE ***** BOUNDARY LINES LEFT AND TOP		INTERSECT IN CELL (	2,	21).
THIS CELL WILL TAKE ITS VALUE FROM THE LATTER LINE				
***** NOTE ***** BOUNDARY LINES RIGHT AND TOP		INTERSECT IN CELL (	16,	21).
THIS CELL WILL TAKE ITS VALUE FROM THE LATTER LINE				

PAGE 1

FILE NAME -- BOUNDARY	BOUNDARY DEFINITION AND CONCENTRATIONS																VERT.	LEVEL 0	START DATE/TIME	76195	5.00 STOP DATE/TIME	76195	5.00
MAP OF BOUNDARY LINE NUMBERS ON THE REGIONAL GRID	1	2	3	4	5	6	7	P	9	10	11	12	13	14	15	16	17						
GRID CELL	22	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	
21	0	1	1	1	1	1	1	1	1	1	1	1	1	1	1	1	0						
20	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
19	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
18	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
17	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
16	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
15	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
14	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
13	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
12	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
11	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
10	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
9	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
8	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
7	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
6	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
5	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
4	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
3	0	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						
2	0	2	2	2	2	2	2	2	2	2	2	2	2	2	2	2	0						
1	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0						

## BOUNDARY LINE IDENTIFICATION

NUMBER	NAME	TIME INTERVAL	76195	500	76195	600	DATE = 76195	TIME = 5.00 **
1	TOP	** READING DIFFBREAK FILE :						
2	BOTTOM	** READING REGTOP FILE :						
3	LEFT	** READING TOPCONC FILE :						
4	RIGHT	METHOD	NO	NO	NO	NO	TIME = 5.00 **	
ALL	NO	NO	NO	NO	NO	NO		
ALL	NO2	NO2	NO2	NO2	NO2	NO2		
ALL	03	03	03	03	03	03		
ALL	RHC	RHC	RHC	RHC	RHC	RHC		
ALL	PAN	PAN	PAN	PAN	PAN	PAN		
ALL	CO	CO	CO	CO	CO	CO		
ALL	ETH	ETH	ETH	ETH	ETH	ETH		
RHC								
ALL	OLE	OLE	OLE	OLE	OLE	OLE		
RHC								
ALL	PAR	PAR	PAR	PAR	PAR	PAR		
RHC								
ALL	CARB	CARB	CARB	CARB	CARB	CARB		
RHC								
ALL	ARO	ARO	ARO	ARO	ARO	ARO		
RHC								
ALL	BZA	BZA	BZA	BZA	BZA	BZA		
RHC								
END								
VERTICAL METHOD								
ALL	ALL	RELPROFRAT	.0000	100.0000	0	0		
END								

BOUNDARYREADINGS

ALL	PATI	.0000
TOP	O3	.0230
RIGHT	O3	.0260
BOTTOM	O3	.0260
LEFT	O3	.0230
TOP	CO	.1210
RIGHT	CO	.1920
BOTTOM	CO	.1920
LEFT	CO	.1210
TOP	RHC	.0790
RIGHT	RHC	.0500
BOTTOM	RHC	.0500
LEFT	RHC	.0790
TOP	NO	.0030
RIGHT	NO	.0020
BOTTOM	NO	.0020
LEFT	NO	.0030
TOP	NO2	.0070
RIGHT	NO2	.0020
BOTTOM	NO2	.0020
LEFT	NO2	.0070
END		.0000

VERTICAL PROFILE

ALL	ALL	35296.0000 5036.0000
9.		.0000
102.		.0000
122.		1.0000
5000.		1.0000
END		.0000
ENDTIME		0

FILE NAME -- BOUNDARY      BOUNDARY DEFINITION AND CONCENTRATIONS  
 BOUNDARY CONCENTRATIONS (PARTS PER MILLION) FOR NO VERT. LEVEL 1 START DATE/TIME 76195 5.00 STOP DATE/TIME 76195 6.00  
 GRID CELL    1    2    3    4    5    6    7    8    9    10    11    12  
 4 -9.0000 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010  
 3 -9.0000 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010  
 2 -9.0000 .0020 .0030 .0030 .0030 .0030 .0030 .0030 .0030 .0030 .0030 .0030 .0030  
 1 -9.0000 .0020 .0030 .0030 .0030 .0030 .0030 .0030 .0030 .0030 .0030 .0030 .0030

## FILE NAME -- BOUNDARY BOUNDARY DEFINITION AND CONCENTRATIONS

	GRID CELL	13	14	15	16	17	18	19	20	21	22
4	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
3	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
2	.0030	.0030	.0030	.0030	.0030	.0030	.0030	.0030	.0030	.0030	.0030
1	.0030	.0030	.0030	.0030	.0030	.0030	.0030	.0030	.0030	.0030	.0030

Exhibit VII-4 (Continued)

Exhibit VII-4 (Continued)

PAGE 1  
FILE NAME -- BOUNDARY      BOUNDARY DEFINITION AND CONCENTRATIONS  
BOUNDARY CONCENTRATIONS (PARTS PER MILLION) FOR NO VERT. LEVEL 2 START DATE/TIME 76195 5.00 STOP DATE/TIME 76195 6.00  
GRID CELL    1    2    3    4    5    6    7    8    9    10    11    12  
4    -9.0000    .0010    .0010    .0010    .0010    .0010    .0010    .0010    .0010    .0010    .0010    .0010  
3    -9.0000    .0010    .0010    .0010    .0010    .0010    .0010    .0010    .0010    .0010    .0010    .0010  
2    -9.0000    .0020    .0020    .0020    .0020    .0020    .0020    .0020    .0020    .0020    .0020    .0020  
1    -9.0000    .0020    .0020    .0020    .0020    .0020    .0020    .0020    .0020    .0020    .0020    .0020

PAGE 2

FILE NAME -- BOUNDARY		BOUNDARY DEFINITION AND CONCENTRATIONS									
		BOUNDARY CONCENTRATIONS (PARTS PER MILLION) FOR NO				VERT.				LEVEL 2 START DATE/TIME 76195 5.00 STOP DATE/TIME 76195 6.00	
GRID CELL		13	14	15	16	17	18	19	20	21	22
4	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
3	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
2	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0030	.0030
1	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0020	.0030	.0030

FILE NAME -- BOUNDARY  
 BOUNDARY CONCENTRATIONS (PARTS PER MILLION) FOR NO VERT. LEVEL 3 START DATE/TIME 76195 5.00 STOP DATE/TIME 76195 6.00  
 GRID CELL 1 2 3 4 5 6 7 8 9 10 11 12  
 4 -9.0000 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010  
 3 -9.0000 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010 .0010  
 2 -9.0000 .0020 .0020 .0020 .0020 .0020 .0020 .0020 .0020 .0020 .0020 .0020  
 1 -9.0000 .0020 .0020 .0020 .0020 .0020 .0020 .0020 .0020 .0020 .0020 .0020

PAGE 1

Exhibit VII-4 (Continued)

PAGE 2

FILE NAME -- BOUNDARY	BOUNDARY DEFINITION AND CONCENTRATIONS				LEVEL 3 START DATE/TIME 76195 5.00	STOP DATE/TIME 76195 6.00
GRID CELL	13	14	15	NO	VERT.	
4	.0010	.0010	.0010	.0010	.0010	-9.0000
3	.0010	.0010	.0010	.0010	.0010	-9.0000
2	.0020	.0020	.0020	.0020	.0020	-9.0000
1	.0020	.0020	.0020	.0020	.0020	-9.0000

FILE NAME -- BOUNDARY                    BOUNDARY DEFINITION AND CONCENTRATIONS  
 BOUNDARY CONCENTRATIONS (PARTS PER MILLION) FOR NO                    VERT.                    LEVEL 4 START DATE/TIME 76195 5.00 STOP DATE/TIME 76195 6.00  
 GRID CELL            1            2            3            4            5            6            7            8            9            10            11            12  
 4            -9.0000            .0010            .0010            .0010            .0010            .0010            .0010            .0010            .0010            .0010            .0010  
 3            -9.0000            .0010            .0010            .0010            .0010            .0010            .0010            .0010            .0010            .0010            .0010  
 2            -9.0000            .0030            .0030            .0030            .0030            .0030            .0030            .0030            .0030            .0030            .0030  
 1            -9.0000            .0030            .0030            .0030            .0030            .0030            .0030            .0030            .0030            .0030            .0030

Exhibit VII-4 (Continued)

FILE NAME -- BOUNDARY      BOUNDARY DEFINITION AND CONCENTRATIONS  
 BOUNDARY CONCENTRATIONS (PARTS PER MILLION) FOR NO      VERT.      LEVEL 4 START DATE/TIME 76195 5.00 STOP DATE/TIME 76195 6.00  
 GRID CELL      13      14      15      16      17  
 4      .0010      .0010      .0010      .0010      -9.0000  
 3      .0010      .0010      .0010      .0010      -9.0000  
 2      .0030      .0030      .0030      .0030      -9.0000  
 1      .0030      .0030      .0030      .0030      -9.0000

## C. DIFFBREAK

The DIFFBREAK file contains time-varying two-dimensional matrices of diffusion break heights. The diffusion break is used to define the distribution of cells in the vertical direction and to delimit layers of the atmosphere with different diffusion characteristics. The diffusion break is usually set to the mixing height or the height of the inversion base. The program DFSNBK is used to create the DIFFBREAK file. DFSNBK requires subroutines from the libraries UTILITY and FILUTIL. Figure VII-5 is an information flow diagram illustrating the job control to be used. In the following sections we discuss the input to the DFSNBK program.

### 1. Variables

The output variable for the DIFFBREAK file is also named DIFFBREAK. This is an "implicit output variable" and need not be referred to anywhere in the CONTROL packet. The internal units for DIFFBREAK are meters (m). If the input values for this variable are to be in any other units, DIFFBREAK must be specified in the UNITS packet.

### 2. Methods

The methods that can be used to generate the DIFFBREAK file are:

- > CONSTANT
- > GRID VALUE
- > STATINTERP
- > POISSON
- > E-WINTERP
- > N-SINTERP
- > USER.

These methods are discussed in detail in Section VI.A.6.

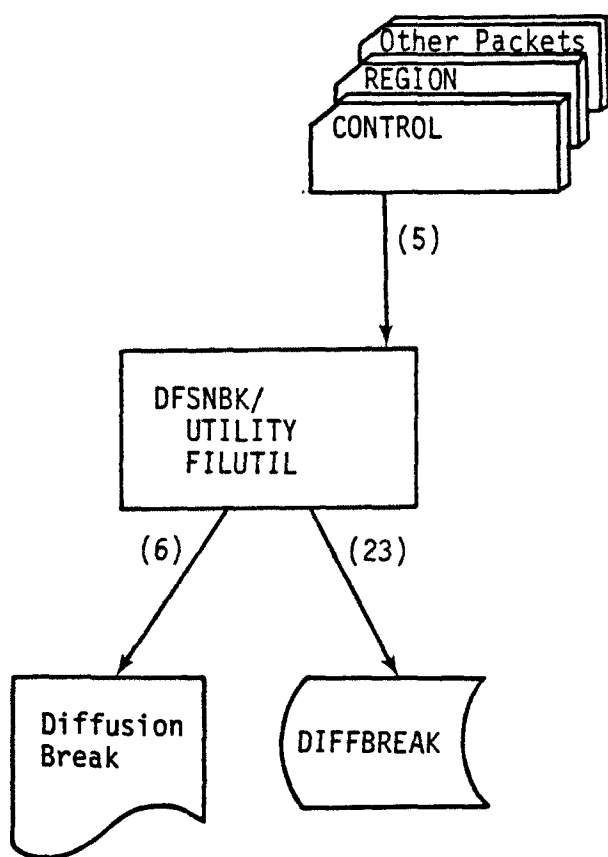


Figure VII-5. Information Flow Diagram for the DIFFBREAK File

### 3. Time Variation

The diffusion break values on the DIFFBREAK file are considered to apply at the beginning of the time interval. Because the Airshed Simulation Program calculates diffusion break values continuously over time by linear interpolation, it also requires values at the end of the time interval. These are read by the program as the values at the beginning of the next time interval. Thus, the last time interval on the file must begin at or after the ending simulation time. For example, if a simulation is to be made from 0500-1700, and diffusion break values are input hourly, the values used between 500 and 600 are calculated by interpolating between values input for the 0500-0600 time interval and those input for the 0600-0700 time interval. Similarly, to calculate values between 1600 and 1700, the simulation program requires values for the interval 1600-1700 as well as another set for an interval beginning at 1700.

### 4. Packets

Figure VII-6 shows the input deck structure in terms of the packets to be used. Each of these packets is described in detail in Section VI.B. Following are special input packet considerations for the DIFFBREAK file:

> CONTROL

- The file name on Card 2 must be DIFFBREAK.
- The control variables to be specified on Cards 4 to 8 for DIFFBREAK are shown in Table VII-3.
- The number of species should be zero.
- If there are input variables that do not appear as output variables, their number must appear as the number of user-defined variables. All such variables must also be named in the UNITS packet.

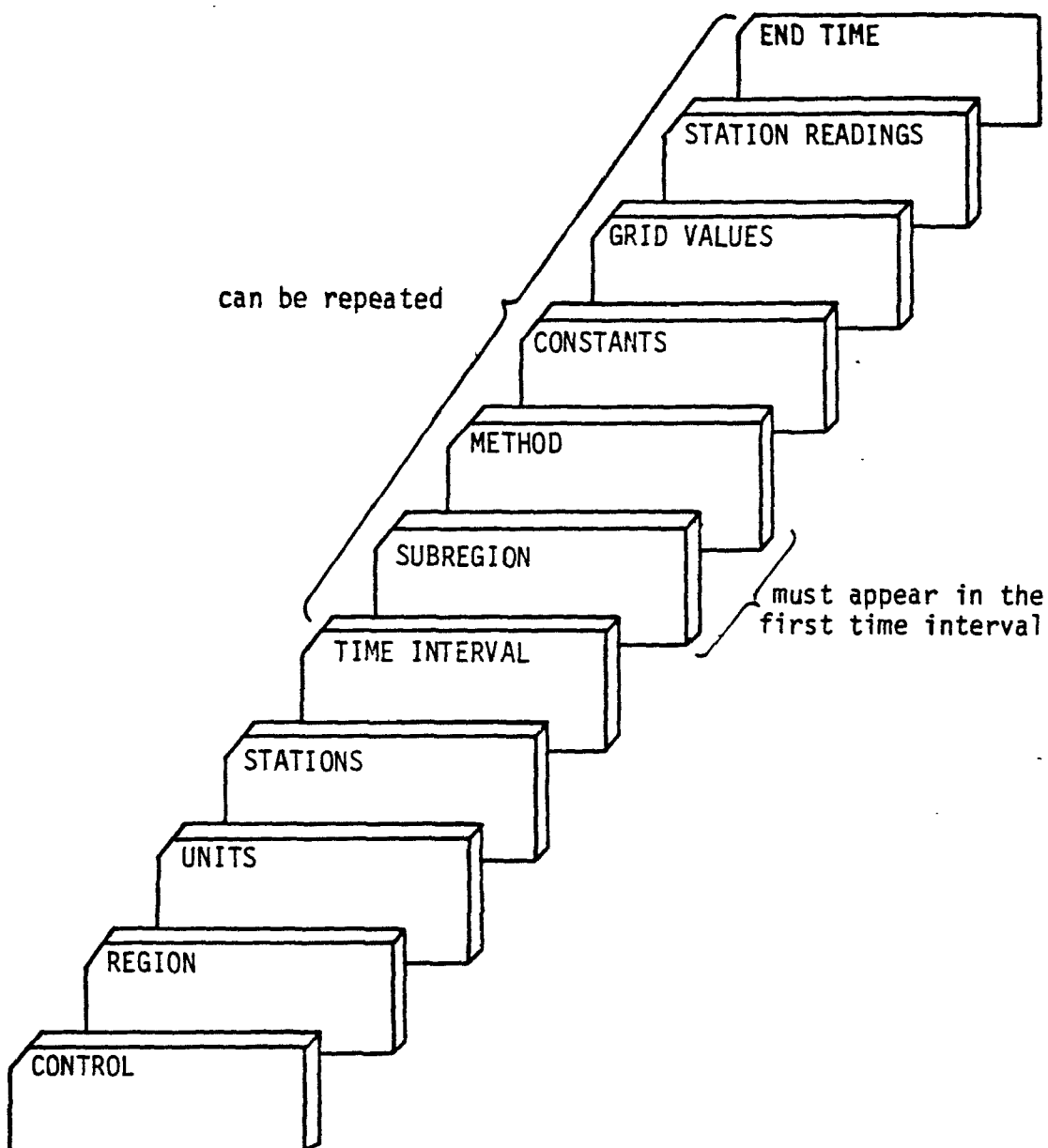


Figure VII-6. Input Card Deck Structure for Preparing the DIFFBREAK FILE

**Table VII-3. CONTROL Packet Entries for the DIFFBREAK File**

<u>Number</u>	<u>Entry</u>
4	Number of species (=0) Number of user-defined variables Number of stations Number of subregions Number of parameters Spare
5	Output file number Print input cards Print output grid Spare Spare Spare
6	Print units table Print station locations table Print regional grid Print methods table Print station values table Spare
7	Spare Spare Spare Spare Spare Spare
8	Spare Spare Spare Spare Spare Spare

- If data from measuring stations are to be used (methods STATINTERP or POISSON), the maximum number of such stations must be given.
  - The number of subregions must be at least one.
  - The maximum number of parameters must be sufficient to include all specifications of all parameters.
  - The vertical controls card (No. 7) should be left blank.
  - The file unit assignment card (No. 8) should be left blank.
  - The beginning and ending dates and times should reflect the time variation considerations discussed in Section 3 above.
  - A set of output species names is not required; if they are present, their number must be the same as the entry in the first control parameter on Card 4, but they will be ignored by the program.
- > REGION. This packet must follow the CONTROL packet. The vertical parameters will be ignored for the DIFFBREAK file.
- > UNITS. This packet, if present, must follow the REGION packet. The UNITS packet must be provided if:
- Any input variable will be input in other than internal units.
  - Any user-defined variables are specified.
  - COORD or HEIGHT unit conversions are to be used.
- The number of user-defined variables must not exceed the maximum specified in the CONTROL packet.
- > STATIONS. This packet is required if either of the methods STATINTERP or POISSON is specified. The number of stations listed must not exceed the maximum specified in the CONTROL packet.
- > TIME INTERVAL. Two or more TIME INTERVAL packets must be present. The first time interval must begin at or before the beginning of the time span specified on Card 10 of the CONTROL packet. The last time interval must begin at or

after the ending time of any simulation run. All time intervals must be contiguous and of nonzero length. Each TIME INTERVAL packet contains one or more of the following packets and ends with an ENDTIME card. Following the first time interval, only those data that are to be changed need be specified.

- > SUBREGION. The first time interval must contain a SUBREGION packet; the inclusion of this packet in other time intervals is optional. The number of subregions must not exceed the maximum specified in the CONTROL packet.
- > METHOD. A method must be provided for every variable-- including user-defined variables--in every subregion in the first time interval. Methods can be changed in subsequent TIME INTERVAL packets if desired. Note that each parameter entry contributes to the overall parameter count; the total number of parameters must not exceed the maximum specified in the CONTROL packet.
- > CONSTANTS. If the method CONSTANT is assigned to any variable in the METHOD packet, the first time interval must contain a CONSTANTS packet. More than one CONSTANTS packet can appear in any time interval.
- > GRID VALUES. If the method GRID VALUE is assigned to any variable in the METHOD packet, the first time interval must contain a GRID VALUES packet. More than one GRID VALUES packet can appear in any time interval.
- > STATION READINGS. If either the.POISSON or STATINTERP method is assigned to any variable in the METHOD packet, the first time interval must contain a STATION READINGS packet. More than one STATION READINGS packet can appear in any time interval.

## 5. Additional Input Files

No additional input files are required.

6. Examples

Exhibit VII-5 shows a sample input deck for DIFFBREAK file generation. The output from a DFSNBK run using that input deck is shown in Exhibit VII-6.

CONTROL  
 DIFFBREAK  
 HEIGHT OF THE MIXED LAYER  
 23 1 12 1 1 10

END  
 REGION  
 725000. 4252000. 15  
 -19000. -16000.  
 4000. 4000.  
 17 22 4  
 2 0. 50.  
 200.

END  
 STATIONS  
 141 17949. 27053.  
 141A 14000. 23000.  
 141B 14000. 27000.  
 141C 14000. 31000.  
 141D 18000. 31000.  
 141E 18000. 23000.  
 141F 22000. 23000.  
 141G 22000. 27000.  
 141H 22000. 31000.  
 142 -9294. 14115.  
 143 35296. 5036.  
 144 30861. 56067.

END  
 TIME INTERVAL  
 76195 500 76195 600  
 SUBREGION  
 A 1 1 -1  
 END

<b>METHOD</b>	<b>DIFFBREAK POISSON</b>	<b>0.</b>	<b>2000.</b>
<b>A</b>	<b>RMAXIT</b>	<b>200.</b>	
	<b>TLRNE</b>	<b>0.5</b>	
	<b>RO1EGA</b>	<b>1.4</b>	
<b>END</b>			
<b>STATION READINGS</b>			
141	DIFFBREAK	402.	
141A	DIFFBREAK	402.	
141B	DIFFBREAK	402.	
141C	DIFFBREAK	402.	
141D	DIFFBREAK	402.	
141E	DIFFBREAK	402.	
141F	DIFFBREAK	402.	
141G	DIFFBREAK	402.	
141H	DIFFBREAK	402.	
142	DIFFBREAK	102.	
143	DIFFBREAK	102.	
144	DIFFBREAK	102.	
<b>END</b>			
<b>ENDTIME</b>			
<b>TIME</b>	<b>INTERVAL</b>		
76195	600	76195	700
<b>STATION READINGS</b>			
141	DIFFBREAK	418.	
141A	DIFFBREAK	418.	
141B	DIFFBREAK	418.	
141C	DIFFBREAK	418.	
141D	DIFFBREAK	418.	
141E	DIFFBREAK	418.	
141F	DIFFBREAK	418.	
141G	DIFFBREAK	418.	
141H	DIFFBREAK	418.	
142	DIFFBREAK	167.	
143	DIFFBREAK	167.	
144	DIFFBREAK	167.	
<b>END</b>			
<b>ENDTIME</b>			

Exhibit VII-5 (Concluded)

PROCESSING OF INPUT DATA FOR THE PREPARATION OF FILE DIFFBREAK  
 HEIGHT OF THE MIXED LAYER  
  
 NUMBER OF STATIONS = 12  
 NUMBER OF SUBREGIONS = 1  
 NUMBER OF METHOD PARAMETERS = 10  
 PROGRAM OUTPUT WILL BE PLACED ON FILE NUMBER 23  
 INPUT CARDS WILL BE PRINTED  
 OUTPUT GRID WILL BE PRINTED  
 UNIT TABLE WILL NOT BE PRINTED  
 STATION LOCATIONS WILL NOT BE PRINTED  
 SUBREGION GRID WILL NOT BE PRINTED  
 METHOD TABLE WILL NOT BE PRINTED  
 STATION VALUES WILL NOT BE PRINTED  
 BEGINNING DATE OF FILE 76195 BEGINNING TIME OF FILE 5.00  
 ENDING DATE OF FILE 76196 ENDING TIME OF FILE .17  
  
 REGION  
 725000. 4252000. 15  
 -19000. -16000.  
 4000.00 4000.00  
 17 22 4  
 2 2 .00 50.00 200.00  
 END

**Exhibit VII-6.** Sample Output from the DFNSBK Program

## FILE NAME -- DIFFBREAK

## HEIGHT OF THE MIXED LAYER

GRID CELL	DIFFUSION BREAK VALUES (IN METERS ABOVE GROUND LEVEL)				VERT.	LEVEL 1	START DATE/TIME	76195 5.00	STOP DATE/TIME	76195 6.00	PAGE 1
	1	2	3	4							
22	102.0000	102.0000	102.0000	102.0000	102.0000	102.0000	102.0000	102.0000	102.0000	102.0000	102.0000
21	102.0000	104.0061	105.9784	107.8764	109.6442	111.2035	112.4517	113.2621	113.4869	112.9678	111.5785
20	102.0000	106.0492	110.0392	113.8977	117.5229	120.7653	123.4199	125.2271	125.8663	124.9496	122.0040
19	102.0000	108.1576	114.2490	120.1900	125.8482	131.0156	135.3671	138.5582	140.0230	139.1342	134.8865
18	102.0000	110.3394	116.6380	126.8301	134.7796	142.2370	148.8003	153.8979	156.8324	156.8625	153.3217
17	102.0000	112.5715	123.1758	133.8129	144.3856	154.6312	164.0430	171.8205	176.9206	176.3270	175.4780
16	102.0000	114.7858	127.7453	141.0099	154.5690	168.2632	181.4332	192.9575	201.1329	204.1504	201.2360
15	102.0000	116.8553	132.1119	148.1159	165.0816	182.9509	201.1351	218.0985	230.9451	235.9049	231.8983
14	102.0000	118.5852	135.8983	154.5894	175.1980	198.0348	222.8621	246.0858	269.0660	276.4828	270.2190
13	102.0000	119.7123	138.5788	159.6205	183.7796	211.9917	245.0963	283.0558	321.0929	330.4662	321.8958
12	102.0000	119.9191	139.5127	162.1920	169.1754	222.0373	263.4067	318.6224	402.0000	402.0000	318.1304
11	102.0000	118.8524	137.9964	161.3214	169.7096	224.6137	268.7601	326.5473	402.0000	402.0000	324.6042
10	102.0000	116.1240	133.1719	156.4365	164.8351	218.9656	261.2449	317.1956	402.0000	402.0000	313.3499
9	102.0000	111.3740	123.2244	147.5830	175.3284	206.0785	240.6791	279.2575	317.1860	325.2433	314.3020
8	102.0000	105.3165	102.0000	136.5030	163.7951	190.0691	216.5829	242.1357	262.1583	267.2108	257.2377
7	102.0000	107.2511	114.7116	133.6508	154.0439	174.3307	193.7312	210.6377	222.0515	224.0679	215.5090
6	102.0000	108.3177	117.0373	130.1234	144.9114	159.7828	173.5247	184.6783	191.3240	191.4567	183.9997
5	102.0000	108.1039	115.7802	125.3862	135.9776	146.5089	155.9673	163.2427	167.1079	166.4374	160.5847
4	102.0000	107.0870	113.0524	119.9097	127.2197	134.3517	140.6021	145.2149	147.4289	146.6159	142.4751
3	102.0000	105.6063	109.6386	114.9663	118.6635	123.0710	126.6604	129.5737	130.7749	130.1351	127.5416
2	102.0000	103.8637	105.8902	108.0631	110.2633	112.3872	114.1739	115.4294	115.9563	115.6151	114.3737
1	102.0000	102.0000	102.0000	102.0000	102.0000	102.0000	102.0000	102.0000	102.0000	102.0000	102.0000

PAGE 2

FILE NAME -- DIFFBREAK	HEIGHT OF THE MIXED LAYER						VERT.	LEVEL 1	START DATE/TIME 76195 5.00	STOP DATE/TIME 76195 6.00
GRID CELL	13	14	15	16	17					
22	102.0000	102.0000	102.0000	102.0000	102.0000					
21	107.2683	107.4373	107.3662	105.8242	102.0000					
20	109.6837	111.9638	112.2625	109.0659	102.0000					
19	102.0000	116.8009	116.0606	112.6005	102.0000					
18	135.3789	132.9462	127.3580	117.1122	102.0000					
17	158.9566	149.4731	137.5922	121.9777	102.0000					
16	180.1957	165.2262	147.5758	126.6674	102.0000					
15	201.3668	180.3163	156.8746	130.8992	102.0000					
14	223.0223	194.5563	165.1207	134.4611	102.0000					
13	244.1440	206.8965	171.6151	137.0459	102.0000					
12	260.6950	214.9730	175.1758	138.1938	102.0000					
11	263.9149	215.4856	174.4594	137.4240	102.0000					
10	253.7866	207.5854	168.9463	134.5254	102.0000					
9	229.6791	191.6240	158.8958	129.6244	102.0000					
8	200.6825	170.1863	145.3731	123.1938	102.0000					
7	171.0517	143.1141	129.3504	115.9792	102.0000					
6	143.7772	102.0000	113.1119	109.5702	102.0000					
5	133.6663	115.8389	111.6856	107.3430	102.0000					
4	125.6634	116.0916	110.5570	106.2150	102.0000					
3	117.7668	112.3746	106.3042	105.0166	102.0000					
2	109.8678	107.3805	105.3094	103.5776	102.0000					
1	102.0000	102.0000	102.0000	102.0000	102.0000					

## D. EMISSIONS

The EMISSIONS file contains time-varying matrices of ground-level emission fluxes. At present, there is no general-purpose program for creating the EMISSIONS file because its generation is so highly dependent on the form of the emissions data available: traffic, stationary sources, area sources, airports, and so on. There is a program structure, similar to that of the other data preparation programs, that includes the reading of control packets, unit conversions, and writing the proper output records. This program can be tailored to particular input requirements as necessary. The skeleton program is named EMISSN. EMISSN requires subroutines from the libraries UTILITY and FILUTIL. Figure VII-7 is an information flow diagram illustrating the job control to be used. In the following sections we discuss the card input to the EMISSN program.

### 1. Variables

The output variables for the EMISSIONS file are the species named in the CONTROL packet. Additional user-defined input variables (e.g. "reactive hydrocarbons") can be specified in the UNITS packet. The internal units for ground-level emissions of all species except AEROSOLS are gram-moles per hour (g-mol/hr); for AEROSOLS, the units are grams per hour (g/hr). The standard names for reactive species recognized by the Airshed Simulation Program are listed in Section IX.A. If any of these species does not appear on the EMISSIONS file, the emissions values default to zero. If any additional species appear on EMISSIONS, they will be ignored.

### 2. Methods

The methods that can be used to generate the EMISSIONS file are:

- > CONSTANT
- > GRID VALUE
- > SPLIT/COMB
- > USER.

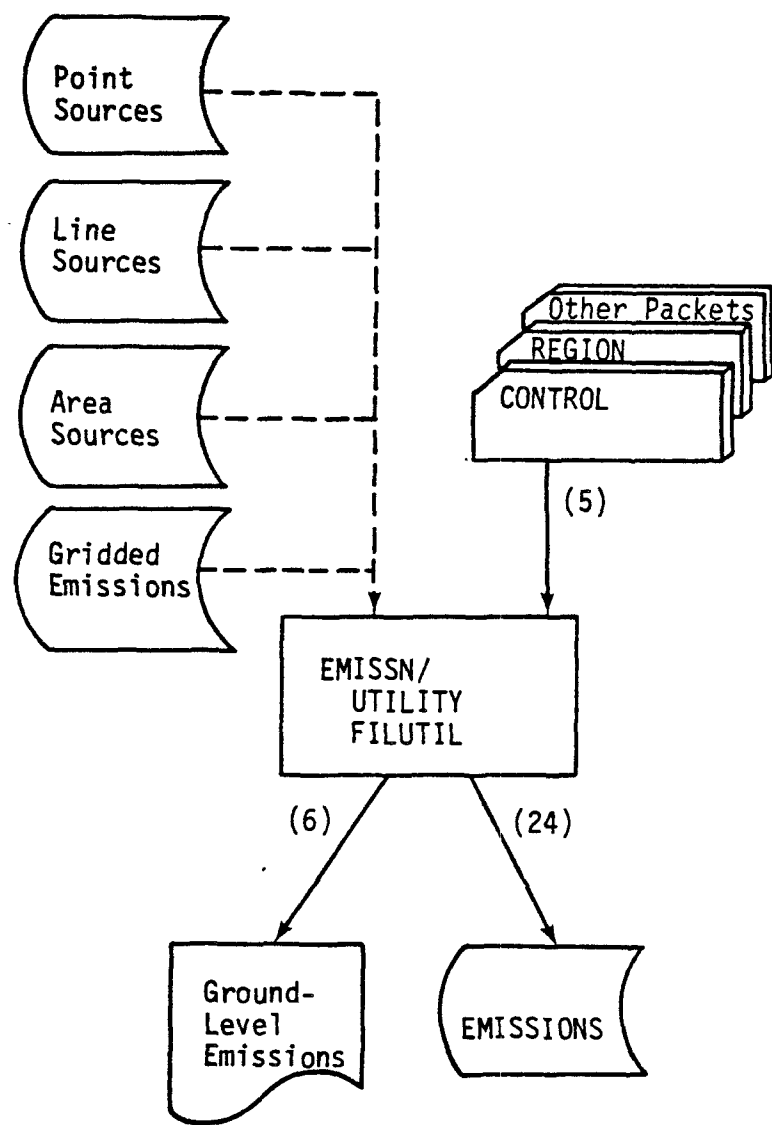


Figure VII-7. Information Flow Diagram for the EMISSIONS File

The actual methods for generating output values from specific input data files must be written for each new application.

### 3. Time Variation

The time span of the EMISSIONS file must include the entire time span of the simulation runs for which it is to be used. Ground-level emissions are considered to be constant during each time interval.

### 4. Packets

The packets to be used for EMISSN will depend on the particular version that is developed for a given set of input data files. Information on control parameters, units, time intervals, computation methods, and data entry will have to be provided when the program has been implemented. Figure VII-8 shows a possible input structure in terms of the packets to be used.

### 5. Additional Input Files

Point source, line source, area source, or gridded area emissions might be input to EMISSN.

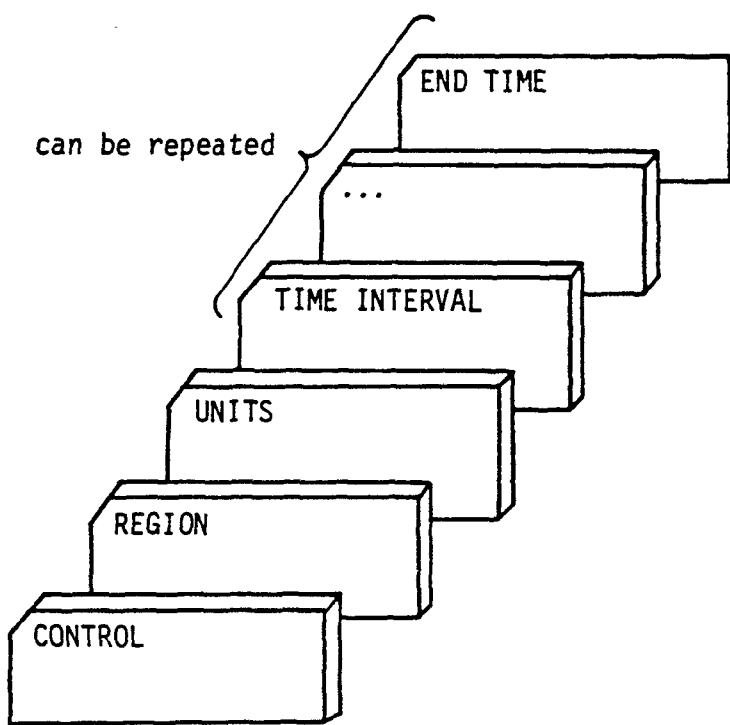


Figure VII-8. A Possible Input Card Deck Structure for Preparing the EMISSIONS File

## E. METSCALARS

The METSCALARS file contains a set of time-varying scalars to be used in meteorological calculations. It has no spatial dependence. The program METSCL is used to create the METSCALARS file. METSCL requires subroutines from the libraries UTILITY and FILUTIL. Figure VII-9 is an information flow diagram illustrating the job control to be used. In the following sections we discuss the input to the METSCL program.

### 1. Variables

There are six output variables for the METSCALARS file. Their names and internal units are:

- > TGRADBELOW--temperature gradient below the diffusion break ( $^{\circ}\text{K}/\text{m}$ ).
- > TGRADABOVE--temperature gradient above the diffusion break ( $^{\circ}\text{K}/\text{m}$ ).
- > EXPCLASS--exposure class, an integer scale (+3 to -2) of the near ground-level atmospheric stability due to surface heating or cooling. It can be estimated from insolation as follows:

$$\text{exposure class} = \begin{cases} 3, \text{ strong} \\ 2, \text{ moderate} \\ 1, \text{ slight} \\ 0, \text{ heavy overcast} \\ -1, \geq \frac{4}{8} \text{ cloud cover} \\ -2, \leq \frac{3}{8} \text{ cloud cover} \end{cases} \begin{array}{l} \text{daytime insolation ,} \\ \text{day or night ,} \\ \text{nighttime cloudiness .} \end{array}$$

- > RADFACTOR--the  $\text{NO}_2$  photolysis rate constant,  $k_1$  ( $\text{min}^{-1}$ ).
- > CONCWATER--average concentration of water (ppm).
- > ATMOSPRESS--atmospheric pressure (atm).

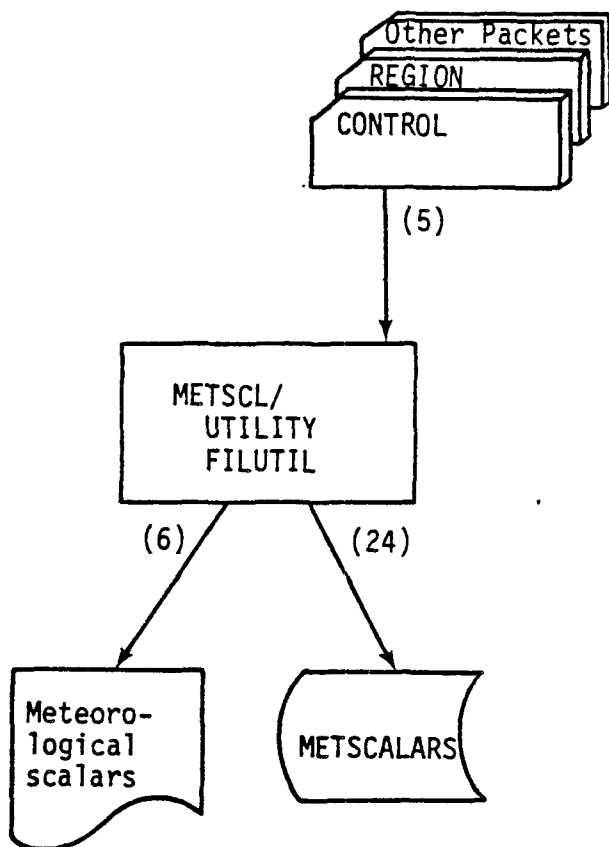


Figure VII-9. Information Flow Diagram for the METSCALARS File

## 2. Methods

Since the METSCALARS variables do not vary spatially, there is no need to define a method for determining spatial distribution. It is assumed that the variables will be explicitly entered in SCALARS packets and that only simple unit conversions, which are defined in the UNITS packet, need be performed.

## 3. Time Variation

Five of the METSCALARS variables, TGRADBELOW, TGRADABOVE, EXPCLASS, CONCWATER, and ATMOSPRESS, are considered to be constant during each time interval. The sixth variable, RADFACTOR, however, is considered to apply at the end of the time interval. Because the Airshed Simulation Program calculates radiation factor values continuously over time by linear interpolation, it also requires values at the beginning of the time interval. These are saved by the program as the values read for the end of the previous time interval. Thus the first time interval on the file must end at or before the beginning simulation time. For example, if a simulation is to be made from 0500-1700, and radiation factor values are input hourly, the values used between 500 and 600 are calculated by interpolating between values input for the 0400-0500 time interval and those input for the 0500-0600 time interval.

## 4. Packets

Figure VII-10 shows the input deck structure in terms of the packets to be used. Each of these packets is described in detail in Sections VI.B. Following are special input packet considerations for the file:

- > CONTROL
  - The file name on Card 2 must be METSCALARS.
  - The control variables to be specified on Cards 4 to 8 for METSCALARS are shown in Table VII-4.

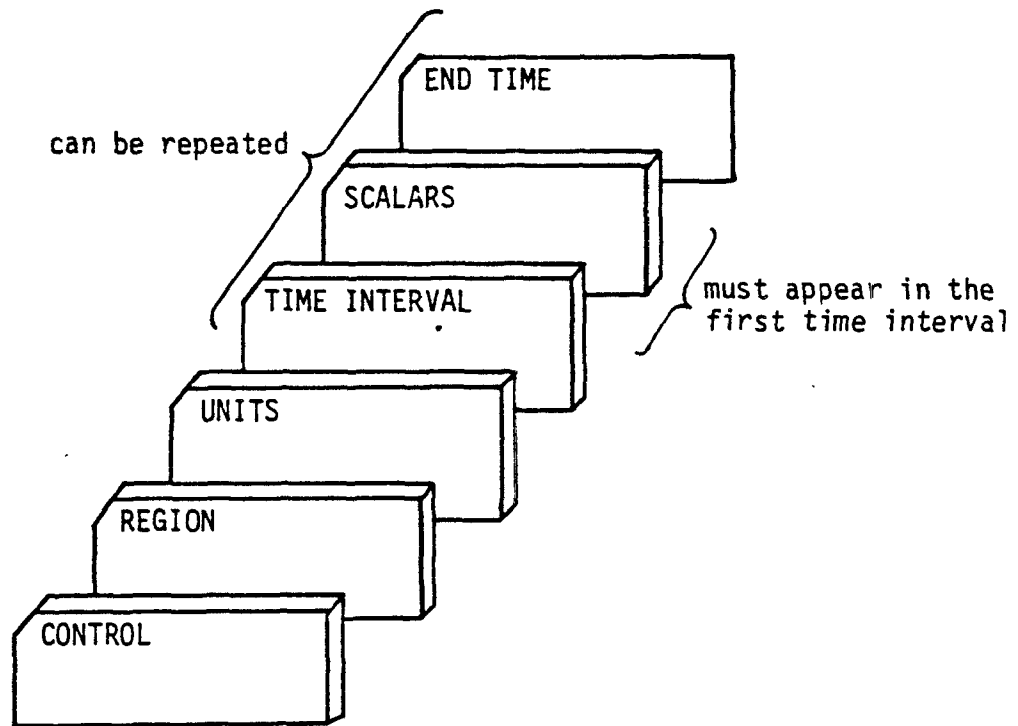


Figure VII-10. Input Card Deck Structure for Preparing the METSCALARS File

Table VII-4. CONTROL Packet Entries for the METSCALARS File

<u>Card Number</u>	<u>Entry</u>
4	Number of species (=0) Spare Spare Spare Spare Spare
5	Output file number Print input cards Print output values Spare Spare Spare
6	Print units table Spare Spare Spare Spare Spare
7	Spare Spare Spare Spare Spare Spare
8	Spare Spare Spare Spare Spare Spare

- The number of species should be zero.
  - The vertical controls card (No. 7) should be left blank.
  - The file unit assignment card (No. 8) should be left blank.
  - The beginning and ending dates and times should reflect the time variation considerations discussed in Section 3 above.
  - A set of output species names is not required; if they are present, their number must be the same as the entry in the first control parameter on Card 4, but they will be ignored by the program.
- > REGION. This packet must follow the CONTROL packet. The vertical parameters will be ignored for the METSCALARS file.
- > UNITS. This packet, if present, must follow the REGION packet. The UNITS packet must be provided if any input variable will be input in other than internal units. The units of RADFACTOR must be  $\text{min}^{-1}$  and cannot be changed.
- > TIME INTERVAL. Two or more TIME INTERVAL packets must be present. The first time interval must end at or before the beginning time of any simulation run. All time intervals must be contiguous and of nonzero length. Each TIME INTERVAL packet contains a SCALARS packet and ends with an ENDTIME card. Following the first time interval, only those data that are to be changed need be specified.
- > SCALARS. The first time interval must contain a SCALARS packet. Following the first time interval, the SCALARS packet can be omitted if no values are to be changed.

## 5. Additional Input Files

No additional input files are required.

## 6. Examples

Exhibit VII-7 shows a sample input deck for METSCALARS file generation. The output from a METSCL run using that input deck is shown in Exhibit VII-8.

```

CONTROL
MET SCALARS
      VALUES FOR METEOROLOGICAL SCALARS
          0   0   0   1   1   3
          24  1
          1

        76195  400  76195  2400
END
REGION
    725000.  4252000.  15
    -19000.  -16000.
    4000.    4000.
    17       22       4
    2       2       0.
    50.     200.

TIME INTERVAL
    76195  400  76195  500
SCALARS
TGRADBELOW  -0.01
TGRADABOVE  0.0039
EXPCLASS    -1.
ATMOSPRESS  0.99
RADFACTOR   .008
CONCWATER   12687.
END
END TIME
TIME INTERVAL
    76195  500  76195  600
SCALARS
RADFACTOR   .084
EXPCLASS    0.0
END
END TIME
TIME INTERVAL
    76195  600  76195  700
SCALARS
EXPCLASS    1.
CONCWATER   12687.
RADFACTOR   .180
END
END TIME

```

**PROCESSING OF INPUT DATA FOR THE PREPARATION OF FILE METSCALARS  
VALUES FOR METEOROLOGICAL SCALARS**

PROGRAM OUTPUT WILL BE PLACED ON FILE NUMBER 24  
INPUT CARDS WILL BE PRINTED  
OUTPUT GRID WILL BE PRINTED  
UNIT TABLE WILL BE PRINTED  
  
BEGINNING DATE OF FILE 76195 BEGINNING TIME OF FILE 4.00  
ENDING DATE OF FILE 76195 ENDING TIME OF FILE 24.00  
  
REGION  
725000. 4252000. 15  
-19000. -16000.  
4000.00 4000.00  
17 22 4  
2 2 .00 50.00 200.00  
  
END  
  
TIME INTERVAL  
76195 400 76195 500  
  
SCALARS  
TGADBELOW -.0100  
TGADABOVE .0039  
EXPCLASS -1.0000  
ATMOSPRESS .9900  
RADFACTOR .0080  
CONCWATER 12667.0000  
.0000  
END  
ENDTIME

**Exhibit VII-8. Sample Output from the METSCL Program**

FILE NAME --METSCALARS		VALUES FOR METEOROLOGICAL SCALARS		PAGE 01
METEOROLOGICAL SCALARS (VARIABLE UNITS -- CONSULT MANUAL) START DATE/TIME ,76195 4.00 STOP DATE/TIME 76195 5.00				
VARIABLE NAME	VALUE			
TGRAD BELOW	- .0100			
TGRAD ABOVE	.0039			
EXPCLASS	-1.0000			
RADFACTOR	.0080			
CONCWATER	12667.0000			
ATMOSPRESS	.9900			
TIME INTERVAL				
SCALARS	500	76195	600	
RADFACTOR	.0840			
EXPCLASS	.0000			
END	.0000			
ENDTIME				

FILE NAME --METSCALARS		VALUES FOR METEOROLOGICAL SCALARS		PAGE 01
METEOROLOGICAL SCALARS (VARIABLE UNITS -- CONSULT MANUAL) START DATE/TIME ,76195 5.00 STOP DATE/TIME 76195 6.00				
VARIABLE NAME	VALUE			
TGRAD BELOW	- .0100			
TGRAD ABOVE	.0039			
EXPCLASS	.0000			
RADFACTOR	.0640			
CONCWATER	12667.0000			
ATMOSPRESS	.9900			
TIME INTERVAL				
SCALARS	600	76195	700	
EXPCLASS	1.0000			
CONCWATER	12667.0000			
RADFACTOR	.1800			
END	.0000			
ENDTIME				

## F. PTSOURCE

The PTSOURCE file contains a set of time-invariant locations of elevated point sources and time-varying emissions fluxes from each source into specific vertical cells. The file can be omitted from a simulation run. The program PTSRCE is used to create the PTSOURCE file. PTSRCE requires subroutines from the libraries UTILITY and FILUTIL. Figure VII-11 is an information flow diagram illustrating the job control to be used. In the following sections we discuss the input to the PTSRCE program.

### 1. Variables

The output variables for the PTSOURCE file are the species named in the CONTROL packet. Additional user-defined input variables (e.g., "reactive hydrocarbons") can be specified in the UNITS packet. The internal units for the emissions of all species except AEROSOLS are gram-moles per hour (g-mol/hr); for AEROSOLS, the units are grams per hour (g/hr). The standard names for reactive species recognized by the Airshed Simulation Program are listed in Section IX.A. If any of these species does not appear on the PTSOURCE file, the emissions will default to zero. If any additional species appear on PTSOURCE, they will be ignored.

There are five other implicit variables used in the PTSRCE program:

- > HEIGHT--stack height
- > DIAMETER--stack exit diameter
- > STACKTEMP--stack exit temperature
- > STACKVEL--stack exit velocity
- > FLOWRATE--flow rate.

The internal units for these variables are shown in Table VI-5; input unit conversions for any of these variables can be specified in the UNITS packet without adding to the count of "user-defined variables." Values

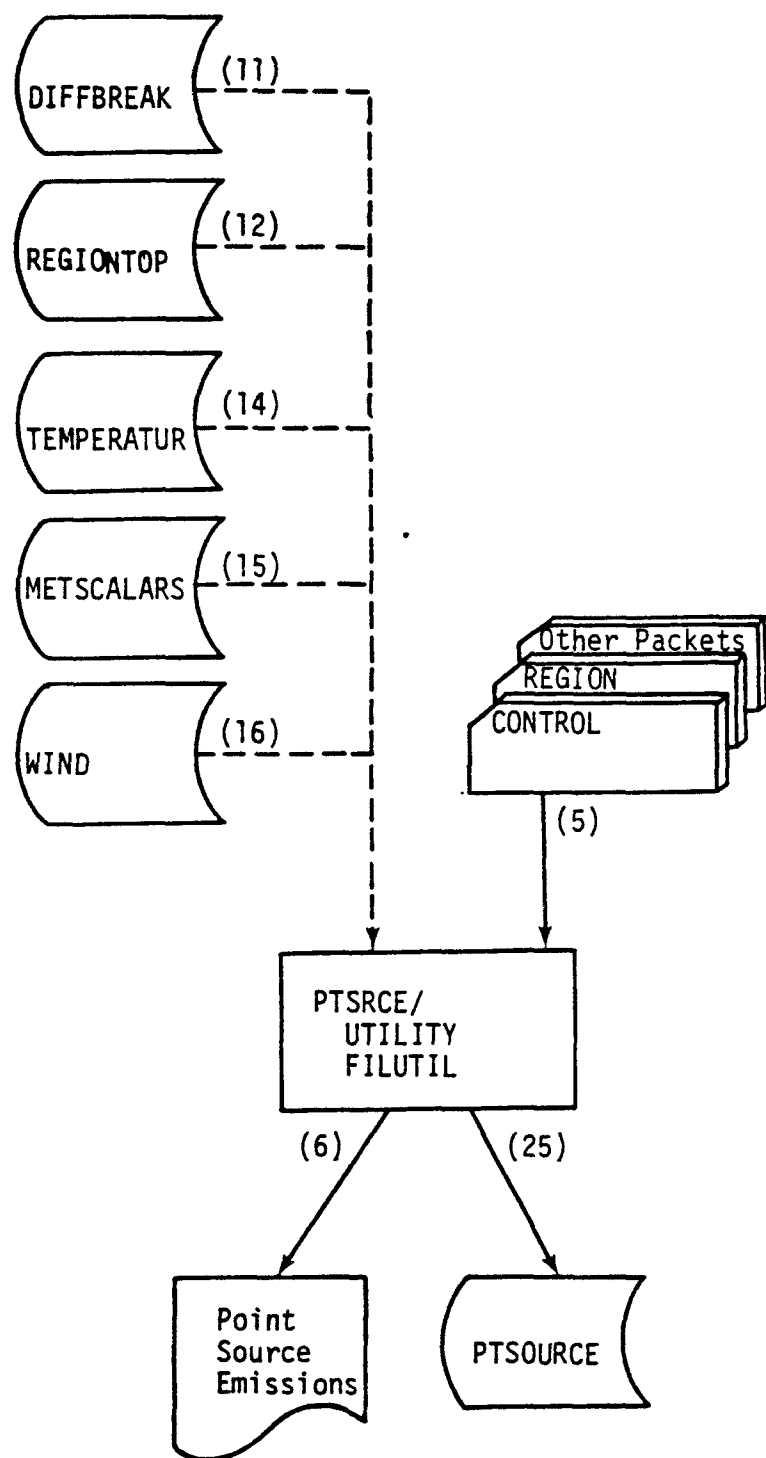


Figure VII-11. Information Flow Diagram for the PTSOURCE File

for the first four variables are entered in the POINT SOURCES packet and are considered time-invariant. FLOWRATE values, if specified, appear in the EMISSIONS VALUES packet and are modified, along with species emissions, by the emissions factors.

## 2. Methods

The methods that can be used to generate the PTSOURCE file are:

- > EMVALUES
- > EMFACTORS
- > SPLIT/COMB
- > USER.

These methods are discussed in detail in Section VI.A.6.

Since the PTSOURCE file varies in the vertical direction, a vertical method must also be specified for each output variable in each subregion. The vertical methods that can be used are:

- > STACKHGT
- > PLUMERISE
- > VERTUSER.

These vertical methods are discussed in detail in Section VI.A.7.

## 3. Time Variation

The time span of the PTSOURCE file must include the entire time span of the simulation runs for which it is to be used. Point source emissions are considered to be constant during each time interval.

#### 4. Packets

Figure VII-12 shows the input deck structure in terms of the packets to be used. Each of these packets is described in detail in Section VI.B. Following are special input packet considerations for the PTSOURCE file:

- > CONTROL
  - The file name on Card 2 must be PTSOURCE.
  - The control variables to be specified on Cards 4 to 8 for PTSOURCE are shown in Table VII-5.
  - The number of species must be greater than zero.
  - If there are input variables that do not appear as output variables, their number must appear as the number of user-defined variables. All such variables must also be named in the UNITS packet.
  - The number of point sources defined in the POINT SOURCES packet must be specified.
  - The number of point source types must be at least one.
  - The maximum number of parameters must be sufficient to include all specifications of all parameters.
  - The vertical controls card (No. 7) must specify the maximum number of vertical parameters as applicable.
  - The file unit assignment card (No. 8) must specify the DIFFBREAK and REGIONTOP files. In addition, it must specify TEMPERATUR, METSCALARS, and WIND if the vertical method PLUMERISE is selected.
  - The beginning and ending dates and times should reflect the time variation considerations discussed in Section 3 above.
  - A set of output species names is required; their number must be the same as the entry in the first control parameter on Card 4.
- > REGION. This packet must follow the CONTROL packet. The vertical parameters must be provided for the PTSOURCE file.

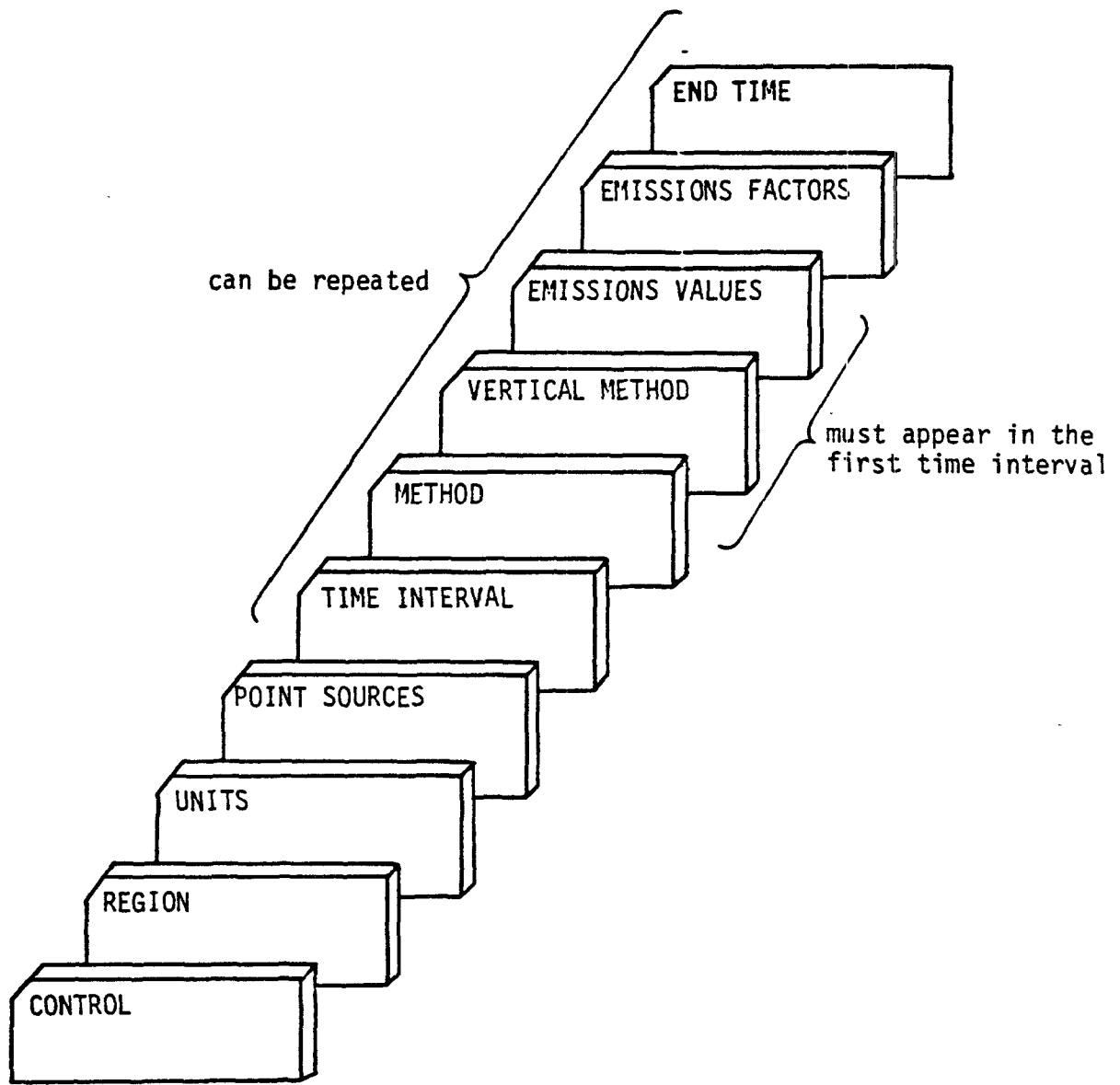


Figure VII-12. Input Card Deck Structure for Preparing the PTSOURCE File

Table VII-5. CONTROL Packet Entries for the PTSOURCE File

<u>Card Number</u>	<u>Entry</u>
4	Number of species Number of user-defined variables Number of point sources Number of point source types Number of parameters Spare
5	Output file number Print input cards Print output grid Spare Spare Spare
6	Print units table Print point source locations table Print regional point source distribution grid Print methods table Print point source values table Spare
7	Number of vertical parameters Spare Print vertical methods table Spare Spare Spare
8	DIFFBREAK file number REGIONTOP file number Spare TEMPERATUR file number METSCALARS file number WIND file number

- > UNITS. This packet, if present, must follow the REGION packet. The UNITS packet must be provided if:
  - Any input variable will be input in other than internal units.
  - Any user-defined variables are specified.
  - COORD or HEIGHT unit conversions are to be used.The number of user-defined variables must not exceed the maximum specified in the CONTROL packet.
- > POINT SOURCES. This packet is required. It names the point sources, assigns to each a type and location, and describes certain time-invariant stack properties. The number of point sources specified must equal the number specified in the CONTROL packet. Each point source must be given a type name. Point sources are grouped by type at the time that methods and vertical methods are assigned, and emission factors can be applied by point source type. The number of different types specified must not exceed the maximum defined in the CONTROL packet.
- > TIME INTERVAL. One or more TIME INTERVAL packets must be present. The first time interval must begin at or before the beginning of the time span specified on Card 10 of the CONTROL packet. All time intervals must be contiguous and of nonzero length. Each TIME INTERVAL packet contains one or more of the following packets and ends with an ENDTIME card. Following the first time interval, only those data that are to be changed need be specified.
- > METHOD. A method must be provided for every variable--including user-defined variables--for every point source type in the first time interval. Methods can be changed in subsequent TIME INTERVAL packets if desired. Note that each parameter entry contributes to the overall parameter count; the total number of parameters must not exceed the maximum specified in the CONTROL packet.

- > VERTICAL METHOD. A vertical method must be provided for every variable--including user-defined variables--for every point source type in the first time interval. Vertical methods can be changed in subsequent TIME INTERVAL packets if desired. Note that each vertical parameter entry contributes to the overall vertical parameter count; the total must not exceed the maximum specified in the CONTROL packet.
- > EMISSIONS VALUES. The first time interval must contain an EMISSIONS VALUES packet. More than one EMISSIONS VALUES packet can appear in any time interval.
- > EMISSIONS FACTORS. If the method EMFACTORS is selected in a subsequent time interval, the EMISSIONS FACTORS packet must appear. This packet can be used to multiply all emissions from a given point source or type by a time-varying factor. More than one EMISSIONS FACTORS packet can appear in any time interval.

## 5. Additional Input Files

The DIFFBREAK and REGIONTOP files must be input to PTSRCE. In addition, if the vertical method PLUMERISE was selected, the TEMPERATUR, METSCALARS, and WIND files must also be input.

## 6. Examples

Exhibit VII-9 shows a sample input deck for PTSOURCE file generation. The output from a PTSRCE run using that input deck is shown in Exhibit VII-10.

CONTROL	PTSOURCE	POINT SOURCES FOR ST.	LOUIS	76195	4X4 GRID
	NO	9	2	217	10
	NO2	25	0	1	
	PAR	1	0	0	1
	OLE				
	CARB				
	ARO				
	SO2				
	AERS				
	CO				
	END	76195	0500	76195	2400
	REGION				
		725000.	4252000.	15	
		-19000.	-16000.		
		4000.	4000.		
		17	22	4	
		2	2	0.	200.
	END				
	UNITS				
	NO	KG/HR			
	NO2	KG/HR			
	PAR				
	OLE				
	CARB				
	ARO	KG/HR			
	SO2	KG/HR			
	AER	KG/HR			1.
	CO	KG/HR			
	FLOWRATE	M3/MIN			
	NOX	KG/HR	60.	0.	1.
	SOX	KG/HR	1000.		46.
	END		1000.		62.
	POINT SOURCES				
	14144001C1ST.LOUIS		76900.	32100.	
		4			
	14144002C1ST.LOUIS		77713.	26692.	
		24			
	1414400401ST.LOUIS		87700.	23100.	
		12			
	1446800101ST.LOUIS		23710.	53630.	
		76			
	1446800102ST.LOUIS		23750.	53600.	
		76			
			421		

Exhibit VII-9 (Continued)

1446800103ST.LOUIS	23770.	53500.
106	417	
1446800201ST.LOUIS	31720.	38680.
9	758	
1446800307ST.LOUIS	20520.	31130.
72	438	
1446800401ST.LOUIS	27330.	52550.
48	471	
1446800407ST.LOUIS	27220.	51650.
48	563	
1446800409ST.LOUIS	27190.	52030.
33	755	
1446800414ST.LOUIS	27180.	52010.
33	568	
1446800417ST.LOUIS	27170.	52310.
23	644	
1446800421ST.LOUIS	27130.	51570.
45	1068	
1446800422ST.LOUIS	26880.	52130.
23	1032	
1446800423ST.LOUIS	27180.	52300.
15	921	
1446800424ST.LOUIS	27020.	52010.
15	286	
1446800436ST.LOUIS	27920.	51120.
17	286	
1446800464ST.LOUIS	26960.	51900.
52	574	
1446800502ST.LOUIS	27770.	50150.
24	532	

• • •

• • •

1446800304ST.LOUIS 72	20520.	31070.
1446800303ST.LOUIS 72	455	455
1446800603ST.LOUIS 4	20520.	31050.
1446900105ST.LOUIS 4	19300.	57400.
END	286	286
TIME INTERVAL 76195	500	76195
METHOD		600
ST.LOUIS ALL	EMVALUES	0.
ST.LOUIS NOX	SPLIT/COMB	50000.
NOX	0.9	1
ST.LOUIS NO2	SPLIT/COMB	50000.
NOX	0.1	1
ST.LOUIS SO2	SPLIT/COMB	50000.
SOX	1.0	1
END		
VERTICAL METHODS	PLUME RISE	0.0
ST.LOUIS ALL	10000.	
END		
EMISSIONS VALUES		
ALL ALL	0.0	
1414400104AERS	.34	
1414400201FLOW RATE	57.77	
1414400201INDX	3.12	
1414400201SOX	.17	
1414400201CO	.85	
1414400201AERS	.12	
1414400201PAR	29.27	
1414400201OLE	.37	
1414400201CARB	.37	
1414400201ARO	.12	
1414400401FLOW RATE	28.32	
1414400401INOX	20.97	
1414400401CO	2.65	
1414400401AERS	.06	
1414400401PAR	617.23	

```

CONTROL
PTSOURCE
POINT SOURCES FOR ST. LOUIS 76195 4X4 GRID
   9      2      217     1     10     0
   25     0      1       0     0     0
   1      0       0     0       1     0
   0      0       0     0       0     0
   11     12      0       0     14     15    16
NO
NO2
PAR
OLE
CARB
ARO
SO2
AERS
CO      76195      500      76195      2400
END
PROCESSING OF INPUT DATA FOR THE PREPARATION OF FILE PTSOURCE
POINT SOURCES FOR ST. LOUIS 76195 4X4 GRID

REGION
 725000.  4252000.      15
 -19000.  -16000.
 4000.00  4000.00
 17      22      4
          2      2     .00     50.00    200.00
END
KCRNT = 111 NSCLFT = 23390 IN SCRSET
KCRNT = 171 NSCLFT = 23330 IN SCRSET
KCRNT = 341 NSCLFT = 23160 IN SCRSET
KCRNT = 358 NSCLFT = 23143 IN SCRSET
KCRNT = 375 NSCLFT = 23126 IN SCRSET

```

KCRNT =	NSCLFT =	23109	IN SCRSET
KCRNT =	NSCLFT =	23103	IN SCRSET
KCRNT =	25668	NSCLFT = 20913	IN SCRSET
KCRNT =	25768	NSCLFT = 20923	IN SCRSET
KCRNT =	2795	NSCLFT = 20706	IN SCRSET
KCRNT =	30112	NSCLFT = 20489	IN SCRSET
KCRNT =	3229	NSCLFT = 20272	IN SCRSET
KCRNT =	5616	NSCLFT = 17885	IN SCRSET
KCRNT =	8003	NSCLFT = 15498	IN SCRSET
KCRNT =	8014	NSCLFT = 15487	IN SCRSET
KCRNT =	8025	NSCLFT = 15476	IN SCRSET
KCRNT =	8036	NSCLFT = 15465	IN SCRSET
KCRNT =	8047	NSCLFT = 15454	IN SCRSET
KCRNT =	8058	NSCLFT = 15443	IN SCRSET
KCRNT =	8158	NSCLFT = 15343	IN SCRSET
KCRNT =	8168	NSCLFT = 15333	IN SCRSET
KCRNT =	8179	NSCLFT = 15322	IN SCRSET
KCRNT =	8190	NSCLFT = 15311	IN SCRSET
KCRNT =	8201	NSCLFT = 15300	IN SCRSET
KCRNT =	8212	NSCLFT = 15289	IN SCRSET
KCRNT =	8223	NSCLFT = 15278	IN SCRSET
KCRNT =	8240	NSCLFT = 15061	IN SCRSET
KCRNT =	8657	NSCLFT = 14844	IN SCRSET
KCRNT =	8874	NSCLFT = 14627	IN SCRSET
KCRNT =	9091	NSCLFT = 14410	IN SCRSET
KCRNT =	9465	NSCLFT = 14036	IN SCRSET
KCRNT =	9839	NSCLFT = 13662	IN SCRSET
KCRNT =	10213	NSCLFT = 13288	IN SCRSET
KCRNT =	11709	NSCLFT = 11792	IN SCRSET
KCRNT =	13205	NSCLFT = 10296	IN SCRSET
KCRNT =	13579	NSCLFT = 9922	IN SCRSET
KCRNT =	13796	NSCLFT = 9705	IN SCRSET
KCRNT =	14013	NSCLFT = 9488	IN SCRSET
KCRNT =	14230	NSCLFT = 9271	IN SCRSET
KCRNT =	14447	NSCLFT = 9054	IN SCRSET
KCRNT =	14664	NSCLFT = 8837	IN SCRSET
KCRNT =	17051	NSCLFT = 6450	IN SCRSET
UNITS			

UNIT CONVERSION TABLE

VARIABLE NAME	UNIT NAME	MULTIPLICATIVE FACTOR	ADITIVE FACTOR
NO	KG/HR	1000.0000000	-9.0000000
NO2	KG/HR	1000.0000000	-9.0000000
PAR		-9.0000000	-9.0000000
OLE		-9.0000000	-9.0000000
CARB		-9.0000000	-9.0000000
ARO		-9.0000000	-9.0000000
SO2	KG/HR	1000.0000000	-9.0000000
AERS	KG/HR	1000.0000000	-9.0000000
CO	KG/HR	1000.0000000	-9.0000000
NOX	KG/HR	1000.0000000	-9.0000000
SOX	KG/HR	1000.0000000	-9.0000000
COORD		-9.0000000	-9.0000000
HEIGHT		-9.0000000	-9.0000000
DIAHETER		-9.0000000	-9.0000000
STACKTEMP		-9.0000000	-9.0000000
STACKVEL		-9.0000000	-9.0000000
FLOWRATE	M3/MIN	60.0000000	-9.0000000
			1.0000000

POINT SOURCES  
TIME INTERVAL  
76195      500      76195      600

METHOD  
VERTICAL METHODS  
EMISSIONS VALUES  
ENDTIME

\*\*\*\*\* NOTE \*\*\*\*\* EMISSIONS FROM POINT SOURCE 1414400104 IN CELL ( 25, 13)  
ARE OUTSIDE HORIZONTAL DIMENSIONS OF MODEL DOMAIN. EMISSIONS FROM THIS SOURCE HAVE BEEN SET TO ZERO

\*\*\*\*\* NOTE \*\*\*\*\* EMISSIONS FROM POINT SOURCE 1414400201 IN CELL ( 25, 11)  
ARE OUTSIDE HORIZONTAL DIMENSIONS OF MODEL DOMAIN. EMISSIONS FROM THIS SOURCE HAVE BEEN SET TO ZERO

\*\*\*\*\* NOTE \*\*\*\*\* EMISSIONS FROM POINT SOURCE 1414400401 IN CELL ( 27, 10)  
ARE OUTSIDE HORIZONTAL DIMENSIONS OF MODEL DOMAIN. EMISSIONS FROM THIS SOURCE HAVE BEEN SET TO ZERO

\*\*\*\*\* NOTE \*\*\*\*\* EMISSIONS FROM POINT SOURCE 1446803301 IN CELL ( 21, 15)  
ARE OUTSIDE HORIZONTAL DIMENSIONS OF MODEL DOMAIN. EMISSIONS FROM THIS SOURCE HAVE BEEN SET TO ZERO

\*\*\*\*\* NOTE \*\*\*\*\* EMISSIONS FROM POINT SOURCE 1464600101 IN CELL ( 18, 1)  
ARE OUTSIDE HORIZONTAL DIMENSIONS OF MODEL DOMAIN. EMISSIONS FROM THIS SOURCE HAVE BEEN SET TO ZERO

\*\*\*\*\* NOTE \*\*\*\*\* EMISSIONS FROM POINT SOURCE 1464600103 IN CELL ( 16, 1)  
ARE OUTSIDE HORIZONTAL DIMENSIONS OF MODEL DOMAIN. EMISSIONS FROM THIS SOURCE HAVE BEEN SET TO ZERO

Exhibit VII-10 (Continued)

## POINT SOURCE EMISSIONS (GRAM-MOLESS/HOUR) SPECIES NO

PT. SOURCE NAME	PT. SOURCE TYPE NAME	X-CELL	Y-CELL	Z-CELL	FLOW RATE (MM*3/HOUR)	PLUME HGT. (METERS)	EMS. VALUE (G-MOL/HR)	EMS. FACTOR (G-MOL/HR)	EMS. FLUX
1446800101	ST.LOUIS	11	18	3	224607.	209.2825	.00	-9.00	2778.85
1446800102	ST.LOUIS	11	18	3	143602.	186.3772	.00	-9.00	173.93
1446800103	ST.LOUIS	11	18	3	940271.	310.3101	.00	-9.00	13977.59
1446800201	ST.LOUIS	13	14	1	10610.	37.2358	.00	-9.00	57.72
1446800307	ST.LOUIS	10	12	2	708572.	267.5112	.00	-9.00	8499.52
1446800401	ST.LOUIS	12	18	3	168270.	177.9466	.00	-9.00	3082.70
1446800407	ST.LOUIS	12	17	3	232827.	214.4082	.00	-9.00	655.63
1446800409	ST.LOUIS	12	18	3	93777.	180.0519	.00	-9.00	578.15
1446800414	ST.LOUIS	12	16	2	34609.	106.4060	.00	-9.00	135.20
1469003401	ST.LOUIS	12	12	1	21472.	25.0110	.00	-9.00	19.37
26222800101	ST.LOUIS	6	1	1	7767.	41.9041	.00	-9.00	373.11
26416000101	ST.LOUIS	6	19	4	804461.	391.2513	.00	-9.00	123442.24
2642800301	ST.LOUIS	10	10	1	53908.	157.5535	.00	-9.00	1091.15
2642800302	ST.LOUIS	10	10	1	1274.	77.3614	.00	-9.00	20.35
2642800305	ST.LOUIS	10	10	1	52349.	85.0143	.00	-9.00	795.13
2642800601	ST.LOUIS	9	13	1	28226.	127.1465	.00	-9.00	328.30
2642901201	ST.LOUIS	9	9	2	88058.	216.5848	.00	-9.00	6.46
2642801202	ST.LOUIS	9	9	2	103633.	202.7230	.00	-9.00	6.46
2642801701	ST.LOUIS	10	12	1	26015.	104.6543	.00	-9.00	355.89
2642801704	ST.LOUIS	10	12	1	797.	40.1493	.00	-9.00	9.96
2642801705	ST.LOUIS	10	12	1	21491.	49.4752	.00	-9.00	444.13
2642802502	ST.LOUIS	10	11	1	41191.	104.3999	.00	-9.00	654.02
2642802401	ST.LOUIS	10	13	2	177407.	291.4135	.00	-9.00	393.07
2642802501	ST.LOUIS	9	10	2	177407.	291.7539	.00	-9.00	393.07
2642803801	ST.LOUIS	10	12	1	61006.	139.7895	.00	-9.00	722.15
2642804001	ST.LOUIS	9	11	1	9312.	85.9915	.00	-9.00	196.24
2642805601	ST.LOUIS	9	12	1	1099.	16.8529	.00	-9.00	42.26
2642805702	ST.LOUIS	10	13	1	16990.	57.9199	.00	-9.00	52.63
2642805801	ST.LOUIS	10	12	1	43367.	91.3986	.00	-9.00	498.52
2642805901	ST.LOUIS	9	11	1	16990.	65.4218	.00	-9.00	.96
2642806001	ST.LOUIS	9	13	1	16990.	65.0096	.00	-9.00	1.96
2643000101	ST.LOUIS	7	15	1	407762.	258.4892	.00	-9.00	2244.33
2643000206	ST.LOUIS	6	9	1	4499.	57.4342	.00	-9.00	368.80
2643000701	ST.LOUIS	6	15	1	5097.	28.8824	.00	-9.00	216.39
2643000704	ST.LOUIS	6	15	1	7287.	42.0511	.00	-9.00	102.91
2643000708	ST.LOUIS	7	15	1	3309.	34.5166	.00	-9.00	65.30
2643000801	ST.LOUIS	9	14	3	679603.	299.2691	.00	-9.00	4935.72
2643000901	ST.LOUIS	9	9	1	57008.	88.8184	.00	-9.00	620.02
2643000903	ST.LOUIS	9	9	1	71429.	93.7199	.00	-9.00	1331.02
2643000905	ST.LOUIS	9	9	1	23143.	54.6988	.00	-9.00	126.78
2643000910	ST.LOUIS	9	9	1	47572.	75.3069	.00	-9.00	68.43
2643000911	ST.LOUIS	9	9	1	110435.	82.1603	.00	-9.00	88.43
2643000913	ST.LOUIS	9	9	1	118931.	96.3039	.00	-9.00	88.43
2643001001	ST.LOUIS	7	5	3	244627.	205.0220	.00	-9.00	11451.13
2643001003	ST.LOUIS	7	5	3	832438.	311.3381	.00	-9.00	32980.50
2643001401	ST.LOUIS	8	8	1	4078.	78.8097	.00	-9.00	37.96
2643001601	ST.LOUIS	6	12	1	8860.	65.0652	.00	-9.00	156.33
2643003101	ST.LOUIS	9	4	1	0.	17.9998	.00	-9.00	101.74

FILE NAME -- PTSOURCE	POINT SOURCE EMISSIONS (GRAM-MOLES/HOUR)	SPECIES NO	POINT SOURCES FOR ST. LOUIS	76195	4X4 GRID	VERT.	LEVEL 0	START DATE/TIME	76195	5.00	STOP DATE/TIME	76195	6.00
GRID CELL	1	2	3	4	5	6	7	8	9	10	11	12	
22	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	
21	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	
20	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	
19	.00	.00	.00	.00	.00	.00	123442.24	.00	.00	21.33	.00	.00	
18	.00	.00	.00	.00	.00	.00	.00	.00	.00	639.59	22346.61	4501.76	
17	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	5216.20	
16	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	
15	.00	.00	.00	.00	.00	319.30	85.30	.00	.00	.00	.00	.00	
14	.00	.00	.00	.00	.00	.00	.00	.00	4935.72	.00	19.76	.00	
13	.00	.00	.00	.00	.00	.00	.00	.00	330.26	445.70	6952.89	.00	
12	.00	.00	.00	.00	.00	.00	.00	156.33	42.26	10530.20	292.11	19.37	
11	.00	.00	.00	.00	.00	.00	.00	.00	197.22	854.02	1329.46	.00	
10	.00	.00	.00	.00	.00	.00	.00	.00	393.07	1906.63	2464.83	9.76	
9	.00	.00	470.54	.00	.00	.00	.00	.00	2356.04	.00	.00	.00	
8	.00	.00	.00	.00	.00	2244.33	37.96	.00	.00	.00	.00	.00	
7	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	
6	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	
5	.00	.00	.00	.00	.00	.00	44431.63	.00	.00	.00	.00	.00	
4	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	
3	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	
2	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	.00	
1	.00	.00	.00	.00	.00	373.11	.00	.00	.00	.00	.00	.00	

GRID CELL	POINT SOURCE EMISSIONS (GRAM-MOLLES/HOUR)	SPECIES NO	VERT.	LEVEL 0 START DATE/TIME 76195 5.00 STOP DATE/TIME 76195 6.00
22	.00	13	14	.00
21	.00	14	15	.00
20	.00	15	16	.00
19	.00	16	17	.00
18	.00	17	18	.00
17	1761.46	18	19	.00
16	.00	19	20	.00
15	.00	20	21	.00
14	57.72	21	22	.00
13	.00	22	23	.00
12	.00	23	24	.00
11	.00	24	25	.00
10	.00	25	26	.00
9	.00	26	27	191.74
8	.00	27	28	.00
7	.00	28	29	.00
6	63.39	29	30	.00
5	.00	30	31	.00
4	.00	31	32	.00
3	.00	32	33	.00
2	.00	33	34	.00
1	.00	34	35	.00

## G. REGIONTOP

The REGIONTOP file contains time-varying two-dimensional matrices of region top heights. The region top height can be linked to the diffusion break or can be independently specified. The program REGNTP is used to create the REGIONTOP file. REGNTP requires subroutines from the libraries UTILITY and FILUTIL. Figure VII-13 is an information flow diagram illustrating the job control to be used. In the following sections we discuss the input to the REGNTP program.

### 1. Variables

The output variable for the REGIONTOP file is also named REGIONTOP. This is an "implicit output variable" and need not be referred to anywhere in the CONTROL packet. The internal units for REGIONTOP are meters (m). If the input values for this variable are to be in any other units, REGIONTOP must be specified in the UNITS packet.

### 2. Methods

The methods that can be used to generate the REGIONTOP file are:

- > CONSTANT
- > GRID VALUE
- > STATINTERP
- > POISSON
- > FIXDHEIGHT
- > SAMEHEIGHT
- > E-WINTERP
- > N-SINTERP
- > USER.

These methods are discussed in detail in Section VI.A.6.

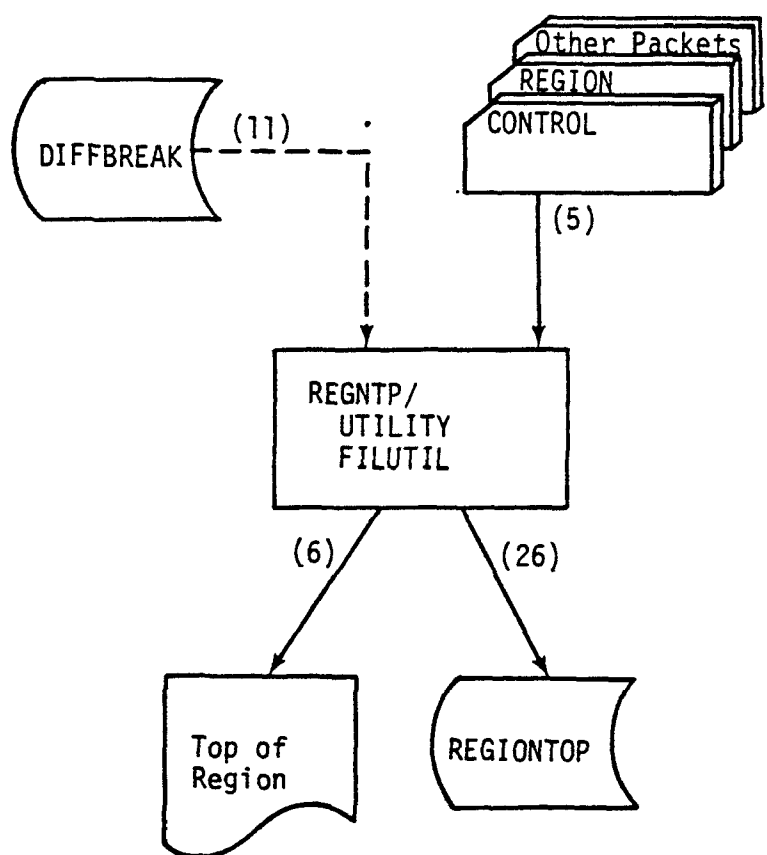


Figure VII-13. Information Flow Diagram for the REGIONTOP File

### 3. Time Variation

The top of the region values on the REGIONTOP file are considered to apply at the beginning of the time interval. Because the Airshed Simulation Program calculates region top values continuously over time by linear interpolation, it also requires values at the end of the time interval. These are read by the program as the values at the beginning of the next time interval. Thus, the last time interval on the file must begin at or after the ending simulation time. For example, if a simulation is to be made from 0500-1700 and region top values are input hourly, the values used between 500 and 600 are calculated by interpolating between values input for the 0500-0600 time interval and those input for the 0600-0700 time interval. Similarly, to calculate values between 1600 and 1700, the simulation program requires values for the interval 1600-1700, and another set for an interval beginning at 1700.

### 4. Packets

Figure VII-14 shows the input deck structure in terms of the packets to be used. Each of these packets is described in detail in Sections VI.B. Following are special input packet considerations for the REGIONTOP file:

- > CONTROL
  - The file name on Card 2 must be REGIONTOP.
  - The control variables to be specified on Cards 4 to 8 for REGIONTOP are shown in Table VII-6.
  - The number of species should be zero.
  - If there are input variables that do not appear as output variables, their number must appear as the number of user-defined variables. All such variables must also be named in the UNITS packet.
  - If data from measuring stations are to be used (methods STATINTERP or POISSON), the maximum number of such stations must be given.
  - The number of subregions must be at least one.

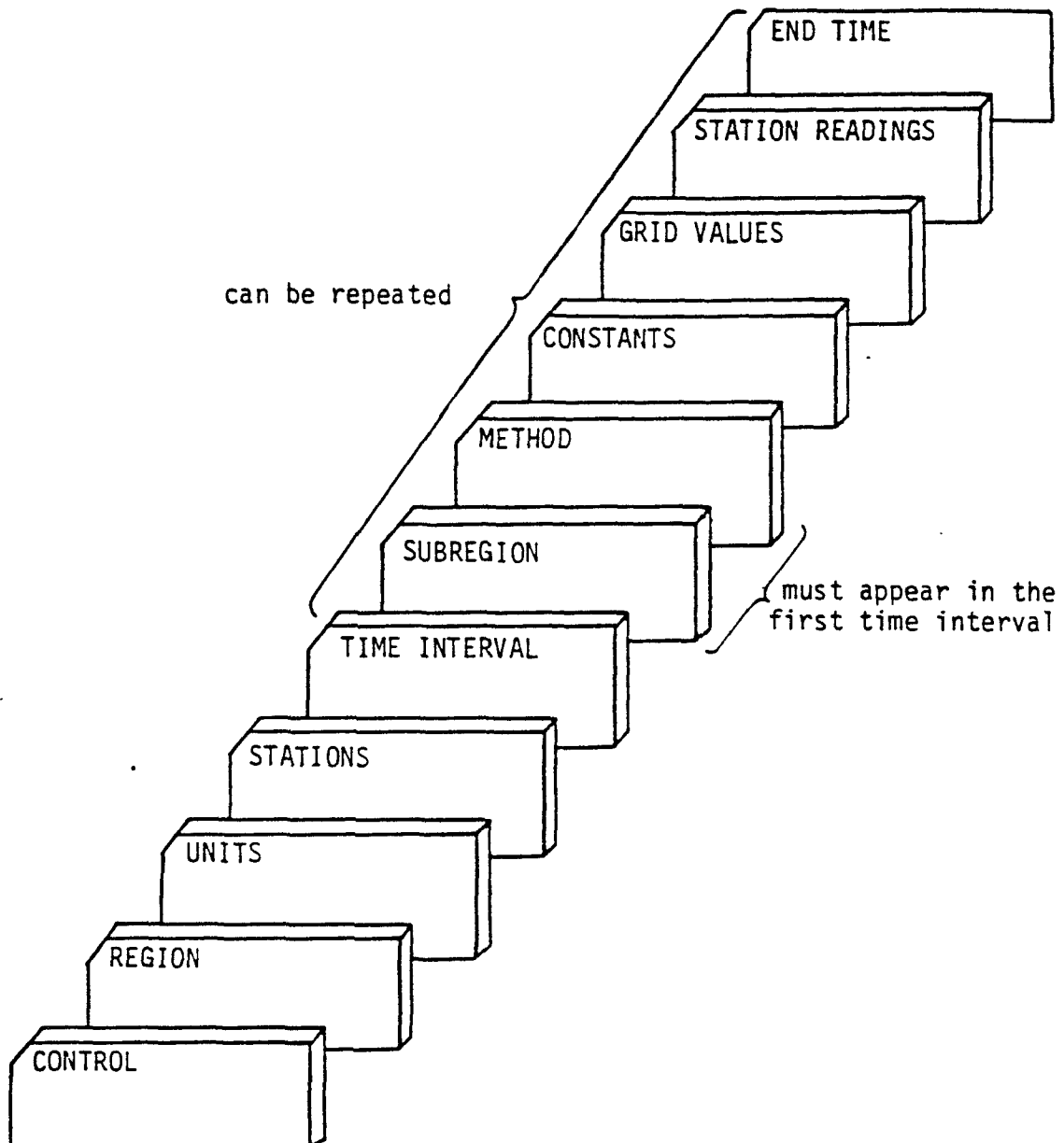


Figure VII-14. Input Card Deck Structure for Preparing the REGIONTOP File

Table VII-6. CONTROL Packet Entries for the REGIONTOP File

<u>Card Number</u>	<u>Entry</u>
4	Number of species (=0) Number of user-defined variables Number of stations Number of subregions Number of parameters Spare
5	Output file number Print input cards Print output grid Spare Spare Spare
6	Print units table Print station locations table Print regional grid Print methods table Print station values table Spare
7	Spare Spare Spare Spare Spare Spare
8	DIFFBREAK file unit number Spare Spare Spare Spare Spare

- The maximum number of parameters must be sufficient to include all specifications of all parameters.
  - The vertical controls card (No. 7) should be left blank.
  - The file unit assignment card (No. 8) must specify DIFFBREAK if the method FIXDHEIGHT or SAMEHEIGHT is selected. Otherwise, it should be left blank.
  - The beginning and ending dates and times should reflect the time variation considerations discussed in Section 3 above.
  - A set of output species names is not required; if they are present, their number must be the same as the entry in the first control parameter on Card 4, but they will be ignored by the program.
- > REGION. This packet must follow the CONTROL packet. The vertical parameters must be provided if FIXDHEIGHT or SAMEHEIGHT is selected. Otherwise, they will be ignored.
- > UNITS. This packet, if present, must follow the REGION packet. The UNITS packet must be provided if:
- Any input variable will be input in other than internal units.
  - Any user-defined variables are specified.
  - COORD or HEIGHT unit conversions are to be used.
- The number of user-defined variables must not exceed the maximum specified in the CONTROL packet.
- > STATIONS. This packet is required if either of the methods STATINTERP or POISSON is specified. The number of stations listed must not exceed the maximum specified in the CONTROL packet.
- > TIME INTERVAL. Two or more TIME INTERVAL packets must be present. The first time interval must begin at or before the beginning of the time span specified on Card 10 of the CONTROL packet. The last time interval must begin at or after the ending time of any simulation run. All time intervals must be contiguous and of nonzero length. Each TIME INTERVAL packet contains one or more of the following

packets and ends with an ENDTIME card. Following the first time interval, only those data that are to be changed need be specified.\*

- > SUBREGION. The first time interval must contain a SUBREGION packet; the inclusion of this packet in other time intervals is optional. The number of subregions must not exceed the maximum specified in the CONTROL packet.
- > METHOD. A method must be provided for every variable--including user-defined variables--in every subregion in the first time interval. Methods can be changed in subsequent TIME INTERVAL packets if desired. Note that each parameter entry contributes to the overall parameter count; the total number of parameters must not exceed the maximum specified in the CONTROL packet.
- > CONSTANTS. If the method CONSTANT is assigned to any variable in the METHOD packet, the first time interval must contain a CONSTANTS packet. More than one CONSTANTS packet can appear in any time interval.
- > GRID VALUES. If the method GRID VALUE is assigned to any variable in the METHOD packet, the first time interval must contain a GRID VALUES packet. More than one GRID VALUES packet can appear in any time interval.
- > STATION READINGS. If either the POISSON or STATINTERP method is assigned to any variable in the METHOD packet, the first time interval must contain a STATION READINGS packet. More than one STATION READINGS packet can appear in any time interval.

## 5. Additional Input Files

If the method FIXDHEIGHT or SAMEHEIGHT was selected, the DIFFBREAK file must be input to REGNTP. Otherwise no additional input files are required.

---

\* If FIXDHEIGHT or SAMEHEIGHT is used, the time intervals specified here must be exactly the same as those on the DIFFBREAK file; after the first time interval, subsequent packets may contain no other packets.

6. Examples

Exhibit VII-11 shows a sample input deck for REGIONTOP file generation. The output from a REGNTP run using that input deck is shown in Exhibit VII-12.

```

CONTROL
REGIONTOP
HEIGHT: OF THE TOP OF THE MODELING REGION
      26      1      1      1      1      5
      1      1      1      1      0
      11
END
REGION
      725000.  4252000.    15
      -19000.  -15000.
      4000.    4000.
      17      22      4
      2       2       0.
      50.     200.
TIME INTERVAL
      76195    500    76195    600
SUBREGION
      A       1       1      -1
END
METHOD
      A      REGIONTOP FIXHEIGHT
      500.   2038.      1
UPCELLHT
      200.
END
ENDTIME
TIME INTERVAL
      76195    600    76195    700
ENDTIME
TIME INTERVAL
      76195    700    76195    800
ENDTIME
TIME INTERVAL
      76195    800    76195    900
ENDTIME
TIME INTERVAL
      76195    900    76195    1000
ENDTIME

```

TIME INTERVAL	76195	1000	76195	1100
ENDTIME				
TIME INTERVAL	76195	1100	76195	1200
ENDTIME				
TIME INTERVAL	76195	1200	76195	1300
ENDTIME				
TIME INTERVAL	76195	1300	76195	1400
ENDTIME				
TIME INTERVAL	76195	1400	76195	1500
ENDTIME				
TIME INTERVAL	76195	1500	76195	1600
ENDTIME				
TIME INTERVAL	76195	1600	76195	1700
ENDTIME				
TIME INTERVAL	76195	1700	76195	1800
ENDTIME				
TIME INTERVAL	76195	1800	76195	1900
ENDTIME				
TIME INTERVAL	76195	1900	76195	2000
ENDTIME				
TIME INTERVAL	76195	2000	76195	2100
ENDTIME				
TIME INTERVAL	76195	2100	76195	2200
ENDTIME				
TIME INTERVAL	76195	2200	76195	2300
ENDTIME				
TIME INTERVAL	76195	2300	76195	2400
ENDTIME				
TIME INTERVAL	76195	2400	76196	10
ENDTIME				

PROCESSING OF INPUT DATA FOR THE PREPARATION OF FILE REGIONTOP

HEIGHT OF THE TOP OF THE MODELING REGION

NUMBER OF STATIONS = 0  
NUMBER OF SUBREGIONS = 1  
NUMBER OF METHOD PARAMETERS = 5  
PROGRAM OUTPUT WILL BE PLACED ON FILE NUMBER 26  
INPUT DATA FOR DIFFUSION BREAK WILL BE OBTAINED FROM FILE NUMBER 11  
INPUT CARDS WILL BE PRINTED  
OUTPUT GRID WILL BE PRINTED  
UNIT TABLE WILL BE PRINTED  
STATION LOCATIONS WILL BE PRINTED  
SUBREGION GRID WILL BE PRINTED  
METHOD TABLE WILL BE PRINTED  
STATION VALUES WILL NOT BE PRINTED  
BEGINNING DATE OF FILE 76195 BEGINNING TIME OF FILE 5.00  
ENDING DATE OF FILE 76196 ENDING TIME OF FILE .17  
  
REGION  
725000. 4252000. 15  
-19000. -16000.  
4000.00 4000.00  
17 22 4  
2 2 .00 50.00 200.00  
END

Exhibit VII-12. Sample Output from the REGNTP Program

FILE NAME -- REGIONTOP		HEIGHT OF THE TOP OF THE MODELING REGION											
		REGION 10P:VALUES (IN METERS ABOVE GROUND LEVEL)											
GRID CELL		1	2	3	4	5	6	7	8	9	10	11	12
22	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000
21	502.0000	504.0061	505.9784	507.8764	509.6442	511.2035	512.4517	513.2621	513.4869	512.9678	511.5785	509.4017	
20	502.0000	506.0492	510.0391	513.0977	517.5229	520.7653	523.4199	525.2271	525.8663	524.9496	522.0040	516.6038	
19	502.0000	508.1576	514.2490	520.1900	525.8482	531.0156	535.3871	538.5582	540.0230	539.1342	534.8865	524.9782	
18	502.0000	510.3394	516.6380	526.0301	534.7796	542.2370	548.8003	553.8979	556.8324	556.8625	553.3217	545.7940	
17	502.0000	512.5715	523.1758	533.0129	544.3856	554.6312	564.0430	571.8205	576.9206	578.3270	575.4780	568.5566	
16	502.0000	514.7858	527.7453	541.0099	554.5890	568.2632	581.4332	592.9575	601.1329	604.1504	601.2368	592.7235	
15	502.0000	516.8553	532.1119	548.1159	565.0816	582.9509	601.1151	618.0985	630.9451	635.9049	631.8983	619.3304	
14	502.0000	518.5852	535.8983	554.5894	575.1980	590.0248	622.8621	648.0858	669.0660	676.4828	670.2190	649.5590	
13	502.0000	519.7123	538.5708	559.6205	583.7796	611.9117	645.0963	683.0550	721.0929	730.4662	721.0950	683.8507	
12	502.0000	519.9191	539.5127	562.1920	589.1754	622.0373	663.4087	718.6224	802.0000	802.0000	802.0000	718.1304	
11	502.0000	518.8524	537.9964	561.3214	589.7096	624.6137	668.7601	726.5473	802.0000	802.0000	802.0000	724.6042	
10	502.0000	516.1240	533.1719	556.4365	586.8351	618.9656	661.2449	717.1955	802.0000	802.0000	802.0000	713.3699	
9	502.0000	511.3740	523.2244	547.5830	575.3284	606.0785	640.6791	679.2575	717.1860	725.2433	714.3020	672.3603	
8	502.0000	505.3165	502.0000	536.5030	563.7951	590.0691	616.5829	642.1357	662.1583	667.2108	657.2377	631.7732	
7	502.0000	507.2511	514.7118	533.6508	554.0439	574.3307	593.7312	610.6377	622.0515	624.0679	615.5090	596.6989	
6	502.0000	508.3177	517.0373	530.1234	544.9114	559.7628	573.5247	584.6783	591.3240	591.4567	583.9997	568.4714	
5	502.0000	508.1039	515.7802	525.3862	535.9775	546.5089	555.9673	563.2427	567.1079	566.4374	560.5847	549.4631	
4	502.0000	507.0670	513.0524	519.9097	527.2197	534.3517	540.6021	545.2149	547.4289	546.6159	542.4751	535.1694	
3	502.0000	505.6063	509.6386	514.0663	510.6635	523.0710	526.8604	529.5737	530.7749	530.1351	527.5416	523.2044	
2	502.0000	503.8637	505.8902	508.0631	510.2833	512.3872	514.1739	515.4294	515.9563	515.6151	514.3737	512.3549	
1	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	

	REGION TOPVALUES (IN METERS ABOVE GROUND LEVEL)						VERT. LEVEL 1 START DATE/TIME 76195 5.00 STOP DATE/TIME 76195 6.00
GRID CELL	13	14	15	16	17		
22	502.0000	502.0000	502.0000	502.0000	502.0000	502.0000	
21	507.2663	507.4373	507.3662	505.8242	502.0000		
20	509.6637	511.9636	512.2625	509.0659	502.0000		
19	502.0000	516.0000	516.0606	512.6005	502.0000		
18	535.3789	532.9462	527.3580	517.1122	502.0000		
17	558.9566	549.4731	537.5922	521.9777	502.0000		
16	580.1956	565.2262	547.5758	526.6674	502.0000		
15	601.3668	580.3163	556.8746	530.8992	502.0000		
14	623.0223	594.5563	565.1207	534.4611	502.0000		
13	644.1440	606.8965	571.6151	537.0459	502.0000		
12	660.6950	614.9730	575.1758	538.1938	502.0000		
11	663.9149	615.4656	574.4594	537.4240	502.0000		
10	653.7866	607.5654	568.9463	534.5254	502.0000		
9	629.6791	591.6240	558.8958	529.6244	502.0000		
8	600.6625	570.1863	545.3731	523.1938	502.0000		
7	571.0517	543.1141	529.3504	515.9792	502.0000		
6	543.7772	502.0000	513.1119	509.5702	502.0000		
5	533.6663	515.0389	511.6856	507.3430	502.0000		
4	525.6634	516.0918	510.5570	506.2150	502.0000		
3	517.7668	512.3746	508.3042	505.0166	502.0000		
2	509.8677	507.3805	505.3094	503.5776	502.0000		
1	502.0000	502.0000	502.0000	502.0000	502.0000		
TIME INTERVAL	76195	600	76195	700			
EENDTIME							

## H. ROADWAY (Not recommended for use at this time)

The ROADWAY file contains time-varying matrices of emission fluxes along roadways and a descriptive parameter for automobile traffic. The ROADWAY file can be omitted from a simulation run if surface layer micro-scale calculations are not to be performed. At present, there is no general purpose program for creating the ROADWAY file because its generation is highly dependent on the form of the traffic data available. There is a program structure, similar to that of the other data preparation programs, that includes the reading of control packets, unit conversions, and writing the proper output records. This program can be tailored to particular input requirements as necessary. The skeleton program is named ROADWY. It requires subroutines from the libraries UTILITY and FILUTIL. Figure VII-15 is an information flow diagram illustrating the job control to be used. In the following sections we discuss the card input to the ROADWY program.

### 1. Variables

The output variables for the ROADWAY file are the species named in the CONTROL packet and the parameter CARM, which is to be calculated from the traffic data. Additional user-defined input variables (e.g. "reactive hydrocarbons") can be specified in the UNITS packet. The internal units for the roadway emissions of all species except AEROSOLS are gram-moles per hour (g-mol/hr); for AEROSOLS, the units are grams per hour (g/hr). The internal units for CARM are meters per hour (m/hr). The standard names for reactive species recognized by the Airshed Simulation Program are listed in Chapter IX. If any of these species does not appear on the ROADWAY file, the roadway emissions will default to zero. If any additional species appear on ROADWAY, they will be ignored.

### 2. Methods

The methods that can be used to generate the ROADWAY file are:

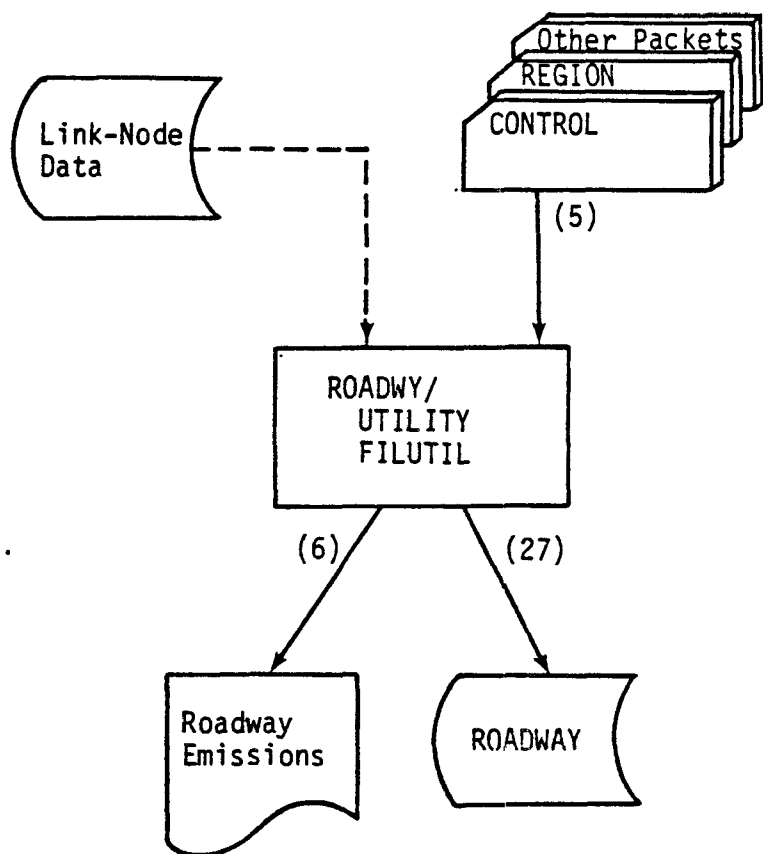


Figure VII-15. Information Flow Diagram for the ROADWAY File

- > CONSTANT
- > ROADVALUE
- > SPLIT/COMB
- > USER.

The methods ROADVALUE and USER have not been defined. The actual methods for generating output values from input traffic data must be written for each new application.

### 3. Time Variation

The time span of the ROADWAY file must include the entire time span of the simulation runs for which it is to be used. Roadway emissions and the parameter CARM are considered to be constant during each time interval.

### 4. Packets

The packets to be used for ROADWY will depend on the particular version that is developed for a given set of traffic data. Information on control parameters, definition of link-node data, units, time intervals, computation methods, and data entry will have to be provided when the program has been implemented. Figure VII-16 shows a possible input structure in terms of the packets to be used.

### 5. Additional Input Files

In general, a traffic data file in link-node form must be provided.

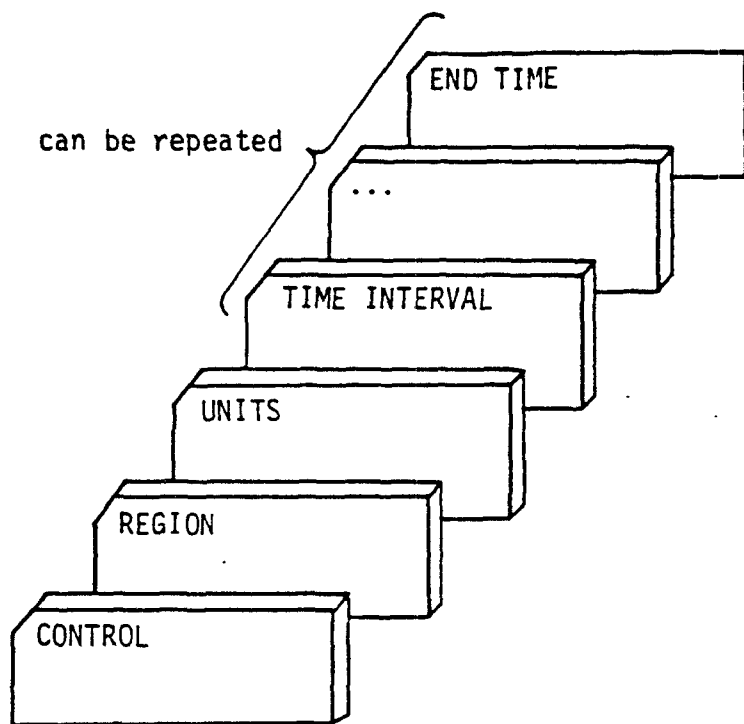


Figure VII-16. A Possible Input Card Deck Structure for Preparing the ROADWAY File

## I. TEMPERATUR

The TEMPERATUR file contains time-varying matrices of temperatures at ground level. If it is omitted from a simulation run, temperature-dependent calculations will not be performed (see Section IX.B). The program TMPTR is used to create the TEMPERATUR file. TMPTR requires subroutines from the libraries UTILITY and FILUTIL. Figure VII-17 is an information flow diagram illustrating the job control to be used. In the following sections we discuss the input to the TMPTR program.

### 1. Variables

The output variable for the TEMPERATUR file is also named TEMPERATUR. This is an "implicit output variable" and need not be referred to anywhere in the CONTROL packet. The internal units for TEMPERATUR are °K. If the input values for this variable are to be in any other units, TEMPERATUR must be specified in the UNITS packet..

### 2. Methods

The methods that can be used to generate the TEMPERATUR file are:

- > CONSTANT
- > GRID VALUE
- > STATINTERP
- > POISSON
- > E-WINTERP
- > N-SINTERP
- > USER.

These methods are discussed in detail in Section VI.A.6.

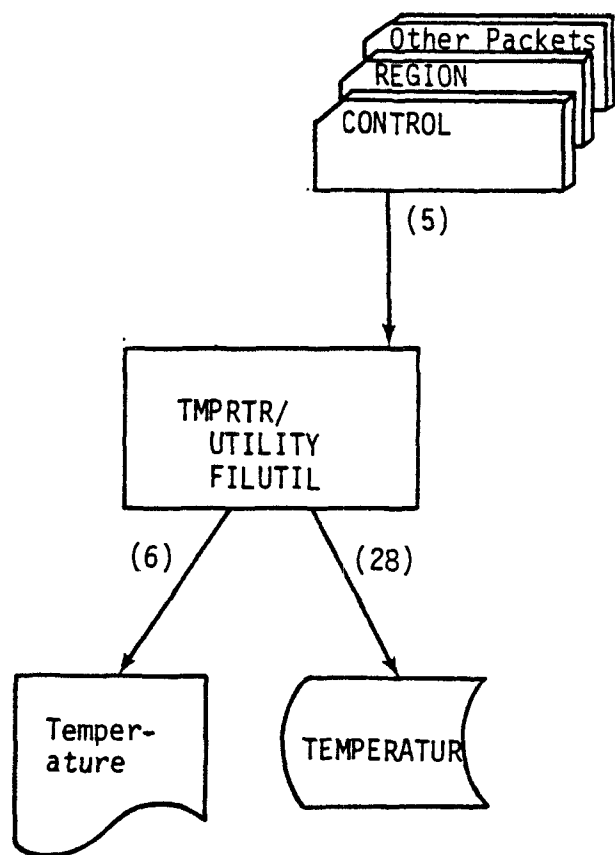


Figure VII-17. Information Flow Diagram for the TEMPERATUR File

### 3. Time Variation

The time span of the TEMPERATUR file must include the entire time span of the simulation runs for which it is to be used. Ground-level temperatures are considered to be constant during each time interval.

### 4. Packets

Figure VII-18 shows the input deck structure in terms of the packets to be used. Each of these packets is described in detail in Section VI.B. Following are special input packet considerations for the TEMPERATUR file:

- > CONTROL
  - The file name on Card 2 must be TEMPERATUR.
  - The control variables to be specified on Cards 4 to 8 for TEMPERATUR are shown in Table VII-7.
  - The number of species should be zero.
  - If there are input variables that do not appear as output variables, their number must appear as the number of user-defined variables. All such variables must also be named in the UNITS packet.
  - If data from measuring stations are to be used (methods STATINTERP or POISSON), the maximum number of such stations must be given.
  - The number of subregions must be at least one.
  - The maximum number of parameters must be sufficient to include all specifications of all parameters.
  - The vertical controls card (No. 7) should be left blank.
  - The file unit assignment card (No. 8) should be left blank.
  - The beginning and ending dates and times should reflect the time variation considerations discussed in Section 3 above.

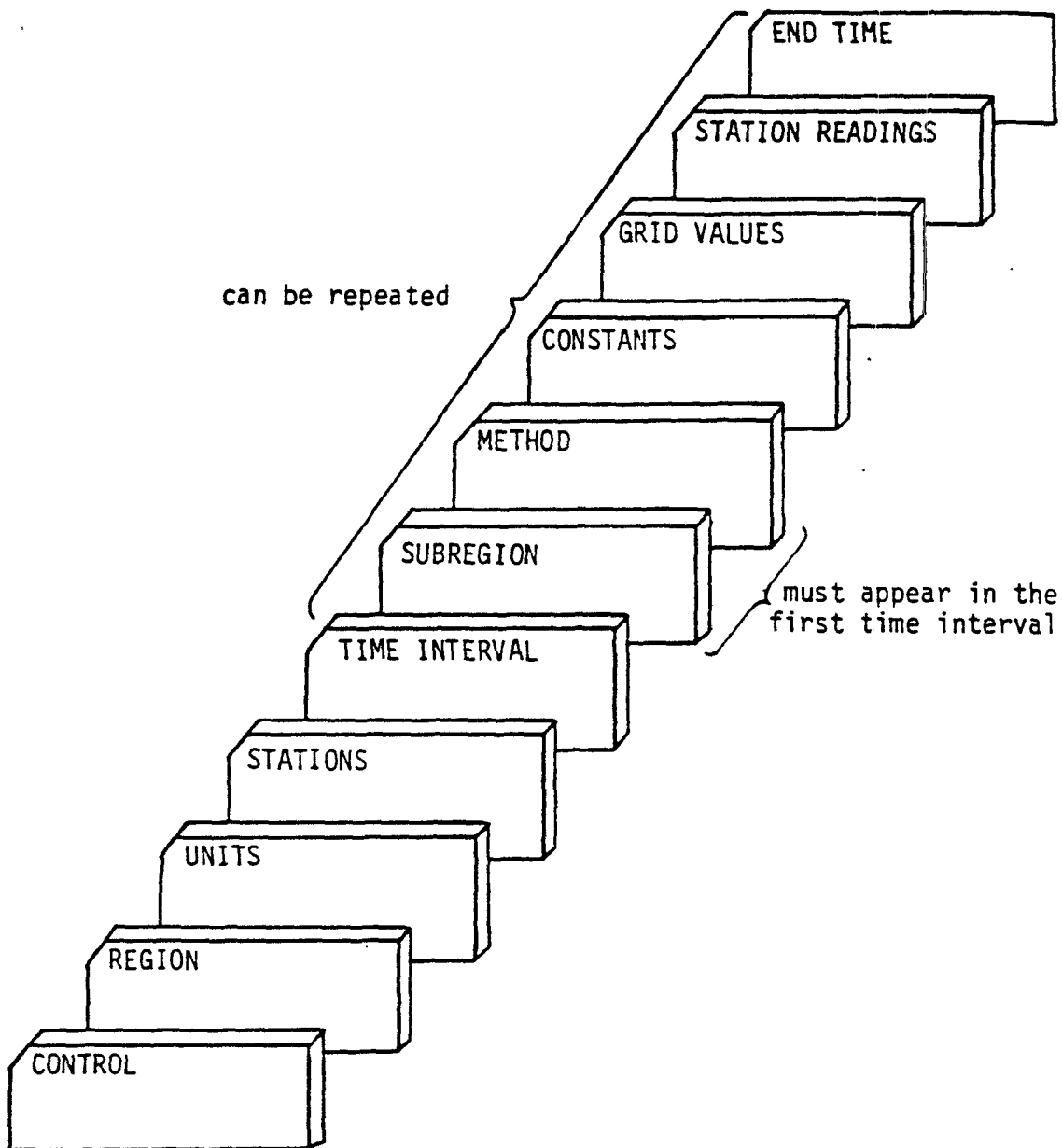


Figure VII-18. Input Card Deck Structure for Preparing the TEMPERATUR File

Table VII-7. CONTROL Packet Entries for the TEMPERATUR File

<u>Card Number</u>	<u>Entry</u>
4	Number of species (=0) Number of user-defined variables Number of stations Number of subregions Number of parameters Spare
5	Output file number Print input cards Print output grid Spare Spare Spare
6	Print units table Print station locations table Print regional grid Print methods table Print station values table Spare
7	Spare Spare Spare Spare Spare Spare
8	Spare Spare Spare Spare Spare Spare

- A set of output species names is not required; if they are present, their number must be the same as the entry in the first control parameter on Card 4, but they will be ignored by the program.
- > REGION. This packet must follow the CONTROL packet. The vertical parameters will be ignored for the TEMPERATUR file.
- > UNITS. This packet, if present, must follow the REGION packet. The UNITS packet must be provided if:
  - Any input variable will be input in other than internal units.
  - Any user-defined variables are specified.
  - COORD or HEIGHT unit conversions are to be used.The number of user-defined variables must not exceed the maximum specified in the CONTROL packet.
- > STATIONS. This packet is required if either of the methods STATINTERP or POISSON is specified. The number of stations listed must not exceed the maximum specified in the CONTROL packet.
- > TIME INTERVAL. One or more TIME INTERVAL packets must be present. The first time interval must begin at or before the beginning of the time span specified on Card 10 of the CONTROL packet. All time intervals must be contiguous and of nonzero length. Each TIME INTERVAL packet contains one or more of the following packets and ends with an ENDTIME card. Following the first time interval, only those data that are to be changed need be specified.
- > SUBREGION. The first time interval must contain a SUBREGION packet; the inclusion of this packet in other time intervals is optional. The number of subregions must not exceed the maximum specified in the CONTROL packet.
- > METHOD. A method must be provided for every variable--including user-defined variables--in every subregion in the first time interval. Methods can be changed in subsequent TIME INTERVAL packets if desired. Note that each parameter

entry contributes to the overall parameter count; the total number of parameters must not exceed the maximum specified in the CONTROL packet.

- > CONSTANTS. If the method CONSTANT is assigned to any variable in the METHOD packet, the first time interval must contain a CONSTANTS packet. More than one CONSTANTS packet can appear in any time interval.
- > GRID VALUES. If the method GRID VALUE is assigned to any variable in the METHOD packet, the first time interval must contain a GRID VALUES packet. More than one GRID VALUES packet can appear in any time interval.
- > STATION READINGS. If either the POISSON or STATINTERP method is assigned to any variable in the METHOD packet, the first time interval must contain a STATION READINGS packet. More than one STATION READINGS packet can appear in any time interval.

## 5. Additional Input Files

No additional input files are required.

## 6. Examples

Exhibit VII-13 shows a sample input deck for TEMPERATUR file generation. The output from a TMPTR run using that input deck is shown in Exhibit VII-14.

**CONTROL**  
**TEMPERATURE FIELD FOR ST. LOUIS DAY 195**

0	0	25	1	10
28	1	1		

**END**  
**REGION**  
725000. 4252000. 15  
-19000. -16000.  
4000. 4000.  
17 22 4  
2 2 0.0 50. 200.  
**END**  
**UNITS**  
**TEMPERATUREDEGREES C**  
1. 273. 1.  
**END**

**Exhibit VII-13.** Sample Input Deck for the TMPRTTR Program

STATIONS	101	19183.	27862.	10.
	102	17518.	34065.	10.
	103	22588.	30467.	10.
	104	22312.	25304.	10.
	105	18706.	24453.	10.
	106	13660.	25566.	10.
	107	15179.	30610.	10.
	108	23407.	39102.	10.
	109	30802.	27886.	10.
	110	22209.	20826.	10.
	111	13812.	20479.	10.
	112	8938.	28913.	10.
	113	12738.	37820.	10.
	114	19320.	45466.	10.
	115	32111.	45799.	10.
	116	37777.	38083.	10.
	117	35560.	20818.	10.
	118	18065.	11256.	10.
	119	4759.	18547.	10.
	120	-1921.	33909.	10.
	121	7414.	50376.	10.
	122	16631.	77223.	10.
	123	52320.	34378.	10.
	124	24475.	-15463.	10.
	125	-27555.	30240.	10.
END	TIME INTERVAL			
	76195	500	76195	600
SUBREGION	A	1	1	-1
END	METHOD	TEMPERATURPOISSON		
	A	MAXITER	100	
ERRORTOL	0.05			
ONEGA	1.4			
END				

Exhibit VII-13 (Continued)

**Exhibit VII-13 (Continued)**

<b>STATION READINGS</b>	
101	TEMPERATUR
102	TEMPERATUR
103	TEMPERATUR
104	TEMPERATUR
105	TEMPERATUR
106	TEMPERATUR
107	TEMPERATUR
108	TEMPERATUR
109	TEMPERATUR
110	TEMPERATUR
111	TEMPERATUR
112	TEMPERATUR
113	TEMPERATUR
114	TEMPERATUR
115	TEMPERATUR
116	TEMPERATUR
117	TEMPERATUR
118	TEMPERATUR
119	TEMPERATUR
120	TEMPERATUR
121	TEMPERATUR
122	TEMPERATUR
123	TEMPERATUR
124	TEMPERATUR
125	TEMPERATUR
	END

**STATION READINGS**

101	TEMPERATUR	22.0
102	TEMPERATUR	21.3
103	TEMPERATUR	20.3
104	TEMPERATUR	20.6
105	TEMPERATUR	21.6
106	TEMPERATUR	22.2
107	TEMPERATUR	22.0
108	TEMPERATUR	20.4
109	TEMPERATUR	20.0
110	TEMPERATUR	20.2
111	TEMPERATUR	21.2
112	TEMPERATUR	21.6
113	TEMPERATUR	20.9
114	TEMPERATUR	20.0
115	TEMPERATUR	20.3
116	TEMPERATUR	19.4
117	TEMPERATUR	19.6
118	TEMPERATUR	20.0
119	TEMPERATUR	21.1
120	TEMPERATUR	21.9
121	TEMPERATUR	20.4
122	TEMPERATUR	19.5
123	TEMPERATUR	19.1
124	TEMPERATUR	19.1
125	TEMPERATUR	21.2

END

EIDOTIME  
TIME INTERVAL

76195      600

TEMP FOR 600

## PROCESSING OF INPUT DATA FOR THE PREPARATION OF FILE TEMPERATURE

TEMPERATURE FIELD FOR ST. LOUIS DAY 195

NUMBER OF STATIONS = 25  
NUMBER OF SUBREGIONS = 1  
NUMBER OF METHOD PARAMETERS = 10  
PROGRAM OUTPUT WILL BE PLACED ON FILE NUMBER 28  
INPUT CARDS WILL BE PRINTED  
OUTPUT GRID WILL BE PRINTED  
UNIT TABLE WILL NOT BE PRINTED  
STATION LOCATIONS WILL NOT BE PRINTED  
SUBREGION GRID WILL NOT BE PRINTED  
METHOD TABLE WILL NOT BE PRINTED  
STATION VALUES WILL NOT BE PRINTED  
BEGINNING DATE OF FILE 76195 BEGINNING TIME OF FILE 5.00  
ENDING DATE OF FILE 76195 ENDING TIME OF FILE 24.00

REGION	725000.	4252000.	15
	-19000.	-16000.	
	4000.00	4000.00	
17	22	4	
2	2	.00	50.00 200.00
END			

Exhibit VII-14. Sample Output from the TMRTR Program

TEMPERATURE VALUES (IN DEGREES KELVIN AT GROUND LEVEL)												VERT.	LEVEL 1	START DATE/TIME	76195 5.00	STOP DATE/TIME	76195 6.00	PAGE 1
GRID CELL	1	2	3	4	5	6	7	8	9	10	11	12						
22	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	
21	292.1000	292.1427	292.1843	292.2232	292.2577	292.2863	292.3070	292.3189	292.3246	292.3269	292.3273	292.3226						
20	292.1000	292.1868	292.2711	292.3501	292.4201	292.4774	292.5169	292.5343	292.5378	292.5351	292.5304	292.5169						
19	292.1000	292.2338	292.3639	292.4858	292.5964	292.6837	292.7434	292.7542	292.7425	292.7251	292.7116	292.6927						
18	292.1000	292.2853	292.4657	292.6351	292.7871	292.9168	293.0133	292.9875	292.9389	292.8914	292.8729	292.8534						
17	292.1000	292.3425	292.5791	292.8013	293.0000	293.1791	293.4000	293.2338	293.1201	293.0101	293.0106	293.0120						
16	292.1000	292.4059	292.7066	292.9893	293.2285	293.3913	293.4620	293.4164	293.2820	293.0000	293.1240	293.1685						
15	292.1000	292.4745	292.8510	293.2182	293.5277	293.6664	293.7068	293.6523	293.5701	293.4127	293.2920	293.2051						
14	292.1000	292.5407	293.0027	293.5015	293.9912	294.0301	294.0137	293.9000	293.9121	293.7621	293.4000	293.2309						
13	292.1000	292.5841	293.1153	293.7695	294.9000	294.4395	294.4049	294.3564	294.3960	294.3000	293.5706	293.2619						
12	292.1000	292.5780	293.0821	293.6364	294.2126	294.4150	294.8000	294.7025	295.0000	294.4038	293.3000	293.2116						
11	292.1000	292.5431	292.9955	293.4575	293.8931	294.2008	294.4509	294.6431	295.2000	295.0000	293.8000	293.2757						
10	292.1000	292.4966	292.8972	293.3018	293.6969	294.0384	294.1521	294.2098	294.2000	293.9881	293.2000	293.0827						
9	292.1000	292.4447	292.7934	293.1538	293.5514	294.1000	293.9037	293.8375	293.7469	293.5457	293.1952	292.9836						
8	292.1000	292.3687	292.6779	292.9673	293.2525	293.5040	293.5219	293.4857	293.4003	293.2491	293.0489	292.8701						
7	292.1000	292.3330	292.5633	292.7851	292.9863	293.1394	293.1922	293.1811	293.1178	293.0000	292.8803	292.7466						
6	292.1000	292.2812	292.4584	292.6247	292.7690	292.8751	292.9258	292.9278	292.8885	292.8151	292.7257	292.6252						
5	292.1000	292.2352	292.3666	292.4881	292.5915	292.6676	292.7088	292.7160	292.6931	292.6458	292.5825	292.5090						
4	292.1000	292.1951	292.2873	292.3720	292.4416	292.4968	292.5276	292.5357	292.5229	292.4929	292.4503	292.3992						
3	292.1000	292.1600	292.2183	292.2718	292.3170	292.3509	292.3714	292.3778	292.3713	292.3537	292.3275	292.2954						
2	292.1000	292.1288	292.1569	292.1827	292.2046	292.2211	292.2313	292.2349	292.2322	292.2241	292.2117	292.1963						
1	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000						

FILE NAME -- TEMPERATURE		TEMPERATURE FIELD FOR ST. LOUIS DAY 195					VERT. LEVEL 1 START DATE/TIME 76195 5.00 STOP DATE/TIME 76195 6.00		
TEMPERATURE VALUES (IN DEGREES KELVIN AT GROUND LEVEL)		13	14	15	16	17			
GRID CELL									
22	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000			
21	292.3098	292.2851	292.2458	292.1874	292.1000	292.1000			
20	292.4899	292.4375	292.3584	292.2488	292.1000	292.1000			
19	292.6503	292.5670	292.4482	292.2942	292.1000	292.1000			
18	292.8037	292.6838	292.5207	292.3274	292.1000	292.1000			
17	292.9857	292.7933	292.5759	292.3480	292.1000	292.1000			
16	293.3000	292.8876	292.5972	292.3500	292.1000	292.1000			
15	293.0997	292.8293	292.5434	292.3241	292.1000	292.1000			
14	293.0325	292.7598	292.4000	292.2829	292.1000	292.1000			
13	293.0157	292.7565	292.4927	292.2955	292.1000	292.1000			
12	292.9955	292.7442	292.5077	292.2993	292.1000	292.1000			
11	293.0000	292.7093	292.4890	292.2909	292.1000	292.1000			
10	292.8676	292.6000	292.4457	292.2739	292.1000	292.1000			
9	292.7846	292.5855	292.4193	292.2589	292.1000	292.1000			
8	292.7009	292.5377	292.3870	292.2424	292.1000	292.1000			
7	292.6112	292.4777	292.3489	292.2237	292.1000	292.1000			
6	292.5197	292.4131	292.3076	292.2034	292.1000	292.1000			
5	292.4297	292.3478	292.2651	292.1825	292.1000	292.1000			
4	292.3428	292.2834	292.2227	292.1615	292.1000	292.1000			
3	292.2593	292.2208	292.1811	292.1407	292.1000	292.1000			
2	292.1787	292.1599	292.1402	292.1202	292.1000	292.1000			
1	292.1000	292.1000	292.1000	292.1000	292.1000	292.1000			

## J. TERRAIN

The TERRAIN file contains time-invariant matrices of surface roughness and vegetation factors that are used by the Airshed Simulation Program to calculate vertical diffusivity and surface deposition. If the TERRAIN file is omitted from a simulation run, single values for surface roughness and vegetation factors will be used (see Section IX.B). The program TERRAIN is used to create the TERRAIN file. TERRAIN requires subroutines from the libraries UTILITY and FILUTIL. Figure VII-19 is an information flow diagram illustrating the job control to be used. In the following sections we discuss the input to the TERRAIN program.

### 1. Variables

The TERRAIN program has two implicit output variables: ROUGHNESS and VEGFACTOR. The internal units for ROUGHNESS are meters (m); VEGFACTOR is the fraction of surface uptake capability compared to that of alfalfa. If the input values for ROUGHNESS are to be in any other units, that variable must be specified in the UNITS packet. VEGFACTOR is unitless; therefore, its units cannot be changed.

### 2. Methods

The methods that can be used to generate the TERRAIN file are:

- > CONSTANT
- > GRID VALUE
- > STATINTERP
- > POISSON
- > E-WINTERP
- > N-SINTERP
- > USER.

These methods are discussed in detail in Section VI.A.6.

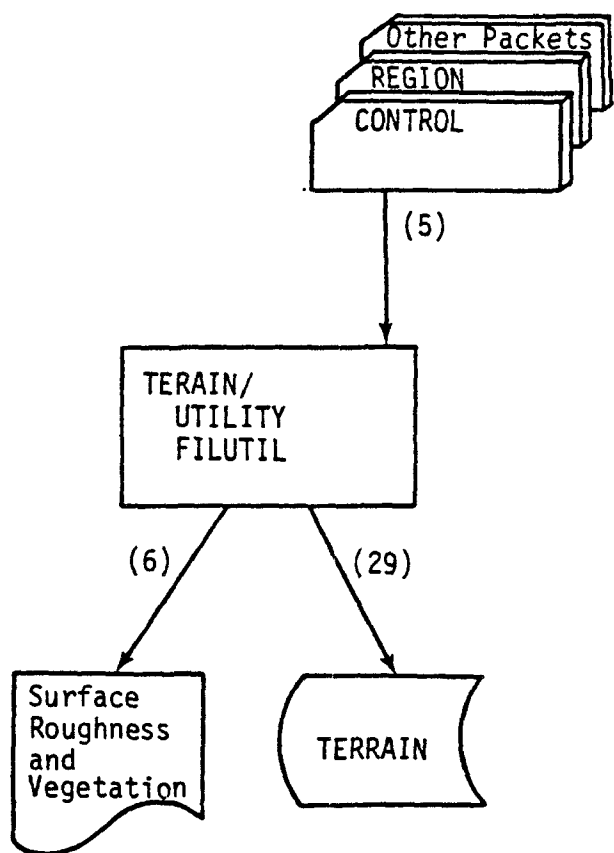


Figure VII-19. Information Flow Diagram for the TERRAIN File

### 3. Time Variation

The TERRAIN file contains values for surface characteristics that are considered to be invariant over time.

### 4. Packets

Figure VII-20 shows the input deck structure in terms of the packets to be used. Each of these packets is described in detail in Section VI.B. Following are special input packet considerations for the TERRAIN file:

- > CONTROL
  - The file name on Card 2 must be TERRAIN.
  - The control variables to be specified on Cards 4 to 8 for TERRAIN are shown in Table VII-8.
  - The number of species should be zero.
  - If there are input variables that do not appear as output variables, their number must appear as the number of user-defined variables. All such variables must also be named in the UNITS packet.
  - If data from measuring stations are to be used (methods STATINTERP or POISSON), the maximum number of such stations must be given.
  - The number of subregions must be at least one.
  - The maximum number of parameters must be sufficient to include all specifications of all parameters.
  - The vertical controls card (No. 7) should be left blank.
  - The file unit assignment card (No. 8) should be left blank.
  - The beginning and ending dates and times can be any valid numbers. They are ignored by the Airshed Simulation Program.
  - A set of output species names is not required; if they are present, their number must be the same as the entry in the first control parameter on Card 4, but they will be ignored by the program.

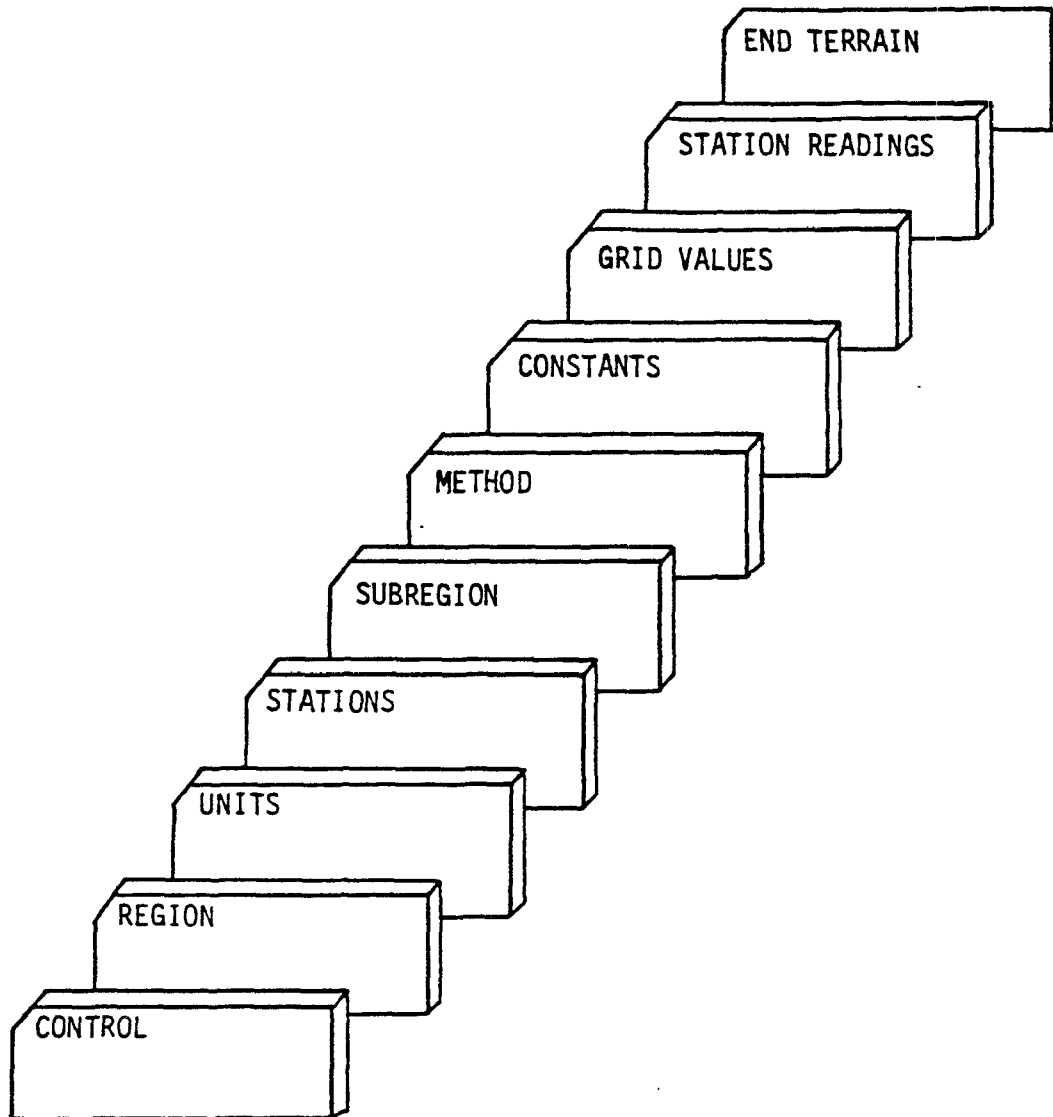


Figure VII-20. Input Card Deck Structure for Preparing the TERRAIN File

Table VII-8. CONTROL Packet Entries for the TERRAIN File

<u>Card Number</u>	<u>Entry</u>
4	Number of species (=0) Number of user-defined variables Number of stations Number of subregions Number of parameters Spare
5	Output file number Print input cards Print output grid Spare Spare Spare
6	Print units table Print station locations table Print regional grid Print methods table Print station values table Spare
7	Spare Spare Spare Spare Spare Spare
8	Spare Spare Spare Spare Spare Spare

- > REGION. This packet must follow the CONTROL packet. The vertical parameters will be ignored for the TERRAIN file.
- > UNITS. This packet, if present, must follow the REGION packet. The UNITS packet must be provided if:
  - Any input variable will be input in other than internal units.
  - Any user-defined variables are specified.
  - COORD or HEIGHT unit conversions are to be used.The number of user-defined variables must not exceed the maximum specified in the CONTROL packet.
- > STATIONS. This packet is required if either method STATINTERP or POISSON is specified. The number of stations listed must not exceed the maximum specified in the CONTROL packet.
- > SUBREGION. A SUBREGION packet is required. The number of subregions must not exceed the maximum specified in the CONTROL packet.
- > METHOD. A method must be provided for every variable--including user-defined variables--in every subregion. Note that each parameter entry contributes to the overall parameter count; the total number of parameters must not exceed the maximum specified in the CONTROL packet.
- > CONSTANTS. If the method CONSTANT is assigned to any variable in the METHOD packet, there must be one or more CONSTANTS packets.
- > GRID VALUES. If the method GRID VALUE is assigned to any variable in the METHOD packet, there must be one or more GRID VALUES packets.
- > STATION READINGS. If either the POISSON or STATINTERP method is assigned to any variable in the METHOD packet, there must be one or more STATION READINGS packets.
- > END TERRAIN. This card must appear to indicate that no more packets are to be read.

5. Additional Input Files

No additional input files are required.

6. Examples

Exhibit VII-15 shows a sample input deck for TERRAIN file generation. The output from a TERRAIN run using that input deck is shown in Exhibit VII-16.

### **Exhibit VII-15. Sample Input Deck from the TERRAIN Program**

Exhibit VII-15 (Concluded)

6 .21  
ROUGNESS 9 .19  
ROUGNESS 10 2 .29  
ROUGNESS 11 2 .29  
ROUGNESS 12 2 .28  
ROUGNESS 13 2 .28  
ROUGNESS 14 2 .27  
VEGFACTOR 2 21 1.00  
VEGFACTOR 3 21 1.00  
VEGFACTOR 4 21 1.00  
VEGFACTOR 5 21 1.00  
VEGFACTOR 6 21 1.00  
VEGFACTOR 7 21 1.00  
VEGFACTOR 8 21 1.00  
VEGFACTOR 9 21 1.00  
VEGFACTOR 10 21 1.00  
VEGFACTOR 11 21 1.00  
VEGFACTOR 12 21 1.00  
VEGFACTOR 13 21 1.00  
VEGFACTOR 14 21 1.00  
VEGFACTOR 15 21 1.00  
VEGFACTOR 16 21 1.00  
VEGFACTOR 17 21 1.00  
VEGFACTOR 1 22 1.00  
VEGFACTOR 2 22 1.00  
VEGFACTOR 3 22 1.00  
VEGFACTOR 4 22 1.00  
VEGFACTOR 5 22 1.00  
VEGFACTOR 6 22 1.00  
VEGFACTOR 7 22 1.00  
VEGFACTOR 8 22 1.00  
VEGFACTOR 9 22 1.00  
VEGFACTOR 10 22 1.00  
VEGFACTOR 11 22 1.00  
VEGFACTOR 12 22 1.00  
VEGFACTOR 13 22 1.00  
VEGFACTOR 14 22 1.00  
VEGFACTOR 15 22 1.00  
VEGFACTOR 16 22 1.00  
VEGFACTOR 17 22 1.00

END  
END TERRAIN

PROCESSING OF INPUT DATA FOR THE PREPARATION OF FILE TERRAIN  
 SURFACE ROUGHNESS AND VEGETATIVE FACTORS

```

REGION          725000.  425000.      15
                -19000.  -16000.
                4000.00  4000.00
                17       22       4     .00   50.00  200.00
                2       2       .00
END
KCRNT = 41   NSCLFT = 26960 IN SCRSET
KCRNT = 61   NSCLFT = 26920 IN SCRSET
KCRNT = 85   NSCLFT = 26916 IN SCRSET
KCRNT = 89   NSCLFT = 26912 IN SCRSET
KCRNT = 93   NSCLFT = 26908 IN SCRSET
KCRNT = 103  NSCLFT = 26898 IN SCRSET
KCRNT = 477  NSCLFT = 26896 IN SCRSET
KCRNT = 479  NSCLFT = 26524 IN SCRSET
KCRNT = 481  NSCLFT = 26522 IN SCRSET
KCRNT = 483  NSCLFT = 26520 IN SCRSET
KCRNT = 485  NSCLFT = 26518 IN SCRSET
KCRNT = 487  NSCLFT = 26516 IN SCRSET
KCRNT = 861  NSCLFT = 26514 IN SCRSET
KCRNT = 1235 NSCLFT = 26626 IN SCRSET
KCRNT = 1983 NSCLFT = 26252 IN SCRSET
SUBREGION
METHOD

```

SUBREGION	VARIABLE	METHOD	MIN. VALUE	MAX. VALUE	PARAMETER	PAR. VALUE
A	ROUGHNESS	GRIDVALUE	.0000	100.0000		
A	VEGFACTOR	GRIDVALUE	.0000	100.0000		
GRIDVALUES						
ENDTERRAIN						

Exhibit VII-16. Sample Output from the TERRAIN Program

## FILE NAME -- TERRAIN

## SURFACE ROUGHNESS AND VEGETATIVE FACTORS

PAGE 1

GRID CELL	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22
	TERRAIN VALUES (IN METERS OR ALFALFA EQUIVALENTS) ROUGHNESS VERT.										LEVEL 1 START DATE/TIME		0 .00 STOP DATE/TIME		0 .00							
	.2300	.2300	.2300	.2300	.2300	.2300	.2300	.2300	.2900	.2600	.2900	.2800	.2500	.2500	.2800	.2800	.2800	.2800	.2800	.2800	.2800	.2800
21	.2300	.2300	.2300	.2300	.2300	.2300	.2300	.2300	.2900	.2600	.2900	.2800	.2500	.2500	.2800	.2800	.2800	.2800	.2800	.2800	.2800	.2800
20	.2300	.2300	.2300	.2300	.2300	.2300	.2300	.2300	.2900	.2600	.2900	.2800	.2500	.2500	.2800	.2800	.2800	.2800	.2800	.2800	.2800	.2800
19	.2200	.2200	.2200	.2200	.2200	.2200	.2200	.2200	.2800	.2500	.2700	.2400	.2100	.2100	.2400	.2400	.2400	.2400	.2400	.2400	.2400	.2400
18	.2100	.2100	.2100	.2100	.2100	.2100	.2100	.2100	.2600	.2300	.2700	.2400	.2100	.2100	.2400	.2400	.2400	.2400	.2400	.2400	.2400	.2400
17	.2200	.2200	.2200	.2200	.2200	.2200	.2200	.2200	.2600	.2300	.2700	.2400	.2100	.2100	.2400	.2400	.2400	.2400	.2400	.2400	.2400	.2400
16	.2900	.2900	.2900	.2900	.2900	.2900	.2900	.2900	.3800	.5200	.4400	.3600	.2900	.2900	.3800	.3600	.3600	.3600	.3600	.3600	.3600	.3600
15	.4400	.4400	.4400	.4400	.4400	.4400	.4400	.4400	.5900	.5900	.6700	.5700	.5100	.5100	.6700	.5700	.5700	.5700	.5700	.5700	.5700	.5700
14	.6400	.6400	.6400	.6400	.6400	.6400	.6400	.6400	.7800	.7400	.6600	.6900	.2300	.2300	.7800	.6600	.6600	.6600	.6600	.6600	.6600	.6600
13	.6700	.6700	.6700	.6700	.6700	.6700	.6700	.6700	.8000	.7400	.9400	.1.5500	.4200	.4200	.8000	.9400	.9400	.9400	.9400	.9400	.9400	.9400
12	.5900	.5900	.5900	.5900	.5900	.5900	.5900	.5900	.6300	.6600	2.0100	3.1700	1.7800	1.7800	.6300	2.0100	2.0100	2.0100	2.0100	2.0100	2.0100	2.0100
11	.5800	.5800	.5800	.5800	.5800	.5800	.5800	.5800	.5800	.6700	1.2000	2.2000	1.2000	1.2000	.5800	1.2000	1.2000	1.2000	1.2000	1.2000	1.2000	1.2000
10	.6200	.6200	.6200	.6200	.6200	.6200	.6200	.6200	.6600	.6400	1.8600	2.2400	.5400	.5400	.6600	1.8600	1.8600	1.8600	1.8600	1.8600	1.8600	1.8600
9	.5400	.5400	.5400	.5400	.5400	.5400	.5400	.5400	.5300	.5200	.5600	.5500	.3800	.3800	.5300	.5600	.5600	.5600	.5600	.5600	.5600	.5600
8	.4700	.4700	.4700	.4700	.4700	.4700	.4700	.4700	.5400	.5400	.5600	.4500	.2300	.4100	.5400	.5600	.5600	.5600	.5600	.5600	.5600	.5600
7	.4700	.4700	.4700	.4700	.4700	.4700	.4700	.4700	.4900	.4800	.3700	.2300	.2800	.3400	.4700	.4900	.4900	.4900	.4900	.4900	.4900	.4900
6	.4900	.4900	.4900	.4900	.4900	.4900	.4900	.4900	.3800	.3300	.2300	.2100	.3400	.3400	.3800	.3800	.3800	.3800	.3800	.3800	.3800	.3800
5	.4800	.4800	.4800	.4800	.4800	.4800	.4800	.4800	.3600	.3100	.2100	.1900	.2900	.2900	.3600	.3600	.3600	.3600	.3600	.3600	.3600	.3600
4	.4800	.4800	.4800	.4800	.4800	.4800	.4800	.4800	.3600	.3100	.2100	.1900	.2900	.2900	.3600	.3600	.3600	.3600	.3600	.3600	.3600	.3600
3	.4800	.4800	.4800	.4800	.4800	.4800	.4800	.4800	.3600	.3100	.2100	.1900	.2900	.2900	.3600	.3600	.3600	.3600	.3600	.3600	.3600	.3600
2	.4800	.4800	.4800	.4800	.4800	.4800	.4800	.4800	.3600	.3100	.2100	.1900	.2900	.2900	.3600	.3600	.3600	.3600	.3600	.3600	.3600	.3600
1	.4800	.4800	.4800	.4800	.4800	.4800	.4800	.4800	.3600	.3100	.2100	.1900	.2900	.2900	.3600	.3600	.3600	.3600	.3600	.3600	.3600	.3600

## FILE NAME -- TERRAIN

## SURFACE ROUGHNESS AND VEGETATIVE FACTORS

GRID CELL	1	2	3	4	5	6	7	8	9	10	11	12	0 .00 STOP DATE/TIME	0 .00
22	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
21	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
20	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
19	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
18	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
17	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
16	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
15	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
14	1.0000	1.0000	1.0000	1.0000	1.0000	.5000	.5000	.5000	.5000	.5000	.5000	.5000	.5000	1 .00
13	1.0000	1.0000	1.0000	1.0000	1.0000	.5000	.5000	.5000	.5000	.5000	.5000	.5000	.5000	1 .00
12	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	.5000	.5000	.5000	.5000	.5000	.5000	1 .00
11	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	.5000	.5000	.5000	.5000	.5000	.5000	1 .00
10	.5000	.5000	.5000	.5000	.5000	.5000	.5000	.5000	.5000	.5000	.5000	.5000	.5000	1 .00
9	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
8	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
7	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
6	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
5	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
4	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
3	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
2	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00
1	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000	0 .00

## K. TOPCONC

The TOPCONC file contains time-varying matrices of pollutant concentrations at the top of the region. The program TPCONC is used to create the TOPCONC file. TPCONC requires subroutines from the libraries UTILITY and FILUTIL. Figure VII-21 is an information flow diagram illustrating the job control to be used. The following sections discuss the input to the TPCONC program.

### 1. Variables

The output variables for the TOPCONC file are the species named in the CONTROL packet. Additional user-defined input variables (e.g. "reactive hydrocarbons") can be specified in the UNITS packet. The internal units for the concentrations of all species except AEROSOLS are parts per million (ppm); for AEROSOLS, the units are micrograms per cubic meter ( $\mu\text{g}/\text{m}^3$ ). The standard names for reactive species recognized by the Airshed Simulation Program are listed in Section IX.A. If any of these species does not appear on the TOPCONC file, the top concentrations will default to a value defined in the CHEMPARAM file. If any additional species appear on TOPCONC, they will be ignored.

### 2. Methods

The methods that can be used to generate the TOPCONC file are:

- > CONSTANT
- > GRID VALUE
- > STATINTERP
- > POISSON
- > SPLIT/COMB
- > ABSTOPCONC
- > RELTOPCONC

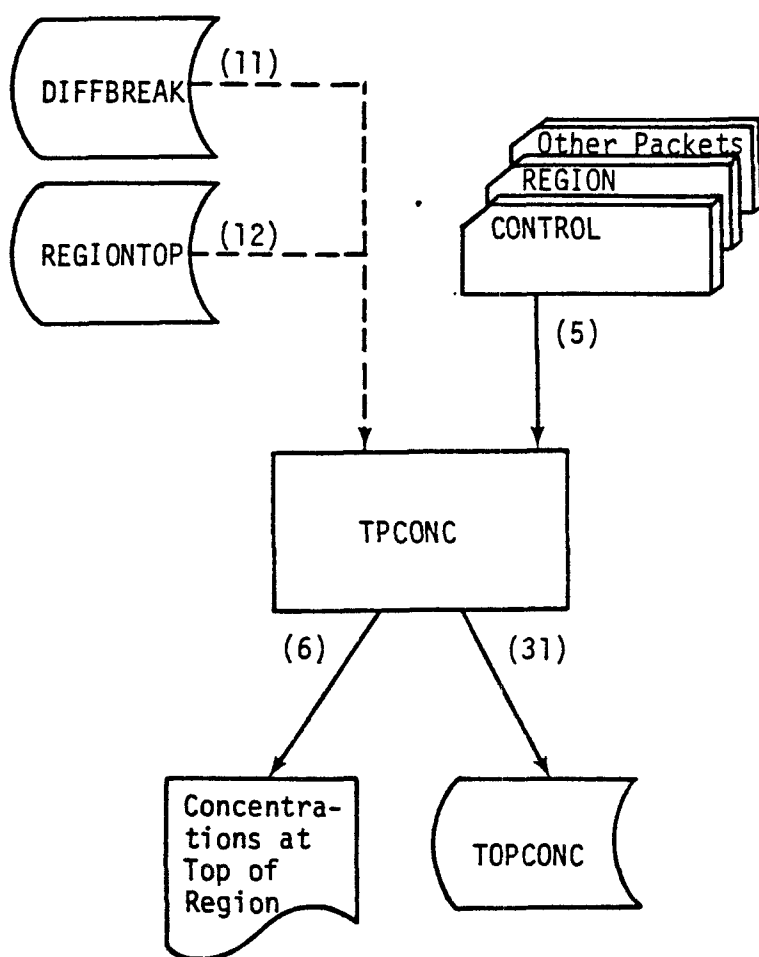


Figure VII-21. Information Flow Diagram for the TOPCONC File

- > E-WINTERP
- > N-SINTERP
- > USER.

These methods are discussed in detail in Section VI.A.6.

### 3. Time Variation

The time span of the TOPCONC file must include the entire time span of the simulation runs for which it is to be used. Concentrations at the top of the region are considered to be constant during each time interval.

### 4. Packets

Figure VII-22 shows the input deck structure in terms of the packets to be used. Each of these packets is described in detail in Section VI.B. Following are special input packet considerations for the TOPCONC file.

- > CONTROL
  - The file name on Card 2 must be TOPCONC.
  - The control variables to be specified on Cards 4 to 8 for TOPCONC are shown in Table VII-9.
  - The number of species must be greater than zero.
  - If there are input variables that do not appear as output variables, their number must appear as the number of user-defined variables. All such variables must also be named in the UNITS packet.
  - If data from measuring stations are to be used (methods STATINTERP or POISSON), the maximum number of such stations must be given.
  - The number of subregions must be at least one.
  - The maximum number of parameters must be sufficient to include all specification of all parameters.

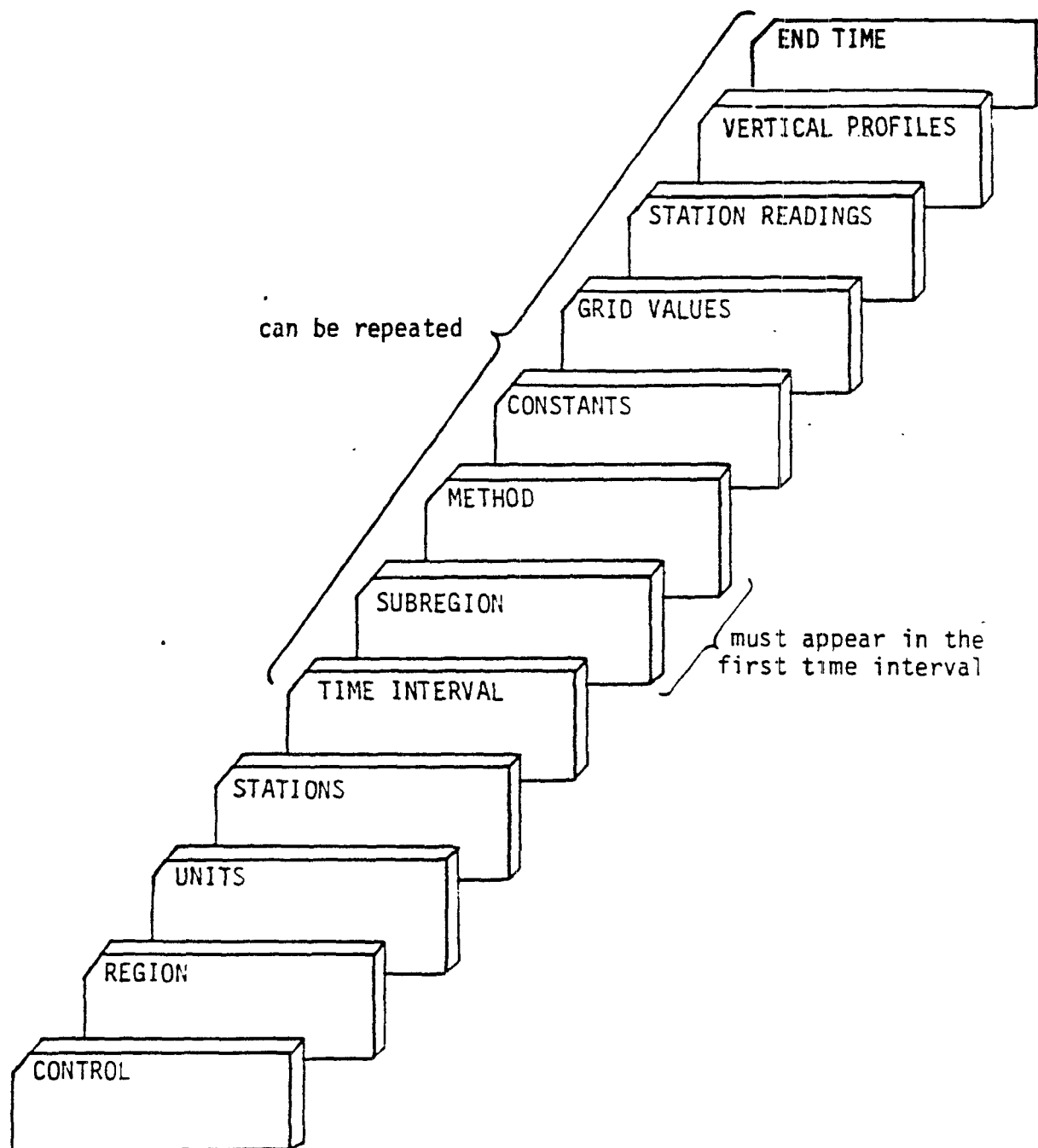


Figure VII-22. Input Card Deck Structure for Preparing the TOPCONC File

Table VII-9. CONTROL Packet Entries for the TOPCONC File

<u>Card Number</u>	<u>Entry</u>
4	Number of species Number of user-defined variables Number of stations Number of subregions Number of parameters Spare
5	Output file number Print input cards Print output grid Spare Spare Spare
6	Print units table Print station locations table Print regional grid Print methods table Print station values table Spare
7	Spare Number of heights in profile Spare Print vertical profile tables Spare Spare
8	DIFFBREAK file number REGIONTOP file number Spare Spare Spare Spare

- The vertical controls card (No. 7) must contain the maximum number of profile heights if ABSTOPCONC or RELTOPCONC is used. Otherwise, this card should be blank.
  - The file unit assignment card (No. 8) must provide an entry for REGIONTOP if the method ABSTOPCONC or RELTOPCONC is selected. It must also provide an entry for DIFFBREAK if RELTOPCONC is selected.
  - The beginning and ending dates and times should reflect the time variation considerations discussed in Section 3 above.
  - A set of output species names is required; their number must be the same as the entry in the first control parameter on Card 4.
- > REGION. This packet must follow the CONTROL packet. The vertical parameters will be ignored for the TOPCONC file.
- > UNITS. This packet, if present, must follow the REGION packet. The UNITS packet must be provided if:
- Any input variable will be input in other than internal units.
  - Any user-defined variables are specified.
  - COORD or HEIGHT unit conversions are to be used.
- The number of user-defined variables must not exceed the maximum specified in the CONTROL packet.
- > STATIONS. This packet is required if the method STATINTERP or POISSON is specified. The number of stations listed must not exceed the maximum specified in the CONTROL packet.
- > TIME INTERVAL. One or more TIME INTERVAL packets must be present. The first time interval must begin at or before the beginning of the time span specified on Card 10 of the CONTROL packet. All time intervals must be contiguous and of nonzero length. Each TIME INTERVAL packet contains one or more of the following packets and ends with an ENDTIME card. Following the first time interval, only those data that are to be changed need be specified.

- > SUBREGION. The first time interval must contain a SUBREGION packet; the inclusion of this packet in other time intervals is optional. The number of subregions must not exceed the maximum specified in the CONTROL packet.
- > METHOD. A method must be provided for every variable--including user-defined variables--in every subregion in the first time interval. Methods can be changed in subsequent TIME INTERVAL packets if desired. Note that each parameter entry contributes to the overall parameter count; the total number of parameters must not exceed the maximum specified in the CONTROL packet.
- > CONSTANTS. If the method CONSTANT is assigned to any variable in the METHOD packet, the first time interval must contain a CONSTANTS packet. More than one CONSTANTS packet can appear in any time interval.
- > GRID VALUES. If the method GRID VALUE is assigned to any variable in the METHOD packet, the first time interval must contain a GRID VALUES packet. More than one GRID VALUES packet can appear in any time interval.
- > STATION READINGS. If either the POISSON or STATINTERP method is assigned to any species in the METHOD packet, the first time interval must contain a STATION READINGS packet. More than one STATION READINGS packet can appear in any time interval.
- > VERTICAL PROFILES. If the method ABSTOPCONC or RELTOPCONC is assigned to any species in the METHOD packet, the first time interval must contain a VERTICAL PROFILES packet. There must be a vertical profile defined (or implied by means of ALL) for every variable in every subregion for which the profile method was specified. The number of height-value pairs in any single profile must not exceed the maximum specified in the CONTROL packet. More than one VERTICAL PROFILES packet can appear in any time interval.

5. Additional Input Files

If the method ABSTOPCONC or RELTOPCONC was selected, the REGIONTOP file must be input to TPCONC. In addition, if the method RELTOPCONC was selected, the DIFFBREAK file must also be input.

6. Examples

Exhibit VII-17 shows a sample input deck for TOPCONC file generation. The output from a TPCONC run using that input deck is shown in Exhibit VII-18.

```

CONTROL
TOPCONC POLLUTANT CONCENTRATIONS ABOVE THE MODELING REGION
    NO      0      0      0      0      0
    NO2     0      0      0      0      0
    O3      1      1      1      1      0
    ETH     1      1      1      1      0
    OLE     1      1      1      1      0
    PAR     1      1      1      1      0
    CARB    1      1      1      1      0
    ARO     0      0      0      0      0
    BZA     0      0      0      0      0
    PAN     0      0      0      0      0
    CO      0      0      0      0      0
END
REGION
    725000. 425000. 15
    -19000. -16000.
    4000. 4000.
    17      22      4
    2       2       0.
    2       2       50.
    2       2       200.
END
TIME INTERVAL
    76195 000 76195 2400
SUBREGION
    A      1      1      -1
END
METHOD
    A      NO      CONSTANT .0001   .01
    A      NO2     CONSTANT .0001   .01
    A      O3      CONSTANT .001    .2
    A      ETH     CONSTANT .0001   .01
    A      OLE     CONSTANT .0001   .01
    A      PAR     CONSTANT .0.01   1.
    A      CARB    CONSTANT .0001   0.1
    A      ARO     CONSTANT .0001   0.1
    A      BZA     CONSTANT .000000 .0001
    A      PAN     CONSTANT .000001 .0001
    A      CO      CONSTANT .01     1.
END

```

**Exhibit VII-17 (Concluded)**

**CONSTANTS**

A	ND	.001
A	NO2	.002
A	O3	.078
A	ETH	.00109
A	OLE	.00220
A	PAR	.03333
A	CARB	.00405
A	ARO	.00124
A	BZA	
A	PAN	.00001
A	CO	0.1

**END**

**ENDTIME**

PROCESSING OF INPUT DATA FOR THE PREPARATION OF FILE TOPCONC  
POLLUTANT CONCENTRATIONS ABOVE THE MODELING REGION

NUMBER OF USER-DEFINED VARIABLES = 0  
NUMBER OF STATIONS = 0  
NUMBER OF SUBREGIONS = 1  
NUMBER OF METHOD PARAMETERS = 0  
NUMBER OF VERTICAL PROFILE HEIGHTS = 0  
PROGRAM OUTPUT WILL BE PLACED ON FILE NUMBER 31  
INPUT CARDS WILL BE PRINTED  
OUTPUT GRID WILL BE PRINTED  
UNIT TABLE WILL BE PRINTED  
STATION LOCATIONS WILL BE PRINTED  
SUBREGION GRID WILL BE PRINTED  
METHOD TABLE WILL BE PRINTED  
STATION VALUES WILL NOT BE PRINTED  
VERTICAL PROFILE VALUES WILL NOT BE PRINTED  
BEGINNING DATE OF FILE 76195 BEGINNING TIME OF FILE .00  
ENDING DATE OF FILE 76195 ENDING TIME OF FILE 24.00  
A TOTAL OF 11 SPECIES NAMES WERE READ, THE SPECIES NAMES ARE --  
NO  
NO2  
O3  
ETH  
OLE  
PAR  
CARB  
ARO  
BZA  
PAN  
CO  
REGION  
725000. 4252000. 15  
-19000. -16000.  
4000.76 4000.00  
17 22 4  
2 2 .00 50.00 200.00  
END

## FILE NAME -- TOPCONC POLLUTANT CONCENTRATIONS ABOVE THE MODELING REGION

GRID CELL	SPECIES NO												VERT. LEVEL 1	START DATE/TIME 76195 0.00 STOP DATE/TIME 76195 24.00
	1	2	3	4	5	6	7	8	9	10	11	12		
22	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
21	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
20	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
19	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
18	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
17	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
16	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
15	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
14	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
13	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
12	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
11	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
10	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
9	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
8	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
7	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
6	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
5	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
4	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
3	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
2	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
1	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010

TOP CONCENTRATIONS (IN PARTS PER MILLION) SPECIES NO  
VERT. LEVEL 1 START DATE/TIME 76198 ••.00 STOP DATE/TIME 76198 24.00

		POLLUTANT CONCENTRATIONS ABOVE THE MODELING REGION									
GRID CELL	13	14	15	16	17	18	19	20	21	22	23
22	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
21	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
20	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
19	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
18	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
17	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
16	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
15	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
14	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
13	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
12	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
11	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
10	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
9	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
8	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
7	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
6	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
5	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
4	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
3	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
2	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010
1	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010	.0010

## L. WIND

The WIND file contains time-varying matrices of horizontal wind vector values in three dimensions and scalars representing overall maxima and average values at the boundaries. The program WNDSET is used to create the WIND file. WNDSET requires subroutines from the libraries UTILITY and FILUTIL. Figure VII-23 is an information flow diagram illustrating the job control to be used. In the following sections we discuss the input to the WNDSET program.

### 1. Variables

The WNDSET program has four implicit variables: SPEED, DIRECTION, WINDX, and WINDY. The first two represent a wind vector in polar form, and the latter two represent the vector in terms of its x- and y-components. A three-dimensional matrix is maintained for each variable. In general, the input to the program will be SPEED and DIRECTION, whereas the variables written to the WIND file are WINDX and WINDY. The WNDSET program calculates values for the undefined variables from those for variables to which values have been assigned.

### 2. Methods

The methods that can be used to generate the WIND file are:

- > CONSTANT
- > GRID VALUE
- > STATINTERP
- > POISSON
- > DIVERGENCE
- > VECTORIZE
- > POLARIZE

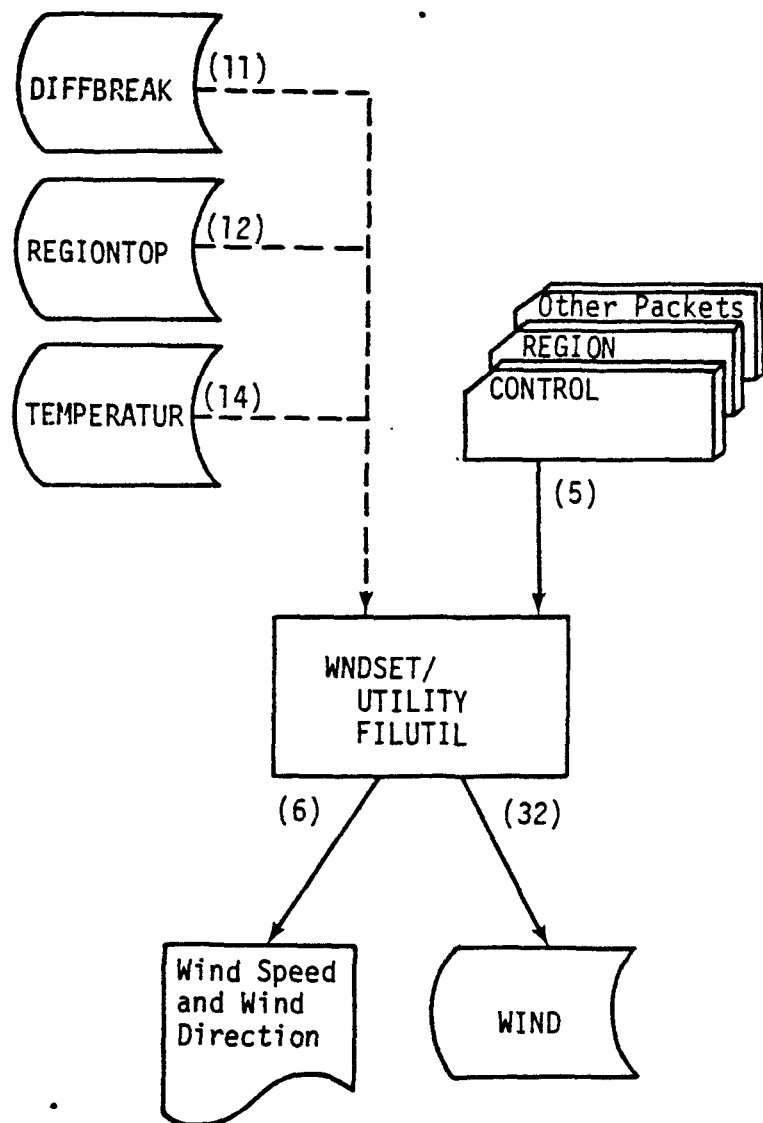


Figure VII-23. Information Flow Diagram for the WIND File

- > E-WINTERP
- > N-SINTERP
- > USER.

These methods are discussed in detail in Section VI.A.6.

Since the WIND file varies in the vertical direction, a vertical method must also be specified for each output variable in each subregion. The vertical methods that can be used are:

- > CONSTANT
- > ABSPROFILE
- > RELPROFILE
- > DIVERGENCE
- > VECTORIZE
- > POLARIZE
- > E-WINTERP
- > N-SINTERP
- > VERTUSER.

These vertical methods are discussed in detail in Section VI.A.6.

### 3. Time Variation

The time span of the WIND file must include the entire time span of the simulation runs for which it is to be used. Wind vectors and scalar values are considered to be constant during each time interval.

### 4. Packets

Figure VII-24 shows the input deck structure in terms of the packets to be used. Each of these packets is described in detail in Section VI.B. Following are special input packet considerations for the WIND file:

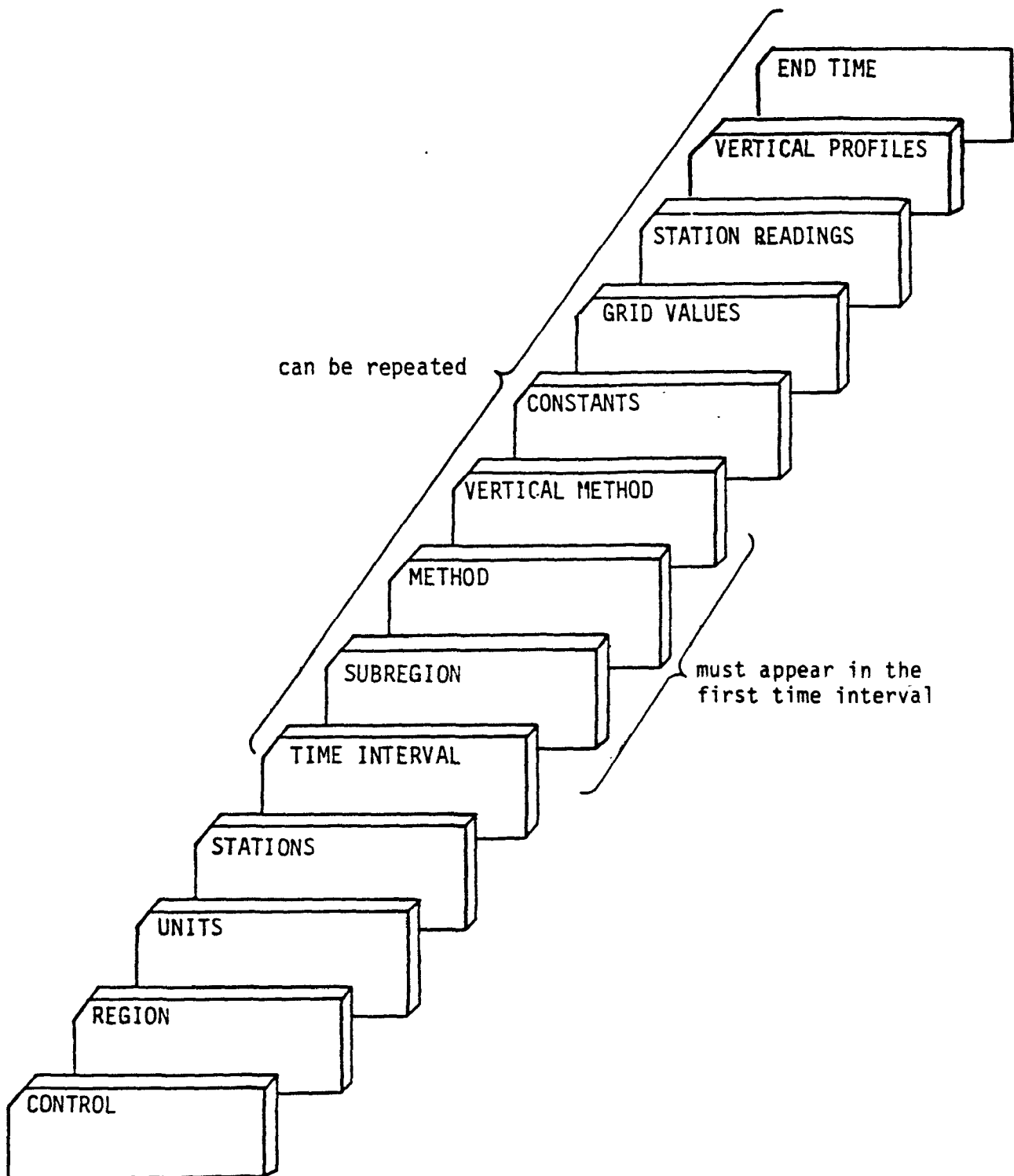


Figure VII-24. Input Card Deck Structure for Preparing the WIND File

> CONTROL

- The file name on Card 2 must be WIND.
- The control variables to be specified on Cards 4 to 8 for WIND are shown in Table VII-10.
- The number of species should be zero.
- If there are input variables other than the four implicit variables, their number must appear as the number of user-defined variables. All such variables must also be named in the UNITS packet.
- If data from measuring stations are to be used (methods STATINTERP, POISSON, or DIVERGENCE), the maximum number of such stations must be given.
- The number of subregions must be at least one.
- The maximum number of parameters must be sufficient to include all specifications of all parameters.
- The vertical controls card (No. 7) must include maximum vertical parameter and profile entries as applicable.
- The file unit assignment card (No. 8) must provide entries for DIFFBREAK and REGIONTOP if any vertical method besides CONSTANT is selected. In addition, entries for TEMPERATUR must appear if DIVERGENCE is used.
- The beginning and ending dates and times should reflect the time variation considerations discussed in Section 3 above.
- A set of output species names is not required; if they are present, their number must be the same as the entry in the first control parameter on Card 4, but they will be ignored by the program.

> REGION. This packet must follow the CONTROL packet. The vertical parameters must be provided for the WIND file.

> UNITS. This packet, if present, must follow the REGION packet. The UNITS packet must be provided if:

- Any input variable will be input in other than internal units.

Table VII-10. CONTROL Packet Entries for the WIND File

<u>Card Number</u>	<u>Entry</u>
4	Number of species (=0) Number of user-defined variables Number of stations Number of subregions Number of parameters Spare
5	Output file number Print input cards Print grids for x- and y-components Print grids for speed and direction Spare Spare
6	Print units table Print station locations table Print regional grid Print methods table Print station values table Spare
7	Number of vertical parameters Number of heights in profile Print vertical methods table Print vertical profile tables Spare Spare
8	DIFFBREAK file number REGIONTOP file number Spare TEMPERATUR file number Spare Spare

- Any user-defined variables are specified.
  - COORD or HEIGHT unit conversions are to be used.
- The number of user-defined variables must not exceed the maximum specified in the CONTROL packet.
- > STATIONS. This packet is required if any of the methods STATINTERP, POISSON, or DIVERGENCE is specified. The number of stations listed must not exceed the maximum specified in the CONTROL packet.
  - > TIME INTERVAL. One or more TIME INTERVAL packets must be present. The first time interval must begin at or before the beginning of the time span specified on Card 10 of the CONTROL packet. All time intervals must be contiguous and of nonzero length. Each TIME INTERVAL packet contains one or more of the following packets and ends with an ENDTIME card. Following the first time interval, only those data that are to be changed need be specified.
  - > SUBREGION. The first time interval must contain a SUBREGION packet; the inclusion of this packet in other time intervals is optional. The number of subregions must not exceed the maximum specified in the CONTROL packet.
  - > METHOD. A method must be provided for every variable--including user-defined variables--in every subregion in the first time interval. Methods can be changed in subsequent TIME INTERVAL packets if desired. Note that each parameter entry contributes to the overall parameter count; the total number of parameters must not exceed the maximum specified in the CONTROL packet.
  - > VERTICAL METHOD. A vertical method must be provided for every variable--including user-defined variables--in every subregion in the first time interval. Vertical methods can be changed in subsequent TIME INTERVAL packets if desired. Note that each vertical parameter entry contributes to the overall vertical parameter count; the total must not exceed the maximum specified in the CONTROL packet.

- > CONSTANTS. If the method CONSTANT is assigned to any variable in the METHOD packet, the first time interval must contain a CONSTANTS packet. More than one CONSTANTS packet can appear in any time interval.
- > GRID VALUES. If the method GRID VALUE is assigned to any variable in the METHOD packet, the first time interval must contain a GRID VALUES packet. More than one GRID VALUES packet can appear in any time interval.
- > STATION READINGS. If either the POISSON, DIVERGENCE, or STATINTERP method is assigned to any variable in the METHOD packet, the first time interval must contain a STATION READINGS packet. More than one STATION READINGS packet can appear in any time interval.
- > VERTICAL PROFILES. If any of the profile methods are assigned to any variable in the VERTICAL METHOD packet, the first time interval must contain a VERTICAL PROFILES packet. There must be a vertical profile defined (or implied by means of ALL) for every variable in every subregion for which a profile method was specified. The number of height-value pairs in any single profile must not exceed the maximum specified in the CONTROL packet. More than one VERTICAL PROFILES packet can appear in any time interval.

Typically, the wind values input will be in terms of SPEED and DIRECTION at measuring stations, whereas the variables output to the file are WINDX and WINDY. In this case, the methods and vertical methods should be specified as follows:

- > For SPEED and DIRECTION, use the method and vertical method POLARIZE if speed and direction grids are to be printed, or CONSTANT if they are not to be printed. If CONSTANT is chosen, supply a dummy CONSTANTS packet to give an arbitrary initial value to the SPEED and DIRECTION grids.
- > For WINDX and WINDY, specify one of the methods using station readings--STATINTERP, POISSON, or DIVERGENCE--and the vertical method desired for computation.

In each time interval, missing values of WINDX and WINDY for station readings and vertical profiles will be calculated from SPEED and DIRECTION, and vice versa. To ensure that old values are replaced with the appropriate new values, the variables WINDX and WINDY must be cleared to -9. before reading in the data for a new time interval. For station readings, the following can be done:

STATION READINGS			STATION READINGS		
ALL	WINDX	-9.	or	ALL	ALL -9.
ALL	WINDY	-9.			
END					

A shorthand method for clearing an entire vertical profile to -9. has also been provided--simply specify -9 as the number of profile heights, as shown in the following example:

#### VERTICAL PROFILES

A ----- ALL ----- 4200. ----- 5000. ----- -9  
END.

#### 5. Additional Input Files

If any vertical method besides CONSTANT was selected, the DIFFBREAK and REGIONTOP files must be input to WNDSET. In addition, if DIVERGENCE was selected the TEMPERATUR file must also be input.

#### 6. Examples

Exhibit VII-19 shows a sample input deck for WIND file generation. The output from a WNDSET run using that input deck is shown in Exhibit VII-20.

```

CONTROL
WIND    WIND FIELD FOR ST. LOUIS      DIVERGENCEF      ALGORITHM USED
      0          0           25          1          10
      32         0           1          0
      1          0           0          0          1
      10         20          0          0
      11         12          0          0
      76195      500          14          2400
END
REGION
      725000.   4252000.      15
      -19000.   -16000.
      4000.     4000.
      17         22          4
      2          2          0.0      50.
      200.      200.
END

```

Exhibit VII-19. Sample Input Deck for the WINDSET Program

UNITS	WINDX	M/SEC	3600.	1.
WINDY	M/SEC	3600.	1.	
SPEED	M/SEC	3600.	1.	
END				
<b>STATIONS</b>				
101	19103.	27862.	30.	
102	17518.	34045.	30.	
103	22588.	30467.	30.	
104	22312.	25304.	30.	
105	16706.	24453.	30.	
106	13660.	25566.	30.	
107	15179.	30610.	30.	
108	23407.	39102.	10.	
109	30802.	27886.	30.	
110	22209.	20826.	10.	
111	13812.	20479.	30.	
112	8938.	28913.	30.	
113	12738.	37820.	30.	
114	19320.	45456.	10.	
115	32111.	45799.	10.	
116	37777.	38083.	10.	
117	35560.	20818.	10.	
118	18065.	11256.	10.	
119	4759.	16547.	30.	
120	-1921.	33909.	30.	
121	7414.	50376.	10.	
122	16631.	77223.	30.	
123	52320.	34378.	30.	
124	24275.	-15663.	30.	
125	-27555.	30240.	30.	
<b>END</b>				
<b>TIME INTERVAL</b>				
76195	500	76195	600	
<b>SUBREGION</b>				
A	1	1	-1	
<b>END</b>				
<b>METHOD</b>				
A	WINDX	DIVERGENCE	-20.	20.
RMAXIT	1000.			3
TLRNE	10000.			
ROMEGA	1.4			
A	WINDY	DIVERGENCE	-20.	20.
RMAXIT	1000.			3
TLRNE	10000.			
ROMEGA	1.4			
A	SPEED	CONSTANT	0.	20.
A	DIRECTION	CONSTANT	-7.	7.
<b>END</b>				

VERTICAL METHOD		WINDY		DIVERGENCE		-20.		-20.	
A	A	WINDX	WINDY	DIVERGENCE	DIVERGENCE	-20.	-20.	20.	20.
A	A	SPEED	SPEED	CONSTANT	CONSTANT	0.	0.	7.	7.
END		CONSTANTS		DIRECTION		3.0		3.14	
CONSTANTS		A		SPEED		3.0		3.14	
CONSTANTS		A		DIRECTION		3.0		3.14	
<b>STATION READINGS</b>									
ALL ALL									
101	101	SPEED	DIRECTION	1.63	3.51				
102	102	SPEED	DIRECTION	2.21	3.26				
103	103	SPEED	DIRECTION	2.16	3.17				
104	104	SPEED	DIRECTION	1.96	3.17				
105	105	SPEED	DIRECTION	3.24	1.72				
106	106	SPEED	DIRECTION	3.75	1.77				
107	107	SPEED	DIRECTION	2.96	1.77				
108	108	SPEED	DIRECTION	1.93	2.96				
109	109	SPEED	DIRECTION	2.06	1.98				
110	110	SPEED	DIRECTION	2.94	1.62				
111	111	SPEED	DIRECTION	1.75	2.38				
111	111	SPEED	DIRECTION	1.61	1.81				
112	112	SPEED	DIRECTION	2.83	1.77				
112	112	SPEED	DIRECTION	1.82	1.92				
113	113	SPEED	DIRECTION	3.03	2.16				
113	113	SPEED	DIRECTION	2.10	1.86				
114	114	SPEED	DIRECTION	2.67	2.70				
114	114	SPEED	DIRECTION	1.92	1.60				
115	115	SPEED	DIRECTION	2.16	2.67				
115	115	SPEED	DIRECTION	2.10	1.97				
116	116	SPEED	DIRECTION	2.67	2.16				
117	117	SPEED	DIRECTION	1.97	2.16				
117	117	SPEED	DIRECTION	2.16	2.16				

Exhibit VII-19 (Continued)

116	SPEED	-9.00
116	DIRECTION	-9.00
116	SPEED	3.24
119	DIRECTION	1.04
119	SPEED	3.21
120	DIRECTION	1.98
120	SPEED	2.68
121	DIRECTION	1.84
121	SPEED	2.51
122	DIRECTION	2.37
122	SPEED	2.26
123	DIRECTION	2.04
123	SPEED	1.90
124	DIRECTION	-9.00
124	SPEED	3.12
125	DIRECTION	1.76
125	SPEED	1.76
END		
	VERTICAL PROFILE	
A	ALL	42948.0
		79053.0
		2
	STA	
A	SPEED	42948.0
		79053.0
		20
	STA 141	
A	0	2.7
34.5		4.9
116.1		6.6
197.0		7.5
276.7		6.0
355.4		4.7
433.2		5.1
510.0		5.5
586.0		6.9
660.9		9.2
734.2		9.2
807.2		11.6
880.2		10.5
953.2		10.5
1026.2		10.5
1099.2		9.6
1172.2		9.3
1245.6		8.9
1319.0		8.7
1392.0		8.2
A	DIRECTION	42948.0
		79053.0
		20
	STA 141	
A	0	1.5
34.5		1.7
116.1		1.9
197.0		2.0
276.7		2.0
355.4		2.2
433.2		4.7
510.0		5.4
586.0		5.5
660.9		5.6

END		ENDTIME		TIME INTERVAL		STATION READINGS		WINDS FOR 600	
734.2	5.6	76195	600	76195	700	ALL	ALL	-9.0	
807.2	5.6					101	SPEED	3.29	
880.2	5.6					101	DIRECTION	2.07	
953.2	5.7					102	SPEED	3.17	
1026.2	5.7					102	DIRECTION	2.15	
1099.2	5.7					103	SPEED	2.56	
1172.2	5.7					103	DIRECTION	2.16	
1245.6	5.7					104	SPEED	2.54	
1319.0	5.6					104	DIRECTION	2.28	
1392.0	5.6					105	SPEED	2.69	
						105	DIRECTION	2.34	
						106	SPEED	2.90	
						106	DIRECTION	2.16	
						107	SPEED	2.96	
						107	DIRECTION	2.13	
						108	SPEED	2.64	
						108	DIRECTION	2.15	
						109	SPEED	2.87	
						109	DIRECTION	2.04	
						110	SPEED	2.05	
						110	DIRECTION	2.25	
						111	SPEED	2.45	
						111	DIRECTION	2.20	
						112	SPEED	3.16	
						112	DIRECTION	2.01	
						113	SPEED	3.92	
						113	DIRECTION	2.13	
						114	SPEED	1.95	
						114	DIRECTION	2.05	
						115	SPEED	2.50	
						115	DIRECTION	1.92	
						116	SPEED	2.97	
						116	DIRECTION	1.89	
						117	SPEED	3.31	
						117	DIRECTION	2.16	
						118	SPEED	2.99	
						118	DIRECTION	2.10	
						119	SPEED	2.70	
						119	DIRECTION	2.18	
						120	SPEED	3.20	
						120	DIRECTION	2.31	
						121	SPEED	3.08	

PROCESSING OF INPUT DATA FOR THE PREPARATION OF FILE WIND

REGION	WIND FIELD FOR ST. LOUIS	DIVERGENCE	ALGORITHM USED
KCRNT =	725000.	4252000.	15
	-1900.	-16000.	
	4000.00	4000.00	
KCRNT =	17	22	4
KCRNT =	2	2	.00
<b>END</b>			<b>50.00</b>
			<b>200.00</b>
KCRNT =	61	NSCLFT = 11040	IN SCRSET
KCRNT =	121	NSCLFT = 11780	IN SCRSET
KCRNT =	127	NSCLFT = 11774	IN SCRSET
KCRNT =	133	NSCLFT = 11768	IN SCRSET
KCRNT =	139	NSCLFT = 11762	IN SCRSET
KCRNT =	389	NSCLFT = 11512	IN SCRSET
KCRNT =	489	NSCLFT = 11412	IN SCRSET
KCRNT =	514	NSCLFT = 11387	IN SCRSET
KCRNT =	539	NSCLFT = 11362	IN SCPSET
KCRNT =	564	NSCLFT = 11337	IN SCRSET
KCRNT =	574	NSCLFT = 11327	IN SCRSET
KCRNT =	948	NSCLFT = 10953	IN SCRSET
KCRNT =	952	NSCLFT = 10949	IN SCRSET
KCRNT =	956	NSCLFT = 10945	IN SCRSET
KCRNT =	960	NSCLFT = 10941	IN SCRSET
KCRNT =	964	NSCLFT = 10937	IN SCRSET
KCRNT =	968	NSCLFT = 10933	IN SCRSET
KCRNT =	1068	NSCLFT = 10833	IN SCRSET
KCRNT =	1078	NSCLFT = 10823	IN SCRSET
KCRNT =	1082	NSCLFT = 10819	IN SCRSET
KCRNT =	1086	NSCLFT = 10815	IN SCRSET
KCRNT =	1090	NSCLFT = 10811	IN SCRSET
KCRNT =	1094	NSCLFT = 10807	IN SCRSET
KCRNT =	1098	NSCLFT = 10803	IN SCRSET
KCRNT =	1198	NSCLFT = 10703	IN SCRSET
KCRNT =	1208	NSCLFT = 10693	IN SCRSET
KCRNT =	1212	NSCLFT = 10689	IN SCRSET
KCRNT =	1216	NSCLFT = 10685	IN SCRSET
KCRNT =	1296	NSCLFT = 10605	IN SCRSET
KCRNT =	1376	NSCLFT = 10525	IN SCRSET
KCRNT =	1750	NSCLFT = 10151	IN SCRSET
KCRNT =	2124	NSCLFT = 9777	IN SCRSET
KCRNT =	2498	NSCLFT = 9403	IN SCRSET
KCRNT =	2872	NSCLFT = 9029	IN SCRSET
KCRNT =	3246	NSCLFT = 8655	IN SCRSET
KCRNT =	9230	NSCLFT = 2671	IN SCRSET
KCPNT =	9604	NSCLFT = 2297	IN SCRSET
			UNITS
			STATIONS

## STATION IDENTIFICATION TABLE

STATION NAME	X-LOCATION(GRID SQ)	Y-LOCATION (GRID SQ)	STATION HEIGHT(METERS)
101	9.55	10.97	30.00
102	9.13	12.51	30.00
103	10.40	11.62	30.00
104	10.33	10.33	30.00
105	9.43	10.11	30.00
106	8.16	10.39	30.00
107	8.54	11.65	30.00
108	10.60	13.76	10.00
109	12.45	10.97	30.00
110	10.30	9.21	10.00
111	8.20	9.12	30.00
112	6.98	11.23	30.00
113	7.93	13.45	30.00
114	9.58	15.36	10.00
115	12.78	15.45	10.00
116	14.19	13.52	10.00
117	13.64	9.20	10.00
118	9.27	6.81	10.00
119	5.94	8.64	30.00
120	4.27	12.48	30.00
121	6.60	16.59	10.00
122	8.91	23.31	30.00
123	17.83	12.59	30.00
124	10.82	.13	30.00
125	-2.14	11.56	30.00
TIME INTERVAL	76195	500	600

SUBREGION  
METHOD  
VERTICAL METHOD

METHODS TABLE

SUBREGION	VARIABLE	METHOD	MIN. VALUE	MAX. VALUE	PARAMETER	PAR. VALUE
-----------	----------	--------	------------	------------	-----------	------------

A	WINDX	DIVERGENCE	-72000.0000	72000.0000		
A	WINDY	DIVERGENCE	-72000.0000	72000.0000		
A	SPEED	CONSTANT	.0000	72000.0000		
A	DIRECTION	CONSTANT	-7.0000	7.0000		
CONSTANTS						
STATION READINGS						
VERTICAL PROFILE						
ENDTIME						
KCRNT = 21	NSCLFT = 2277	IN SCRSET				
KCRNT = 41	NSCLFT = 2257	IN SCRSET				
KCRNT = 45	NSCLFT = 2253	IN SCRSET				
KCRNT = 49	NSCLFT = 2249	IN SCRSET				
KCRNT = 69	NSCLFT = 2229	IN SCRSET				
KCRIT = 89	NSCLFT = 2209	IN SCRSET				
KCRNT = 93	NSCLFT = 2205	IN SCRSET				
KCRNT = 467	NSCLFT = 1831	IN SCRSET				
KCRNT = 841	NSCLFT = 1457	IN SCRSET				

WIND SCALAR VALUES BETWEEN DATE 76195 / TIME 5.00 AND DATE 76195 / TIME 6.00

AVERAGE STATION REFERENCE HEIGHT	23.60 METERS
MAXIMUM SPEED IN X-DIRECTION	22453.10 METERS/HOUR
MAXIMUM SPEED IN Y-DIRECTION	9602.99 METERS/HOUR
AVERAGE SPEED AT WEST BORDER	14801.04 METERS/HOUR
AVERAGE SPEED AT EAST BORDER	14801.11 METERS/HOUR
AVERAGE SPEED AT SOUTH BORDER	14799.13 METERS/HOUR
AVERAGE SPEED AT NORTH BORDER	14798.96 METERS/HOUR

FILE NAME -- WIND	WIND FIELD FOR ST. LOUIS				DIVERGENCE				ALGORITHM USED				PAGE 1
WIND VELOCITIES (IN METERS PER HOUR)	COMPONENT WINDX				VERT.	LEVEL 1	START DATE/TIME	76195 5.00	STOP DATE/TIME	76195 6.00			
GRID CELL	1	2	3	4	5	6	7	8	9	10	11	12	
22	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	
21	-9608.41	-9597.95	-9579.41	-9568.13	-9560.89	-9556.42	-9554.05	-9553.15	-9553.21	-9553.97	-9555.34	-9557.28	
20	-9629.52	-9617.58	-9590.74	-9570.93	-9556.77	-9547.54	-9542.64	-9540.86	-9541.03	-9542.76	-9545.91	-9550.42	
19	-9639.70	-9629.26	-9598.60	-9572.82	-9552.85	-9539.38	-9532.62	-9530.53	-9530.89	-9533.54	-9538.42	-9545.61	
18	-9643.83	-9635.78	-9603.36	-9573.39	-9548.70	-9531.51	-9524.08	-9522.62	-9522.87	-9526.22	-9532.48	-9540.82	
17	-9644.58	-9639.11	-9606.02	-9573.32	-9545.04	-9524.29	-9517.11	-9516.83	-9515.88	-9520.07	-9528.19	-9537.93	
16	-9643.51	-9640.49	-9607.57	-9573.79	-9544.33	-9522.76	-9512.11	-9508.57	-9507.45	-9513.54	-9524.24	-9535.49	
15	-9641.41	-9640.50	-9608.27	-9574.50	-9545.05	-9523.00	-9509.68	-9503.90	-9502.55	-9506.55	-9516.48	-9530.33	
14	-9638.67	-9639.30	-9608.03	-9574.90	-9546.36	-9524.56	-9509.46	-9502.23	-9499.21	-9499.19	-9507.85	-9524.57	
13	-9635.44	-9636.88	-9606.73	-9574.58	-9547.76	-9527.51	-9512.23	-9502.66	-9495.58	-9491.19	-9498.89	-9519.23	
12	-9631.80	-9633.17	-9604.48	-9573.99	-9546.09	-9526.70	-9514.25	-9503.75	-9491.93	-9483.51	-9492.50	-9516.89	
11	-9627.67	-9627.86	-9600.59	-9571.54	-9546.27	-9527.42	-9514.08	-9506.30	-9493.14	-9483.70	-9490.80	-9516.09	
10	-9623.07	-9620.88	-9595.05	-9567.46	-9542.58	-9524.25	-9511.06	-9499.94	-9490.60	-9486.79	-9496.40	-9521.06	
9	-9618.20	-9612.45	-9588.23	-9563.00	-9538.46	-9521.13	-9509.01	-9498.25	-9491.60	-9492.15	-9503.50	-9525.90	
8	-9613.28	-9603.53	-9579.32	-9559.94	-9536.94	-9519.30	-9506.79	-9497.73	-9493.74	-9497.33	-9509.89	-9531.00	
7	-9608.43	-9600.24	-9578.86	-9557.41	-9536.29	-9519.35	-9507.22	-9499.57	-9497.39	-9502.89	-9516.07	-9535.96	
6	-9603.65	-9596.32	-9575.93	-9555.66	-9536.86	-9521.61	-9510.75	-9504.55	-9503.75	-9509.49	-9521.85	-9540.34	
5	-9598.84	-9591.76	-9572.87	-9554.95	-9539.02	-9526.22	-9517.22	-9512.42	-9512.33	-9517.57	-9528.28	-9544.21	
4	-9503.75	-9586.84	-9570.19	-9555.47	-9543.00	-9533.22	-9526.46	-9523.02	-9523.23	-9527.42	-9535.76	-9548.15	
3	-9587.01	-9581.43	-9568.03	-9557.36	-9548.92	-9542.55	-9518.25	-9516.14	-9536.39	-9539.18	-9544.63	-9552.80	
2	-9579.70	-9576.95	-9566.50	-9560.63	-9556.77	-9553.88	-9552.01	-9551.13	-9551.29	-9552.56	-9555.02	-9558.76	
1	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24	

PAGE 2

FILE NAME -- WIND	WIND FIELD FOR ST. LOUIS			DIVERGENCE	ALGORITHM USED
	WIND VELOCITIES (IN METERS PER HOUR)			COMPONENT WINDX	VERT.
GRID CELL	13	14	15	16	17
22	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24
21	-9559.75	-9562.87	-9567.58	-9574.91	-9578.55
20	-9556.05	-9561.82	-9570.62	-9582.49	-9586.79
19	-9556.27	-9562.33	-9574.61	-9589.70	-9593.28
18	-9551.24	-9563.53	-9579.54	-9597.14	-9598.94
17	-9549.96	-9565.47	-9584.86	-9604.72	-9604.12
16	-9549.26	-9567.40	-9590.16	-9612.26	-9609.00
15	-9547.77	-9569.70	-9595.53	-9619.70	-9613.69
14	-9545.83	-9571.59	-9600.60	-9626.83	-9618.20
13	-9544.25	-9573.41	-9605.34	-9633.48	-9622.57
12	-9544.28	-9575.70	-9609.89	-9639.59	-9626.94
11	-9545.81	-9578.68	-9614.32	-9645.04	-9631.46
10	-9550.51	-9583.39	-9618.79	-9649.68	-9636.27
9	-9554.72	-9587.83	-9622.96	-9653.00	-9641.28
8	-9558.72	-9590.80	-9624.42	-9653.57	-9645.85
7	-9561.93	-9590.70	-9621.48	-9650.46	-9649.26
6	-9563.83	-9582.33	-9613.42	-9644.51	-9650.73
5	-9564.38	-9585.54	-9610.81	-9640.02	-9649.32
4	-9564.13	-9582.82	-9605.26	-9632.82	-9643.69
3	-9563.68	-9577.48	-9595.45	-9619.94	-9631.59
2	-9563.98	-9571.18	-9581.71	-9598.90	-9608.83
1	-9566.24	-9566.24	-9566.24	-9566.24	-9566.24

FILE NAME -- WIND	WIND FIELD FOR ST. LOUIS				DIVERGENCE	ALGORITHM USED		PAGE 1
GRID CELL	1	2	3	4	5	6	7	5.00 STOP DATE/TIME 76195 6.00
	WIND VELOCITIES (IN METERS PER HOUR)				COMPONENT WINDY	VERT.	LEVEL 1 START DATE/TIME 76195	
22	3119.40	3161.56	3180.31	3185.97	3183.71	3176.66	3167.01	3156.30 3145.39 3134.84 3125.04 3116.40
21	3119.40	3152.35	3170.68	3178.39	3176.33	3172.83	3163.96	3153.45 3142.37 3131.53 3121.70 3113.60
20	3119.40	3136.33	3149.15	3155.83	3156.53	3152.41	3145.25	3137.09 3126.60 3120.44 3113.50 3106.56
19	3119.40	3127.43	3134.67	3138.66	3136.49	3134.46	3128.06	3122.24 3116.30 3110.66 3106.48 3104.55
18	3119.40	3122.24	3125.32	3126.90	3125.90	3121.98	3115.80	3112.40 3108.25 3103.97 3101.59 3100.84
17	3119.40	3119.21	3119.65	3119.87	3119.29	3117.67	3114.93	3111.19 3106.77 3102.02 3100.30 3099.76
16	3119.40	3117.45	3116.27	3115.66	3116.23	3117.51	3116.89	3115.10 3111.41 3107.45 3103.42 3100.82
15	3119.40	3116.33	3114.04	3113.03	3113.59	3115.98	3116.37	3117.57 3116.92 3114.48 3107.52 3102.53
14	3119.40	3115.54	3112.34	3110.55	3110.55	3113.38	3116.92	3119.12 3119.69 3116.66 3108.76 3103.24
13	3119.40	3114.90	3110.92	3108.45	3107.91	3109.45	3112.72	3116.09 3117.01 3112.68 3105.14 3101.36
12	3119.40	3114.30	3109.46	3105.90	3103.96	3102.56	3102.94	3104.28 3105.10 3103.27 3099.26 3097.51
11	3119.40	3113.73	3107.95	3102.78	3098.29	3094.01	3091.04	3088.53 3085.43 3086.46 3088.70 3091.05
10	3119.40	3113.45	3107.16	3100.63	3094.62	3088.26	3063.76	3078.84 3073.46 3074.10 3080.12 3085.96
9	3119.40	3113.63	3107.79	3101.13	3095.10	3089.57	3084.33	3079.58 3076.41 3077.14 3082.73 3087.52
8	3119.40	3114.19	3110.99	3102.97	3097.85	3094.21	3089.48	3085.84 3084.18 3085.28 3089.66 3094.02
7	3119.40	3114.09	3109.30	3104.12	3100.40	3098.02	3095.84	3094.74 3095.31 3097.67 3101.28 3104.05
6	3119.40	3114.01	3109.23	3105.34	3103.09	3102.51	3103.10	3104.97 3108.15 3112.24 3115.83 3118.64
5	3119.40	3113.86	3109.33	3106.50	3105.88	3107.48	3110.69	3115.74 3121.48 3127.25 3131.76 3134.23
4	3119.40	3113.33	3108.99	3107.24	3108.45	3112.47	3118.76	3126.59 3135.01 3142.83 3148.68 3151.41
3	3119.40	3111.87	3107.55	3107.09	3110.40	3117.00	3126.15	3136.87 3147.98 3158.09 3165.74 3169.60
2	3119.40	3108.20	3104.13	3105.58	3111.39	3120.62	3132.32	3145.44 3158.81 3171.12 3180.99 3187.15
1	3119.40	3105.94	3102.63	3105.45	3112.28	3121.77	3132.94	3144.98 3157.18 3168.81 3179.08 3187.07

FILE NAME -- WIND	WIND FIELD FOR ST. LOUIS	DIVERGENCE	ALGORITHM USED		
MIND VELOCITIES (IN METERS PER HOUR)	COMPONENT WINDY	VERT.	LEVEL 1 START DATE/TIME 76195 5.00 STOP DATE/TIME 76195 6.00		
GRID CELL	13	14	15	16	17
22	3109.38	3104.65	3103.30	3107.09	3119.40
21	3107.72	3103.95	3103.60	3108.33	3119.40
20	3106.23	3104.36	3105.66	3111.02	3119.40
19	3107.51	3104.43	3106.53	3112.11	3119.40
18	3101.76	3103.00	3106.54	3112.51	3119.40
17	3100.65	3102.43	3106.46	3112.65	3119.40
16	3100.28	3102.09	3106.34	3112.68	3119.40
15	3100.35	3101.99	3106.29	3112.70	3119.40
14	3101.00	3102.28	3106.41	3112.76	3119.40
13	3100.44	3102.14	3106.37	3112.75	3119.40
12	3098.46	3101.21	3105.91	3112.58	3119.40
11	3094.44	3099.34	3105.00	3112.27	3119.40
10	3090.97	3097.04	3104.09	3112.05	3119.40
9	3092.24	3097.71	3104.85	3112.50	3119.40
8	3098.26	3102.90	3106.25	3113.98	3119.40
7	3108.07	3111.13	3113.63	3116.35	3119.40
6	3120.16	3120.13	3119.94	3119.37	3119.40
5	3133.91	3130.73	3127.52	3123.30	3119.40
4	3150.19	3145.19	3138.15	3129.02	3119.40
3	3168.74	3163.01	3152.85	3137.91	3119.40
2	3188.54	3184.34	3173.40	3153.15	3119.40
1	3191.59	3190.99	3182.66	3161.99	3119.40

## VIII SEGMENTATION OF M.E.B. FILES

If the entire modeling region is too large to fit within the core limitations of the Airshed Simulation Program, the region must be subdivided into rectangular subregions called "segments" that will fit into core. The segmentation process involves (1) determining the size and layout of the segments and (2) running the file segmentation program to create segmented versions of each spatially varying data file. This chapter first describes how to determine the core requirements for a given segmentation scheme and then how to use the file segmentation program.

### A. CALCULATING CORE REQUIREMENTS

The maximum horizontal dimensions allowable for a single segment depend on the number of segments, the number of vertical levels and chemical species, and the number of point sources within the modeling region; the program options selected; and the amount of core available on the host computer. When the Airshed Simulation Program is installed on a particular computer, it will contain a vector of available space. The length of this vector is specified when the program is compiled and should be as large as possible. The vector of available space is used to hold the following four tables:

- > Segment
- > Local
- > Boundary
- > Segment scratch.

The detailed contents of these tables appear in Chapter VII of the Systems Manual. Only the methods of calculating the length of each table are described here.

To determine whether a segment of a given size will fit in core, the calculated sizes of the tables must be summed and that total compared with the amount of available core. The calculations for each table are discussed next, followed by Exhibit VIII-1, which is a worksheet to be used for the calculations. Finally, a sample problem is presented showing the calculation of core requirements for different segmentation schemes for the same region.

### 1. Segment Table

The Segment Table contains 65 entries for each segment.

$$S.T. = 65 * \text{seg} ,$$

where seg = number of segments.

### 2. Local Table

The Local Table contains vectors used by the numerical integration routines.

$$L.T. = 5 + 32z + zs + 5s$$

$$+ 4r + 2r^2$$

$$+ 6x_{\max} ,$$

where

$z$  = number of vertical levels,

$s$  = number of chemical species,

$r$  = number of reactive species,

$x_{\max}$  = maximum dimension in  $x$  or  $y$  direction for any segment.

Fixed Dimensions      Simulation Options (1 = yes, 0 = no)

$z =$ _____	TEMPERATUR = _____
$s =$ _____	TERRAIN = _____
$r =$ _____	ROADWAY = _____
	VARIATION = _____

	Segment			
	1	2	3	4
x				
y				
p				
interface				

$$65 \times \frac{\text{seg}}{\text{seg}} = \text{_____} \text{ (S.T.)}$$

$$\left[ \begin{array}{l} 5 + \frac{32z}{zs} + \frac{zs}{5s} \\ + \frac{4r}{2r^2} \\ + \frac{6(x,y)_{\max}}{} \end{array} \right] = \text{_____} \text{ (L.T.)}$$

$$\left[ \begin{array}{l} 8(x+y) \\ + \frac{zs}{\text{interface}} \end{array} \right] = \text{_____} \text{ (B.T.)}$$

$$\left[ \begin{array}{l} 8 + \frac{2\text{terr}}{\text{temp}} + \frac{\text{temp}}{\text{road}} \\ + \frac{2z}{s} \\ + \frac{s}{zs} (2 + \frac{\text{road}}{\text{road}} + \frac{\text{var}}{\text{var}}) \end{array} \right] \rightarrow \boxed{[ ]} \times \frac{xy}{p} + \frac{2+s}{2+s} \times \frac{xy}{p} = \text{_____} \text{ (S.S.T.)}$$

Sum of all tables = \_\_\_\_\_

Available storage = \_\_\_\_\_

### 3. Boundary Table

The Boundary Table contains the boundary cell definitions and the concentrations in the interface cells for a single segment. The "interface cells" of a segment are those that lie within the two-cell overlap into the adjacent segment. The number of interface cells in a segment is two times the sum of the lengths of each nonexternal edge. For non-symmetrical segmentation schemes (such as that shown in Figure VIII-3), the Boundary Table length for each segment must be calculated to determine the maximum storage requirement.

$$B.T. = 8(x + y) + zs \text{ (interface)} ,$$

where

$z$  = number of vertical levels,

$s$  = number of chemical species,

$(x+y)$  = sum of  $x$  and  $y$  dimensions for the segment,

$(\text{interface})$  = number of interface cells in the segment,

i.e., 2 times the length of all nonexternal edges.

### 4. Segment Scratch Table

The Segment Scratch Table contains all the other spatially varying data arrays, including predicted concentrations and variations, and the point source data, for one segment.

$$\begin{aligned} S.S.T. = xy & \left\{ 8 + 2(\text{terr}) + 1(\text{temp}) + 1(\text{road}) \right. \\ & + 2z + s[2 + 1(\text{road}) + 1(\text{var})] + zs \left. \right\} \\ & + p(2 + s) , \end{aligned}$$

where

```
x = number of cells in x direction;  
y = number of cells in y direction;  
p = number of point sources;  
z = number of vertical levels;  
s = number of chemical species;  
terr = 1, if TERRAIN file input,  
      0, otherwise;  
temp = 1, if TEMPERATUR file input,  
      0, otherwise;  
road = 1, if ROADWAY file input,  
      0, otherwise;  
var = 1, if concentration variations are calculated,  
      0, otherwise.
```

## 5. A Sample Problem

As an example, consider the region shown in Figure VIII-1, which is 17 x 22 cells, with the outer edge used as the boundary; the area modeled is thus 15 x 20 cells. There are 4 cells in the vertical direction, 13 chemical species to be modeled, and 200 elevated point sources. There are spatially varying terrain and temperature data but no roadway emissions file and, therefore, no surface layer variations. The core requirement calculations for this unsegmented region, which are shown in Exhibit VIII-2, indicate that the vector of available space must be at least 40373 words long. If it is not that long, the region must be divided into segments small enough to fit in core.

First, the user must try dividing the region in half, as shown in Figure VIII-2. (Recall that at each segment interface there is a two-cell overlap into the adjacent segment.) The resulting two segments are each 17 x 13 and the area modeled within each is 15 x 10. Each has one non-external edge 17 cells long. If the point sources are distributed fairly

<u>Segment</u>	<u>Inner Area</u>	<u>Total Area</u>	<u>Interface Cells</u>
1	15 x 20	17 x 22	0

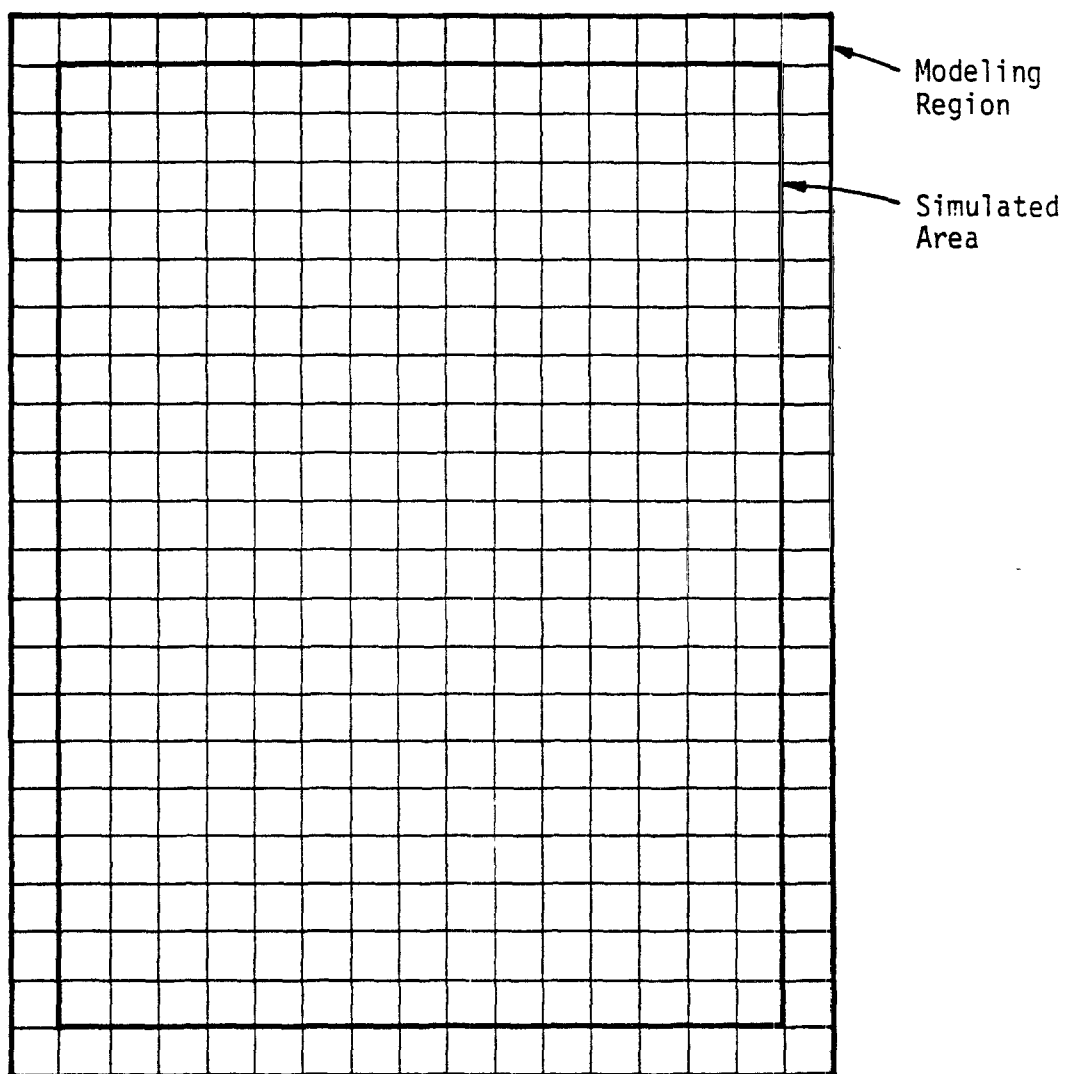


Figure VIII-1. Sample Region--Unsegmented

Fixed Dimensions · Simulation Options (1 = yes, 0 = no)

z = 4  
s = 13  
r = 12

TEMPERATUR = 1  
TERRAIN = 1  
ROADWAY = 0  
VARIATION = 0

	Segment			
	1	2	3	4
x	17			
y	22			
p	200			
interface	-			

$$65 \times \frac{1}{\text{seg}} = \underline{\quad 65 \quad} (\text{S.T.})$$

$$\left[ 5 + \frac{128}{32z} + \frac{52}{zs} + \frac{65}{5s} + \frac{48}{4r} + \frac{288}{2r^2} + \frac{132}{6(x,y)_{\max}} \right] = \underline{\quad 718 \quad} (\text{L.T.})$$

$$\left[ \frac{8 \cdot 37}{8(x+y)} + \frac{0}{zs \text{ (interface)}} \right] = \underline{\quad 312 \quad} (\text{B.T.})$$

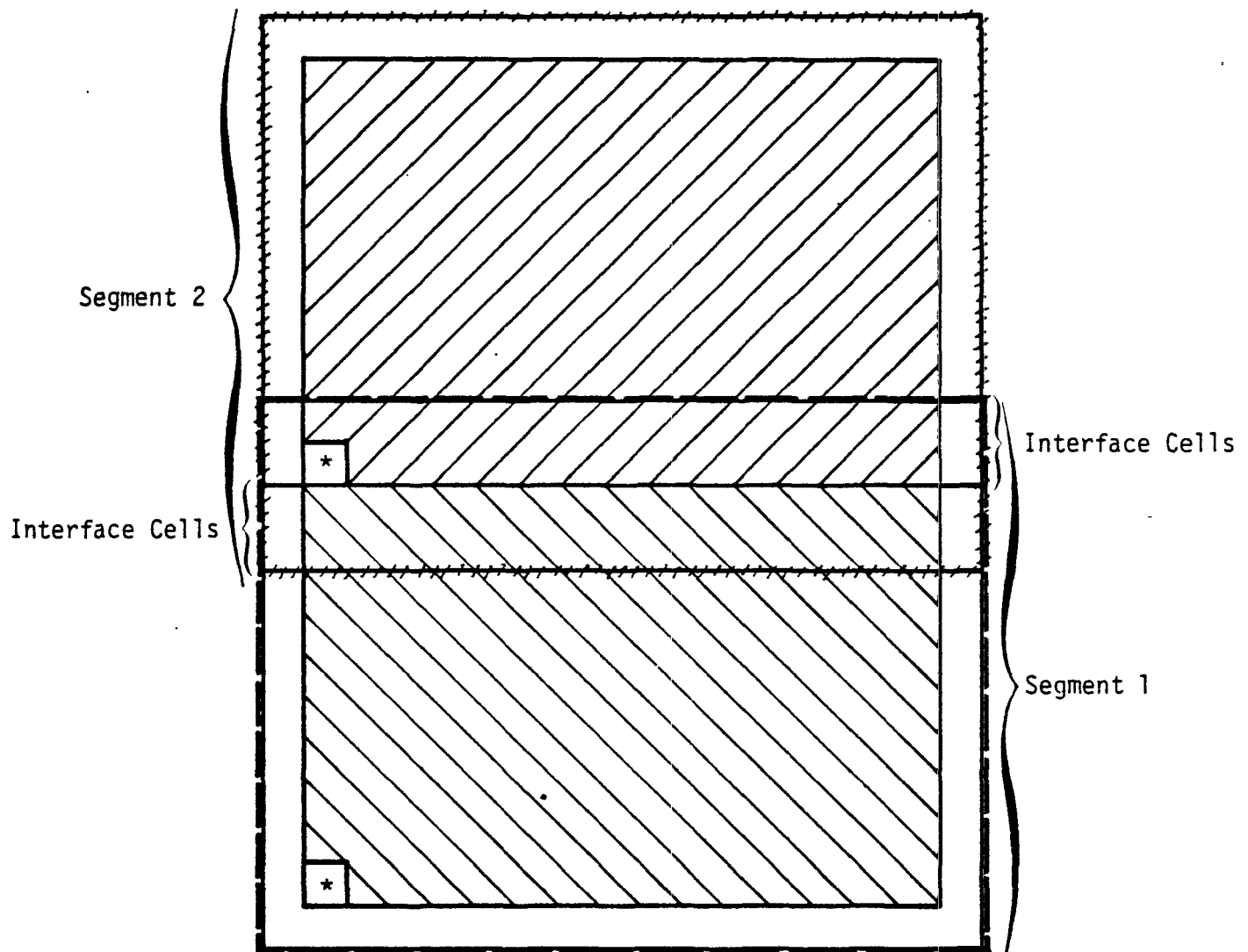
$$\left[ 8 + \frac{2}{2\text{terr}} + \frac{1}{\text{temp}} + \frac{0}{\text{road}} + \frac{8}{2z} + \frac{13}{s} (2 + \frac{0}{\text{road}} + \frac{0}{\text{var}}) + \frac{52}{zs} \right] \xrightarrow{\frac{97}{[\ ]} \times \frac{374}{xy}} + \frac{15}{2+s} \times \frac{200}{p} = \underline{\quad 39278 \quad} (\text{S.S.T.})$$

$$\text{Sum of all tables} = \underline{\quad 40373 \quad}$$

$$\text{Available storage} = \underline{\quad 30,000 \quad}$$

Exhibit VIII-2. Worksheet Filled Out for an Unsegmented Region

<u>Segment</u>	<u>Inner Area</u>	<u>Total Area</u>	<u>Interface Cells</u>
1	15 x 10	17 x 13	2 x 17
2	15 x 10	17 x 13	2 x 17



\* Segment origins input to the file segmentation program.

Figure VIII-2. Sample Region Divided into Two Segments

evenly throughout the region, there will be approximately 100 in each segment (count only the point sources within the modeled area, not those in the interface cells). Exhibit VIII-3 shows the computation of a core requirement of 25763 for this two-segment scheme.

If the two-segment scheme is still too large, the region can be further divided. Figure VIII-3 illustrates a three-segment scheme, without the actual segment edges drawn in. Size computations are shown in Exhibit VIII-4.

The objective of any segment layout design is to minimize the length of the segment boundaries to obtain segments that are as nearly square and equal sized as possible. Thus, the arrangement of the three segments shown in Figure VIII-3 represents a more efficient utilization of core space than that which would be obtained by merely cutting the region into three slices along either axis.

## B. THE FILE SEGMENTATION PROGRAM

The spatially varying data files must be converted into segmented form according to the region segmentation scheme selected. If segmentation is not necessary, the files may be used in their unconverted form since unsegmented files are structured like segmented files that contain only one segment. The file segmentation program (SEGDEF) is used to create a segmented file from an unsegmented file. It will operate on any of the 11 spatially varying M.E.B. files, but only on one at a time. SEGDEF requires subroutines from the libraries UTILITY and FILUTIL.

The input to SEGDEF consists of a card deck and one unsegmented file, as illustrated in Figure VIII-4. The CONTROL and REGION packets are described in detail in Section V.A. The SEGMENTDEF and SEGMENT FILE packets are defined here in Sections B.1 and B.2.

Fixed Dimensions    Simulation Options (1 = yes, 0 = no)

$$\begin{array}{l} z = \underline{\underline{4}} \\ s = \underline{\underline{13}} \\ r = \underline{\underline{12}} \end{array}$$

$$\begin{array}{l} \text{TEMPERATUR} = \underline{\underline{1}} \\ \text{TERRAIN} = \underline{\underline{1}} \\ \text{ROADWAY} = \underline{\underline{6}} \\ \text{VARIATION} = \underline{\underline{0}} \end{array}$$

	Segment			
	1	2	3	4
x	17	17		
y	13	13		
p	100	100		
interface	34	34		

$$65 \times \frac{2}{\text{seg}} = \underline{\underline{130}} \text{ (S.T.)}$$

$$\left[ \begin{array}{l} 5 + \frac{128}{32z} + \frac{52}{zs} + \frac{65}{5s} \\ + \frac{48}{4r} + \frac{288}{2r^2} \\ + \frac{102}{6(x,y)_{\max}} \end{array} \right] = \underline{\underline{688}} \text{ (L.T.)}$$

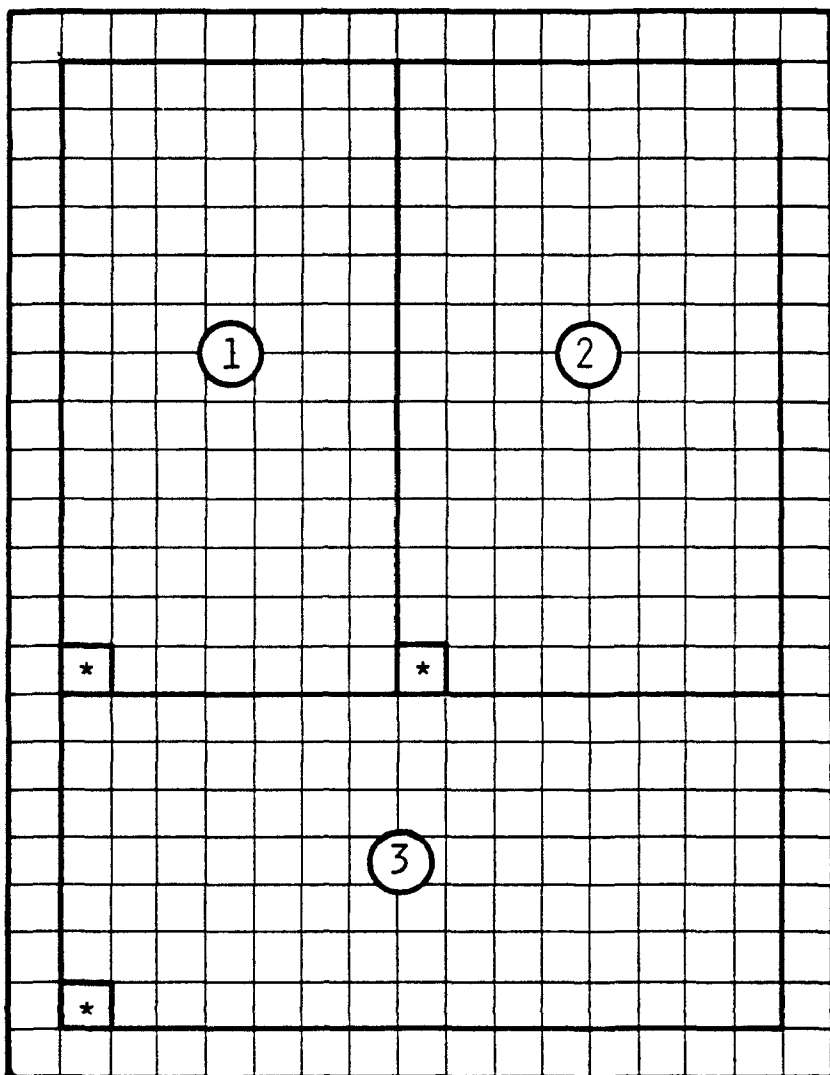
$$\left[ \frac{240}{8(x+y)} + \frac{52 \cdot 34}{zs \text{ (interface)}} \right] = \underline{\underline{2008}} \text{ (B.T.)}$$

$$\left[ \begin{array}{l} 8 + \frac{2}{2\text{terr}} + \frac{1}{\text{temp}} + \frac{0}{\text{road}} \\ + \frac{8}{2z} \\ + \frac{13}{s} (2 + \frac{0}{\text{road}} + \frac{0}{\text{var}}) \\ + \frac{52}{zs} \end{array} \right] \xrightarrow{\text{[ ]}} \frac{97}{2+s} \times \frac{221}{xy}, + \frac{15}{2+s} \times \frac{100}{p} = \underline{\underline{22937}} \text{ (S.S.T.)}$$

Sum of all tables = 25763

Available storage = 30000

<u>Segment</u>	<u>Inner Area</u>	<u>Total Area</u>	<u>Interface Cells</u>
1	7 x 13	10 x 16	2 x (10 + 16)
2	8 x 13	11 x 16	2 x (11 + 16)
3	15 x 7	17 x 10	2 x 17



\* Segment origins input to the file segmentation program.

Figure VIII-3. Sample Region Divided into Three Segments

Fixed Dimensions    Simulation Options (1 = yes, 0 = no)

$z = \underline{4}$   
 $s = \underline{13}$   
 $r = \underline{12}$

TEMPERATUR = /  
 TERRAIN = /  
 ROADWAY = 0  
 VARIATION = 0

	Segment			
	1	2	3	4
x	10	11	17	
y	16	16	10	
p				
interface	52	54	34	

$$65 \times \frac{3}{\text{seg}} = \underline{195} \text{ (S.T.)}$$

$$\left[ \begin{array}{l} 5 + \frac{128}{32z} + \frac{52 + 65}{zs} + \frac{65}{5s} \\ + \frac{48}{4r} + \frac{288}{2r^2} \\ + \frac{102}{6(x,y)_{\max}} \end{array} \right] = \underline{688} \text{ (L.T.)}$$

$$\left[ \frac{8 \cdot 27}{8(x+y)} + \frac{52 \cdot 54}{zs \text{ (interface)}} \right] = \underline{3024} \text{ (B.T.)}$$

$$\left[ \begin{array}{l} 8 + \frac{2}{2\text{terr}} + \frac{1}{\text{temp}} + \frac{0}{\text{road}} \\ + \frac{8}{2z} \\ + \frac{13}{s} (2 + \frac{-}{\text{road}} + \frac{-}{\text{var}}) \\ + \frac{52}{zs} \end{array} \right] \xrightarrow{\left[ \begin{array}{l} 97 \\ 176 \\ xy \end{array} \right]} + \frac{15}{2+s} \times \frac{80}{p} = \underline{18272} \text{ (S.S.T.)}$$

$$\text{Sum of all tables} = \underline{22179}$$

$$\text{Available storage} = \underline{30000}$$

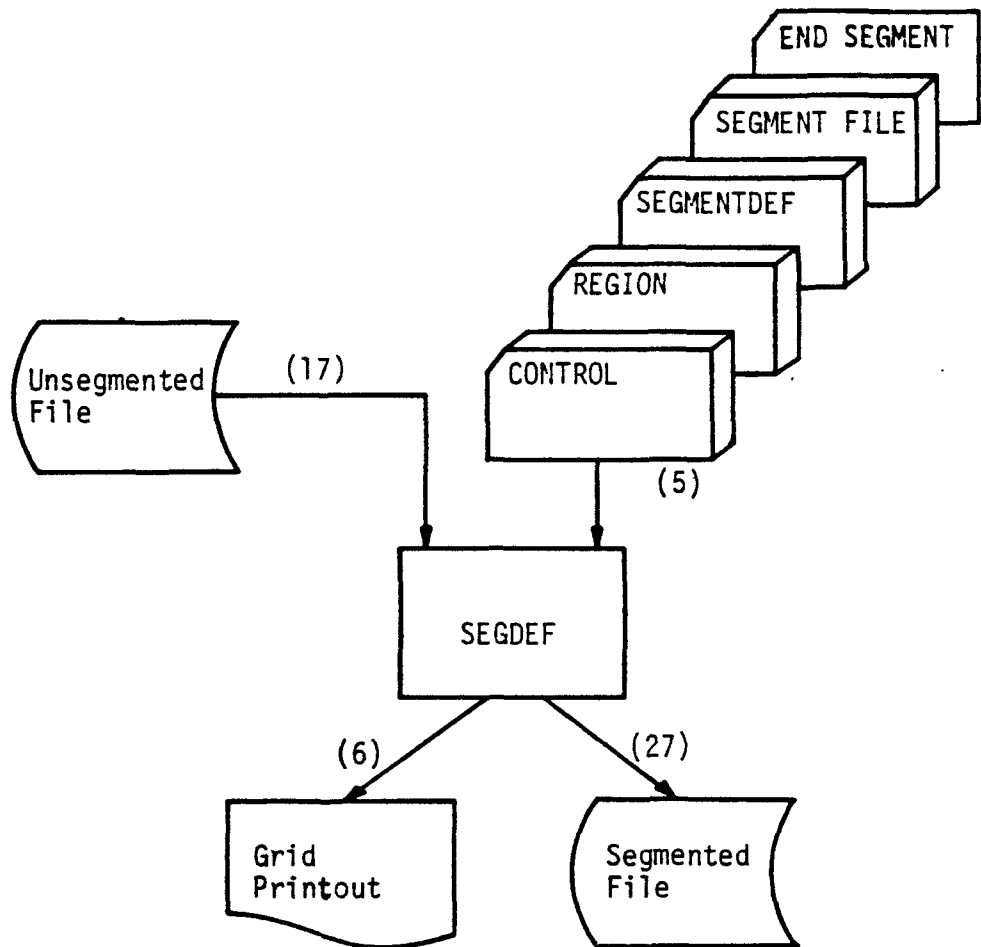


Figure VIII-4. Information Flow Diagram for the File Segmentation Program

The CONTROL packet entries for SEGDEF are shown in Table VIII-1. The following are special considerations for the CONTROL and REGION packets that pertain to this program:

> CONTROL

- The file name provided in Card 2 must be SEGMENTDEF.
- The only options recognized by the file segmentation program are the number of species, the number of segments, and the print flags for the input cards and the segmented regional grid.
- The beginning and ending dates can be the same as or contained within those of the file to be segmented.
- The list of species may be a subset of the list in the file; their number must be the same as the entry in the control parameter above. If the file to be segmented contains no species, the list can be omitted.

> REGION

- This packet is standard in all respects. The region can be a subregion of the region described in the file; if the file is three-dimensional, vertical parameters in the REGION packet must match those on the file.

## 1. The SEGMENTDEF Packet

The SEGMENTDEF packet provides the definition of the segmentation scheme. The contents of this packet are described in Table VIII-2, and the format is shown in Exhibit VIII-5. For every segment there should be one Card 2 containing parameters defining the inner, modeled area of the grid. Thus, in the Figure VIII-2, a two-segment scheme, the origin of Segment 1 is defined as Cell (2,2) and its dimensions are 15 x 10. The origin of Segment 2 is defined as Cell (2,12) and its dimensions are also 15 x 10. The file segmentation program adds extra cells as necessary at external boundaries and at segment interfaces, and adjusts the origins and dimensions accordingly. In Figure VIII-3, the segment origins to be specified in the SEGMENTDEF packet are indicated by stars in the grid cells.

Table VIII-1. CONTROL Packet Entries for File Segmentation

<u>Card Number</u>	<u>Entry</u>
4	Number of species Spare Spare Number of segments Spare Spare
5	Spare Print input cards Print regional grid Spare Spare Spare
6	Spare Spare Spare Spare Spare Spare
7	Spare Spare Spare Spare Spare Spare
8	Spare Spare Spare Spare Spare Spare

**Table VIII-2. Contents of SEGMENTDEF Packet Cards**

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<b>1</b>	Packet Header	Header 1-10	60A1	--
	Segment Definition	x-origin 1-10 y-origin 11-20	I10 I10	Grid units x-index of cell in lower left corner of the modeled area of the segment y-index of cell above
	Number of x-cells 21-30	I10	--	Length in x-direction of the modeled area of the segment
	Number of y-cells 31-40	I10	--	Length in y-direction of the modeled area of the segment
<b>2+</b>	Packet Terminator	Terminator 1-3	10A1	--
				Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	1-10	11-20	21-30	31-40	41-50	51-60	<u>FORTRAN Format</u>
1	Packet Header	SEGMENTDEF						60A1
2+	Segment Definition	x-origin	y-origin	x-size	y-size			4110
3	Packet Terminator	END						10A1

Exhibit VIII-5. SEGMENTDEF Packet Format

## 2. The SEGMENT FILE Packet

The SEGMENT FILE packet provides the name of the file to be segmented and the standard input and output units for the unsegmented and segmented files, respectively. The contents of this packet are described in Table VIII-3, and the format is shown in Exhibit VIII-6. The file name on the file designation card must be one of the following:

- > AIRQUALITY
- > BOUNDARY
- > DIFFBREAK
- > EMISSIONS
- > PTSOURCE
- > REGIONTOP
- > ROADWAY
- > TEMPERATUR
- > TERRAIN
- > TOPCONC
- > WIND

The input file unit number must always be 17, and the output file unit number must always be 27.

Exhibit VIII-7 shows sample input to the file segmentation program; Exhibit VIII-8 shows sample output from this program.

Table VIII-3. Contents of SEGMENT FILE Packet Cards

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	60A1	--	Must be SEGMENT FILE, left adjusted
2 File Designation				
	File name 1-10	10A1	--	File name
	Input unit 11-20	I10	--	Unit number of unsegmented file-- must be 17
	Output unit 21-30	I10	--	Unit number of segmented file-- must be 27
3 Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	SEGMENT FILE					
2	File Designation	file name	input unit	output unit			
3	Packet Terminator	END					

FORTRAN Format

60A1	10A1, 2I10	10A1
------	------------	------

Exhibit VIII-6. SEGMENT FILE Packet Format

CONTROL  
SEGMENT DEFINITION  
SEGMENTATION OF ST. LOUIS FILES FOR 4 X 4 GRID

11	0	0	2
1	1		

NO  
NO2  
O3  
ETH  
OLE  
PAR  
CARB  
ARO  
BZA  
PAN  
CO      76195      500      76195      2400  
END  
REGION  
  725000.    4252000.      15  
  -19000.    -16000.  
  4000.    4000.  
  17      22      4  
  END      2      2      0.0      50.  
SEGMENTDEF  
  1      2      2      15      10  
  2      2      12      15      10  
END  
SEGMENT FILE  
TEMPERATUR      17      27  
END

## REGIONAL DEFINITION

UTM ORIGIN X-COORDINATE	725000.
UTM ORIGIN Y-COORDINATE	4252000.
UTM ORIGIN ZONE NUMBER	15
REGION ORIGIN X-COORDINATE	-19000.
REGION ORIGIN Y-COORDINATE	-16000.
CELL LENGTH IN X-DIRECTION	4000.00
CELL LENGTH IN Y-DIRECTION	4000.00
NUMBER OF CELLS IN X-DIRECTION	17
NUMBER OF CELLS IN Y-DIRECTION	22
NUMBER OF CELLS IN Z-DIRECTION	4
NUMBER OF CELLS IN LOWER VERTICAL LAYER	2
NUMBER OF CELLS IN UPPER VERTICAL LAYER	2
HEIGHT OF SURFACE LAYER	.00
MINIMUM CELL HEIGHT IN LOWER LAYER	50.00
MINIMUM CELL HEIGHT IN UPPER LAYER	200.00
KCRNT = 113 NSCLFT = 35798 IN SCRSET	
KCRNT = 115 NSCLFT = 35796 IN SCRSET	
KCRNT = 117 NSCLFT = 35794 IN SCRSET	
KCRNT = 119 NSCLFT = 35792 IN SCRSET	
KCRNT = 130 NSCLFT = 35781 IN SCRSET	
KCRNT = 150 NSCLFT = 35761 IN SCRSET	
KCRNT = 524 NSCLFT = 35387 IN SCRSET	
SEGMENTDEF	
1                   2                   2                   15                   10	
2                   2                   12                  15                  10	
END                   0                   0                   0                   0	

PAGE 1

SEGMENTATION OF ST. LOUIS FILES FOR 4 X 4 GRID

FILE NAME -- SEGMENTDEF	SEGMENTDEF FOR DATA FILE CONTAINING,	SEGMENTDEF VERT.	LEVEL 1	START DATE/TIME 76195 5.00	STOP DATE/TIME 76195 24.00
GRID CELL	1	2	3	4	5
22	0	0	0	0	0
21	0	2	2	2	2
20	0	2	2	2	2
19	0	2	2	2	2
18	0	2	2	2	2
17	0	2	2	2	2
16	0	2	2	2	2
15	0	2	2	2	2
14	0	2	2	2	2
13	0	2	2	2	2
12	0	2	2	2	2
11	1	1	1	1	1
10	1	1	1	1	1
9	0	1	1	1	1
8	0	1	1	1	1
7	1	1	1	1	1
6	6	1	1	1	1
5	5	0	1	1	1
4	4	0	1	1	1
3	3	0	1	1	1
2	2	0	0	1	1
1	1	0	0	0	0

Exhibit VIII-8 (Continued)

## SEGMENT DEFINITION TABLE

SEGMENT NAME	X-ORIGIN CELL	Y-ORIGIN CELL	NO. X-CELLS	NO. Y-CELLS
1	0	0	17	13
2	0	9	17	13

SEGMENT FILE  
TEMPERATUR  
KCRNT = 375 NSCLFT = 35013 IN SCRSET  
KCRNT = 749 HSCLFT = 34639 IN SCRSET  
KCRNT = 1123 NSCLFT = 34265 IN SCRSET  
END 0 0

Exhibit VIII-8 (Continued).

PROCESSING OF INPUT DATA FOR THE PREPARATION OF FILE SEGMENTDEF  
SEGMENTATION OF ST. LOUIS FILES FOR 4 X 4 GRID

NUMBER OF SPECIES = 11  
NUMBER OF SEGMENTS = 2  
INPUT CARDS WILL BE PRINTED  
BEGINNING DATE OF FILE 76195 BEGINNING TIME OF FILE 5.00  
ENDING DATE OF FILE 76195 ENDING TIME OF FILE 24.00  
A TOTAL OF 11 SPECIES NAMES WERE READ.  
THE SPECIES NAMES ARE --  
HO  
NO2  
O3  
ETH  
OLE  
PAR  
CARB  
ARO  
BZA  
PAN  
CO  
REGION  
725000. 4252000. 15  
-190000. -16000.  
4000.00 4000.00  
17 22 4  
2 2 .00 50.00 200.00  
END

## IX PREPARATION OF CONTROL DATA

Of the 14 files input to the Airshed Simulation Program, two are classified as control data files: the Chemistry Parameters file (CHEMPARAM), and the Simulation Controls file (SIMCONTROL). This chapter describes in detail the preparation of these files and includes for each a job control illustration, packet item definitions and packet formats, worksheets, and sample input and output listings.

### A. THE CHEMISTRY PARAMETERS FILE

The Chemistry Parameters (CHEMPARAM) file names the species to be simulated and specifies values for species-related and reaction-related parameters. Figure IX-1 is an information flow diagram for use in setting up the job control for a CHEMPARAM file creation run. The program that creates the CHEMPARAM file is called CPREP; it requires as input a CONTROL packet, a SPECIES packet, and if there is to be chemistry, a REACTIONS packet and a COEFFICIENTS packet (fully described below). The printed output from CPREP consists of input numbers, error messages, if any, and the values written to the CHEMPARAM file (see Section A.5 of this chapter for sample input and output listings). The file itself is written to FORTRAN Unit 2, and the file format is given in Chapter IX of the Systems Manual.

#### 1. CONTROL Packet Rules for the CHEMPARAM File

An abbreviated CONTROL packet is used to create the CHEMPARAM file. The packet names and identifies the file and specifies for it the number of species, reactions, and coefficients. This packet must be entered before any other packet. The contents of each card in the CONTROL packet are described in Table IX-1 and the format is shown in Exhibit IX-1.

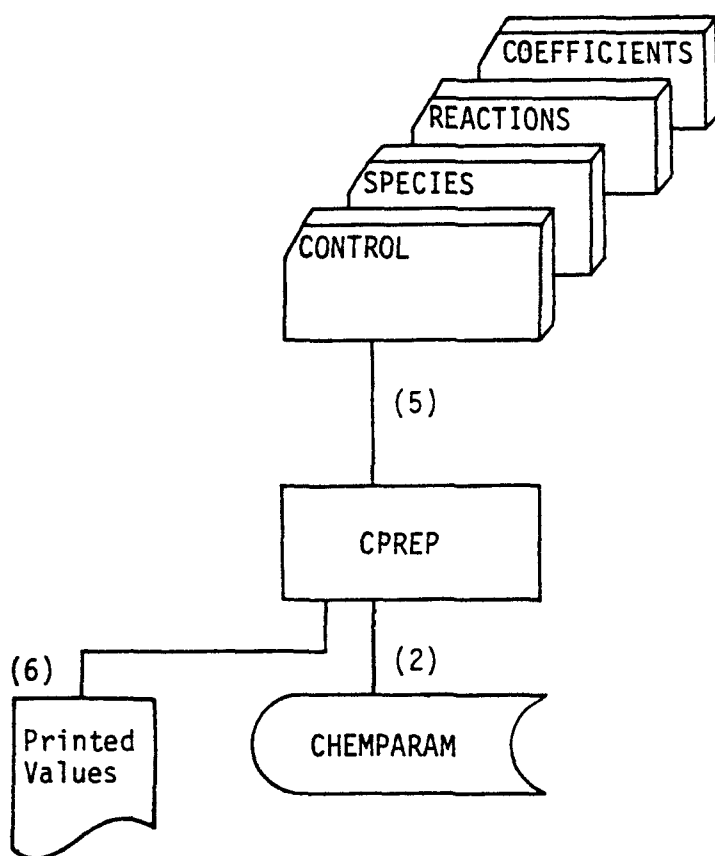


Figure IX-1. Information Flow Diagram for Use in  
Creation of the CHEMPARAM File

Table IX-1. CONTROL Packet for the CHEMPARAM File

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<b>1</b> Packet Header	Header 1-10	10A1	--	Must be CONTROL, left adjusted
<b>2</b> File Name	File name 1-10	10A1	--	Must be CHEMPARAM, left adjusted
<b>3</b> File Identifier	Identifier 1-60	60A1	--	Any 60 characters; the file identifier will appear on the File Description Header Record
<b>4</b> Control Options	Number of species 1-10	I10	--	Must be at least 1; the number of species that will appear in the SPECIES packet and that will thus be included in the airshed simulation; at present, this number must be $\leq 20$
	Number of reactions 11-20	I10	--	The number of reactions that will be described in the REACTIONS packet; for the Carbon-Bond Mechanism, this number should be 69; at present, this number must be $\leq 80$ .
	Number of coefficients 21-30	I10	--	The number of coefficients that will be defined in the COEFFICIENTS packet; if inert species only are to be simulated, set this number to zero and omit the COEFFICIENTS packet; for the Carbon-Bond Mechanism, this number should be 4; at present, this number must be $\leq 10$
<b>5</b> Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>
1	Packet Header
2	File Name
3	File Identifier
4	Control Parameters
5	Packet Terminator

<u>FORTRAN Format</u>	<u>1-10</u>	<u>11-20</u>	<u>21-30</u>	<u>31-40</u>	<u>41-50</u>	<u>51-60</u>
CONTROL						10A1
CHEMPARAM						10A1
# species	# reacs	# coeffs				60A1
END						3I10
						10A1

Exhibit IX-1. CONTROL Packet Format for the CHEMPARAM File

## 2. SPECIES Packet Rules for the CHEMPARAM File

The SPECIES packet must follow the CONTROL packet in the input deck used to create the CHEMPARAM file. The SPECIES packet consists of a packet header card, one pair of Cards 2 and 3 for each species to be simulated, and a packet terminator card. The contents of each card in the SPECIES packet are described in Table IX-2, and the format is shown in Exhibit IX-2.

The chemical species can appear in any order on the CHEMPARAM file, provided that all the reactive species precede all the unreactive species. The output files AVERAGE, INSTANT, and VARIATION will contain these species in this order.

If any reactive species are to be simulated, their names must correspond to the species names used in the chemical mechanism that is built into the program. For the Carbon-Bond II Mechanism (currently built into the program), the following species names are required:

- > NO
- > NO2
- > O3
- > ETH
- > OLE
- > PAR
- > CARB
- > ARO
- > BZA
- > PAN
- > CO.

All the first 10 species (NO through PAN) must appear (though not necessarily in the order listed) and all must be flagged as reactive. CO must

Table IX-2. SPECIES Packet for the CHEMPARAM File

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1	Packet Header	Header 1-10	10A1	--
	Species Name and Flags	Species name 1-10	10A1	--
	Reactive flag	A1, 9X 11	--	An R in Col. 11 indicates that the species is reactive; anything else means unreactive
	Steady-state initial condi- tions flag	9X, L1 30	--	T or F in Col. 30--a blank means false; if true, steady-state computations will be performed in the Airshed Simulation Program to set initial concentrations; at present, these computations are implemented in the Carbon-Bond Mechanism for O <sub>3</sub> only; when computations are performed for O <sub>3</sub> , concentrations of NO and NO <sub>2</sub> are also adjusted
	Steady-state boundary condi- tions flag	9X, L1 40	--	T or F in Col. 40--a blank means false; if true, steady-state computations will be performed at each time slice in the airshed simulation to set boundary concentrations on the vertically varying west, east, south, and north planes (from file BOUNDARY) and on the x-y plane at the top of the region (from file TOPCONC); at present, these computations are implemented in the Carbon-Bond Mechanism for O <sub>3</sub> only; when computations are performed for O <sub>3</sub> , concentrations of NO and NO <sub>2</sub> are also adjusted

Table IX-2 (Concluded)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<sup>3</sup> Other Parameters	Surface uptake resistance 1-10	F10.0	(m/hr) <sup>-1</sup>	To eliminate surface deposition, set this number to a high value, e.g., 1.E6
	Steady-state lower bound 11-20	F10.0	ppm ( $\mu\text{g}/\text{m}^3$ for AEROSOLS)	Computed steady-state concentrations will not be allowed to go below this value, which will also be used for initial or boundary concentrations if this species is missing from an input data file (AIRQUALITY, BOUNDARY, or TOPCONC)
	Steady-state upper bound 21-30	F10.0	ppm ( $\mu\text{g}/\text{m}^3$ for AEROSOLS)	This number must be greater than the steady-state lower bound; computed steady-state concentrations will not be allowed to exceed this value.
Numerical lower bound 31-40	F10.0	ppm ( $\mu\text{g}/\text{m}^3$ for AEROSOLS)	Computed concentrations will not be allowed to go below this value in the horizontal transport steps of the simulation	
Numerical upper bound 41-50	F10.0	ppm ( $\mu\text{g}/\text{m}^3$ for AEROSOLS)	This number must be greater than the numerical lower bound; computed concentrations will not be allowed to exceed this value in the horizontal transport steps of the simulation (at present, this test is not implemented)	
<sup>4</sup> Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	<u>FORTRAN Format</u>
1	Packet Header	10A1
2	Species Name and Flags	10A1, A1, 9X, 2L10
3	Other Species Parameters	5F10.0
4	Packet Terminator	10A1

Exhibit IX-2. SPECIES Packet Format for the CHEMPARAM File

also appear, without a reactive flag, and must follow the listing of all the reactive species. SO<sub>2</sub> and AEROSOLS each can be named as reactive or unreactive, or they can be omitted from the simulation altogether. Any other reactive species are illegal, since there have been no reactions defined to deal with them. However, any other unreactive species can be specified without limitation other than core availability.

The species names on the species-varying data files must also correspond to the names in the CHEMPARAM file. If a name that is not on the CHEMPARAM file appears on a data file, the data for that species will be ignored. If a name is on the CHEMPARAM file but does not appear on a data file, the following default values apply:

- > For concentration data (from AIRQUALITY, BOUNDARY, or TOPCONC), the steady-state lower bound values (see Card 3 of the SPECIES packet) will be used.
- > For emissions data (from EMISSIONS, ROADWAY, or PTSOURCE), the value zero will be used.

Exhibit IX-3(a) is a worksheet that can be used to aid in the preparation of the SPECIES packet. Exhibit IX-3(b) is a sample of that worksheet filled in with typical values used for the Carbon-Bond II Mechanism.

Species	Reactive?	Initial Steady-State Boundary	Steady-State Boundary	Surface Uptake Resistance	Steady-State Lower Bound	Steady-State Upper Bound	Numerical Lower Bound	Numerical Upper Bound
					1-10	21-30	41-50	41-50
Column Format	A1	L10	L10	F10.0	F10.0	F10.0	F10.0	F10.0

(a) Format

Exhibit IX-3. SPECIES Definition Worksheet

Species	Reactive?	Steady-State Initial			Steady-State Boundary			Surface Uptake Resistance			Steady-State			Steady-State		
		11	A1	L10	31-30	L10	F10.0	1-10	F10.0	F10.0	Lower Bound	Upper Bound	Numerical Lower Bound	Numerical Upper Bound	41-50	
Column Format																
NO	R	F	F	F	0.28	1. E-5	1. E+6	1. E-5	1. E-5	1. E+6	1. E-5	1. E+6	1. E-5	1. E+6		
NO2	R	F	F	F	0.011	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4		
O3	R	T	T	T	0.0147	1. E-7	1. E-7	1. E-7	1. E-7	1. E-7	1. E-7	1. E-7	1. E-7	1. E-7		
ETH	R	F	F	F	1. E+10	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4		
OCE	R	F	F	F	1. E+10	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4		
PAR	R	F	F	F	1. E+10	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4		
CARB	R	F	F	F	1. E+10	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4		
ARO	R	F	F	F	1. E+10	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4	1. E-4		
PAN	R	F	F	F	1. E+10	1. E-8	1. E-8	1. E-8	1. E-8	1. E-8	1. E-8	1. E-8	1. E-8	1. E-8		
BZA	R	F	F	F	1. E+10	1. E-8	1. E-8	1. E-8	1. E-8	1. E-8	1. E-8	1. E-8	1. E-8	1. E-8		
AEROSOL	R	F	F	F	1. E+10	2.4.0	2.4.0	2.4.0	2.4.0	2.4.0	2.4.0	2.4.0	2.4.0	2.4.0		
SO2	R	F	F	F	0.0097	1. E-6	1. E-6	1. E-6	1. E-6	1. E-6	1. E-6	1. E-6	1. E-6	1. E-6		
CO	C	F	F	F	1. E+10	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01		

(b) Worksheet Filled In with Typical Values Used for the Carbon-Bond II Mechanism

Exhibit IX-3 (Concluded)

### 3. REACTIONS Packet Rules for the CHEMPARAM File

If the number of reactions specified in the CONTROL packet is greater than zero, there must be a REACTIONS packet following the SPECIES packet. If there are no reactions, the REACTIONS packet must be omitted. This packet consists of a header card, a mechanism name, one card for each reaction, and a packet terminator. The contents of each card in the REACTIONS packet are listed in Table IX-3, and the format is shown in Exhibit IX-4.

For simulation purposes, a reaction is considered to be photolysis, or temperature-dependent, or neither of these. If the photolysis flag is "on" (i.e., a value of P is punched in Column 21), the reaction is photolysis. If the photolysis field is not P and the reaction rate constant and activation energy are nonzero, the reaction is temperature-dependent, and a nonzero value for the reference temperature must be provided. If the reaction rate constant or the activation energy is zero, the reaction is considered to be not temperature dependent. Exhibit IX-5(a) is a worksheet that can be used to aid in the preparation of the REACTIONS packet. Exhibit IX-5(b) is a sample of that worksheet filled in with typical values used for the Carbon-Bond II Mechanism. Since temperature-dependent reaction rates are recalculated with an exponential expression at each time step in each grid cell, it is suggested that the temperature-dependence parameters--activation energy and reference temperatures--be specified for only those reactions for which the temperature range experienced during the simulation period will produce significant differences in predicted concentrations. For the Carbon-Bond Mechanism, the activation energies considered important at present are indicated in the example of Exhibit IX-5(b).

Table IX-3. REACTIONS Packet for the CHAMPARAM File

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
1 Packet Header	Header 1-10	10A1	--	Must be REACTIONS, left adjusted
2 Mechanism Name	Mechanism name 1-60	60A1	--	Any 60-character string; this name will appear in the file and on any simulation runs using the file
3 Reaction Parameters	Reaction number 1-10	I10	--	Must be 1 card for each reaction
				This number is used to place the reaction parameters in their respective vectors, as well as to aid in visual identification; although the scheme described allows reaction parameter cards to appear in any sequence, strict adherence to numerical order is suggested to avoid omission or duplication of reactions
				For photolysis reactions, this number is multiplied by the parameter RADFACTOR on the METSCALARS file (see Chapter VII on METSCALARS); RADFACTOR is the NO <sub>2</sub> photolysis rate constant expressed in min <sup>-1</sup> ; therefore, this number is not a rate constant but rather is a dimensionless factor for relating each photolysis rate constant to the temporally varying NO <sub>2</sub> photolysis rate constant and for converting the rate from min <sup>-1</sup> to hour <sup>-1</sup>
				ppm <sup>-n</sup> hours <sup>-1</sup> * for nonphotolysis reactions
				Nominal reaction rate constant 11-20
			F10.0	

Table IX-3. (Concluded)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
3 (cont'd)				
	Reaction type 21	A1, 9X	--	If Col. 21 is set to P, it will be considered a photolysis reaction
	Activation energy 31-40	F10.0	°K	If this field is left blank, temperature dependence calculations will not be performed
	Reference temperature 41-50	F10.0	°K	This field must be set to nonzero if both the reaction rate and the activation energy are nonzero
4	Packet Terminator Terminator 1-3	10A1	--	Must be END, left adjusted

\* The value of n is determined by the order of the reaction. For example, n takes on values of 0, 1, and 2 for first-, second-, and third-order reactions, respectively.

<u>Card Number</u>	<u>Card Identifier</u>	<u>FORTRAN Format</u>
1	Packet Header	10A1
2	Mechanism Name	60A1
3	Reaction Parameters	I10, F10.0, A1, 9X, 2F10.0
4	Packet Terminator	10A1

Exhibit IX-4. REACTIONS Packet Format for the CHEMPAR11 File

Column Format	Reaction	Number	Rate Constant	Type	Activation Energy	Reference Temperature
		1-10 110	11-20 F10.0	21 A1	31-40 F10.0	41-50 F10.0

(a) Format

Exhibit IX-5. REACTIONS Definition Worksheet

Column Format	Reaction	Number	Rate Constant	Type	Activation Energy		Reference Temperature
					1-10	11-20	
		110	F10.0	A1	F10.0	F10.0	
	$\text{NO}_2 + \text{H}_2 \rightarrow \text{NO} + \text{O}$	1	60.	P			
	$\text{O}_2 + \text{H} \rightarrow \text{O}_3$	2	2.64E + 8				298.
	$\text{O}_3 + \text{NO} \rightarrow \text{NO}_2 + \text{O}_2$	3	1.434E + 3		1450.		
	$\text{O}_3 + \text{NO}_2 \rightarrow \text{NO}_3 + \text{O}_2$	4	2.88		2450.		
	$\text{O} + \text{NO}_2 \rightarrow \text{NO} + \text{O}_2$	5	8.04E + 5				298.
	$\text{O}_3 + \text{OH} \rightarrow \text{HO}_2 + \text{O}_2$	6	4.62E + 3		1000.		
	$\text{O}_3 + \text{HO}_2 \rightarrow \text{OH} + 2\text{O}_2$	7	3.00E + 2		1525.		
	$\text{NO}_2 + \text{OH} \rightarrow \text{HO}_2$	8	8.40E + 5				
	$\text{CO} + \text{OH} \xrightarrow{?} \text{HO}_2 + \text{CO}_2$	9	2.64E + 4				
	$\text{NO} + \text{NO} \xrightarrow{?} 2\text{NO}_2$	10	9.00E - 3				
	$\text{NO}_3 + \text{NO} \rightarrow 2\text{NO}_2$	11	1.68E + 6				

Column Format	Reaction	Number	Rate Constant	Type	Activation Energy	Reference Temperature
		1-10	11-20	21	31-40	41-50
		110	F10.0	A1	F10.0	F10.0
	$\text{NO}_3 + \text{HO}_2 + \text{H}_2\text{O} \rightarrow 2\text{HNO}_3$	12	9.00E - 3		-10600.	298.
	$\text{HO}_2 + \text{NO} \rightarrow \text{NO}_2 + \text{OH}$	13	7.20E + 5			
	$\text{HO}_2 + \text{HO}_2 \rightarrow \text{H}_2\text{O}_2 + \text{O}_2$	14	9.00E + 5			
	$\text{PAR} + \text{O} \rightarrow \text{MEO}_2 + \text{OH}$	15	1.20E + 3			
	$\text{PAR} + \text{OH} \rightarrow \text{MEO}_2$	16	9.00E + 4			
	$\text{OLE} + \text{O} \rightarrow \text{MEO}_2 + \text{ACO}_3 + \text{x}$	17	1.62E + 5			
	$\text{OLE} + \text{O} \rightarrow \text{CARB} + \text{PAR}$	18	1.62E + 5			
	$\text{OLE} + \text{OH} \rightarrow \text{RAO}_2$	19	2.52E + 6			
	$\text{OLE} + \text{O}_3 \rightarrow \text{CARB} + \text{CRIG}$	20	4.80E - 1			
	$\text{OLE} + \text{O}_3 \rightarrow \text{CARR} + \text{MCRG} + \text{x}$	21	4.80E - 1			
	$\text{ETH} + \text{O} \rightarrow \text{MEO}_2 + \text{HO}_2 + \text{CO}$	22	3.60E + 4			
	$\text{ETH} + \text{O} \rightarrow \text{CARR} + \text{PAR}$	23	3.60E + 4			
	$\text{ETH} + \text{OH} \rightarrow \text{RB}\text{O}_2$	24	7.20E + 5			

Column Format	Reaction	Number	Rate Constant	Type	Activation Energy		Reference Temperature
					1-10	11-20	
	ETH + O <sub>3</sub> + CARB + CRIG	25	1.44E - 1				
	ACO <sub>3</sub> + NO + NO <sub>2</sub> + ME0 <sub>2</sub> + CO <sub>2</sub>	26	2.28E + 5				
	RB0 <sub>2</sub> + NO + NO <sub>2</sub> + 2 CARB + HO <sub>2</sub>	27	7.20E + 5				
	RA0 <sub>2</sub> + NO + NO <sub>2</sub> + 2 CARB + HO <sub>2</sub>	28	7.20E + 5				
	ME0 <sub>2</sub> + NO + NO <sub>2</sub> + CARB + ME0 <sub>2</sub> + X.	29	2.30E + 5				
	ME0 <sub>2</sub> + NO + NO <sub>2</sub> + CARB + HO <sub>2</sub>	30	4.60E + 5				
	ME0 <sub>2</sub> + NO + Nitrates	31	3.00E + 4				
	RB0 <sub>2</sub> + O <sub>3</sub> + 2 CARB + HO <sub>2</sub>	32	3.00E + 2				
	RA0 <sub>2</sub> + O <sub>3</sub> + 2 CARB + HO <sub>2</sub>	33	1.20E + 4				
	ME0 <sub>2</sub> + O <sub>3</sub> + CARB + HO <sub>2</sub>	34	3.00E + 2				
	CARB + OH + a (HO <sub>2</sub> + CO) + (1 - a) (ACO <sub>3</sub> + X)	35	1.14E + 6				
	CARB + hv + CO	36	1.35E - 3	P			
	CARB + (1 + a) NO <sub>2</sub> + (1 - a) (ME0 <sub>2</sub> + X) + CO	37	1.215E - 1	P			

Column Format	Reaction	Number	Rate Constant	Type	Activation Energy	Reference Temperature
		1-10	11-20	21	31-40	41-50
		110	F10.0	A1	F10.0	F10.0
	X + PAR + Products	38	6.00E + 6			
	ACO <sub>3</sub> + NO <sub>2</sub> + PAN	39	1.20E + 5			
	PAN + ACO <sub>3</sub> + NO <sub>2</sub>	40	1.68E + 0		12500.	298.
	ACO <sub>3</sub> + NO <sub>2</sub> + Products	41	2.40E + 5			
	NO <sub>2</sub> + HO <sub>2</sub> + Products	42	2.40E + 5			
	CRIG + NO + NO <sub>2</sub> + CARB	43	7.20E + 5			
	CRIG + NO <sub>2</sub> + NO <sub>3</sub> + CARB	44	4.80E + 5			
	CRIG + CARB + Ozonide	45	1.20E + 5			
	MCRIG + NO + NO <sub>2</sub> + CARB + PAR	46	7.20E + 5			
	MCRIG + NO <sub>2</sub> + NO <sub>3</sub> + CARR + PAR	47	4.80E + 5			
	MCRIG + CARB + Ozonide	48	2.00E + 3			
	CRIG + CO	49	4.02E + 4			
	CRIG + Products	50	1.44E + 4			

Column Format	Reaction	Number	Rate Constant	Type	Activation Energy	Reference Temperature
	CRIG + 2HO <sub>2</sub> + CO <sub>2</sub>	51	5.40E + 3			
	MCRG + Products	52	9.00E + 3			
	MCRG + ME0 <sub>2</sub> + OH + CO	53	2.04E + 4			
	MCRG + ME0 <sub>2</sub> + HO <sub>2</sub> + CO <sub>2</sub>	54	2.55E + 4			
	MCRG + CARB + 2HO <sub>2</sub> + CO	55	5.10E + 3			
	ARO + OH + HO <sub>2</sub> + 3 ARPI	56	3.60E + 5			
	ARO + OH + HO <sub>2</sub> + BZA + X	57	9.60E + 4			
	ARO + OH + BZA + W	58	9.00E + 5			
	CARB + W + Products	59	6.00E + 6			
	NO + ARPI + PAR + CARB + NO	60	1.80E + 3			
	NO + ARPI + NO <sub>2</sub> + Aerosol	61	9.00E + 2			
	NO <sub>3</sub> + ARPI + 2 CARB	62	2.10E + 6			

Column Format	Reaction	Number	Rate Constant	Type	Activation Energy	Reference Temperature
	$O_3 + APR1 + Aerosol$	63			$3.60E + 1$	
	$BZA + OH + HO_2 + CO + 3 APR1$	64			$6.00E + 5$	
	$BZA + NO_2 + HO_2 + 3 APR1$	65		P	$2.16E + 0$	
	$SO_2 + OH \xrightarrow{O_3} HSO_5$	66			$5.40E + 1$	
	$HSO_5 + NO + NO_2 + HSO_4$	67			$4.80E + 4$	
	$HSO_4 + NO_2 + H_2SO_4 + HNO_3$	68			$5.40E + 5$	
	$SO_2 + Aerosol$	69			$3.00E - 2$	

#### **4. COEFFICIENTS Packet Rules for the CHEMPARAM File**

If the number of coefficients specified in the CONTROL packet is greater than zero, there must be a COEFFICIENTS packet, and it must follow the REACTIONS packet (or the SPECIES packet, if there is no REACTIONS packet). If there are no coefficients, this packet must be omitted. The COEFFICIENTS packet consists of a header card, one card for each coefficient, and a packet terminator. The contents of each card are described in Table IX-4, and the format is shown in Exhibit IX-6.

Exhibit IX-7(a) is a worksheet that can be used to aid in the preparation of the COEFFICIENTS packet. Exhibit IX-7(b) is a sample of that worksheet filled in with typical values used for the Carbon-Bond II Mechanism.

**Table IX-4. COEFFICIENTS Packet for the CHEMPARAM File**

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<b>1</b> Packet Header	Header 1-10	10A1	--	Must be COEFFICIENTS, left adjusted
<b>2</b> Coefficient	Coefficient name 1-10	10A1	--	10 characters, left adjusted; must be identical to a coefficient name required by the built-in chemical mechanism
	Coefficient value 11-20	F10.0	--	
<b>3</b> Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	<u>Column Number</u>					<u>FORTRAN Format</u>
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	COEFFICIENTS					10A1
2	Coefficients	Name					10A1,
3	Packet Terminator	Value					F10.0
		END					

Exhibit IX-6. COEFFICIENTS Packet Format for the CHEMPARAM File

Meaning	Name	Value
Column	1-10	11-20
Format	10A1	F10.0

(a) Format

Exhibit IX-7. COEFFICIENTS Definition Worksheet

Column	Name	Value
Meaning		
Format	1-10	11-20
	10A1	F10.0
	<i>ALPHA</i>	0.5
	<i>BETA</i>	.48
	<i> GAMMA</i>	10
	<i> ETA</i>	0.1

(b) Worksheet Filled In with Typical Values Used for the Carbon-Bond Mechanism

Exhibit IX-7 (Concluded)

5. CPREP Sample Input and Output

Exhibit IX-8 is a listing of an input deck used to create a CHEMPARAM file; it contains the values from the sample worksheets [Exhibits IX-3(b), IX-5(b), and IX-7(b)]. Exhibit IX-9 is the printout that was generated by CPREP; the first two pages list the values on the input cards as they are read in, and the last two show the values put out to the CHEMPARAM file.

```

CONTROL
CHEMPARAM
  CHEMISTRY PARAMETERS NEW CHEM HIGHER PARAFINS LOWER LIMIT
    11      6.9        4
END
SPECIES
HO      R      F      F      1.0E+6      1.0E-5      1.0E+6
NO2     R      F      F      1.E+6      1.E-4      1.E+6
O3      R      T      T      1.E+6      1.E-9      1.E+6
ETH      R      F      F      1.E+6      1.E-9      1.E+6
OLE      R      F      F      1.E+6      1.E-6      1.E+6
PAR      R      F      F      1.E+6      1.E-6      1.E+6
CARB     R      F      F      1.E+6      1.E-4      1.E+6
ARO      R      F      F      1.E+6      1.E-6      1.E+6
BZA      R      F      F      1.E+6      1.E-6      1.E+6
PAN      R      F      F      1.E+6      1.E-6      1.E+6
CO       R      F      F      1.E+6      1.E-6      1.E+6
C0      1.E10   .01    .01    1.E+6      .01    1.E+6
END
REACTIONS
CARBON BOND MECHANISM -- NOVEMBER 78
  1 60.00E+0P
  2 264.E+6
  3 1434.
  4 288.E-2
  5 80.4E+4
  6 462.E+1
  7 300.
  8 84.E+4
  9 264.E+2
 10 90.E-4
 11 168.E+4
 12 9.0E-3
 13 72.E+4
 14 90.E+4
 15 120.E+1
 16 90.E+3
 17 162.E+3
 18 162.E+3
 19 252.E+4
 20 480.E-3
 21 480.E-3
 22 360.E+2
 23 360.E+2
 24 72.E+4

```

25	144.E-3
26	228.E+3
27	72.E+4
28	72.E+4
29	2.3E+5
30	4.6E+5
31	300.E+2
32	300.
33	120.E+2
34	300.
35	114.E+4
36	81.E-3P
37	121.5E-3P
38	60.E+5
39	120.E+3
40	168.E-2
41	240.E+3
42	240.E+3
43	72.E+4
44	480.E+3
45	120.E+3
46	72.E+4
47	480.E+3
48	120.E+3
49	402.E+2
50	144.E+2
51	540.E+1
52	90.E+2
53	204.E+2
54	255.E+2
55	510.E+1
56	3.60E+5
57	9.6E+4
58	9.0E+5
59	6.0E+6
60	1.0E+3
61	9.0E+2
62	2.1E+6
63	3.6E+1
64	6.0E+5
65	2.160E+0P
66	540.00E+2
67	480.00E+2
68	540.00E+3
69	3.00E-2

END  
**COEFFICIENTS**  
 ALPHA      0.5  
 BETA        .48  
 GAMMA      1.0  
 ETA        0.1  
 END

CONTROL  
 CHEMPARAM  
 CHEMISTRY PARAMETERS NEW CHEM HIGHER PARAFFINS LOWER LIMIT  
 11 69 4  
 END  
 SPECIES NO R F F F F  
 NO2 R 2.8000-001 1.0000-005 1.0000+006 1.0000-005 1.0000+006  
 NO2 R 1.1100-002 1.0000-004 1.0000+006 1.0000-004 1.0000+006  
 O3 R 1.4700-002 1.0000-009 T T 1.0000+006 1.0000-009 1.0000+006  
 ETH R 1.0000+010 F F F F  
 OLE R 1.0000+010 1.0000-006 1.0000+006 1.0000-006 1.0000+006  
 PAR R 1.0000+010 1.0000-006 1.0000+006 1.0000-006 1.0000+006  
 CARB R 1.0000+010 1.0000-003 F F F F  
 APO R 1.0000+010 1.0000-006 1.0000+006 1.0000-006 1.0000+006  
 BZA R 1.0000+010 1.0000-006 1.0000+006 1.0000-006 1.0000+006  
 PAN R 1.0000+010 1.0000-006 F F F F  
 CO R 1.0000+010 1.0000-002 F F F F  
 END  
 REACTIONS  
 CARBON BOND MECHANISM -- NOVEMBER 78  
 1 6.0000+001 P .0000 .0000  
 2 2.6400+008 .0000 .0000  
 3 1.4340+003 1.4500+003 2.9800+002  
 4 2.8800+000 2.4500+003 2.2800+002  
 5 8.0400+005 .0000 .0000  
 6 4.6200+003 1.0000-003 2.9300+002  
 7 3.0000+002 1.5250+003 2.9500+002  
 8 8.4000+002 .0000 .0000  
 9 2.6400+004 .0000 .0000  
 10 9.0000-003 .0000 .0000  
 11 1.6300+006 .0000 .0000  
 12 9.0000-003 -1.0600+004 2.9800+002  
 13 7.2000+005 .0000 .0000  
 14 9.0000+005 .0000 .0000  
 15 1.2000+003 .0000 .0000  
 16 9.0000+004 .0000 .0000  
 17 1.6200+005 .0000 .0000  
 18 1.6200+005 .0000 .0000  
 19 2.5200+006 .0000 .0000  
 20 4.8000-001 .0000 .0000  
 21 4.8000-001 .0000 .0000  
 22 3.6000+004 .0000 .0000  
 23 3.6000+004 .0000 .0000  
 24 7.2000+005 .0079 .0000  
 25 1.4400-001 .0000 .0000

ETID	COEFFICIENT	ALPHA	BETA	GAMMA	ETA
26	2.2800+005	.0000	.0000	.0000	.0000
27	7.2000+005	.0000	.0000	.0000	.0000
28	7.2000+005	.0000	.0000	.0000	.0000
29	2.3000+005	.0000	.0000	.0000	.0000
30	4.6000+005	.0000	.0000	.0000	.0000
31	3.0000+004	.0000	.0000	.0000	.0000
32	3.0000+002	.0000	.0000	.0000	.0000
33	1.2000+004	.0000	.0000	.0000	.0000
34	3.0000+002	.0000	.0000	.0000	.0000
35	1.1400+006	.0000	.0000	.0000	.0000
36	8.1000-002	P	.0000	.0000	.0000
37	1.2150-001	P	.0000	.0000	.0000
38	6.0000+006	.0000	.0000	.0000	.0000
39	1.2000+005	.0000	.0000	.0000	.0000
40	1.6600+000	.0000	.0000	.0000	.0000
41	2.4000+005	.0000	.0000	.0000	.0000
42	2.4000+005	.0000	.0000	.0000	.0000
43	7.2000+005	.0000	.0000	.0000	.0000
44	4.8000+005	.0000	.0000	.0000	.0000
45	1.2000+005	.0000	.0000	.0000	.0000
46	7.2000+005	.0000	.0000	.0000	.0000
47	4.8000+005	.0000	.0000	.0000	.0000
48	1.2000+005	.0000	.0000	.0000	.0000
49	4.0200+004	.0000	.0000	.0000	.0000
50	1.4400+004	.0000	.0000	.0000	.0000
51	5.4000+003	.0000	.0000	.0000	.0000
52	9.0050+003	.0000	.0000	.0000	.0000
53	2.0400+004	.0000	.0000	.0000	.0000
54	2.5500+004	.0000	.0000	.0000	.0000
55	5.1000+003	.0000	.0000	.0000	.0000
56	3.6000+005	.0000	.0000	.0000	.0000
57	9.6000+004	.0000	.0000	.0000	.0000
58	9.0000+005	.0000	.0000	.0000	.0000
59	6.0000+006	.0000	.0000	.0000	.0000
60	1.0000+003	.0000	.0000	.0000	.0000
61	9.0000+002	.0000	.0000	.0000	.0000
62	2.1000+006	.0000	.0000	.0000	.0000
63	3.6000+001	.0000	.0000	.0000	.0000
64	6.0000+005	.0000	.0000	.0000	.0000
65	2.1600+000	P	.0000	.0000	.0000
66	5.4000+004	.0000	.0000	.0000	.0000
67	4.0000+004	.0000	.0000	.0000	.0000
68	5.4000+005	.0000	.0000	.0000	.0000
69	3.0000-002	.0000	.0000	.0000	.0000

Exhibit IX-9 (Continued)

## \*\*\*\*\* CHEMPARAM FILE \*\*\*\*\*

## FILE DESCRIPTION HEADER RECORD

FILE TYPE = CHEMPARAM  
 FILE ID = CHEMPARAM  
 NO OF SEGMENTS = 0  
 NO OF SPECIES = 11  
 BEG DATE = 0.  
 BEG TIME = 0.  
 END DATE = 0.  
 END TIME = 0.

## SPECIES DESCRIPTION HEADER RECORD

HO  
 NO2  
 O3  
 ETH  
 OLE  
 PAR  
 CARB  
 ARO  
 BZA  
 PAN  
 CO

## CHEMISTRY DEFINITION RECORD

MECHANISM ID = CARBON BOND MECHANISM -- NOVEMBER 78  
 NO OF REACTIONS = 69  
 NO OF COEFFS = 4

## SPECIES PARAMETERS RECORD

SPEC NAME	REACT	SSIC	SSBC	DVRES	LOW BD SS	UP BD SS	LOW BD NUM	UP BD NUM
HO	T	F	F	2.8000+001	1.0000+005	1.0000+006	1.0000+005	1.0000+006
NO2	T	F	F	1.1100+002	1.0000+004	1.0000+006	1.0000+004	1.0000+006
O3	T	T	T	1.4700+002	1.0000+009	1.0000+006	1.0000+009	1.0000+006
ETH	T	F	F	1.0000+010	1.0000+006	1.0000+006	1.0000+006	1.0000+006
OLE	T	F	F	1.0000+010	1.0000+006	1.0000+006	1.0000+006	1.0000+006
PAR	T	F	F	1.0000+010	1.0000+003	1.0000+006	1.0000+004	1.0000+006
CARB	T	F	F	1.0000+010	1.0000+006	1.0000+006	1.0000+006	1.0000+006
ARO	T	F	F	1.0000+010	1.0000+006	1.0000+006	1.0000+006	1.0000+006
BZA	T	F	F	1.0000+010	1.0000+006	1.0000+006	1.0000+006	1.0000+006
PAN	T	F	F	1.0000+010	1.0000+006	1.0000+006	1.0000+006	1.0000+006
CO	T	F	F	1.0000+010	1.0000+002	1.0000+006	1.0000+002	1.0000+006

## REACTION PARAMETERS RECORD

RATE CONST	PHOTOL	TEMP	ACT ENERGY	REF TEMP
1 6.0000+001	T	F	.0000	.0000
2 2.6400+008	F	F	.0000	.0000
3 1.4340+003	F	T	1.4500+003	2.9600+002
4 2.2800+000	F	T	2.4500+003	2.9600+002
5 8.0400+005	F	F	.0000	.0000
6 4.6200+003	F	T	1.0000+003	2.9600+002
7 3.0000+002	F	T	1.5250+003	2.9600+002
8 8.6000+005	F	F	.0000	.0000
9 2.6100+004	F	F	.0000	.0000

10	9.0000-003	.0000
11	1.6000+006	.0000
12	9.0000-003	-1.0600+002
13	7.2000+005	2.9800+004
14	9.0000+005	.0000
15	1.2000+003	.0000
16	9.0000+004	.0000
17	1.6200+005	.0000
18	1.6200+005	.0000
19	2.5200+006	.0000
20	7.2000+005	.0000
21	4.8000-001	.0000
22	3.6000+004	.0000
23	3.6000+004	.0000
24	2.3000+005	.0000
25	1.4400-001	.0000
26	2.2800+005	.0000
27	7.2000+005	.0000
28	7.2000+005	.0000
29	2.3000+005	.0000
30	4.6000+005	.0000
31	3.0000+004	.0000
32	3.0000+002	.0000
33	1.2000+004	.0000
34	3.0000+002	.0000
35	1.1400+006	.0000
36	8.1000-002	.0000
37	1.2150-001	.0000
38	6.0000+006	.0000
39	1.2000+005	.0000
40	1.6800+000	1.2500+004
41	2.4000+005	2.9800+002
42	2.4000+005	.0000
43	7.2000+005	.0000
44	4.6000+005	.0000
45	1.2000+005	.0000
46	7.2000+005	.0000
47	4.8000+005	.0000
48	1.2000+005	.0000
49	4.0200+004	.0000
50	1.4400+004	.0000
51	5.4000+003	.0000
52	9.0000+003	.0000
53	2.0400+004	.0000
54	2.5500+004	.0000
55	5.1000+003	.0000
56	3.6000+005	.0000
57	9.6000+004	.0000
58	9.0000+005	.0000
59	6.0000+006	.0000
60	1.8000+003	.0000

Exhibit IX-9 (Continued)

**Exhibit IX-9 (Concluded)**

61	9.0000+002	F	F	.0000
62	2.1000+006	F	F	.0000
63	3.6000+001	F	F	.0000
64	6.0000+005	F	F	.0000
65	2.1600+000	T	F	.0000
66	5.4000+004	F	F	.0000
67	4.8000+004	F	F	.0000
68	5.4000+005	F	F	.0000
69	3.0000-002	F	F	.0000

**STOICHIOMETRIC COEFFICIENTS RECORD**

ALPHA	5.0000-001
BETA	4.8000-001
Gamma	1.0000-000
ETA	1.0000-001

## B. THE SIMULATION CONTROLS FILE

The Simulation Controls (SIMCONTROL) file, generally the last to be prepared, is the input file likely to be varied the most frequently. As previously mentioned, this file can routinely be prepared as a separate job step within each airshed simulation run.

Figure IX-2 is an information flow diagram for use in setting up the job control for a SIMCONTROL file creation run. The program that creates the SIMCONTROL file is called SPREP. It requires as input a CONTROL packet and a SIMULATION packet, which are described in detail in the next section. The printed output from SPREP consists of input numbers, error messages, if any, and the values written to the SIMCONTROL file (see Section B.3 of this chapter for sample input and output listings). The file itself is written to FORTRAN Unit 1, and the file format is given in Chapter IX of the Systems Manual.

### 1. CONTROL Packet Rules for the SIMCONTROL File

An abbreviated CONTROL packet is used to create the SIMCONTROL file. The packet names and identifies the type of file to be created, and it must be entered before any other packet. The contents of each card in the CONTROL packet are described in Table IX-5, and the format is shown in Exhibit IX-10.

### 2. SIMULATION Packet Rules for the SIMCONTROL File

The SIMULATION packet must follow the CONTROL packet in the input deck used to create the SIMCONTROL file. The contents of each card in the SIMULATION packet are described in Table IX-6 and the format is shown in Exhibit IX-11.

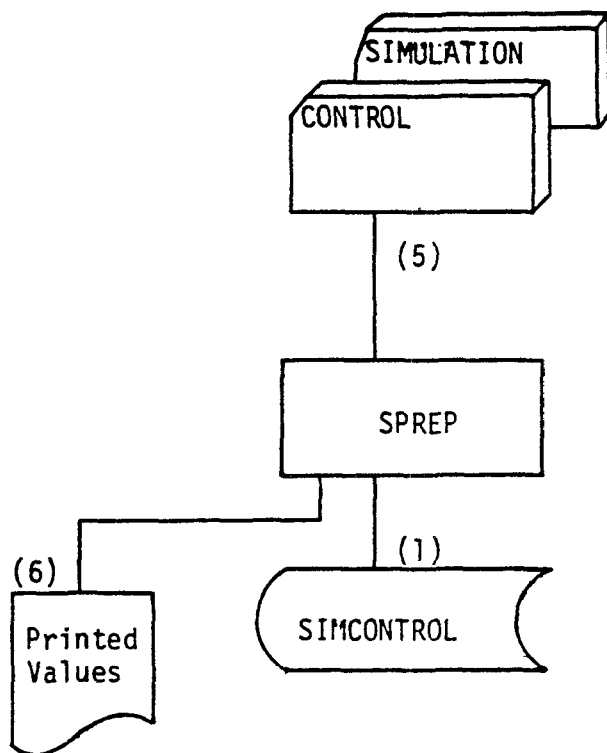


Figure IX-2. Information Flow Diagram for Use in  
Creation of the SIMCONTROL File

**Table IX-5. CONTROL Packet for the SIMCONTROL File**

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<b>1</b> Packet Header	Header 1-10	10A1	--	Must be CONTROL, left adjusted
<b>2</b> File Name	File name 1-10	10A1	--	Must be SIMCONTROL, left adjusted
<b>3</b> File Identifier	Identifier 1-60	60A1	--	Any 60 characters; the file identifier will appear on the File Description Header Record
<b>4</b> Packet Terminator	Terminator 1-3	10A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	<u>Column Number</u>				<u>FORTRAN Format</u>	
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	CONTROL					10A1
2	File Name	SIMCONTROL					10A1
3	File Identifier						60A1
4	Packet Terminator	END					10A1

Exhibit IX-10. CONTROL Packet Format for the SIMCONTROL File

Table IX-6. SIMULATION Packet for the SIMCONTROL File

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
<sup>1</sup> Packet Header	Header 1-10	10A1	--	Must be SIMULATION, left adjusted
<sup>2</sup> File Identifier	Identifier 1-60	60A1	--	Any 60-characters; to provide a unique identifier, change this card for each run
<sup>3</sup> <b>Time Limits of the Run</b>				
Beginning date 1-10		I10	Julian date (yyddd)	Beginning date and time must appear within a time interval on the AIRQUALITY file, and the entire time span shown here must be included within the time span of every other time-varying data file; when restarting a run from an intermediate stopping point, these time limits must be changed
Beginning time 11-20		I10	Military time (hhmm)	yy is the last two digits of the year and ddd is the three-digit Julian date hh is hours and mm is minutes--this time will be internally converted to hours and fraction
End date 21-30		I10	Julian date (yyddd)	Same format as beginning date
End time 31-40		I10	Military time (hhmm)	Same format as beginning time
<sup>4</sup> <b>Simulation Options</b>				
Restart flag 1-10		L10	--	For each flag; .TRUE. means "yes, do it," and .FALSE. means "no, don't do it" Set to TRUE if this is a restart; the simulation program will not perform steady-state computations on initial concentrations if the restart flag is TRUE

Table IX-6 (Continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
4 (cont'd)	Surface removal flag 11-20	L10	--	Set to TRUE if the effects of pollutant uptake at the surface are to be calculated; when the surface removal flag is TRUE, there must be either a TERRAIN file or a nonzero vegetation factor specified on Card 6 of this packet
	PTSOURCE flag 21-30	L10	--	Set to TRUE if elevated point source emissions from a PTSOURCE file are to be included in the simulation; if flag is TRUE, there must be a PTSOURCE file; if it is FALSE, the PTSOURCE file, if present, will be ignored
	ROADWAY flag 31-40	L10	--	Set to TRUE if there is a ROADWAY file; flag may be reset to FALSE by the simulation program if there is no surface layer or if no reactive species are being simulated; if flag is TRUE, there must be a ROADWAY file; if flag is set to FALSE or reset to FALSE under the conditions described above, the ROADWAY file, if present, will be ignored. ( <u>use FALSE</u> ).
	TEMPERATUR flag 41-50	L10	--	Set to TRUE if there is a TEMPERATUR file; if flag is TRUE, there must be a TEMPERATUR file; if it is FALSE, the TEMPERATUR file, if present, will be ignored; when flag is FALSE, an ambient temperature of 25°C will be assumed, and the nominal input values of the reaction rate constants will be employed in the chemistry calculations (i.e., temperature effects will be ignored)

Table IX-6 (Continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
4 (cont'd)	TERRAIN flag 51-60	L10	--	Set to TRUE if there is a TERRAIN file; if flag is TRUE, the TERRAIN file must be included; if it is false, the TERRAIN file, if present, will be ignored; when flag is FALSE, a default value for surface roughness must be specified on Card 6 of this packet, and a default vegetation factor must also be specified on Card 6 if surface uptake effects are to be calculated
5	More Simulation Options	Concentration variation flag 1-10	L10	Set to TRUE if subgrid scale concentration variation is to be calculated; if flag is TRUE, concentration variation will be calculated at each time step and averaged over the same period that concentrations are averaged; the average concentration variations will be output on the VARIATION file; the flag will be reset to FALSE in the program if there is no surface layer or no ROADWAY file
6	Default Terrain Values	Default surface roughness 1-10	F10.0	m Must be included whether or not the values are required This value will be used to calculate the vertical diffusivity if the TERRAIN file is not input to the run; if the TERRAIN file is input, this value will be ignored

Table IX-6 (Continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
6 (cont'd)	Default vegeta- tion factor 11-20	F10.0	--	Pollutant uptake capability of the surface relative to that of alfalfa; this value will be used to calculate surface deposition if the TERRAIN file is not input to the run; if the TERRAIN file is input or if surface removal effects are not to be calculated, this value will be ignored
7 Integration Controls	Maximum time slice 1-10	F10.0	hr	The largest simulation time interval during which the boundary conditions at the segment interfaces will be held constant; this parameter is used with the next one to determine the time slice size used in the Airshed Simulation Program (see Chapter IV); if the maximum slice is set very large (i.e., 1. or more), the calculated time slice will be limited by the maximum number of steps in a slice (or, if the time steps are very large, by the output time intervals or the intervals on the input files); if the maximum slice is set small relative to the typical time step calculated, it will be the limiting factor in computing the time slice size to be used
	Maximum number of steps in a slice 11-20	I10	--	This parameter is used with the previous one to calculate the time slice size used in the Airshed Simulation Program; if this number is set very large, the slice will be limited by the maximum time slice size; if the number is small, the slice will be limited by

Table IX-6 (Continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
7 (cont'd)				the maximum number of the smallest time steps; if this number is set to 1, the boundary conditions at the segment interfaces will be updated at the end of every time step
8	Chemistry Integration Controls			Provides a lower limit on the chemistry step size; the overall time step size for each segment will be determined by the maximum wind speed in the segment; within each cell, however, the time step can be further reduced for the chemistry step; if the chemistry step fails to converge or gives negative concentrations at this step size, the program will terminate
	Minimum chemistry time step	F10.0 1-10	hr	If the chemistry step does not converge within this number of iterations for any cell at a given time step, the chemistry step size for that cell will be halved and the iterative procedure will begin again
	Maximum number of iterations	I10 11-20	--	Convergence criterion for the chemistry step--when the relative change in concentration for all species is less than this number, the iterative procedure is said to have converged; if the relative change in concentration for any species is greater than this number, another iteration will be performed
	Relative error tolerance	F10.0 21-30	--	

Table IX-6 (Continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
8 (cont'd)	Darkness criterion 31-40	F10.0	Same as for RADFACTOR on the METSCALARS file	If the calculated RADFACTOR, which varies continuously in time, is less than this number, night chemistry will be used; if the calculated RADFACTOR is greater than this number, the normal (daytime) chemistry will be used; if the simulation is performed on inert species only or if the night chemistry mechanism has been disabled, this number is ignored
9	Instantaneous output interval 1-10	F10.0	hr	Instantaneous concentrations are written to the output file INSTANT at the beginning of the simulation and at this time interval thereafter; if the end of a time slice will go past the output time, the slice will be reduced to hit the output time exactly; instantaneous concentrations are also written when time-averaged concentrations are written, enabling a restart from the beginning of any averaging period
	Average output intervals 11-20	F10.0	hr	Starting at the beginning of the simulation, concentrations will be averaged over this interval and written to the AVERAGE file at the end of the interval; if the end of a time slice will go past the end of an averaging interval, the slice will be reduced to correspond to the end of the interval exactly

Table IX-6 (Continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
10 Print Options				Six options are read, but only five are used If this field is set to =1, a one-line message will be printed each time a time interval record is read on any data file; if set to blank or zero, the file read messages will not be printed
	Print file history 1-10	I10	--	If this field is set to =1, all storage allocation pointers and controls will be printed; this information is useful only for program debugging; if set to blank or zero, the information will not be printed
	Print storage allocation 11-20	I10	--	If set positive, this is the number of vertical levels of concentrations that will be printed at the end of each instantaneous output interval (see Card 9); the boundary cells printed may contain irrelevant information and, therefore, the printouts should be used for debugging only; concentration maps will be printed for each segment, for each species, for each level up to the number specified, but not exceeding the number of levels modeled; if set to blank or zero, instantaneous concentrations will not be printed
	Print instantaneous concentrations 21-30	I10	--	If set positive, this is the number of vertical levels of average concentrations that will be printed at the end of each averaging interval (see Card 9); format and order of the average concentration printout is the same as that of the instantaneous concentration option; if set to blank or zero, average concentrations will not be printed
	Print average concentrations 31-40	I10	--	

Table IX-6 (Concluded)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Units</u>	<u>Comments</u>
10 (cont'd)	Print concentra- tion variations 41-50	I10	--	If set positive, concentration variations will be printed after instantaneous and after average concentrations as selected above. If set to blank or zero, if both options above are zero, or if concentra- tion variations are not calculated, con- centration variations are not printed
11	Packet Terminator Terminator 1-3	I0A1	--	Must be END, left adjusted

<u>Card Number</u>	<u>Card Identifier</u>	<u>Column Number</u>				<u>FORTRAN Format</u>	
		1-10	11-20	21-30	31-40	41-50	51-60
1	Packet Header	SIMULATION					10A1
2	Run Identification						60A1
3	Time Limits of the Run	beg date	beg time	end date	end time		4I10
4	Simulation Options	restart	surface removal	pt srces	roadway	temp	6L10
5	More Simulation Options	conc vartn				terrain	
6	Default Terrain Values	surface roughness	vegetation				
7	Integration Time Interval Controls	max time slice	max # steps in slice				F10.0, I10
8	Chemistry Integration Controls	min chem step	max # iter to	rel error	dark crit		F10.0, I10
9	Output intervals	instant	average		conc vartn		2F10.0
10	Print Options	file history	storage allocation	instant conc	average conc		6I10
11	Packet Terminator	END					10A1

3. SPREP Sample Input and Output

Exhibit IX-12 is a listing of an input deck used to create a SIMCONTROL file. Exhibit IX-13 is the printout that was generated by SPREP; it first lists the values on the input cards as they are read in, and then shows these values put out to the SIMCONTROL file.

```
CONTROL  
SIMCONTROL  
RAPS TEST RUN CONTROLS FOR JUNE-JULY 1978  
END  
SIMULATION  
RAPS 4-KM DAY 195 -- 4-HR RUN 07/13/76 C-BOND II 6-79  
76195 0500 76195 0900  
F T T F T T  
.5 1.  
.1 1.  
.003 20 .01 .01  
1. 1. 1 1  
END
```

Exhibit IX-12. Sample SPREP Input for the SIMCONTROL File

```
CONTROL
SIMCONTROL
RAPS TEST RUN CONTROLS FOR JUNE-JULY 1978
END
SIMULATION
RAPS 4-KM DAY 195 -- 4-HR RUN 07/13/76 C-BOND II 8-79
76195   500    76195   900
          F      T      F      T      T
          F
          F
          5.0000-001  1.0000+000
          1.0000-001  1
          3.0000-003  20
          1.0000+000  1.0000-002  1.0000-002
          1           1           1           0           0
END
```

Exhibit IX-13. Sample SPREP Output for  
the SIMCONTROL File

\*\*\*\*\* SIMCONTROL FILE \*\*\*\*\*

FILE DESCRIPTION HEADER RECORD  
FILE TYPE = SIMCONTROL  
FILE ID = RAPS TEST RUN CONTROLS FOR JUNE-JULY 1976  
NO OF SEGMENTS = 0  
NO OF SPECIES = 0  
BEG DATE = 0  
BEG TIME = 0.  
END DATE = 0  
END TIME = 0.

SIMULATION CONTROLS RECORD  
RUN ID = RAPS 4-KM DAY 195 -- 4-HR RUN 07/13/76 C-BOND II 6-79

BEG DATE	=	76195
BEG TIME	=	5.00
END DATE	=	76195
END TIME	=	9.00
RESTART FLAG	=	F
SINK FLAG	=	T
PTS FLAG	=	T
RWY FLAG	=	F
TEMP FLAG	=	T
TERR FLAG	=	T
CONC VAR FLAG	=	F
DEF ROUGHNESS	=	.5000
DEF VEGETATN	=	1.0000
MAX TIME SLICE	=	.1000
MAX STEPS	=	1
MIN STEP SIZE	=	.0030
MAX ITERS	=	20
ERR TOL	=	.010000
DARK CRITERION	=	.0100
INST INTERVAL	=	1.00
AVG INTERVAL	=	1.00
PRINT OPTIONS	=	1
		1
		1
		1
		0
		0

DBRKPT PRINT\$

## X AIRSHED SIMULATION

After all the input data files have been prepared as described in the preceding chapters, running the Airshed Simulation Program is simply a matter of properly assigning all the input files and arranging to save all the desired output files. That is, running the simulation is primarily a job control function, since the bulk of the input and output processing is performed outside of this step.

This chapter discusses job control in terms of input, scratch, and output files; describes and presents samples of the printed output; and discusses the error messages generated by the program. Also included is a worksheet that can be used to keep track of files.

### A. JOB CONTROL

Figure X-1 shows the information flow for the Airshed Simulation Program. (Each group of files is discussed in more detail later.) Dotted lines indicate a possible creation of the SIMCONTROL file in a separate job step, and a possible restart using a previously generated output INSTANT file as an input AIRQUALITY file. Because the actual job control statements required are highly dependent on a particular implementation or computer, they are not specified here, but should be developed by the computer personnel who implement the system.

Tables X-1, X-2, and X-3 show the FORTRAN unit assignments of the Airshed Simulation Program's input, scratch, and output files, respectively. Currently, these unit assignments are compiled in the program by data statements in the subprogram BLKDAT. For instructions on changing the assignments, see Section VII.H. of the Systems Manual. The local file designation that corresponds to each data file should be entered in the tables for each specific implementation:

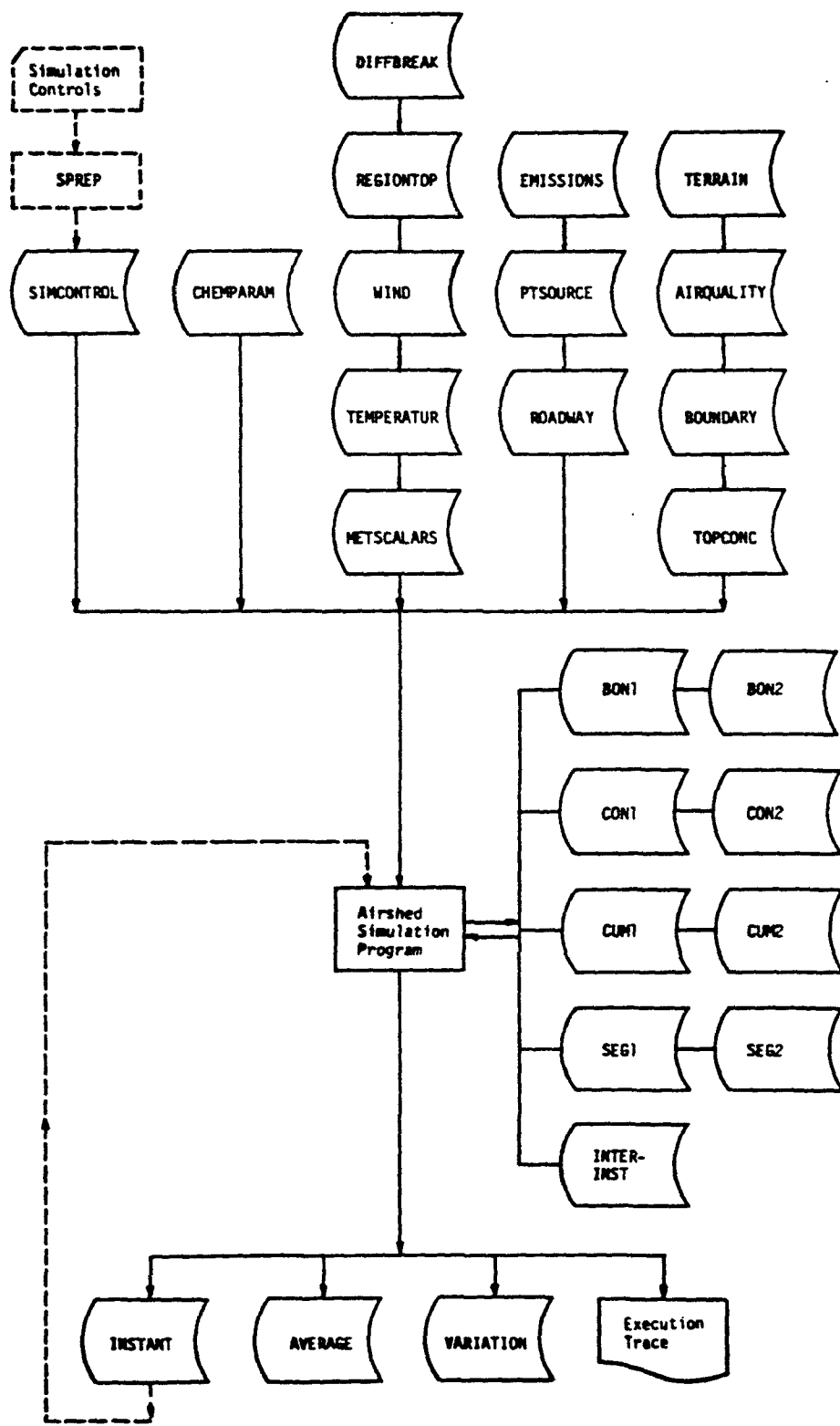


Figure X-1. Information Flow Diagram for the Airshed Simulation Program

Table X-1. Input File Assignments

<u>File Name</u>	<u>Unit No.</u>	<u>LBL lfn*</u>	<u>Optional File</u>	<u>File Closed During Simulation</u>
AIRQUALITY	3	AIRQ		✓
BOUNDARY	11	BOUND		
DIFFBREAK	12	DIFBK		
EMISSIONS	13	EMISS		
METSCALARS	7	METS		
PTSOURCE	14	POINTS	✓	
REGIONTOP	15	REGTOP		
ROADWAY	16	ROAD	✓	
TEMPERATUR	17	TEMP	✓	
TERRAIN	4	TERAIN	✓	✓
TOPCONC	18	TOPCON		
WIND	19	WIND		
CHEMPARAM	2	CPARAM		✓
SIMCONTROL	1	SIMCTL		✓

---

\* "lfn" means local file name.

Table X-2. Scratch File Assignments

<u>File Name</u>	<u>Unit No.</u>	<u>LBL fn</u>	<u>Optional File</u>	<u>File Closed During Simulation</u>
INTERINST	30	TAPE30		
BON1	37	†	*	
BON2	38	†	*	
CON1	33	TAPE33	*	
CON2	34	TAPE34		
CUM1	35	TAPE35	*	
CUM2	36	TAPE36		
SEG1	31	TAPE31	*	
SEG2	32	TAPE32		

\* Not used for runs with only one segment.

† The LBL system utilizes large core memory instead of these files; UNIVAC uses extended memory.

Table X-3. Output File Assignments

<u>File Name</u>	<u>Unit No.</u>	<u>LBL fn</u>	<u>Optional File</u>	<u>File Closed During Simulation</u>
AVERAGE	21	AVERAG	✓	
INSTANT	22	INST	✓	
VARIATION	23	CONVAR	✓	
Execution Trace	6	OUTPUT		

- > For the LBL CDC 7600 system, six-character mnemonic names are used instead of "TAPEnn."
- > For the UNIVAC 1100 systems, the internal file names are the unit numbers themselves.
- > For IBM systems, the DD names are of the form "FTnnF001."

### 1. Input Files

Table X-1 lists fourteen input files--twelve M.E.B. files and two control files. Four of the M.E.B. files are optional: PTSOURCE, ROADWAY, TEMPERATUR, and TERRAIN; all other input files must be present. For more information on selecting input file options and providing default values for the data omitted, see the Section IX.B on preparation of the SIMCONTROL file.

### 2. Scratch Files

Table X-2 shows the nine scratch files that might be used by the simulation. The actual number of files used and the method of assigning them are governed by the particular implementation of the system. For example, at LBL, only seven of the files are used, and the other two are represented by vectors in large core memory. See Section VII.B of the Systems Manual for more detailed information on the scratch files.

### 3. Output Files

As shown in Table X-3, the Airshed Simulation Program can write up to three binary data files, which are described in detail in Chapter IX of the Systems Manual. The presence or absence of these files is determined by parameters on the SIMCONTROL file:

- > AVERAGE will be written if the averaging interval is less than 99.0 hours.
- > INSTANT will be written at the beginning of the simulation, at the instantaneous output interval, and also at the end of each averaging interval if that time does not coincide with an instantaneous output time. If both the instantaneous output interval and the averaging interval are greater than 99.0 hours, INSTANT will not be written at all.
- > VARIATION will be written if the concentration variation option is TRUE and the averaging interval is less than 99.0 hours.

## B. PRINTED OUTPUT

The printed output from the simulation program appears on the Execution Trace file--ordinarily the standard print unit--and always contains at least the following information:

- > A complete display of the contents of the SIMCONTROL and CHEMPARAM files.
- > The region description as specified on the BOUNDARY file. This description serves as the standard for the run: the horizontal grid parameters on all other spatially varying files and the vertical distribution parameters on all other vertically varying files must match those of the BOUNDARY file.
- > The header information from each M.E.B. file. If the files were uniquely identified at the times they were created, this information completely defines all data input to the run.
- > A message when each non-time-varying file is closed.
- > A message at the end of each time slice indicating elapsed CPU time (if the host computer provides the capability).
- > A message at the termination of the run.

As indicated on the SIMCONTROL file, additional printed output options are also available that are primarily useful for debugging:

- > The file history printout consists of a message printed whenever a time interval record is read from any file. During early simulation runs, this information can help the user follow the progress of the run.
- > The storage allocation printout consists of tables listing pointers to scratch storage. Each entry is the relative location of a data array in a single storage vector. To interpret these tables, see Section VII.B of the Systems Manual.
- > Instantaneous and average concentration arrays may be printed at the same time they are written on the files. They are printed by segments, within each segment by species, and within each species by vertical level. These printouts contain values in the boundary cells that do not necessarily represent the actual boundary values at the time; thus, they should be used only for program debugging or for preliminary runs for which the binary output files are not to be saved.

Exhibit X-1 is an annotated excerpt of the output from a successful Airshed Simulation Program run that contains samples of all the normal and optional information printed.

If the Airshed Simulation Program is unable to perform the chemistry step successfully for any cell, it will give a "crash printout," showing the values in all data arrays in the vicinity of the cell where the error occurred. Also included are the values in the local vectors used by the simulation. A sample of the crash printout is shown in Exhibit X-2. The user should examine this printout and look for anomalous values of

\*\*\*\*\* SIMCONTROL FILE \*\*\*\*\*

```
FILE DESCRIPTION HEADER RECORD
FILE TYPE = SIMCONTROL
FILE ID = RAPS TEST RUN CONTROLS FOR JUNE-JULY 1978
NO OF SEGMENTS = 0
NO OF SPECIES = 0
BEG DATE = 0
BEG TIME = 0.
END DATE = 0.
END TIME = 0.

SIMULATION CONTROLS RECORD
RUN ID = RAPS 4-KM DAY 195 -- 4-HR RUN 07/13/76 C-BOND II 8-79
BEG DATE = 76195
BEG TIME = 5.00
END DATE = 76195
END TIME = 9.00

RESTART FLAG = F
SINK FLAG = T
PTS FLAG = T
RWY FLAG = F
TEMP FLAG = T
TERR FLAG = T

CONC VAR FLAG = F

DEF ROUGHNESS = .5000
DEF VEGETAT'N = 1.00000

MAX TIME SLICE= .1000
MAX STEPS = 1

MIN STEP SIZE = .0030
MAX ITERS = 20
ERROR TOL = .010000
DARK CRITERION=.0100

INST INTERVAL = 1.00
AVG INTERVAL = 1.00

FILE HISTORY =
STORAGE ALLOC = 1
LFVELS INST = 1
LEVELS AVG = 1
CONC VAR = 0
OPT J = 0

FILE 1 SIMCONTROL CLOSED.
```

## \*\*\*\*\* CHEMPARAM FILE \*\*\*\*\*

FILE DESCRIPTION HEADER RECORD  
 FILE TYPE = CHEM/PAHM  
 FILE ID = CHEMISTRY PARAMETERS NEW CHEM HIGHER PARAFINS LOWER LIMI  
 NO OF SEGMENTS = 0  
 NO OF SPECIES = 11  
 BEG DATE = 0  
 BEG TIME = 0.  
 END DATE = 0.  
 END TIME = 0.

## SPECIES DESCRIPTION HEADER RECORD

NO  
 NO2  
 O3  
 ETH  
 OLE  
 PAR  
 CARB  
 APO  
 BZA  
 PAH  
 CO

CHEMISTRY DEFINITION RECORD  
 MECHANISM ID = CARBON BOND MECHANISM -- NOVEMBER 78  
 NO OF REACTIONS = 69  
 NO OF COEFFS = 4

## SPECIES PARAMETERS RECORD

SPEC. NAME	REACT	SSIC	SSBC	DYRES	LOW BD SS	HIGH BD SS	UP BD NUM	LOW BD NUM	HIGH BD NUM
NO	T	F	F	2.8000-001	1.0000-005	1.0000+006	1.0000-005	1.0000-005	1.0000+006
N <sub>2</sub> O	T	F	F	1.1100-002	1.0000-004	1.0000+006	1.0000-004	1.0000-004	1.0000+006
O3	T	T	T	1.4700-002	1.0000-009	1.0000+006	1.0000-009	1.0000-009	1.0000+006
ETH	T	F	F	1.0000+010	1.0000-006	1.0000+006	1.0000-006	1.0000-006	1.0000+006
OLE	T	F	F	1.0000+010	1.0000-006	1.0000+006	1.0000-006	1.0000-006	1.0000+006
PAR	T	F	F	1.0000+010	1.0000-003	1.0000+006	1.0000-004	1.0000-004	1.0000+006
CAPB	T	F	F	1.0000+010	1.0000-006	1.0000+006	1.0000-006	1.0000-006	1.0000+006
ARO	T	F	F	1.0000+010	1.0000-006	1.0000+006	1.0000-006	1.0000-006	1.0000+006
BZA	T	F	F	1.0000+010	1.0000-006	1.0000+006	1.0000-006	1.0000-006	1.0000+006
PAH	T	F	F	1.0000+010	1.0000-006	1.0000+006	1.0000-006	1.0000-006	1.0000+006
CO	T	F	F	1.0000+010	1.0000-002	1.0000+006	1.0000-002	1.0000-002	1.0000+006

## REACTION PARAMETERS RECORD

RATE CONST	PHOTOL	TEMP	ACT ENERGY	REF TEMP
1	6.0000+001	T	.0000	.0000
2	2.6400+008	F	.0000	.0000
3	1.4340+003	F	1.4500+003	2.9200+002
4	2.8200+000	F	2.4500+003	2.9200+002
5	8.0400+005	F	.0000	.0000
6	4.6200+003	F	1.0000+003	2.9200+002
7	3.0000+002	F	1.5250+003	2.9200+002
8	6.4000+035	F	.0000	.0000
9	2.6400+004	F	.0000	.0000

10	9.0000-003	.0000
11	1.6800+006	.0000
12	9.0000-003	.0000
13	7.2000+005	2.9300+002
14	9.0000+005	.0000
15	1.2000+003	.0000
16	9.0000+004	.0000
17	1.6200+005	.0000
18	1.6200+005	.0000
19	2.5200+005	.0000
20	4.8000-001	.0000
21	4.8000-001	.0000
22	3.6000+004	.0000
23	3.6000+004	.0000
24	7.2000+005	.0000
25	1.4400-001	.0000
26	2.2800+005	.0000
27	7.2000+005	.0000
28	7.2000+005	.0000
29	2.3000+005	.0000
30	4.6000+005	.0000
31	3.0000+004	.0000
32	3.0000+002	.0000
33	1.2000+004	.0000
34	3.0000+002	.0000
35	1.1400+006	.0000
36	8.1000-002	T
37	1.2150-001	T
38	6.0000+006	F
39	1.2000+005	F
40	1.6800+000	F
41	2.4000+005	F
42	2.4000+005	F
43	7.2000+005	F
44	4.8000+005	F
45	1.2000+005	F
46	7.2000+005	F
47	4.8000+005	F
48	1.2000+005	F
49	4.0200+006	F
50	1.4400+004	F
51	5.4000+003	F
52	9.0000+003	F
53	2.0400+004	F
54	2.5500+004	F
55	5.1000+003	F
56	3.6000+005	F
57	9.6000+004	F
58	9.0000+005	F
59	6.0000+006	F
60	1.0000+003	F
61	9.0000+002	F

Exhibit X-1 (Continued)

62	2.1000+006	F	.0000
63	3.6000+001	F	.0000
64	6.0000+005	F	.0000
65	2.1600+000	T	.0000
66	5.4000+004	F	.0000
67	4.8000+004	F	.0000
68	5.4000+005	F	.0000
69	3.0000-002	F	.0000

STOICHIOMETRIC COEFFICIENTS RECORD  
ALPHA 5.0000-001  
BETA 4.8000-001  
GAMMA 1.0000-000  
ETA 1.0000-001

FILE 2 CHEMPARAM CLOSED.

**LOCAL POINTERS**

1	1
2	5
3	9
4	13
5	17
6	21
7	25
8	36
9	47
10	168
11	185
12	202
13	219
14	236
15	253
16	168
17	172
18	177
19	188
20	232
21	237
22	248
23	253
24	269
25	285
26	289
27	253
28	257
29	261
30	265
31	269
32	273
33	278
34	282
35	253
36	264
37	275
38	286
39	407
40	418

Exhibit X-1 (Continued)

BOUNDARY INDEX TABLE FOR SEGMENT 1		WEST	EAST																
ROW	COL			0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
1	1	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
2	2	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
3	3	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
4	4	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
5	5	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
6	6	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
7	7	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
8	8	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
9	9	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
10	10	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
11	11	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
12	12	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
13	13	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
14	14	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
15	15	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
16	16	2	2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
17	17	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0

**Exhibit X-1 (Continued)**

ROW		WEST			EAST			NORTH									
COL	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17
1	0	2	2	2	2	2	2	2	2	2	2	2	2	2	2	2	0
2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
4	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
5	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
6	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
7	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
8	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
9	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
10	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
11	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
12	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
13	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
14	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
15	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
16	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
17	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0

RAPS 4-KM DAY 195 -- 4-HR RUN 07/13/76 C-BOND II 8-79

\*\*\*\*\* REGION DESCRIPTION \*\*\*\*\*

HORIZONTAL GRID

	X	Y
REFERENCE ORIGIN (UTM - METERS)	725000.	4252000. (ZONE 15)
REGION ORIGIN (M FROM REF. ORG.)	-19000.	-16000.
CELL SPACING (M)	4000.	4000.
NUMBER OF CELLS	17	22

VERTICAL DISTRIBUTION

NO. CELLS IN LOWER LAYER = 2

NO. CELLS IN UPPER LAYER = 2

SURFACE LAYER HT (M) = 0.

MIN HT LOWER LAYER CELLS (M) = .50.

MIN HT UPPER LAYER CELLS (M) = 200.

TOTAL NO. VERTICAL CELLS = 4

SEGMENTS	IORGX	IORGY	NDX	NDY
1	0	0	17	13
2	0	9	17	13

BEGINNING OF TABLES RELATIVE TO BEGINNING OF AVAILABLE SPACE

SEGMENT	1
LOCAL	131
BOUNDARY	559
SEG SCRATCH	2295
LAST WORD	22756

Exhibit X-1 (Continued)

## SEGMENT ALLOCATION TABLE

	1	2
1	0	0
2	0	9
3	17	17
4	13	13
5	80	129
6	0	0
7	221	221
8	886	884
9	9724	9724
10	2431	2431
11	1736	1736
12	1	1737
13	9724	9724
14	0	9724
15	9724	9724
16	0	9724
17	10101	10758
18	0	10101
19	0	0
20	0	0
21	0	0
22	0	0
23	0	0
24	0	0
25	0	0
26	0	0
27	0	0
28	0	0
29	1	1
30	1	1
31	1	1
32	1	0
33	0	1
34	1	1
35	53	53
36	105	105
37	173	173
38	0	0
39	0	0
40	0	241
41	241	0
42	1	1
43	1	1
44	865	865
45	2432	2432

Exhibit X-1 (Continued)

46	2653	
47	2874	2874
48	3095	3095
49	3316	3316
50	3537	3537
51	3758	3758
52	3979	3979
53	4200	4200
54	5084	5084
55	5968	5968
56	6189	6189
57	6410	6410
58	6631	6631
59	0	0
60	0	0
61	9062	9062
62	9142	9191
63	9222	9320
64	10102	10739
65	0	0

FILE = AIRQUALITY ID = INITIAL POLLUTANT CONCENTRATIONS ST LOUIS 500 76195  
 SEGMENTS = 2 SPECIES = 11  
 TIME SPAN = 76195 5.00 TO 76195 5.02  
 725000.42:2000. 15 -19000. -16000. 4000. 4000. 17 22 4 2 2 .00 50.00 200.00

## SEGMENTS

0	0	17	13
0	9	17	13

## SPECIES ON AIRQUALITY FILE

## NO. NAME NO. WITHIN RUN

1	NO	1
2	NO2	2
3	O3	3
4	ETH	4
5	OLE	5
6	PAR	6
7	CARB	7
8	ARO	8
9	BZA	9
10	PAN	10
11	CO	11

FILSKP READS TIME INTERVAL 76195 5.00 76195 5.02 ON UNIT 3

FILE = BOUNDARY ID = BOUNDARY DEFINITION AND CONCENTRATIONS  
 SEGMENTS = 2 SPECIES = 11  
 TIME SPAN = 76195 5.00 TO 76195 24.00  
 725000.425:2000. 15 -19000. -16000. 4000. 4000. 17 22 4 2 2 .00 50.00 200.00

## SEGMENTS

0	0	17	13
0	9	17	13

## SPECIES ON BOUNDARY FILE

## NO. NAME NO. WITHIN RUN

1	NO	1
2	NO2	2
3	O3	3
4	ETH	4
5	OLE	5
6	PAR	6
7	CARB	7
8	ARO	8
9	BZA	9
10	PAN	10
11	CO	11

FILSKP READS TIME INTERVAL 76195 5.00 76195 6.00 ON UNIT 11

Exhibit X-1 (Continued)

```

FILE = DIFFBREAK   ID = HEIGHT OF THE MIXED LAYER
SEGMENTS = 2      SPECIES = 0
TIME SPAN = 76195 5.00 TO 76195 24.00
725000.4252000. 15 -19000. -16000. 4000. 4000. 17 22 0 2 2 .00 50.00 200.00

SEGMENTS
0    0    17   13
0    9    17   13
FILSKP READS TIME INTERVAL 76195 5.00 76195 6.00 ON UNIT 12

```

```

FILE = EMISSIONS  ID = RAPS EMISSIONS (17 X 22) DAY195 07/13/76
SEGMENTS = 2      SPECIES = 8
TIME SPAN = 76195 5.00 TO 76195 24.00
725000.4252000. 15 -19000. -16000. 4000. 4000. 17 22 0 2 2 .00 50.00 200.00

SEGMENTS
0    0    17   13
0    9    17   13
SPECIES ON EMISSIONS FILE
NO. NAME          NO. WITHIN RUN
1  NO            1
2  NO2           2
3  ETH            4
4  OLE            5
5  PAR            6
6  CARB           7
7  ARO            8
8  CO             11
FILSKP READS TIME INTERVAL 76195 5.00 76195 6.00 ON UNIT 13

```

```

FILE = METSCALARS ID = VALUES FOR METEOROLOGICAL SCALARS
SEGMENTS = 0      SPECIES = 0
TIME SPAN = 76195 4.00 TO 76195 24.00
725000.4252000. 15 -19000. -16000. 4000. 4000. 17 22 4 2 2 .00 50.00 200.00

FILSKP READS TIME INTERVAL 76195 4.00 76195 5.00 ON UNIT 7
TGRADBELOW TGRADABOVE EXPCLASS RADFACTOR CONCWTTER ATMOSPRESS
-.010   .004   -1.000   .008   12667.000   .990
FILSKP READS TIME INTERVAL 76195 5.00 76195 6.00 ON UNIT 7

```

FILE = PTSOURCE ID = POINT SOURCES FOR ST. LOUIS 76195 4X4 GRID  
 SEGMENTS = 2 SPECIES = 7  
 TIME SPAN = 76195 5.00 TO 76195 24.00  
 725000.4252000. 15 -19000. -16000. 4000. 4000. 17 22 4 2 2 .00 50.00 200.00

SEGMENTS	0	0	17	13
	0	9	17	13

SPECIES ON PTSOURCE FILE  
 NO. NAME NO. WITHIN RUN  
 1 NO 1  
 2 NO2 2  
 3 OLE 5  
 4 PAR 6  
 5 CARB 7  
 6 ARO 8  
 7 CO 11  
 FILSKP READS TIME INTERVAL 76195 5.00 76195 6.00 ON UNIT 14

FILE = REGIONTOP ID = HEIGHT OF THE TOP OF THE MODELING REGION  
 SEGMENTS = 2 SPECIES = 0  
 TIME SPAN = 76195 5.00 TO 76195 24.00  
 725000.4252000. 15 -19000. -16000. 4000. 4000. 17 22 0 2 2 .00 50.00 200.00

SEGMENTS	0	0	17	13
	0	9	17	13

FILSKP READS TIME INTERVAL 76195 5.00 76195 6.00 ON UNIT 15

FILE = TEMPERATUR ID = TEMPERATURE FIELD FOR ST. LOUIS DAY 195  
 SEGMENTS = 2 SPECIES = 0  
 TIME SPAN = 76195 5.00 TO 76195 24.00  
 725000.4252000. 15 -19000. -16000. 4000. 4000. 17 22 0 2 2 .00 50.00 200.00

SEGMENTS	0	0	17	13
----------	---	---	----	----

FILSKP READS TIME INTERVAL 76195 5.00 76195 6.00 ON UNIT 17

FILE = TERRAIN ID = SURFACE ROUGHNESS AND VEGETATIVE FACTORS  
 SEGMENTS = 2 SPECIES = 0  
 TIME SPAN = 76195 5.00 TO 76195 24.00  
 725000.4252000. 15 -19000. -16000. 4000. 4000. 17 22 0 2 2 .00 50.00 200.00

Exhibit X-1 (Continued)

SEGMENTS	0	0	17	13
	0	9	17	13

FILE = TDPCONC ID = POLLUTANT CONCENTRATIONS ABOVE THE MODELING REGION

SEGMENTS = 2 SPECIES = 11  
 TIME SPAN = 76195 5.00 TO 76195 24.00  
 725000.425:000. 15 -19000. -16000. 4000. 4000. 17 22 0 2 2 .00 50.00 200.00

SEGMENTS	0	0	17	13
	0	9	17	13

SPECIES ON TOPCONC FILE  
 NO. NAME NO. WITHIN RUN

1	NO	1
2	ND2	2
3	O3	3
4	ETH	4
5	OLE	5
6	PAR	6
7	CARB	7
8	ARO	8
9	BZA	9
10	PAN	10
11	CO	11

FILSKP READS TIME INTERVAL 76195 .00 76195 24.00 ON UNIT 18

FILE = WIND ID = WIND FIELD FOR ST. LOUIS DIVERGENCE ALGORITHM USED

SEGMENTS = 2 SPECIES = 0  
 TIME SPAN = 76195 5.00 TO 76195 24.00  
 725000.425:000. 15 -19000. -16000. 4000. 4000. 17 22 4 2 2 .00 50.00 200.00

SEGMENTS	0	0	17	13
	0	9	17	13

FILSKP READS TIME INTERVAL 76195 5.00 76195 6.00 ON UNIT 19

TGRAD BELOW	TGRAD ABOVE	EXPCLASS	RADFACTOR	CONCWATER	ATMOSPRESS
-.010	.004	.000	.084	12687.000	.990

FILE 4 TERRAIN CLOSED.  
 TZERO READS TIME INTERVAL 76195 6.00 76195 7.00 ON UNIT 12 DIFFBREAK  
 TZERO READS TIME INTERVAL 76195 6.00 76195 7.00 ON UNIT 15 REGIONTOP

Exhibit X-1 (Continued)

**FILE 3 AIRQUILITY CLOSED.**

76195 5.00 SEG	WXMAX	WYMAX	DTMAX	DT NDT
1 22453.	.9603.	.089	.089	1
2 .2453.	.9603.	.089	.089	1
TIME = 76195 5.089	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs	
76195 5.09 SEG	WXMAX	WYMAX	DTMAX	DT NDT
1 22453.	.9603.	.089	.089	1
2 22453.	.9603.	.089	.0C9	1
TIME = 76195 5.178	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs	
76195 5.18 SEG	WXMAX	WYMAX	DTMAX	DT NDT
1 22453.	.9603.	.089	.089	1
2 22453.	.9603.	.089	.089	1
TIME = 76195 5.267	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs	
76195 5.27 SEG	WXMAX	WYMAX	DTMAX	DT NDT
1 22453.	.9603.	.089	.089	1
2 22453.	.9603.	.089	.089	1
TIME = 76195 5.356	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs	
76195 5.36 SEG	WXMAX	WYMAX	DTMAX	DT NDT
1 22453.	.9603.	.089	.089	1
2 22453.	.9603.	.089	.089	1
TIME = 76195 5.445	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs	
76195 5.45 SEG	WXMAX	WYMAX	DTMAX	DT NDT
1 22453.	.9603.	.089	.089	1
2 22453.	.9603.	.089	.089	1
TIME = 76195 5.534	SLICE = .0691	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs	
76195 5.53 SEG	WXMAX	WYMAX	DTMAX	DT NDT
1 22453.	.9603.	.089	.089	1
2 22453.	.9603.	.089	.089	1
TIME = 76195 5.624	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs	
76195 5.62 SEG	WXMAX	WYMAX	DTMAX	DT NDT

TIME =	76195	5.713	SEG	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs	
	1	22453.	WXMAX	WYMAX DTMAX DT NDT			
	2	22453.	9603.	.089 .089 1			
	76195	5.71	SEG	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs	
	1	22453.	9603.	.089 .089 1			
	2	22453.	9603.	.089 .089 1			
	TIME =	76195	5.802	SEG	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs
	76195	5.80	SEG	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs	
	1	22453.	WXMAX	WYMAX DTMAX DT NDT			
	2	22453.	9603.	.089 .089 1			
	TIME =	76195	5.891	SEG	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs
	76195	5.89	SEG	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs	
	1	22453.	WXMAX	WYMAX DTMAX DT NDT			
	2	22453.	9603.	.089 .089 1			
	TIME =	76195	5.980	SEG	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs
	76195	5.98	SEG	SLICE = .0891	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs	
	1	22453.	WXMAX	WYMAX DTMAX DT NDT			
	2	22453.	9603.	.089 .089 1			
	TIME =	76195	6.000	SEG	SLICE = .0202	ELAPSED CPU TIME = .000 SECs	TOTAL ELAPSED CPU TIME = .000 SECs

## RAPS 4-KM DAY 195 -- 4-HR RUN 07/13/76 C-BOND II 8-79

Avg	Concentrations At 76195	6.00 For Species 1 NO	Segment= 1	Level=1	
1	2	3	4	5	
13	8.44-004	5.04-003	3.31-003	4.85-003	
12	4.22-004	5.02-004	4.38-004	3.56-004	
11	8.45-004	1.48-002	5.33-003	5.07-003	
10	5.06-004	4.47-004	2.98-004	2.74-004	
9	6.66-004	3.19-004	3.77-004	3.18-004	
8	8.48-004	2.38-003	4.72-003	4.76-003	
7	5.56-004	4.52-004	5.52-004	5.00-004	
6	8.50-004	1.45-003	2.57-003	4.12-003	
5	6.03-004	7.12-004	6.07-004	4.27-004	
4	8.51-004	1.12-003	1.48-003	1.99-003	
3	4.08-004	6.75-004	7.37-004	4.88-004	
2	8.53-004	8.15-004	8.88-004	1.05-003	
1	3.02-004	3.71-004	6.63-004	5.07-004	
	6.854-004	4.91-004	6.41-004	8.83-004	
	3.05-004	3.64-004	4.10-004	4.14-004	
	5	8.56-004	4.07-004	5.30-004	7.01-004
	3.08	3.55-004	3.94-004	4.17-004	3.24-004
	4	8.57-004	3.25-004	4.37-004	5.52-004
	3	2.91-004	3.36-004	3.75-004	3.85-004
	2	8.59-004	2.62-004	3.38-004	4.93-004
	2	2.71-004	3.09-004	3.40-004	3.25-004
	1	3.01-004	2.56-004	3.07-004	3.43-004
	1	2.67-004	2.77-004	2.95-004	3.12-004
	1	2.00-003	3.03-004	3.05-004	3.06-004
	3.20-004	3.22-004	3.23-004	3.25-004	2.00-003

Avg	Concentrations At 76195	6.00 For Species 2 NO2	Segment= 1	Level=1	
1	2	3	4	5	
13	9.16-003	1.76-002	1.69-002	2.41-002	
12	6.64-003	6.24-003	5.03-003	4.01-003	
11	9.15-003	2.27-002	2.03-002	2.24-002	
10	6.68-003	5.65-003	4.26-003	3.67-003	
9	9.15-003	1.69-002	2.26-002	2.40-002	
8	5.95-003	4.84-003	4.75-003	3.89-003	
7	9.15-003	1.29-002	1.63-002	1.99-002	
6	6.87-003	5.82-003	5.71-003	4.63-003	
5	9.15-003	9.62-003	5.74-002	1.72-002	
4	7.30-003	7.43-003	5.89-003	4.29-003	
3	6.95-003	7.96-003	9.63-003	1.14-002	
2	5.85-003	6.96-003	6.49-003	4.41-003	
1	7.91-003	6.18-003	6.84-003	7.98-003	
	7	4.66-003	4.88-003	4.87-003	4.59-003
	6	9.15-003	4.86-003	5.81-003	7.13-003
	5	4.54-003	4.89-003	4.27-003	4.17-003
	4.26-003	4.25-003	4.30-003	4.14-003	3.60-003

4	9.14-003	3.90-003	4.54-003	5.48-003	7.47-003	9.28-003	4.60-003	4.43-003	4.35-003	4.30-003	4.36-003	3.91-003
4	4.05-003	4.18-003	4.16-003	3.99-003	3.68-003	3.68-003	0.91-003	1.40-002	3.55-003	4.04-003	4.16-003	4.20-003
3	9.14-003	3.55-003	4.05-003	5.30-003	5.30-003	8.91-003	1.40-002	1.40-002	3.55-003	4.04-003	4.16-003	4.31-003
3	3.80-003	3.92-003	3.92-003	3.93-003	3.79-003	3.67-003	5.16-003	5.61-003	3.63-003	3.67-003	3.90-003	3.86-003
2	3.70-003	3.52-003	3.79-003	4.15-003	5.16-003	5.16-003	5.16-003	5.16-003	3.67-003	3.67-003	3.90-003	3.86-003
2	3.52-003	3.64-003	3.66-003	3.66-003	3.66-003	3.66-003	4.62-003	4.62-003	4.62-003	4.62-003	4.62-003	3.69-003
1	2.00-003	3.70-003	3.70-003	3.69-003	3.69-003	3.69-003	3.69-003	3.69-003	3.69-003	3.69-003	3.68-003	3.68-003
1	3.68-003	3.68-003	3.68-003	3.68-003	3.68-003	3.68-003	2.00-003	2.00-003	2.00-003	2.00-003	2.00-003	2.00-003

AVG	SEGMENT= 1 LEVEL=1											
	CONCENTRATIONS AT 76195			6.00 FOR SPECIES 3 03			CONCENTRATIONS AT 76195			6.00 FOR SPECIES 4 ETH		
1	2	3	4	5	6	7	8	9	10	11	12	
13	2.08-002	1.10-002	1.33-002	1.21-002	1.82-002	1.75-002	1.69-002	2.16-002	2.58-002	3.18-002		
12	2.08-002	6.92-003	1.14-002	2.50-002	2.43-002	1.51-002	1.44-002	1.43-002	1.96-002	2.04-002	3.36-002	
11	2.08-002	1.06-002	8.51-003	9.60-003	1.32-002	1.40-002	1.17-002	9.47-003	9.39-003	2.07-002	2.99-002	3.69-002
10	2.08-002	1.47-002	1.23-002	1.31-002	1.50-002	1.66-002	1.64-002	1.83-002	2.06-002	2.55-002	2.95-002	3.66-002
9	2.08-002	1.68-002	1.47-002	1.32-002	1.43-002	1.74-002	1.77-002	1.88-002	2.22-002	3.09-002	3.27-002	3.29-002
8	3.22-002	3.10-002	2.83-002	2.50-002	2.43-002	2.50-002	1.48-002	1.59-002	2.20-002	2.62-002	2.90-002	3.20-002
7	2.09-002	1.79-002	1.71-002	1.72-002	1.70-002	1.91-002	2.15-002	2.57-002	2.80-002	3.04-002	3.14-002	3.14-002
6	2.09-002	1.94-002	1.85-002	1.79-002	1.77-002	1.83-002	2.22-002	2.37-002	2.54-002	2.71-002	2.93-002	3.08-002
5	2.09-002	2.94-002	2.31-002	2.35-002	2.43-002	1.88-002	1.83-002	2.24-002	2.41-002	2.54-002	2.67-002	2.81-002
4	2.73-002	2.57-002	2.41-002	2.33-002	2.43-002	1.93-002	1.84-002	2.32-002	2.43-002	2.52-002	2.64-002	2.69-002
3	2.09-002	2.27-002	2.23-002	2.14-002	1.86-002	1.46-002	2.45-002	2.47-002	2.51-002	2.56-002	2.57-002	2.61-002
2	2.43-002	2.33-002	2.32-002	2.30-002	2.23-002	2.20-002	2.43-002	2.44-002	2.47-002	2.50-002	2.52-002	2.50-002
1	2.00-002	2.45-002	2.39-002	2.33-002	2.43-002	2.43-002	2.43-002	2.43-002	2.43-002	2.43-002	2.43-002	2.43-002
0	2.43-002	2.43-002	2.43-002	2.43-002	2.43-002	2.00-002						

AVG	SEGMENT= 1 LEVEL=1											
	CONCENTRATIONS AT 76195			6.00 FOR SPECIES 4 ETH			CONCENTRATIONS AT 76195			6.00 FOR SPECIES 3 03		
1	2	3	4	5	6	7	8	9	10	11	12	
13	1.71-003	5.07-003	6.97-003	9.06-003	9.41-003	6.57-003	5.38-003	4.87-003	2.99-003	2.39-003	1.98-003	
12	1.71-003	1.96-003	1.64-003	1.10-003	1.09-003	6.84-003	6.41-003	6.34-003	5.47-003	4.16-003	2.55-003	2.44-003
11	1.71-003	4.09-003	5.25-003	5.35-003	5.52-003	6.80-003	6.89-003	6.40-003	5.36-003	4.04-003	2.85-003	
10	1.71-003	1.72-003	1.53-003	1.18-003	1.09-003	4.68-003	4.73-003	4.77-003	4.20-003	3.39-003	2.98-003	2.79-003
9	1.71-003	2.44-003	2.94-003	3.73-003	4.38-003	4.05-003	3.85-003	3.35-003	2.40-003	2.11-003	2.18-003	2.37-003
8	2.56-003	2.62-003	2.09-003	1.33-003	1.09-003	2.57-003	3.75-003	3.64-003	2.41-003	1.66-003	1.72-003	1.55-003

Exhibit X-1 (Continued)

AVG CONCENTRATIONS AT 76195 6.00 FOR SPECIES 5 OLE												
SEGMENT= 1 LEVEL=1												
	1	2	3	4	5	6	7	8	9	10	11	12
7	1.64-003	2.40-003	2.31-003	1.39-003	1.09-003	2.27-003	2.53-003	2.16-003	1.78-003	1.61-003	1.61-003	1.32-003
7	1.71-003	1.49-003	1.77-003	1.93-003	1.09-003	2.11-003	1.47-003	1.33-003	1.47-003	1.77-003	1.42-003	1.35-003
1	1.39-003	1.49-003	1.49-003	1.39-003	1.09-003	2.11-003	1.47-003	1.33-003	1.47-003	1.77-003	1.42-003	1.26-003
6	1.71-003	1.37-003	1.50-003	1.84-003	1.23-003	1.09-003	1.27-003	1.21-003	1.99-003	1.28-003	1.19-003	1.26-003
5	1.71-003	1.38-003	1.42-003	1.51-003	1.61-003	1.09-003	1.27-003	1.21-003	1.66-003	1.28-003	1.11-003	1.29-003
4	1.71-003	1.28-003	1.38-003	1.45-003	1.45-003	1.09-003	1.21-003	1.21-003	1.09-003	1.21-003	1.15-003	1.13-003
3	1.71-003	1.14-003	1.22-003	1.32-003	1.63-003	1.09-003	1.17-003	1.07-003	1.13-003	1.16-003	1.22-003	1.33-003
3	1.14-003	1.16-003	1.16-003	1.11-003	1.09-003	1.20-003	1.18-003	1.20-003	1.40-003	1.09-003	1.13-003	1.14-003
2	1.09-003	1.12-003	1.12-003	1.13-003	1.11-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.13-003
1	1.11-003	1.12-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.11-003
1	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003
1	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003	1.09-003

SUBROUTINE CRASH SHOWS YOU WHAT WAS WHAT AT THE TIME OF THE CRASH IN STEP 3. CELL 6, 4, 5, OF SEGMENT 1-SPECIES 1= CO

VALUES IN SPATIAL ARRAYS WILL BE PRINTED IN MAP FORM, I.E.

N  
W E  
S

WITH THE OFFENDING CELL ( 6, 4) IN THE CENTER

( 5, 5)	( 6, 5)	( 7, 5)
( 5, 4)	( 6, 4)	( 7, 4)
( 5, 3)	( 6, 3)	( 7, 3)

CONCENTRATIONS FOR SPECIES 1 CO

LEVELS	1	2	3
1.293E+00	1.220E+00	1.373E+00	1.231E+00
1.218E+00	1.210E+00	1.214E+00	1.216E+00
1.195E+00	1.206E+00	1.195E+00	1.205E+00
LEVELS	4	5	
1.202E+00	1.183E+00	1.020E+00	1.019E+00
1.171E+00	1.183E+00	9.535E-01	1.019E+00
1.165E+00	1.167E+00	9.534E-01	9.528E-01

DIFFUSION BREAK	TEMP	ROUGH	VEG	CARM
BEG	BEG	BEG	BEG	BEG
52.	52.	.52.	54.	199.0
52.	52.	.52.	54.	189.0
52.	52.	.52.	54.	189.0
				189.0
				189.0
				189.0
				189.0
END	END	END	END	END
				198.1
				198.1
				198.1
				198.1
				198.1

LEVEL	WK	WT	WY	
1	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
2	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
3	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
4	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
5	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
	3.000E+03	3.000E+03	3.000E+03	-1.000E+04 -1.000E+04 -1.000E+04
SPECIES	QT	QR	CTOP	
I-CO	1.559E+01	5.772E+00	1.934E+00	1.000E-01 1.000E-01 1.000E-01
	9.193E-01	2.197E+00	1.152E+00	1.000E-01 1.000E-01 1.000E-01
	0.	1.027E+00	3.249E-01	1.000E-01 1.000E-01 1.000E-01
BOUNDARY INDICES	I J	ADJ SEG	I J WITHIN THAT SEGMENT	
BDY	1 2	0 4	0 6	
W				
E	6	4	0	
S	6	2	0	
N	6	11	0	
SPEC	1.00E-01	1.00E-01	1.00E-01	1.00E-01
	1.00E-01	1.00E-01	1.00E-01	1.00E-01
	1.00E-01	1.00E-01	1.00E-01	1.00E-01
	1.00E-01	1.00E-01	1.00E-01	1.00E-01
	1.00E-01	1.00E-01	1.00E-01	1.00E-01

$\nu_1 + \zeta_1 = -1$

#### LOCAL VARIABLES

K	TMOLD	TMNEW	THAVG	HTOLD	HTNEW	HTAVG	AER	FACT	WZOLD	WZNEW	XKZ
1	26.	26.	26.	16.	16.	16.	0.	0.0000	0.	0.	0.
2	26.	26.	26.	20.	30.	30.	0.	0.0000	0.	0.	2.319E+03
3	26.	26.	26.	20.	50.	50.	0.	0.0000	0.	0.	2.432E+03
4	26.	26.	26.	20.	70.	70.	0.	0.0000	0.	0.	2.224E+03
5	109.	110.	114.	134.	139.	137.	0.	0.0000	0.	0.	1.954E+03
6							0.	0.0000	-2.491E+02	-2.700E+02	7.635E+02

#### CONTEN ARRAY (STEPS 3 AND 4)

SPEC	1	1.216E+00	1.217E+00	2	1.198E+00	3	1.169E+00	4	9.529E-01	5

SPEC	1	1.000E-10	0.	SRC	2.197E+00	FLUX

#### STEP 3 VECTORS

K	Y1-1	Y1-2	Y1-3	A	B	C	Y	X	CTEST	Z	AOMEGA

1	0.	0.	0.	0.	0.	0.	0.	0.	0.	0.	0.
2	0.	6.96E-01	3.04E-01	0.	1.30E+00	-3.04E-01	1.222E+00	0.	1.22E+00	9.38E-01	1.30E+00
3	3.04E-01	4.18E-01	2.70E-01	-3.04E-01	1.58E+00	-2.70E-01	1.20E+00	0.	1.23E+00	9.63E-01	1.51E+00
4	2.70E-01	6.46E-01	7.57E-02	-2.70E-01	1.35E+00	-7.07E-02	1.18E+00	0.	1.20E+00	1.12E+00	1.30E+00
5	1.28E-02	9.10E-01	0.	-1.28E-02	1.01E+00	0.	9.50E-01	0.	1.17E+00	1.18E+00	1.91E+00

#### MET SCALARS

TCRADBELOW	TCRADABOVE	EXPCLASS	RADFACTOR	CONCWATER	ATMOSPRESS
-.009	.029	-1.000	.220	13000.000	.840
ATFACT	TMFEC	TMEND	FHKFEC	FRKDEL	FACTRK

1.000E-01  
0.000  
.220  
.055

Exhibit X-2 (Continued)

<b>MISC SCALARS</b>		<b>WYMAX</b>	<b>WAVW</b>	<b>WAVS</b>	<b>WAVN</b>
ZREF	3000.000	10000.000	10440.307	10440.307	10440.307
DTMAX	DT	NDT			
.322	.100	1			
EL	USTAR	DIFRES			
TIME = 62.889	996.072	.014			
TIME = 76210	6.300	SLICE = .1000	ELAPSED CPU TIME = .096 SECs	TOTAL ELAPSED CPU TIME = .096 SECs	TOTAL ELAPSED CPU TIME = 3.736 SECs

Exhibit X-2 (Concluded)

any data. Check that printed values correspond to the data expected from the input files.

If no readily apparent errors can be found in the model inputs, then the available input and output should be examined by someone familiar with the technical components of the model. In undertaking the diagnosis, one should note the cause of the failure (iteration limit exceeded or negative concentration predicted). First, attention should be given to reaction rate constants and stoichiometric coefficients. Next, the predicted concentrations should be examined. Usually, one will note predictions for one or more pollutants that fall outside the expected nominal range of values for the particular contaminants in question. The development of an understanding of how these concentration values occurred will often suggest means for resolving the problem. For example, a very high NO concentration prediction may be causing the model to predict negative O<sub>3</sub> concentrations. The user might subsequently find that the NO emissions rate from a point source was in error by a factor of 100. Skill in diagnosing such problems requires some familiarity with the physical and chemical processes that take place in the atmosphere as well as an understanding of how these processes are treated in the model.

#### C. ERROR CONDITIONS

Table X-4 lists error conditions that could cause the Airshed Simulation Program to terminate before the simulation is completed. These are situations detected by the program itself rather than by the operating system, and the program will exit normally. Included in the table are the names of the subroutines in which the error messages are printed.

Table X-4. Airshed Simulation Program Error Conditions

<u>Error Condition</u>	<u>Subroutine(s) That Print Errors</u>
<b>Control file errors</b>	
File improperly named	SCREAD, CHREAD
Species or coefficient required by built-in chemical mechanism is missing	SPECID
Reactive species not listed in built-in mechanism	SPECID
<b>Allocation errors</b>	
Space allocation exceeded--region dimensions too large	ALLOC
<b>M.E.B. file errors</b>	
File missing or improperly named	HDRCK
Time span on file does not encompass time span of run	HDRCK
Region or segment definition does not match that on BOUNDARY file	HDRCK
Time intervals on file are not contiguous	FILSKP, FILCK
Unexpected end-of-file	Any file-read routine
<b>Chemistry errors</b>	
Negative concentrations at minimum step size	STEP4
Nonconvergence of concentration values at minimum step size	STEP4
Nonconvergence of $[HO_2]$ iteration	CHEM
<b>Other errors</b>	
Program about to run out of time	FINISH

## XI DISPLAY AND ANALYSIS

Concentration values predicted by the Airshed Simulation Program are saved on binary files and are thus available for any display and analysis techniques that might be developed. At present, there is one program available to handle the airshed simulation output files. Currently, it can be used only for the display of the concentration data, but analysis capabilities can be easily added. In this chapter, the general capabilities of the display program are discussed, followed by descriptions and examples of its input and output. A detailed description of the program is given in the Systems Manual.

### A. CAPABILITIES OF THE DISPLAY PROGRAM

The display program provides for two distinct operating modes: prediction and comparison. In the "prediction" mode, concentration predictions are read from one airshed simulation run, and concentration maps and vertical concentration profiles are printed. Capabilities that could be added to the program in this mode include contour mapping of concentration isopleths and calculation of some simple statistics, such as hourly maximum concentrations for each species, frequency of exceeding certain specified levels, and concentrations averaged over time periods longer than that used by the simulation.

In the "comparison" mode, which has been provided for but not yet implemented, one set of concentrations would be subtracted from another, and all further display and analysis would be performed on the resulting array of differences. The comparison mode could be used for validating model predictions against observed data as well as assessing the impact of different model scenarios.

The display program is currently capable of processing a region of up to 30 x 30 grid cells in the horizontal plane, with a maximum of 13 chemical species. These maximum dimensions can be changed by recompiling the program.

## B. INPUT REQUIRED BY THE DISPLAY PROGRAM

As illustrated in Figure XI-1, the input to the display program consists of the following:

- > Card-supplied display options and print controls.
- > Three data files used by the simulation program.
- > One output concentration file from the simulation program.
- > Optionally, a second concentration data file.

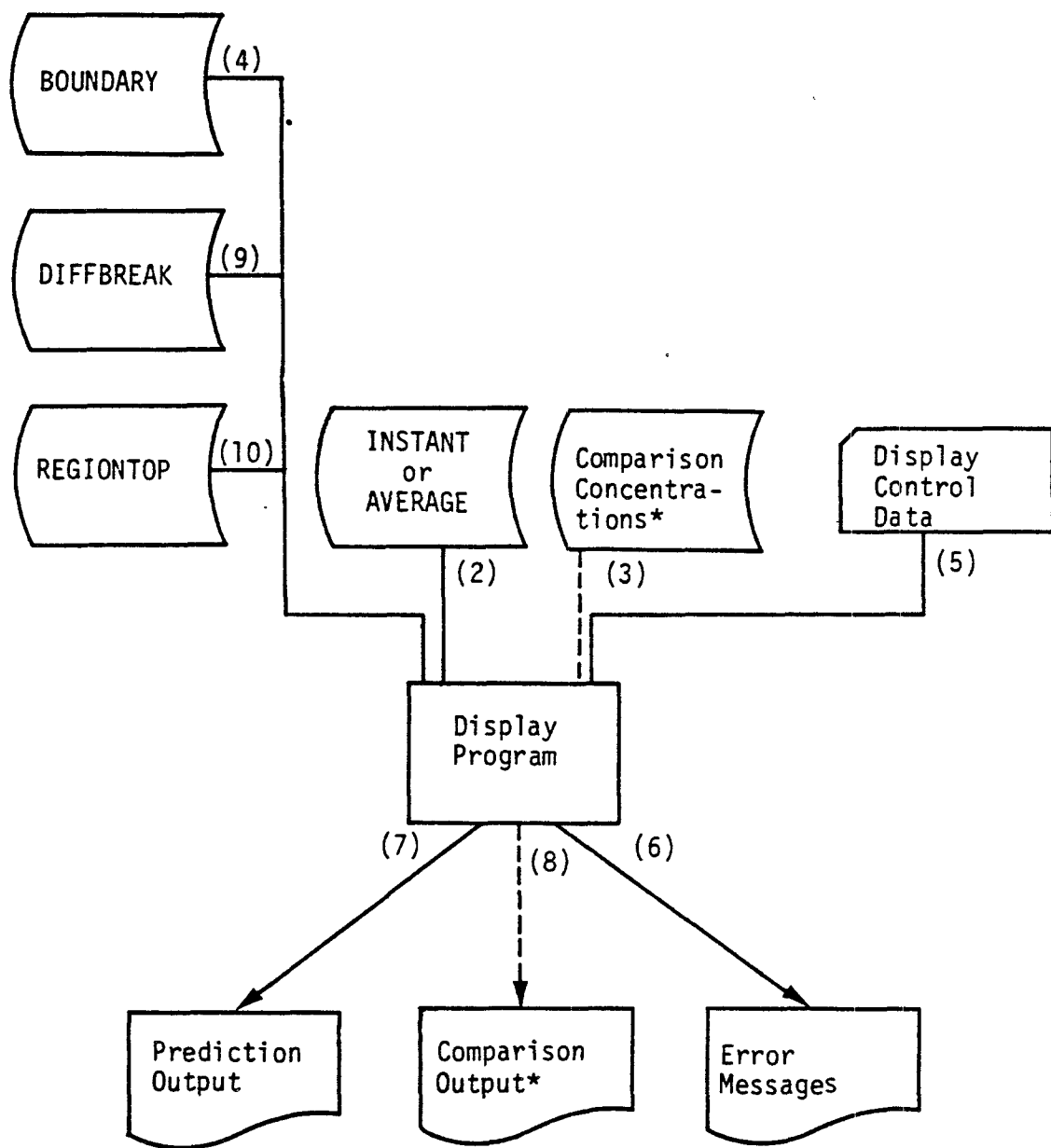
This section discusses first the data files that are to be input, and then details the contents and format of the input cards.

### 1. Data File Input

If concentration maps and vertical profiles are requested (currently the only features available), the BOUNDARY, DIFFBREAK, and REGIONTOP files used for the simulation run must be input. The BOUNDARY file describes the segmentation layout and defines the boundary of the simulated area in the horizontal plane. The DIFFBREAK and REGIONTOP files are used to calculate vertical heights at selected stations. If the simulation has been performed with more than 1 segment, it is the segmented versions of the BOUNDARY, DIFFBREAK, and REGIONTOP files that must be provided to the display program.

One of the Airshed Simulation Program output files containing predicted concentration values must also be input. The concentration values can be instantaneous (INSTANT) or averaged over a time interval (AVERAGE); the output intervals for both files are specified in the SIMCONTROL file. The display program recognizes the file type from the header record, and formats the output accordingly.

If a comparison run is to be made (an option not yet fully implemented), a second concentration file must also be input. This file could contain concentration fields gridded from observed data using the AIRQUALITY data preparation program, or it could contain the results from another airshed simulation run.



\* This option is not yet available.

Figure XI-1. Information Flow Diagram for the Display Program

All input files must contain the same segment structure; that is, all must be either unsegmented or segmented according to the same segment layout.

## 2. Card Input

The display program requires the following data to be input on cards:

- > Output display options
- > Names of species to be examined
- > Vertical level to be considered
- > Print controls
- > Measurement station and landmark names and locations
- > Display units conversion factors
- > User comments to appear on output.

The exact structure of each data card is shown in Table XI-1; Exhibit XI-1 is a sample input deck. Consistent with assumptions used in both the data preparation and simulation programs, the following conventions are generally followed in specifying data fields on each card:

- > Integer input is in 10-character fields, up to six per card.
- > Real number input is floating point in 10-character fields, up to six per card.
- > Alphanumeric input is read and stored as one character per machine word.

Table XI-1. Display Program Control Input Definition

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Comments</u>
<sup>1</sup> <b>Output Options</b>			
KMAP 1-10	I10	Map selection flag: 1, if concentration maps are desired; 0, if no maps are desired	
KCOMP 11-20	I10	Comparison option selection flag: 1, if comparison of two concentration files is required; 0, if no comparison is required	
KSTNFL 21-30	I10	If greater than 0, writes station predictions to unit 12.	
KALLFL 31-40	I10	If greater than 0, writes concentration field predictions to unit 16.	
<sup>2</sup> <b>Species Count</b>			
NSPECP	I10	Number of species to be examined: must be ≤ 13	
<sup>3+</sup> <b>Species Names</b>	((SPNAME(I,J), J = 1, 10), I = 1, NSPECP) 1-60	6(10A1)	Alphanumeric names of species to be examined-- 10 characters per name, left adjusted, six names per card; each name must correspond to a name used in the Airshed Simulation Program run; the species can be named in any order
<sup>4</sup> <b>Vertical Level</b>	NLEVEL	I10	Vertical level to be examined and mapped
<sup>5</sup> <b>Map Print Times (only if KMAP = 1)</b>			Print types must correspond to times for which data on the concentration data files are available
DATPRS	1-10	I10	Print start date (Julian date)
TPRS	F10.2	F10.2	Print start time (hours past midnight)

Table XI-1 (Continued)

Card Number and Name	Item and Columns	Format	Comments
<b>5</b> <b>Map Print Times (con't)</b>			
DTPR	21-30	F10.2	Print interval (hours)
DATPRF	31-40	I10	Print finish date (Julian date)
TPRF	41-50	F10.2	Print finish time (hours past midnight)
The next four cards (Nos. 6 through 9) are to be input only for the comparison mode (not yet implemented)			
<b>6</b> <b>Comparison Times (only if KCOMP = 1)</b>			
DATSCS	1-10	I10	Comparison start date (Julian date)
TSCS	11-20	F10.2	Comparison start time (hours past midnight)
DTSC	21-30	F10.2	Comparison interval (hours)
DATSCF	31-40	I10	Comparison finish date (Julian date)
TSCF	41-50	F10.2	Comparison finish time (hours)
<b>7</b> <b>Comparison Station Count (only if KCOMP = 1)</b>	NSTATC	I10	Number of stations at which concentrations are to be compared; must be $\leq 16$ ; this number is used to determine how many Card 8's are to be read
XI-6	1-10		

Table XI-1 (Continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Comments</u>
8+ Comparison Stations (only if KCOMP = 1)	((STATNA(I,J), J = 1, 20), I = 1, NSTATC) 1-60	3(20A1)	Alphanumeric names of stations at which concentrations are to be compared--20 characters per name, left adjusted, three names per card; each name must match a station name listed on Card 11.
9 Comparison Options (only if KCOMP = 1)	KPLOT 1-10 KSTAT 11-20	I10 I10	Comparison plot flag: 1, if plots are desired; 0, if no plots are needed Statistics option flag: 1, if comparative statistics are to be calculated; 0, if no statistics are to be calculated
10 Point Counts	NOSTN 1-10 NOLAND 11-20	I10 I10	Number of stations at which concentrations are to be calculated--must be $\leq 16$ ; stations are also included on concentration maps Number of landmarks to be included on concentration maps--must be $\leq 8$

Table XI-1 (Continued)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Comments</u>
11+			
Station	Station Name 1-20	20A1	There must be NOSTN of these cards Alphanumeric station name, left adjusted
	x-location 21-30	F10.0	x-coordinate of the station, measured in meters from the reference origin
	y-location 31-40	F10.0	y-coordinate of the station, measured in meters from the reference origin
12+			
Landmark	Landmark name 1-20	20A1	There must be NOLAND of these cards Alphanumeric landmark name, left adjusted
	x-location 21-30	F10.0	x-coordinate of the landmark, measured in meters from the reference origin
	y-location 31-40	F10.0	y-coordinate of the landmark, measured in meters from the reference origin
13+			
Output Units	(IOUNTS(I), I = 1, NSPECP) 1-60	6I10	Output conversion flags, six per card; the number and order must correspond to the species named on Cards 3+; internal units are in $\mu\text{g}/\text{m}^3$ for AEROSOLS and ppm for all other species; different units, however, can be selected for dis- play output using IOUNTS as follows:  1 = ppm; 2 = pptm; 3 = pphm; 4 = ppb; 5 = $\mu\text{g}/\text{m}^3$

Table XI-1 (Concluded)

<u>Card Number and Name</u>	<u>Item and Columns</u>	<u>Format</u>	<u>Comments</u>
14 User Comment	NOTE      1-80	20A4	Note--up to 80 characters--written on header page of display output; it is referred to on that page as "*COMMENTS*"
15 Region Name	REGION      1-80	20A4	Region description note--up to 80 characters--to be used in entitling display output header page

NO CARB	NO2 ARO	O3 CO	ETH	OLE	PAR
1		0			
9					
76195	05.00	1.00	76195	09.01	
21	0				
101		19183.0	27862.0		
102		17518.0	34045.0		
103		22588.0	30467.0		
104		22312.0	25304.0		
105		18706.0	24453.0		
106		13660.0	25566.0		
107		15179.0	30610.0		
108		23407.0	39102.0		
109		30802.0	27886.0		
110		22209.0	20826.0		
111		13812.0	20479.0		
112		8938.0	28913.0		
113		12738.0	37820.0		
114		19320.0	45456.0		
115		32111.0	45799.0		
116		37777.0	38083.0		
117		35560.0	20818.0		
118		18065.0	11256.0		
119		4759.0	18547.0		
120		-1921.0	33909.0		
121		7414.0	50376.0		
4	4	4	4	4	3
4	4	2			
USER COMMENT ST. LOUIS REGION					

Exhibit XI-1. Sample Input for the Display Program

### C. OUTPUT PRODUCED BY THE DISPLAY PROGRAM

The display program first prints a header page, including the alphanumeric region description supplied in the input deck, the dimensions of the simulated area of the grid, the input user comment, and the run identifier from the simulation run that produced the concentration predictions. A sample of the display program header page is shown in Exhibit XI-2.

The display program, operating in prediction mode, currently produces two types of output for each time interval on the input concentration file.

- > Vertical concentration profiles. At each station location input, the concentrations at each vertical level are printed for each designated species. The heights of the diffusion break (mixing depth) and top of the region are also printed at each station. An example of vertical profile output is shown in Exhibit XI-3.
- > Ambient concentration maps. Maps are printed of concentrations in all horizontal grid cells (excluding boundary cells) at the vertical level designated by the user. Printout is confined to those hours and species designated by the user. A separate map is produced for each species at each print time. Station sites and landmarks are also printed on each concentration map. An example of the concentration maps is shown in Exhibit XI-4.

Operating in comparison mode, the display program would calculate residuals for a designated vertical level by subtracting the concentration field on the "comparison" file from the corresponding field on the primary concentration file. Among the statistics calculated for these residuals would be the average value, standard deviation, and correlation coefficient. Calculation of other statistics could be readily incorporated into the program. These statistics would be calculated for each designated species at each print time. Maps of residuals could also be generated for each desired species at each specified print time.

URBAN SCALE AIR POLLUTION SIMULATION MODEL OF  
ST. LOUIS REGION

'INPUT PARAMETERS'

MESH SPACING IN X-DIRECTION 4.0000000+003 MESH SPACING IN Y-DIRECTION 4.0000000+003  
NUMBER OF MESH POINTS IN X-DIRECTION , 15 NUMBER OF MESH POINTS IN Y-DIRECTION , 20  
NUMBER OF VERTICAL MESH POINTS 4

'COMMENTS'

USER COMMENT  
'RUN ID'

RAPS 4-KM DAY 195 -- 4-HR RUN 07/13/76 C-BOND II 6-79

Exhibit XI-2. Standard Display Program Header Page

RUN ID = RAPS 4-KM DAY 195 -- 4-HR RUN 07/13/76 C-BOND II 8-79

VERTICAL AVERAGE CONCENTRATION PROFILES ABOVE SELECTED SITES ON 76195 BETWEEN THE HOURS OF 5 AND 6. LST

		121	.61	MIXING DEPTH= .04	TOP OF REGION=.06	564.0 METERS
NO	(PPB )	1.07	.61			
NO2	(PPB )	6.87	8.26	2.32	2.63	
03	(PPB )	15.20	22.18	77.32	77.13	
ETH	(PPB )	2.43	1.90	1.08	1.08	
OLE	(PPB )	4.10	3.48	2.07	2.07	
PAR	(PPHM)	7.57	5.90	3.32	3.32	
CARB	(PPB )	7.15	6.45	4.29	4.29	
ARO	(PPB )	2.64	2.12	1.22	1.22	
CO	(PPTM)	1.61	1.27	1.02	1.00	

		115	.17	MIXING DEPTH= .06	TOP OF REGION=.06	501.4 METERS
NO	(PPB )	.39	4.17	2.89	2.88	
NO2	(PPB )	5.15	40.40	76.95	76.97	
03	(PPB )	28.81	1.29	1.08	1.08	
ETH	(PPB )	1.53	2.42	2.08	2.08	
OLE	(PPB )	2.78	4.00	3.32	3.32	
PAR	(PPHM)	4.84	4.67	4.28	4.27	
CARB	(PPB )	4.98	1.44	1.22	1.22	
ARO	(PPB )	1.65	1.17	1.01	1.00	
CO	(PPTM)	1.27				

		114	.33	MIXING DEPTH= .05	TOP OF REGION=.06	635.9 METERS
NO	(PPB )	.67	7.04	2.48	2.63	
NO2	(PPB )	8.86	33.67	77.29	76.99	
03	(PPB )	27.19	1.45	1.07	1.08	
ETH	(PPB )	1.70	2.78	2.06	2.07	
OLE	(PPB )	3.14	5.59	3.28	3.32	
PAR	(PPHM)	5.59	4.58	4.30	4.29	
CARB	(PPB )	5.69	5.30	1.22	1.22	
ARO	(PPB )	1.94	1.65	.86	1.00	
CO	(PPTM)	1.18	1.04			

		108	.32	MIXING DEPTH= .06	TOP OF REGION=.07	670.2 METERS
NO	(PPB )	.63	7.61	2.91	3.04	
NO2	(PPB )	10.63	37.10	76.93	76.83	
03	(PPB )	26.26	1.51	1.08	1.08	
ETH	(PPB )	2.01	2.79	2.06	2.07	
OLE	(PPB )	3.35	4.82	3.32	3.32	
PAR	(PPHM)	6.97	5.30	4.30	4.29	
CARB	(PPB )	5.89	1.75	1.22	1.22	
ARO	(PPB )	2.67	1.22	1.00	1.00	
CO	(PPTM)	1.47				

		116	.18	MIXING DEPTH= .06	TOP OF REGION=.06	594.6 METERS
NO	(PPB )	.54	4.10	2.89	2.88	
NO2	(PPB )	5.89	44.22	77.00	76.98	
03	(PPB )	30.56				

Exhibit XI-3.

Sample Vertical Concentration Profile Output--Prediction Mode

RUN ID =RAPS 4-KM DAY 195 -- 4-HR RUN 07/13/76 C-BOND II 8-79

PANEL 1	AVERAGE AMBIENT CONCENTRATIONS(PPB ) OF 03															ON 76195 BETWEEN THE HOURS OF			
*	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	*	*	*	*
21 *	22	21	21	21	21	21	22	22	23	24	24	24	23	23	21	*	21	*	*
20 *	20	19	18	18	18	19	20	21	23	24	24	24	23	24	23	*	20	*	*
19 *	18	17	16	15	16	18	20	20	20	22	24	24	23	24	23	*	19	*	*
18 *	17	15	13	13	14	16	20	21	20	19	20	24	25	25	24	*	18	*	*
17 *	15	13	11	11	12	14	21	25	26	24	22	23	27	25	24	*	17	*	*
16 *	13	10	7	8	13	14	19	25	28	27	27	29	29	27	24	*	16	*	*
15 *	13	19	9	10	12	14	17	20	24	26	28	30	31	28	24	*	15	*	*
14 *	15	14	12	12	15	18	19	20	24	26	30	32	32	30	25	*	14	*	*
13 *	10	13	12	13	17	18	17	16	21	26	33	33	32	30	26	*	13	*	*
12 *	6	11	12	12	16	14	14	14	18	29	34	34	32	30	26	*	12	*	*
11 *	11	9	10	13	14	12	9	9	21	30	37	36	34	30	26	*	11	*	*
10 *	15	12	13	15	17	16	16	21	25	29	37	35	34	29	25	*	10	*	*
9 *	17	15	13	14	17	18	19	22	31	33	33	32	31	29	27	*	9	*	*
8 *	16	17	15	15	14	16	22	26	29	31	32	31	29	27	25	*	8	*	*
7 *	18	17	17	17	19	21	26	28	30	31	31	29	26	24	24	*	7	*	*
6 *	19	18	18	18	18	22	24	25	27	29	31	31	29	23	23	*	6	*	*
5 *	20	20	19	19	18	22	24	25	27	28	28	27	26	24	23	*	5	*	*
4 *	22	21	20	19	18	23	24	25	26	27	27	26	24	23	23	*	4	*	*
3 *	23	22	21	19	15	25	25	26	26	26	26	25	24	23	2	*	3	*	*
2 *	23	23	22	22	24	24	25	25	25	25	25	24	23	2	*	2	*	*	*

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TECHNICAL REPORT DATA <i>(Please read Instructions on the reverse before completing)</i>		
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16. ABSTRACT  This report presents a general view of the Systems Applications, Inc. (SAI) Airshed Model as well as detailed operating instructions for the user. Included are discussions of all the files needed to run the model, the data preparation programs that produce each file, the input formats and data preparation methods, samples of input and output, and information flow diagrams that illustrate the job stream control on any computer. Chapter II presents a technical summary of the SAI Airshed Model; Chapter III contains an overview of the model as a system of computer programs, outlining the five main functions that must be performed by users and programs together. Chapter IV discusses some of the concepts and terms basic to an understanding of the system, followed in Chapter V by descriptions of the job control notations that will be used throughout this manual. Chapters VI through XI examine in detail the five main functions of the system, providing input and output definitions and examples, and job control diagrams for each program that must be exercised.		
17. KEY WORDS AND DOCUMENT ANALYSIS		
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