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## **HYSPLIT\_4 USER's GUIDE**

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## INFORMATION

This document discusses the technical aspects of the installation and operation of the Hysplit4 model version designed to run on Windows 95/98/NT platforms. The executable code can be obtained at: <http://www.arl.noaa.gov/hysplit.html>.

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# HYSPLIT\_4 USER's GUIDE

**ABSTRACT.** The HYSPLIT\_4 (Hybrid Single-Particle Lagrangian Integrated Trajectory) Model installation, configuration, and operating procedures are reviewed. Examples are given for setting up the model for trajectory and concentration simulations, graphical displays, and creating publication quality illustrations. Programs that can be used to create the model's meteorological input data are described.

## 1. MODEL OVERVIEW

### 1.1 Features

The HYSPLIT\_4 (HYbrid Single-Particle Lagrangian Integrated Trajectory) model is a complete system for computing trajectories to complex dispersion and deposition simulations using either puff or particle approaches.<sup>1</sup> It consists of a modular library structure with main programs for each application: trajectories and air concentrations.

Gridded meteorological data, on one of three conformal (Polar, Lambert, Mercator) map projections, are required at regular time intervals. The input data are interpolated to an internal sub-grid to reduce memory requirements and increase computational speed. Calculations may be performed sequentially on multiple meteorological grids, usually specified from fine to coarse resolution.

Air concentration calculations require the definition of the pollutant's emissions and physical characteristics (if deposition is required). When multiple pollutant species are defined, an emission would consist of one particle or puff associated with each pollutant type. Alternately, the mass associated with a single puff may contain several species. The latter approach is used for calculation of chemical transformations when all the species follow the same transport pathway. Chemical transformation subroutines are not part of the model distribution.

The dispersion of a pollutant is calculated by assuming either a Gaussian or Top-Hat horizontal distribution within a puff or from the dispersal of a fixed number of particles. A single released puff will expand until its size exceeds the meteorological grid cell spacing and then it will split into several puffs. An alternate approach combines both puff and particle methods by assuming a puff distribution in the horizontal and particle dispersion in the vertical direction. The resulting calculation may be started with a single particle. As its horizontal distribution expands beyond the grid length scale, it will split into multiple particle-puffs, each with their respective fraction of the pollutant mass. In this way, the greater accuracy of the vertical dispersion parameterization of the particle model is combined with the advantage of having an expanding number of particles represent the pollutant

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<sup>1</sup>Draxler, R.R., and G.D. Hess, 1998, An overview of the Hysplit\_4 modelling system for trajectories, dispersion, and deposition, *Australian Meteorological Magazine*, 47, 295-308.

distribution as the spatial coverage of the pollutant increases.

Air concentrations are calculated at a specific grid point for puffs and as cell-average concentrations for particles. A concentration grid is defined by latitude-longitude intersections. Simultaneous multiple grids with different horizontal resolutions and temporal averaging periods can be defined for each simulation. Each pollutant species is summed independently on each grid.

The routine meteorological data fields required for the calculations may be obtained from existing archives or from forecast model outputs already formatted for input to Hysplit. In addition, pre-processor programs are provided to convert NOAA, NCAR (National Center for Atmospheric Research) re-analysis, or ECMWF (European Centre for Medium-range Weather Forecasts) model output fields to a format compatible for direct input to the model. The model's meteorological data set structure is compressed and in "direct-access" format. Each time period within the data file contains an index record that includes grid definitions to locate the spatial domain, check-sums for each record to ensure data integrity, variable identification, and level information. These data files require no conversion between computing platforms.

The modeling system includes a Graphical User Interface (GUI) to set up a trajectory, air concentration, or deposition simulation. The post-processing part of the model package incorporates graphical programs to generate multi-color or black and white publication quality Postscript printer graphics.

A complete description of all the equations and model calculation methods for trajectories and air concentrations has been published,<sup>2</sup> and is also available on-line (<http://www.arl.noaa.gov/hysplit.html>).

## 1.2 Pre-Installation Preparation

Although the self-installing executable, *hysplit4{x}.exe*, does not require any additional software, it will only provide a command line interface to the model. To enable the model's GUI, the computer should have Tcl/Tk script language installed. It can be obtained over the Internet from: <http://dev.scriptics.com>. The installation of Tcl/Tk will result in the association of the *.tcl* suffix with the *Wish* executable and all Hysplit GUI scripts will then show the Tk icon.

The Hysplit GUI contains options to convert either the trajectory or concentration model output files to Postscript. The Postscript files can also be viewed directly through the GUI if a Postscript viewer, such as Ghostscript has been installed prior to the Hysplit installation. See [www.cs.wisc.edu/~ghost](http://www.cs.wisc.edu/~ghost) for more information on Postscript viewers.

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<sup>2</sup>Draxler, R.R., and G.D. Hess, 1997, Description of the Hysplit\_4 modeling system, NOAA Technical Memorandum ERL ARL-224, December, 24p .

### 1.3 Windows 95/98/NT Installation

Hysplit installation to a computer running Windows (16 bit versions are not supported) is provided through a self installing file. Executables are installed in various directories for trajectories, dispersion, display of results, manipulation of results, and creation of input meteorological data files. The trajectory and dispersion model source code is not provided. However all the Fortran source code to create meteorological data files in a format that the model can read are provided in the \metdata directory. Each subdirectory contains a *Readme.txt* file with more complete information about the contents of that directory.

At the beginning of the installation you will be prompted as to the directory location. It is suggested you select the default location (*C:\hysplit4*). The installation program is very simple and although the selection of a different drive or directory will install the code in the selected directories, the shortcut links will not be placed correctly in the Start Menu or Desktop. In this situation you will need to edit the *\icons\setup.bat* file for the correct drive and directory locations.

The following subdirectories will be created after the installation has completed:

**bdyfiles** - This directory contains an ASCII version of gridded land use, roughness length, and terrain data. The current file resolution is 360 x 180 at 1 degree. The upper left corner starts at 180 W, 90 N. The files are read by both Hysplit executables, *hymodelt* and *hymodelc*, from this directory. If not found, the model uses default constant values for land-use and roughness length. The data structure of these files is defined in the file *ASCDATA.CFG*, which should be located in either the model's startup directory or the *\bdyfiles* directory. This file defines the grid system and gives an optional directory location for the land-use and roughness length files. These files may be replaced by higher resolution customized user-created files. However, regardless of their resolution, the model will only apply the data from these files at the same resolution as the input meteorological data grid. More information on the structure of these files can be found in the local *Readme.txt* file.

**data2arl** - Current forecast or archive meteorological data can be obtained from the ARL ftp server: *ftp://gus.arlhq.noaa.gov/pub/archives (or /forecasts)*. Older archive data can be ordered from the NCDC (National Climatic Data Center). However if you have access to your own meteorological data or data formatted as GRIB (Gridded Binary), this directory contains various example decoder programs to convert meteorological data in various formats to the format (ARL packed) that Hysplit can read. Sample programs include GRIB decoders for ECMWF model fields, NCAR/NCEP (National Centers for Environmental Prediction) re-analysis data, and NOAA Aviation and Regional Spectral Model files. All the required packing and unpacking subroutines can be found in the various subdirectories of *\data2arl*. More information on these programs can be found in Section 5.4.

**concmdl** - The directory contains the Hysplit4 air concentration prediction model (*hymodelc*) and related display programs. Although the model can be run through the GUI, at times it may be desirable to run the model from the command line (e.g. using automated scripts). The example *Control* file should produce some results for viewing. The sequence of

commands would be *hymodelc* to execute the model, and *concpplot* to create a Postscript file called *concpplot.ps*. Command line arguments are required for *concpplot*. Normally the GUI is used to create the *Control* file for the simulation. If the file is missing, the model will prompt you for input from the keyboard. Your inputs are copied to a file called *Startup*. That file may then be edited and renamed to *Control* for subsequent simulations. *Conread* and *con2asc* are provided as examples of how to read concentration files for users to develop other applications.

**document** - This directory contains PDF (Adobe Portable Document Format) versions of the User's Guide and other documentation such as ARL-224, the principal ARL Technical Memorandum describing the model and equations. This User's Guide (this document) provides detailed instructions in setting up the model, modifications to the *Control* file to perform various simulations and output interpretation. The *Readme.txt* file contains additional information about compilation, typical CPU times, and a summary of recent model updates.

**graphics** - There are two types of graphical plotting programs provided in the *\concmdl* and *\trajmdl* directories. Publication quality graphics can be created using the postscript conversion programs, *concpplot* and *trajplot*, which use a Fortran Postscript library created by Kevin Kohler<sup>3</sup>. All graphical routines use the map background file *arlmap* in this directory. The map background file uses a simple ASCII format and contains the world's coastal and political boundaries at relatively coarse resolution. Other higher resolution map background files are available from the Hysplit download web page. All graphical programs search the startup directory first for *arlmap* before going to *\graphics*, therefore customized maps can be created without changing the Hysplit installation structure.

**guicode** - This directory contains a Tcl/Tk GUI interface source code script for Hysplit. The interface is used to set up the input *Control* file as well as run the graphical output display programs. To use the interface you must first install Tcl/Tk. The upper-level Tcl script is called *hysplit4.tcl*, which calls all other Tcl scripts. The Hysplit GUI is started by executing this script. The Desktop short-cut as well as the Start Menu options should point to this script. If the installation program did not properly setup the Desktop, you should copy the shortcut from *\working* or you can manually create a short-cut to the script and edit its properties such that the "Start In" directory is *\working*. You should also select the Hysplit icon from the *\icons* directory before moving it to the Desktop.

**icons** - Normally if you install Hysplit to the default drive *c:\hysplit4*, there should be a desktop icon as well as entries in the Start Programs menu. If these short-cuts were not set up properly, you can edit the *Setup.bat* file to reflect your directory structure. If you still have trouble, then the icons supplied in this directory can be used to replace the default Windows icons if you create shortcuts to Hysplit from the desktop. Use the right mouse button and select: Properties | Shortcut | Change Icon. Note that the *\working\hysplit4.tcl* shortcut should be copied to the desktop.

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<sup>3</sup>PSPLOT libraries can be found at [www.nova.edu/ocean/psplot.html](http://www.nova.edu/ocean/psplot.html) and were created by Kevin Kohler ([kevin@ocean.nova.edu](mailto:kevin@ocean.nova.edu)).



**metdata** - This directory contains the sample meteorological data file: *oct1618.BIN*. It is an extract of the NGM (NOAA's Nested Grid Model) over the US from 0000 UTC 10/16/95 through 0000 UTC of 10/18/95. The file is used for all calculations shown in the User's Guide. In addition, several sample programs are provided that can be used to examine and display the meteorological data files. Source code for some of these routines can be found in the *\source* subdirectory. More information on these programs can be found in Section 5.4.

**trajmdl** - This directory contains the Hysplit4 trajectory model (*hymodelt*) and related display programs. Although the model can be run through the GUI, at times it may be desirable to run the model from the command line (e.g. automated scripts). The example *Control* file should produce some results for viewing. The sequence of commands would be *hymodelt* to execute the model, and *trajplot* to create a Postscript file called *trajplot.ps*. Normally the GUI is used to create the *Control* file for the simulation. If the file is missing, the model will prompt you for input from the keyboard. Your inputs are copied to a file called *Startup*. That file may then be edited and renamed to *Control* for subsequent simulations.

**visualization** - contains Fortran source code and some executables to convert trajectory, air concentration, and meteorological data files for input to GRADS and VIS5D.

**working** - This should be the default working directory when running the model through the GUI. The properties of the Hysplit4 shortcut should point to this as the startup directory. This directory will contain all the user created input and output files unless they are explicitly directed to be read or written from/to another directory in the input *Control* file, such as meteorological data files that might be found in *\metdata*. A sample tcl script, *Auto\_traj.tcl*, is provided as an example of how one might automate multiple trajectory calculations. The script creates the *Control* file and executes the model in a loop, varying specific parameters with each simulation.

## 1.4 Quick Start

The easiest way to run the model is to use the GUI menu to edit the model's input *Control* file. For the purposes of this demonstration appropriate meteorological files are provided. If for some reason the menu system is not available, the *Control* file can be created manually. See the discussion in Section 2.

Step 1 - start the GUI menu system using *\working\hysplit4.tcl* or the desktop shortcut to *Hysplit4*. A widget will appear with the *HYSPLIT* graphic and two button options: *Menu* and *Exit*. On some systems the graphic may be scrambled or the colors may be flat. Switching the PC display to VGA and 256 colors usually solves these problems. However it can be left alone as the faulty graphic will not affect any of the other widgets or display graphics.

Step 2 - from the *HYSPLIT* graphic widget click on *Menu*. The three main menus of Hysplit4 will appear: *Meteorology*, *Trajectory*, *Concentration*, and on some custom installations the optional menu: *Advanced*. An additional small widget underneath the main menu gives the current Hysplit4 version information. Do not delete this widget as it will

terminate the GUI. It provides the reference frame for the model's standard output and messages.

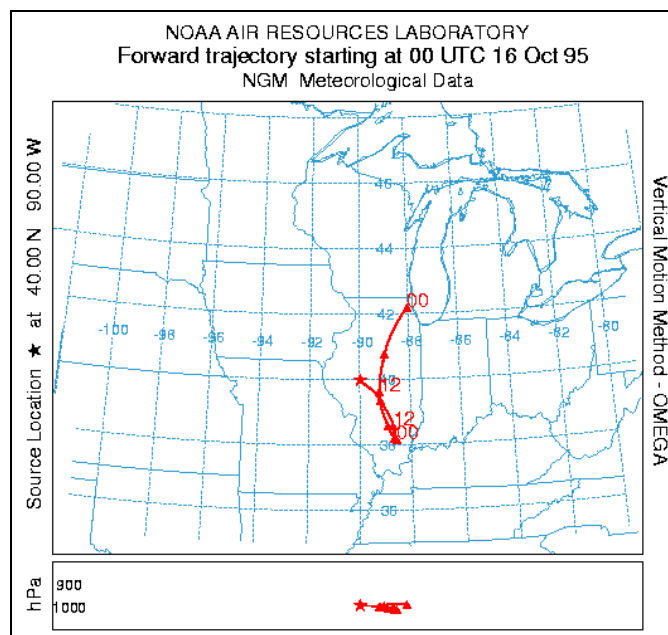
Step 3 - for the first example calculation select the *Trajectory* option. Four options appear under this item: *Trajectory Setup*, *Run Model*, *Trajectory Display*, and *Utility Programs*. Normally these are run in sequence, however any item can be selected and run if the appropriate input files were created during a previous simulation. Currently no utility program options are available.

Step 4 - *Trajectory Setup* is used to enter the basic model simulation parameters: the starting time of the calculation; starting location in terms of latitude, longitude, and height; the run-time or duration of the trajectory calculation; and the names and locations of all required files. When modifications to this menu are complete, click on *Save*. However for this example, you will use the *Retrieve* option for predefined configurations, so do nothing here and go on to Step 5.

Step 5 - for the sample calculation click on *Retrieve*, enter name of the example pre-configured control file: *sample\_traj*, then click on *OK*, then after the data entry widget is closed, click on *Save* and the setup menu will close.

Step 6 - *Run Model* copies the setup configuration to the model's input *Control* file and starts the model calculation. Messages will appear on standard output showing the progress of the calculation. When the simulation is completed, the trajectory end-points output file is ready to be converted to a graphical display. Under Windows 95/98 the standard output widget will not show any output until the end of the calculation and the *Trajectory* menu items will be locked until the calculation completes.

Step 7 - selecting *Display* will run a special program that converts the ASCII file of trajectory end-point positions into a high quality Postscript file (*trajplot.ps*) suitable for printing. The conversion widget provides options for the frequency of the labels on the trajectory, a "HiResolution" (zoomed) display, and color or black and white graphics. If a Postscript viewer (*Ghostscript / Ghostview*) has been installed and associated with the .ps file suffix, then it will be invoked by the GUI. If the viewer does not automatically open, it may be necessary to manually edit *hysplit4.tcl* to change the directory location associated with the program *gsview32.exe*. The sample output from the Postscript file is shown in Fig. 1.



**Figure 1.** Example from Postscript conversion.

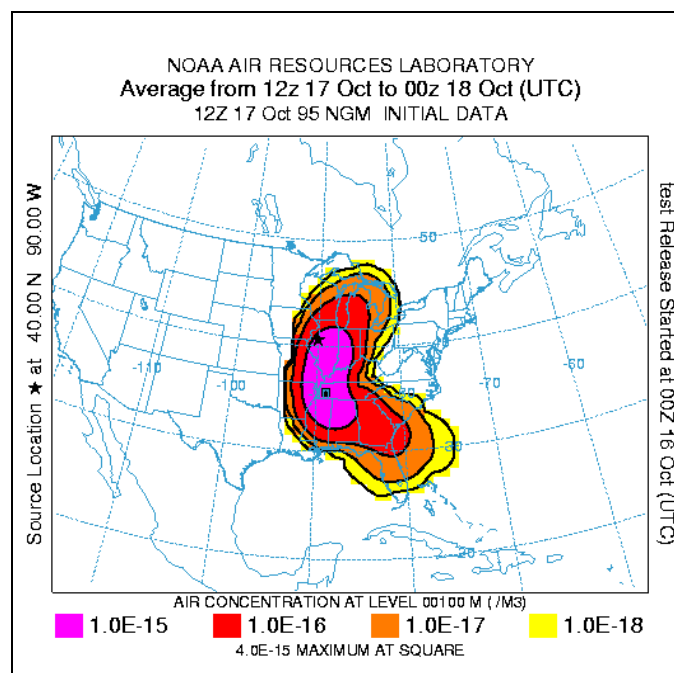
In the example case, trajectory positions are marked at 6-h intervals and the vertical projection is shown in the lower panel with each point below its corresponding point on the horizontal projection..

Step 8 - for the second example calculation select *Concentration*. Under this menu there are also four options: *concentration setup*, *run model*, *concentration display*, and *Utility Programs*. In general they should be executed in sequential order.

Step 9 - selecting *Concentration Setup* brings up similar starting information as with trajectories, but with three additional sub-menus (*Pollutant*) that can be used set the emission rate, duration, and start time of the emission; (*Grids*) to set the location, resolution, levels, and averaging times of the concentration output grid; and (*Deposition*) to set the characteristics of each pollutant. Click on *Retrieve*, enter name of sample pre-configured control file: *sample\_conc*, then click on *OK*, then after the data entry widget is closed, click on *Save* and the setup menu will close.

Step 10 - selecting *Run Model* copies the setup configuration to the model's input *Control* file and starts the model calculation. Messages will appear on standard output showing the progress of the calculation after the calculation has completed. At that point the binary gridded concentration output file is ready to be converted to a graphical display. Be patient as concentration calculations may take considerably longer than trajectory calculations.

Step 11 - selecting *Display* will run a special program that converts the binary concentration file to a Postscript file (*concpplot.ps*) suitable for printing. The display widget contains multiple options for different pollutants (if defined), data grids, levels, and contour options. These are discussed in more detail in the "Graphics" section. For this example accept the defaults and just click on "Execute Display." If the *Ghostview* Postscript viewer has been installed and properly associated with .ps files, then it will be automatically invoked by the GUI. If the viewer does not open, it may be necessary to manually edit the file *hysplit4.tcl* for the directory entry associated with the program *gsview32.exe*. The sample output from the Postscript file is shown in Fig. 2. The output file can be printed directly on any Postscript printer or printed through *Ghostscript*.



**Figure 2.** Example from the Postscript conversion.

Step 12 - selecting *Utility Programs* brings up options for three different utility programs:

"*Convert to ASCII*" which will run a program that converts the binary air concentration file to an ASCII file with one record for each non-zero grid point giving the latitude-longitude position and air concentration. The file can be used as input to GIS or other visualization programs; "*Grid to Station*" will extract a concentration time series at one or more locations to an output text file; and "*Select from Display*" shows a coarse graphic screen display of the concentration field and uses the mouse to selection of points at which the position and concentration value is written to a text file. These programs will be discussed in more detail in the graphics section.

## 1.5 Problems

If Tcl/Tk does not exist on your system or there are other problems with the GUI interface, it is very easy to run the sample cases directly in either the `\trajmdl` or `\concmdl` sub-directories by double-clicking on *hymodelt*, *trajplot* for trajectories and *hymodelc*, *concpplot* for air concentrations. If the sample simulation works well, then it is only necessary to manually edit the *Control* file to try out different simulation variations. The other options are explained in more detail in Section 2.

In general, premature termination during the model initialization phase will result in messages to standard output. However after the model has started, fatal, diagnostic, and progress notification messages are written to a file called *Message*. If the model output is not what you expected, first check the *Control* file to determine if the input setup is what is desired, then check the *Message* file for indication of abnormal performance. These files are always written to the model's startup directory - `\working` if the model is run from the GUI.

## 2. ADVANCED SIMULATION CONTROL

When the model starts it looks for an input file called *Control*. If found, this file is used to read all input parameters. If not found, a prompt will appear on standard input requesting appropriate information. These prompts are described in more detail below and are identical to the comparable entries through the GUI menu. When data entry is through the keyboard, a file named *Startup* is created. This contains a copy of the input, and which later may be renamed to *Control* to permit direct editing and model execution without data entry. If you are unsure as to a value required in an input field, just enter the forward slash (/) character, the indicated default value will be used. This default procedure is valid for all input fields except directory and file names. An automatic default selection procedure is also available when the input fields are read from the *Control* file for certain fields when they are set to zero. Those options are discussed in more detail below. Each input line is numbered (only in this text) according to the order it appears in the file. A number in parenthesis after the line number indicates that there is an input loop and multiple entry lines may be required depending upon the value of the previous entry.

### 2.1 Trajectories

1- Enter starting time (year, month, day, hour) Default: 0 0 0 0

Enter the two digit values for the UTC time that the calculation is to start. Use 0's to start at the beginning (or end) of the file according to the direction of the calculation. Zero's will force the calculation to use the time of the first (or last) record of the meteorological data file.

2- Enter number of starting locations Default: 1

Simultaneous trajectories can be calculated at multiple levels or starting locations. The maximum number depends upon the compilation parameters. The GUI menu can accommodate up to 6 simultaneous starting locations. Specification of additional locations requires manual editing of the *Control* file.

3(1)- Enter starting location (lat, lon, meters) Default: 40.0 -90.0 50.0

Position in degrees and decimal (West and South are negative). Height is entered as meters above ground. See section 4.4 on how to set heights relative to mean sea level.

4- Enter total run time (hours) Default: 48

Specify the duration of the calculation in hours. Backward calculations are entered as negative hours.

5- Vertical motion option (0:data 1:isob 2:isen 3:dens 4:sigma) Default: 0

Indicates the vertical motion calculation method. The default "data" selection will

use the meteorological model's vertical velocity fields; other options include isobaric, isentropic, constant density, and constant sigma (internal model coordinate).

6- Top of model domain (internal coordinates m-agl) Default: 10000.0

Vertical limit of the internal meteorological grid. If calculations are not required above a certain level, fewer meteorological data are processed thus speeding up the computation. Trajectories will terminate when they reach this level.

A secondary use of this parameter is to set the model's internal scaling height – the height at which the internal sigma surfaces go flat relative to terrain. The default internal scaling height is set to 25 km but it is set to the top of the model domain if the entry exceeds 25 km. Further, when meteorological data are provided on terrain sigma surfaces it is assumed that the input data were scaled to a height of 20 km. If a different height is required to decode the input data it should be entered on this line as a negative of the height. Hysplit's internal scaling height remains at 25 km unless the absolute value of the domain top exceeds 25 km.

7- Number of input data grids Default: 1

Number of simultaneous input meteorological files. The following two entries (directory and name) will be repeated this number of times. Always start with the finest resolution grid as number #1. When the computation shifts from grid #1 to #2, it will not return to #1 again. The run will terminate when the computation is off the last grid. Multiple trajectory calculations will switch grids for all trajectories when the first trajectory passes to the new grid. Multiple grid definitions also have the restriction that there should be some overlap between the grids in space. Time overlap, although desirable, is not required. However, without time overlap it is not possible to interpolate in time across grids and hence start trajectories at those times.

8(1)- Meteorological data grid # 1 directory Default: \main\sub\data\

Directory location of the meteorological file on the grid specified. Always terminate with the appropriate slash (\).

9(2)- Meteorological data grid # 1 file name Default: file\_name

Name of the file containing meteorological data. Located in the previous directory.

10- Directory of trajectory output file Default: \main\trajectory\output\

Directory location to which the ASCII trajectory end-points file will be written. Always terminate with the appropriate slash (\).

11- Name of the trajectory endpoints file Default: file\_name

The trajectory end-points output file is named in this entry line. The format of the

file is given in Section 6.

## 2.2 Air Concentration

The entries in the *Control* file for air concentration simulations consist of four groups of input data. The first group is almost identical to the trajectory simulation. The second group defines the pollutant emission characteristics. The third group defines the concentration grid in terms of spacing and integration interval. The fourth group of entries defines pollutant characteristics relevant to computing deposition and removal processes.

### 2.2.1 Meteorological Simulation Entries

1- Enter starting time (year, month, day, hour) Default: 0 0 0 0

Enter the two digit values for the UTC time that the calculation is to start. Use 0's to start at the beginning of the file. Note that the calculation start time may be different from the emission start time that is specified below. However the simulation start time may not occur after the emission start time.

2- Number of starting locations Default: 1

Multiple pollutant sources may be simultaneously tracked. The emission rate specified below is assigned to each source. In addition, the emissions are distributed vertically in a layer between the current emission height and the previous source emission height if the previous source is at the same location. The effective source will be a vertical line source between the two heights. When multiple sources are in different locations, the pollutant is emitted as a point source from each location at the height specified. Point and vertical line sources can be mixed in the same simulation. The GUI menu can accommodate up to 6 simultaneous starting locations. Specification of additional locations requires manual editing of the *Control* file. Area source emissions can be specified from an input file: *emission.txt*. When this file is present in the root directory, the emission parameters in the *Control* file are superseded by the emission rates specified in the file. More information on this file structure can be found in Section 4.2.

3(1)- Enter starting location (lat, lon, meters, Opt-4, Opt-5) Default: 40.0 -90.0 50.0

Position in degrees and decimal (West and South are negative). Height is entered as meters above ground level unless the mean-sea-level flag has been set (see section 4.4).

The optional 4<sup>th</sup> (emission rate - units per hour) and 5<sup>th</sup> (emission area - square meters) columns on this input line can be used to supercede the value of the emission rate (line 12-2) when multiple sources are defined, otherwise all sources have the same rate as specified on line 12-2. The 5<sup>th</sup> column defines the virtual size of the source: point sources default to "0".

4- Enter total run time (hours)

Default: 48

The duration of the calculation in hours. Backward calculations are indicated by a negative run time. See discussion in Section 4 on backward "dispersion" calculations.

5- Vertical (0:data 1:isob 2:isen 3:dens 4:sigma)

Default: 0

Indicates the vertical motion calculation method. The default "data" option uses the meteorological model's vertical velocity fields; other options include isobaric, isentropic, constant density, and constant sigma above terrain.

6- Top of model domain (internal coordinates m AGL)

Default: 10000.0

Vertical limit of the internal grid. If calculations are not required above a certain level, fewer meteorological data are processed, thus speeding up the computation. Particles and puffs are restricted from mixing above this level. Complete reflection is assumed.

A secondary use of this parameter is to set the model's internal scaling height - the height at which the internal sigma surfaces go flat relative to terrain. The default internal scaling height is set to 25 km but it is set to the top of the model domain if the entry exceeds 25 km. Further, when meteorological data are provided on terrain sigma surfaces it is assumed that the input data were scaled to a height of 20 km. If a different height is required to decode the input data it should be entered on this line as a negative of the height. Hysplit's internal scaling height remains at 25 km unless the absolute value of the domain top exceeds 25 km.

7- Number of input data grids

Default: 1

Number of simultaneous meteorological fields to be input. The following two entries (directory and name) will be repeated this number of times. Always start with the finest resolution grid as number #1. When the first puff or particle moves on to the next grid, all subsequent particles are automatically transferred to the new grid. There should be time and space overlap for multiple grids.

8(1)- Enter grid # 1 directory

Default: \main\sub\data\

Directory location of the meteorological files. Terminate with appropriate (\) slash.

9(2)- Enter grid # 1 file name

Default: file\_name

Name of the meteorological data file.



## 2.2.2 Pollutant Definition Entries

### 10- Number of different pollutants

Default: 1

Multiple pollutant species may be defined for emission. Each pollutant is assigned to its own particle or puff and therefore may behave differently due to deposition or other pollutant specific characteristics. Each will be tracked on its own concentration grid. The following four entries are repeated for each pollutant defined. Although the GUI shows that up to seven pollutants can be defined, the compilation default is to permit only two simultaneous definitions.

### 11(1)- Pollutant four Character Identification

Default: TEST

Any four-character label that can be used to identify the pollutant. The label is written with the concentration output grid to identify output records associated with that pollutant and will appear in display labels. Additional user supplied deposition and chemistry calculations may be keyed to this identification string.

### 12(2)- Emission rate (per hour)

Default: 1.0

Mass units released each hour. Units are arbitrary except when specific chemical transformation subroutines are associated with the calculation. Output air concentration units will be in the same units as specified on this line: input kg/hr -> output kg/m<sup>3</sup>; input Bq/hr -> output Bq/m<sup>3</sup>. When multiple sources are defined this rate is assigned to all sources unless optional parameters are present on line 3(1).

### 13(3)- Hours of emission

Default: 1.0

The duration of emission may be defined in fractional hours. Durations of less than one time-step will be emitted over one time-step with a total emission that would yield the requested rate over the emission duration. "Backward" simulations require a negative value for the hours of emission.

### 14(4)- Release start time: year month day hour minute

Default: [simulation start time]

The previously specified hours of emission start at this time. An entry of zero's in the field, when input is read from a file, will also result in the selection of the default values. "Backward" calculations require this field to be set with explicit rather than relative or default values.

## 2.2.3 Concentration Grid Definition

Dispersion calculations are performed on the computational (meteorological) grid without regard to the definition or location of any concentration grid. Therefore it is possible to complete a simulation and have no results to view if the concentration grid was in the

wrong location. In addition, the concentration grid spacing may restrict the model's integration time step to a smaller value for higher resolution concentration grids. This section is used to define the grid system to which the concentrations are summed during the integration and subsequently for postprocessing and display of the model's output.

15- Number of simultaneous concentration grids

Default: 1

Multiple or nested grids may be defined. The concentration output grids are treated independently. The following 10 entries will be repeated for each grid defined. The number of grids permitted depends upon model compilation parameters – the default compilation is usually supports two independent grids.

16(1)- Center Latitude, Longitude (degrees)

Default: [source location]

The center position of the concentration sampling grid in degrees and decimal. Input of zero's will result in selection of the default value: location of the emission source. Sometimes it may be desirable to move the grid center location downwind near the center of the projected plume position.

17(2)- Grid spacing (degrees) Latitude, Longitude

Default: 1.0 1.0

The interval in degrees between nodes of the sampling grid. Puffs must pass over a node to contribute concentration to that point and therefore if the spacing is too wide, they may pass between intersection points. Particle model calculations represent grid-cell averages, where each cell is centered on a node position, with its dimensions equal to the grid spacing. Finer resolution concentration grids require correspondingly finer integration time-steps. This may be mitigated to some extent by limiting fine resolution grids to only the first few hours of the simulation.

18(3)- Grid span (deg) Latitude, Longitude

Default: [max-Y / d-lat] [max-X / d-lon]

The total span of the grid in each direction. For instance, a span of 10 degrees would cover 5 degrees on each side of the center grid location. A default span is computed from the compiled maximum dimensions of the concentration grid divided by the grid spacing requested in the previous entry. The typical default compilation size supports a 300x300 node grid. If a grid resolution of 0.1 deg (about 10 km) was selected, then the maximum grid span would be 30 degrees latitude-longitude. A plume that goes off the grid would have cutoff appearance. This can sometimes be mitigated by moving the grid center further downwind.

19(4)- Enter grid # 1 directory

Default: \main\sub\output\

Directory to which the binary gridded concentration output file for this grid is written. As in other directory entries a terminating (\) slash is required.

20(5)- Enter grid # 1 file name Default: file\_name

Name of the concentration output file for each grid. See Section 6 for a description of the format of the concentration output file.

21(6)- Number of vertical concentration levels Default: 1

The number of vertical levels in the concentration grid including the ground surface level if deposition output is required. The default compilation usually supports up to 10 levels.

22(7)- Height of each level (m) Default: 50

Output grid levels may be defined in any order for the puff model as long as the deposition level (0) comes first (a height of zero indicates deposition output). Air concentrations must have a non-zero height defined. A height for the puff model indicates the concentration at that level. A height for the particle model indicates the average concentration between that level and the previous level (or the ground for the first level). Therefore heights for the particle model need to be defined in ascending order. Note that the default is to treat the levels as above-ground-level (AGL) unless the the MSL (above Mean-Sea-Level) flag has been set (see Section 4.4).

23(8)- Sampling start time: year month day hour minute Default: [simulation start time]

Each concentration grid may have a different starting, stopping, and output averaging time. Zero entry will result in setting the default values. "Backward" calculations require this and the following parameter to be explicitly set and further the stop time should come before the start time.

24(9)- Sampling stop time: year month day hour minute Default: {+1} 12 31 24 60

After this time no more concentration records are written. Early termination of high resolution grids (after the plume has moved away from the source) is an effective way of speeding up the computation for high resolution output because that particular grid resolution is no longer used for time-step computations.

25(10)- Sampling interval: type hour minute Default: 0 24 0

Each grid may have its own sampling or averaging interval. The interval can be of two different types: averaging (type=0) or snapshot (type=1). Averaging will produce output averaged over the specified interval. Snapshot will give the instantaneous output at the output interval. For instance you may want to define a concentration grid that produces 24-hour average air concentrations for the duration of the simulation which for the default case of

a 2-day simulation will result in 2 output maps, one for each day. Each defined grid can have a different output type and interval.

## 2.2.4 Deposition Definitions

26- Number of pollutants depositing      **Default: number of pollutants defined on line # 10**

Deposition parameters must be defined for each pollutant species emitted. Each species may behave differently for deposition calculations. Each will be tracked on its own concentration grid. The following five lines are repeated for each pollutant defined. The number here must be identical to the number on line 10. Deposition is turned off for pollutants by an entry of zero in all fields.

27(1)- Particle: Diameter ( $\mu\text{m}$ ), Density (g/cc), and Shape      **Default: 0.0 0.0 0.0**

These three entries are used to define the pollutant as a particle for gravitational settling and wet removal calculations. A value of zero in any field will cause the pollutant to be treated as a gas. All three fields must be defined ( $>0$ ) for particle deposition calculations. These values need to be correct only if gravitational settling is to be computed by the model, otherwise a nominal value of 1.0 may be assigned as a default for each entry to define the pollutant as a particle. If a dry deposition velocity is specified as the first entry in the next line (28), then that value is also used as the particle settling velocity.

28(2)- Deposition velocity (m/s), Pollutant molecular weight (Gram/Mole), Surface Reactivity Ratio, Diffusivity Ratio, Effective Henry's Constant  
**Default: 0.0 0.0 0.0 0.0 0.0**

Dry deposition calculations are performed in the lowest model layer based upon the relation that the deposition flux equals the velocity times the ground-level air concentration. This calculation is available for gases and particles. The dry deposition velocity can be set directly for each pollutant by entering a non-zero value in the first field or it can be calculated by the model using the resistance method which requires setting the remaining four parameters (molecular weight, surface reactivity, diffusivity, and the effective Henry's constant) - see Table I for more information.

29(3)- Wet Removal: Actual Henry's constant, In-cloud (L/L), Below-cloud (1/s)  
**Default: 0.0 0.0 0.0**  
**Suggested: 0.0  $3.2 \times 10^5$   $5 \times 10^{-5}$**

Henry's constant defines the wet removal process for soluble gases. It is defined only as a first-order process by a non-zero value in the field. Wet removal of particles is defined by non-zero values for the in-cloud and below-cloud parameters. In-cloud removal is defined as a ratio of the pollutant in air (g/liter of air in the cloud layer) to that in rain (g/liter) measured at the ground. Below-cloud removal is defined through a removal time constant.

30(4)- Radioactive decay half-life (days)

Default: 0.0

A non-zero value in this field initiates the decay process of both airborne and deposited pollutants.

31(5)- Pollutant Resuspension (1/m)

Default: 0.0  
Suggested : $10^{-6}$

A non-zero value for the resuspension factor causes deposited pollutants to be re-emitted based upon soil conditions, wind velocity, and particle type. Pollutant resuspension requires the definition of a deposition grid, as the pollutant is re-emitted from previously deposited material. Under most circumstances, the deposition should be accumulated on the grid for the entire duration of the simulation. Note that the air concentration and deposition grids may be defined at different temporal and spatial scales.

Table I. Some typical Pollutant constants from Wesely (1989)<sup>1</sup> and Walmsley and Wesely (1996)<sup>2</sup>.

Chemical	Symbol	$D_{hx}$	$H^*$ (M/atm) effective	H(M/atm) actual	$f_o$
Sulfur dioxide	SO <sub>2</sub>	1.9	$1 \times 10^5$	1.24	0.0
Ozone	O <sub>3</sub>	1.6	0.01	0.013	1.0
Nitrogen dioxide	NO <sub>2</sub>	1.6	0.01	0.01	0.1
Nitric oxide	NO	1.3	$3 \times 10^{-3}$	$1.9 \times 10^{-3}$	0.0
Nitric acid	HNO <sub>3</sub>	1.9	$1 \times 10^{14}$	$2.1 \times 10^5$	0.0
Hydrogen peroxide	H <sub>2</sub> O <sub>2</sub>	1.4	$1 \times 10^5$	$1.0 \times 10^5$	1.0
Ammonia	NH <sub>3</sub>	0.97	$2 \times 10^{14}$	62	0.0
Peroxyacetyl nitrate	PAN	2.6	3.6	5	0.1
Nitrous acid	HNO <sub>2</sub>	1.6	$1 \times 10^5$	$2.1 \times 10^5$	0

$D_{hx}$  - Diffusivity ratio; H - Henry's constant;  $f_o$  - Surface reactivity ratio

---

<sup>1</sup>Wesely, M.L., 1989, Parameterizations of surface resistances to gaseous dry deposition in regional-scale numerical models, *Atmos. Environ.*, 23, 1293-1304.

<sup>2</sup>Walmsley, J. L., and M.L. Wesely, 1996, Modification of coded parameterizations of surface resistances to gaseous dry deposition, *Atmos. Environ.*, 30, 1181-1188.



### 3. GRAPHICAL DISPLAY OPTIONS

The trajectory and concentration models each generate their own output files, which are read by the other programs to produce various displays and other output. The trajectory model generates an ASCII file of end-point positions while the concentration model produces a binary output file (big-endian) on a regular latitude-longitude grid. All mapping programs use the same ASCII map background file, *arlmap*, which normally would be located in the *\graphics* directory. However all the graphics programs search the local start directory first, then the *\graphics* directory. Customized map background files could be placed in the local directory for specialized applications. Some higher resolution map background files are available from the Hysplit download web page. The *Readme.txt* file in the *\graphics* directory has more information about developing custom map background files.

The feature rich Postscript conversion programs can both be accessed through the GUI or run directly from the command line. The Postscript conversion programs for both trajectories and concentrations have a variety of command line options, most of which are also available through the GUI.

#### 3.1 Trajectories

The Postscript conversion program (*trajplot*), found in the *\trajmdl* directory, reads the trajectory endpoints output file, calculates the most optimum map for display, and creates the output file - *trajplot.ps*. When executed from the command line, there are three other optional {} inputs:

```
trajplot [file_name] {Size} {Color} {Labels},
```

where default values are in red and only the *file\_name* is required:

Size	= 0	for standard resolution maps
	= 1	for high resolution maps (zoomed)
Color	= 0	for black and white output
	= 1	for color differentiation of multiple trajectories
Label	= 0	for no labels along the trajectory
	= 6	for labels every 6 hours
	= 12	for labels every 12 hours
	= {X}	any hour selection permitted.

The output example was shown previously in Fig. 1.

## 3.2 Air Concentration

The Postscript conversion program (*concpplot*), found in the `\concmdl` directory, reads the binary concentration output file, calculates the most optimum map for display, and creates the output file *concpplot.ps*. Multiple pollutant species or levels can be accommodated. Most routine variations can be invoked from the GUI or the command line using the following 7 optional { } parameters:

```
concpplot [file_name] {Z1} {Z2} {Type} {Size} {Color} {Value} {Units}.
```

where default values are in red and only the *file\_name* is required:

- |               |              |  |
|---------------|--------------|--|
| <i>Z1, Z2</i> |              | Heights that represent the levels that will be displayed. The heights are always defined as meters and should correspond to the range of values defined in the input section (line 22). The level information is interpreted according to the <i>Type</i> definition. Also $Z2 \geq Z1$ .  |
| <i>Type</i>   | <b>=1</b>    | All output levels between the levels specified on the command line are displayed as individual frames. A single level will be displayed if either both specified levels equal the calculation level or they bracket that level. Deposition plots are produced if available in the file and when a level height is set to 0. If level information is omitted, all levels are displayed. |
|               | <b>=2</b>    | The concentrations at all levels between the specified range are averaged to produce one output frame per time period. If a deposition plot is required then <i>Z1</i> should be set to 0.   |
|               | <b>=3</b>    | A customized exposure output in which all the output concentrations are converted to time-integrated units and vertically averaged for all levels that are found in the file between <i>Z1</i> to <i>Z2</i> . The last frame displayed represents the accumulated deposition through the model simulation.   |
| <i>Size</i>   | <b>= 0</b>   | Standard resolution  |
|               | <b>= 1</b>   | High resolution map (less white space around the concentration pattern)  |
| <i>Color</i>  | <b>= 0</b>   | Uses grey shade patterns for the contour color fill  |
|               | <b>= 1</b>   | Uses the standard four color fill.   |
| <i>Value</i>  | <b>= 0</b>   | Contour intervals are to be optimized for each map   |
|               | <b>= 1</b>   | Contour intervals are the same for all maps  |
|               | <b>= {X}</b> | where {X} represents the integer power of 10 of the maximum contour.   |
| <i>Units</i>  | <b>= {X}</b> | where {X} is the multiplier applied to the input data before output.   |



The output example was shown previously in Fig. 2. One final note is that if multiple pollutant species are defined, a prompt to standard output will appear, requesting the selection of a specific species. Only one pollutant species may be displayed per plot sequence. However, an entry of "0" will cause all species to be summed for display.

### 3.3 Utility Programs

Currently only utility programs to manipulate the binary concentration output files are available. Each of these programs is discussed in more detail below. All are available through the "*Utility Programs*" selection of the GUI as well as through the command line.

#### 3.3.1 con2asc - Convert to ASCII

This program can be used to convert the binary concentration file to a simple ASCII file composed of one record per grid point for all grid points where concentrations at any level are non-zero. Concentrations for multiple levels and pollutant species are all listed on the same record for each grid point. The primary purpose of the conversion is to create a file that can be imported into other applications. The format of each record in the output file is given by:

2I3	- End of Sample: Julian Day and Hour
F7.2, F8.2	- Latitude and Longitude of grid cell
45E9.2	- Concentration data (by level and pollutant)

Each output record is identified by the day (Julian: 1 to 365) and hour (UTC) of the ending time of each sample. In addition, a new output file is created for each sampling period, where the name of the file is composed of the *{input file name}\_{Julian day}\_{hour}*. Only the input file name is required on the command line.

```
con2asc [input file name]
```

The selection of input file names through the GUI is determined by the *Setup* menu.

#### 3.3.2 con2stn - Grid to Station

The purpose of con2stn is to list concentrations at specific latitude-longitude locations. The program can be run from the command line, through interactive prompts from the keyboard, or through the GUI. Command line argument syntax is that there should be no space between the switch and options. No options are available with interactive mode. The output gives the Julian day, month, day, and hour of the sample start; day and hour of sample ending time, and the concentrations for each station (location selected by latitude-longitude). The format of each output record is as follows:

F8.3,6I4	- Starting: Julian day, Year, month, day, hour; Ending: day, hour
XF10	- Concentration value at X stations

Unlike the other programs, the command line arguments can appear in any order and the syntax is as follows:

```
con2stn -i -o -c -s -x -z -p
```

where the default value is indicated in red:

```
-i[input concentration file name: std input]  
-o[output concentration file name: std output]  
-c[input to output concentration multiplier: 1.0]  
-s[station list file name: std input]  
-x[n(neighbor) or i(interpolation): n]  
-z[level index: 1]  
-p[pollutant index: 1]
```

Unspecified file names will result in a standard input prompt. The default interpolation method (-xn) is to use the value at nearest grid point to each latitude-longitude position. The station positions can be read from a file (space delimited) with the first field being an integer that represents the location identification, followed by that location's latitude and longitude.

Examples:

0) con2stn ... Results in prompts -->

```
Enter input concentration file name...  
[name of hysplit output file]  
  
Enter sampler ID#, latitude, longitude ...  
[integer sample ID, real latitude, real longitude]  
0 0 0 (to terminate input)
```

1) Read the model output file 'cdump' and write text output to file 'clist.txt' for station #517 at 53N 85W.

```
con2stn -icdump -oclist.txt  
517 53.0 -85.0  
0 0 0
```

2) As in 1) but multiply all concentrations by 1000.0

```
con2stn -icdump -oclist.txt -c1000.0  
517 53.0 -85.0  
0 0 0
```

3) As in 1) but linear interpolate concentration to station rather than using the nearest grid point

```
con2stn -icdump -oclist.txt -xi
517 53.0 -85.0
0 0 0
```

4) As in 1) but read the station lat-lon from a file "slist.txt"

```
con2stn -icdump -oclist.txt -sslist.txt
```

The format of "slist.txt" is "free form", for example ...

```
401 39.0 -88.5
422 36.5 -87.5
657 36.0 -88.0
004 35.0 -89.0
```

### 3.3.3 wincpick - Select from Display

Windows based concentration plot point registration program is designed to read the binary concentration data file, display the data on the screen, and then use the mouse to select locations at which the position and concentration data are read and written to the text file: *wincpick.txt*. The concentration data are displayed over the entire grid domain. If you want to zoom in on a specific area, then it is necessary to rerun the model with a smaller concentration grid domain. The program is available from the GUI or the command line with the following syntax:

```
wincpick [input file name]
```

Upon startup *Wincpick* will display the domain background map with a summary of the mouse based instructions: left-click registers the lat-lon position of that point to the output file *wincpick.txt*. A right-click of the mouse redraws the map for the next time period and at the end of the input file saves and closes all files; and a CNTL+right-click closes all files and exits the program before the end of the input file. Right-click the mouse to go to the first concentration map. Maps are drawn in sequence of height, pollutant species, and time. Left-clicks register the lat-lon position of the mouse to the output file. If you are interested in only one time period, skip past those times (right-click), and then register the points of interest, then CNTL-right to exit.

### 3.3.4 Timeplot - Time Series Concentration Plot

```
timeplot -i -n
-i[input concentration file name in format as output from con2stn]
-n[sequential station number; repeat for multiple sites; 999 for all]
```

### 3.4 Output Customization

Many of the Postscript graphics programs that have extensive label information can be customized to some extent, primarily the title field (upper center) and the units. This is accomplished by placing a file called *Labels.cfg* in the \working or startup directory which contains the following two entries (all in single quotes terminated by &) replacing the new string with the desired text. A sample file called *Labels.bak* may be found in the relevant directory.

```
'TITLE&', 'NEW TITLE STRING&'
'UNITS&', 'NEW UNITS STRING&'
```

### 3.5 File Formats

#### 3.5.1 Trajectory Endpoints

The format of the ASCII endpoints file written by the trajectory model (*hymodelt*) and read by all trajectory display programs is given below:

Record #1

I6 - Number of meteorological grids used in calculation

Records **Loop** #2 through the number of grids

A8 - Meteorological Model identification

5I6 - Data file starting Year, Month, Day, Hour, Forecast Hour

Record #3

I6 - number of different trajectories in file

A8 - direction of trajectory calculation (FORWARD, BACKWARD)

A8 - vertical motion calculation method (OMEGA, THETA, ...)

Record **Loop** #4 through the number of different trajectories in file

4I6 - starting year, month, day, hour

2F8.3 - starting latitude, longitude

F8.3 - starting level above ground (meters)

Record #5

I6 - number (n) of diagnostic output variables

nA8 - label identification of each variable (PRESSURE, THETA, ...)

Record **Loop** #6 through the number of hours in the simulation

I6 - trajectory number

I6 - meteorological grid number

5I6 - time of point: year month day hour minute

I6 - forecast hour at point

F8.1 - age of the trajectory in hours

2F8.3 - position latitude and longitude

F8.1 - position height in meters above ground

nF8.1 - n diagnostic output variables (1<sup>st</sup> output always pressure)

### 3.5.2 Binary Gridded Concentrations

The output format of the binary concentration file written by *hymodelc* and read by all concentration display programs is as follows:

#### Record #1

CHAR\*4 Meteorological *MODEL* Identification  
INT\*4 Meteorological file starting time (*YEAR, MONTH, DAY, HOUR, FORECAST*)  
INT\*4 *NUMBER* of starting locations

#### Record #2 Loop to record: Number of starting locations

INT\*4 Release starting time (*YEAR, MONTH, DAY, HOUR*)  
REAL\*4 Starting location and height (*LATITUDE, LONGITUDE, METERS*)

#### Record #3

INT\*4 Number of (*LATITUDE-POINTS, LONGITUDE-POINTS*)  
REAL\*4 Grid spacing (*DELTA-LATITUDE, DELTA-LONGITUDE*)  
REAL\*4 Grid lower left corner (*LATITUDE, LONGITUDE*)

#### Record #4

INT\*4 *NUMBER* of vertical levels in concentration grid  
INT\*4 *HEIGHT* of each level (meters above ground)

#### Record #5

INT\*4 *NUMBER* of different pollutants in grid  
CHAR\*4 Identification *STRING* for each pollutant

#### Record #6 Loop to record: Number of output times

INT\*4 Sample start (*YEAR MONTH DAY HOUR MINUTE FORECAST*)

#### Record #7 Loop to record: Number of output times

INT\*4 Sample stop (*YEAR MONTH DAY HOUR MINUTE FORECAST*)

#### Record #8 Loop to record: Number of levels, Number of pollutant types

CHAR\*4 Pollutant type identification *STRING*  
INT\*4 Output *LEVEL* (meters) of this record  
REAL\*4 Concentration output *ARRAY* (number of lat/lon elements)

## 4. SPECIAL APPLICATIONS

This section provides some guidance in configuring the model input to do certain specialized calculations. The default configuration supplied with the test meteorological data is confined to a simple trajectory and inert transport and dispersion calculation. Some other simple configurations will be reviewed in this section. Note some of these configurations may not be possible from default compilation of the distribution version of the code.

### 4.1 Particle or Puff Releases

The concentration model default simulation assumes a particle dispersion in the vertical direction and a top-hat puff dispersion in the horizontal direction. Other options are set with the *INITD* parameter of the *SETUP.CFG* namelist file defined in Section 4.4. Normally changes to the dispersion distribution are straightforward. However there are some considerations with regard to the initial number of particles released. The default release is set to be 500 particles over the duration of the emission cycle (see *NUMPAR* in Section 4.4). A 3-dimensional (3D) particle simulation requires many more particles to simulate the proper pollutant distribution, the number depending upon the maximum downwind distance of the simulation and the duration of the release, longer in each case require more particles. Too few particles result in noisy concentration fields. A 3D puff simulation can be started with one puff as the puff-splitting process in conjunction with the vertical dispersion quickly generates a sufficient number of puffs to represent the complex dispersion process. The default configuration represents a compromise in permitting particle dispersion in the vertical for greater accuracy and puff dispersion in the horizontal to limit the particle number requirements.

### 4.2 Continuous Emissions

As noted in Section 4.1 the default release is 500 particles over the duration of the emission cycle. If continuous emissions are specified (e.g. over the duration of the simulation), then those 500 particles are spread out over that time period. This may easily result in the release of too few particles each hour to provide smooth temporal changes in the concentration field. Imagine a single particle passing in and out of the vertical concentration grid plane due to turbulent diffusion. One solution would be to increase the *NUMPAR* parameter until smoother results are obtained. Another possibility would be to cycle the emissions by emitting 100 particles only for the first time step of each hour. Those particles would contain the total mass for a one-hour release (see how to set *QCYCLE* as described in Section 4.4).

### 4.3 Gridded Area Source Emissions

Normally emissions are assumed to be point or vertical line sources. Virtual point sources (initial source area >0) can be defined two ways: 1) through the definition of an initial area on the source location input line or 2) by the definition of a gridded emissions file. If the model's root startup directory contains the file *Emissions.txt*, then the pollutants are emitted from each grid cell according to the definitions previously set in the *Control* file. Two source

points should be selected, which define the lower left (1<sup>st</sup> point) and upper right (2<sup>nd</sup> point) corner of the emissions grid that will be used in the simulation. This can be a subset of the grid defined in *Emissions.txt*. The release height represents the height from the ground through which pollutants will be initially distributed. The emission file's first record contains information about the internal grid cell size that is used by the dispersion model to accumulate the file's emissions. The emission file defines the emissions at latitude-longitude points, the values at these points are accumulated in an internal grid, the size of which is defined on the first record. This value can be arbitrarily changed according to the desired resolution of the simulation. The pollutant puffs are released with an initial size comparable to the accumulation cell size. Because the emission file data are re-mapped to an internal grid, the file can consist of emissions data on a regular grid or just a collection of individual cells. The emission rate in the *Control* file is used as an additional multiplication factor for the data in the emission file. Also note that previously discussed particle number restrictions still apply. The initial number of particles are spread out over the duration of the emission and the number of grid cells that are defined in the emission domain. The format of the *Emission.txt* file is given below:

Record #1

- I4 - Number (n) of pollutant species in file
- F10.4 - Conversion factor from file emission units to internal model units/hour
- 2F10.4 - Internal grid cell size (latitude & longitude) at which file emissions are accumulated
- nA4 - Character identification of each pollutant (should match control file)

Records Loop #2 to the number of i,j grid point

- 2I4 - I,J grid point index of emission cell (arbitrary units for user identification)
- 2F10.4 - Southwest corner Longitude and Latitude of this emission cell

Record Loop #3 to the number of pollutant species

- 12E10.3 - emissions for pollutant#1 hours 1-12
- 12E10.3 - emissions for pollutant#1 hours 13-24

The model can easily be configured to simulate more complex pollutant episodes with multiple pollutant types or multiple pollutant species on the same particle. This is accomplished by changing either *MAXTYP* or *MAXDIM* in *DEFSIZE.INC* to the appropriate value and recompiling the code. If an external chemistry routine is used that converts mass from one species to another, all tracking together (advecting and dispersing), then *MAXDIM* is raised to the required value. If multiple species are emitted, have no interaction, and may track differently, then *MAXTYP* is adjusted to the required value. This latter situation may represent a volcanic ash plume where each pollutant, a different sized particle, settles at a different rate. Note that multiple species defined by the latter method can be accomplished within the default configuration of the model. However the *MAXDIM* definition always requires an external routine to adjust the mass between species.

#### 4.4 Namelist File: *SETUP.CFG*

Additional simulation options are available through modification of the *Setup.cfg* namelist file. This file is not required, and if not present in the root startup directory, default

values are used. The trajectory model has only five *namelist* options. The concentration model has an additional 10 parameters. These parameters can all be changed without recompilation by modification of the contents of *Setup.cfg* and in some cases their modification will substantially change the nature of the simulation. The file should be present in the root directory (either *\working* for the GUI interface or *\concmdl* or *\trajmdl* for command line) with the following contents:

Options valid for either the Trajectory or Concentration models:

*TRATIO* - valid for trajectories or concentration simulations and defines the fraction of a grid cell that a trajectory is permitted to transit in one advection time step. Reducing this value will reduce the time step and increase computational times. However, smaller time steps result in less integration error. Integration errors can be estimated by computing a backward trajectory from the forward trajectory end position and computing the ratio of the distance between that endpoint and the original starting point divided by the total forward and backward trajectory distance. Default value = 0.75

*DELT* - can be used to set the integration time step to a fixed value in minutes from 1 to 60 and it should be evenly divisible into 60. The default value of 0.0 causes the program to compute the time step each hour according to the maximum wind speed, meteorological and concentration grid spacing, and the *TRATIO* parameter. The option to use a fixed time step should only be used when strong winds in regions not relevant to the dispersion simulation are causing the model to run with unrealistically small time steps. Improper specification of the time step could cause aliasing errors in advection as well as substantial underestimates of air concentrations. Default value = 0.0

*MGMIN* - is the minimum size in grid units of the meteorological sub-grid. The sub-grid is set dynamically during the calculation and depends upon the horizontal distribution of points and the wind speed. Larger sub-grids than necessary will slow down the calculation by forcing the processing of meteorological data in regions where no transport or dispersion calculations are being performed. In some situations, such as when the computation is between meteorological data files that have no temporal overlap, the model may try to reload meteorological data with a new sub-grid. This will result in an error from the *metpos* subroutine. One solution to this error would be to increase the minimum grid size so that the size is sufficient to cross the file boundary with trying to reload data from the old closed file. Default value = 10

*KMSL* - When set to "0" source input heights are assumed to be relative to the terrain height of the meteorological model - hence input heights are specified as AGL. Setting this parameter to "1" forces the model to subtract the local terrain height from source input heights before further processing - hence input heights should be specified as relative to Mean Sea Level (MSL). In concentration simulations this also forces the vertical concentration grid heights to be considered relative to mean sea level. Default value = 0

*AA, BB, CC* - Polynomial parameters that control the resolution of the internal Hysplit terrain



following grid system. Input meteorological data are interpolated to this grid. The polynomial relates height (AGL) to the internal vertical index number where  $k=1$  would be the first level above ground:  $Z = AA*k**2 + BB*k + CC$ . **Default values = AA(30), BB(-25), CC(5)**

Options valid only for the Concentration Model:

*INITD* - determines if the model is configured as a puff or particle model. Valid options: 0 (3D particle); 1 (Gaussian Puff); 2 (Top Hat Puff); 3 (Gaussian Particle) 4 (Top-Hat Particle). **Default value = 4**

*KHMAX* - is the maximum age (hours) that any puff or particle is permitted to attain. All pollutants beyond this age are deleted. **Default value = 9999**

*NUMPAR* - would be the maximum number of particles or puffs permitted during a simulation. This value is different from *MAXPAR* in the include file, in that *NUMPAR* cannot exceed *MAXPAR*. *NUMPAR* has a different meaning for puff and particle simulations. In a full puff simulation, where only one puff per time step is released, the total number of puffs on the grid at any one time cannot exceed *NUMPAR*. In a particle simulation, or mixed particle-puff simulation, *NUMPAR* represents the total number of particles that are released during one release cycle. Multiple release cycles cannot produce more than *MAXPAR* number of particles. For a mixed simulation (particle-puff), *NUMPAR* should be greater than one but does not need to be anything close to what is required for a full 3D particle simulation. In all simulation types, particle or puffs are emitted if the particle count is less than *MAXPAR* **Default value = 500**

*QCYCLE* - are the number of hours between emission start cycles. A zero value means that the emissions are not cycled. When non-zero, the number of emission hours is repeated again at *QCYCLE* hours after the starting emission time specified in the input. **Default value = 0.0**

*KRND* - at this interval in hours, enhanced puff merging takes place. Enhanced merging is less restrictive and will degrade the accuracy of the simulation. Puffs can be further apart and still be merged into the same position. Less frequent merging will improve accuracy, however too many puffs may remain and the simulation time will be substantially increased. The selection of an appropriate value depends if the pollutant release is instantaneous or continuous. Enhanced merging only occurs when the puff number exceeds 25% of *MAXPAR*. **Default value = 6**

*FRME* - is the fraction of the total mass that represents a puff mass at which all puffs with a mass less than that puff value will only account for *FRME* of the total mass. These "Low Mass" puffs will be subject to enhanced merging. **Default value = 0.10**

*FRMR* - is the fraction of the mass that is permitted to be removed at *KRND* intervals. The normal situation is to permit no mass loss. However for certain simulations, such as when a pollutant has a high ambient background relative, a small removal rate will significantly reduce the number of puffs on the grid at no loss in accuracy. **Default value = 0.0**

*ISOT* - is a flag used to set the isotropic turbulence option. The default value of 0 results in the computation of vertical turbulence from vertical stability estimates and the horizontal turbulence from the wind field deformation. In shorter range dispersion simulations (<100 km) the deformation parameterization used in conjunction with larger scale meteorological fields will not reflect the diurnal variations in horizontal turbulence. In these situations isotropy is assumed by setting the *ISOT* parameter to 1, i.e., the horizontal velocity variance is set equal to the vertical velocity variance. In addition, the vertical diffusivity profile is NOT replaced by its boundary layer average value when *ISOT*=1. Default value = 0

*NDUMP* - can be set to dump out all the particle/puff points at selected time intervals to a file called *PARDUMP*. This file can be read from the root directory at the start of a new simulation to continue the previous calculation. Valid *NDUMP* settings are [0] for no I/O or [hours] to set the number of hours at which all the endpoint positions will be written to the file. The file must exist in the root directory and *NDUMP*>0 for the model to initialize pollutant particles from the file. *NDUMP* is used in conjunction with *NCYCL* (see below). Default value = 0

*NCYCL* - sets the cycle interval at which the *PARDUMP* file is to be written after the first write at hours *NDUMP*. Multiple outputs will overwrite the last output. For instance in a multi-day simulation, one application would be to set *NDUMP*=24 and *NCYCL*=24 to output all points at the end of every simulation day. If the model were to crash unexpectedly, the simulation could be restarted from the last *PARDUMP* output. Default value = 0

#### 4.5 Compilation Limits: *DEFSIZE.INC*

A variety of different parameters are set in the compilation "include" files. The value of some of these parameters may effect the results of a simulation and the options that are available in setting up the *Control* file. There is a copy of the *DEFSIZE.INC* include file in each subdirectory: trajectories, air concentrations, and other special applications. The parameters in the local include file are optimized for the application in that directory. Although there is nothing that can be altered to change the model simulation, the file provides information about the compiled model limits and can be used to diagnose certain error messages. A summary of the main features of the primary "include" file *DEFSIZE.INC* follows:

##### Meteorological grid size parameters ...

*NXYD* - The product of the number of points in X and Y of the packed meteorological data grid should not exceed this value.

*NXM*, *NYM* - The horizontal limits of the meteorological sub-grid. Note this value may be larger or smaller than the actual data grid on the input file. It only specifies how much data is read and processed from the input file at any one time. This is the prime dimension controlling memory requirements because all meteorological variables are of this dimension. These values are usually kept small for trajectory calculations; you do not need to process the whole grid to compute advection at only one point. However for concentration simulations, it is best if a

larger grid domain can be loaded, because pollutant plumes usually encompass a much larger area. A small sub-grid for concentration simulations may cause the model to keep reloading the data at the same time but slightly different locations to complete a calculation.

*NZM* - In a similar manner to the horizontal dimension, it limits the scope of the model's internal coordinate system in that only meteorological data from the ground to *NZM* number of levels are processed. In addition, the internal vertical grid to which the meteorological data are interpolated is of the same dimension.

*MLVL, MVAR* - define the maximum data levels and parameters that may be found on an input file. It is necessary that these parameters cover the scope of potential input files as the model must be able to read the index record of the data file. This record will contain information for each level and variable.

*MGRD* - permits the simultaneous definition of multiple meteorological input files. The ability to open multiple files has low memory requirements. However multiple files may slow calculations as only one file's data are permitted in memory at any one time.

#### Additional parameters for concentration calculations...

*MAXPAR* - is the maximum number of particles or puffs that may be followed for the compiled version of the code. Larger values require correspondingly more system memory. The actual particle/puff limit during a particular calculation is set in the *namelist* file.

*MAXDIM* - The maximum number of different species that can be associated with any single particle or puff. There is no routine model configuration that requires this value to be greater than one. Certain optional additional subroutines may be added that may convert mass from one species to another on the same particle or puff.

*MAXTYP* - The number of different pollutant species. Each species is associated with a different puff or particle. *MAXTYP* should always be equal to or greater than *MAXDIM*. Different pollutant types may be defined as part of the normal input procedure for emissions and deposition.

*MAXGRD* - The number of simultaneous concentration output grids. Values larger than one quickly increase memory requirements because each grid will have the dimensions: *MAXXP, MAXYP, MAXZP, MAXTYP, MAXGRD* .

*MAXXP, MAXYP* - The maximum number of points in the West to East direction and the South to North direction on any concentration grid.

*MAXZP* - The maximum number of levels on any concentration grid.

#### To increase the number of sources for trajectories and concentrations ...

*MLOC* - the number of simultaneous trajectories or particle starting locations that can be simultaneously tracked during a computation.

#### 4.6 Optional Features - Advanced GUI Menu

Some Hysplit installations may include the additional main menu item called "*Advanced*". This menu permits interactive modification of the *Setup.cfg* file through the "*Configuration Setup*" tab. Not all the options discussed in Section 4.4 are available through this menu. File display tabs are available for the *MESSAGE* and *PARDUMP* files. The *MESSAGE* file is viewed through a simple text scroll bar widget while the *PARDUMP* file viewer displays the particle end point positions on the screen. The "*FTP Satellite Data*" tab provides access to selected NOAA and NASA satellite archives. The *PARDUMP* file view tab will also display the corresponding satellite image if it is available. These features are still under development and additional instructions are provided through the viewer and ftp menu tabs.

#### 4.7 Configuration for Operational Applications

Most of the discussion in various sections of the User's Guide are tailored to individually configured simulations. However there are several features to the model that can be used to automate the computational environment. For instance, a sample *Auto\_traj.tcl* script is provided in the *\working* directory that can be used as a guide to automate many applications.

```
# Auto_traj.tcl
# the next line restarts using wish \
# exec wish "$0" "$@"
set Start_hgt "10.0"
set Traj_path "../trajmdl"
set Start_time "00 00 00 00"
set Run_hours "24"
set Vert_coord "0"
set Top_model "10000.0"
set Meteo_path "../metdata/"
set Meteo_file "oct1618.BIN"
set Output_path "/"
set Output_base "tdump"
set Output_num 1
foreach {Start_lat Start_lon} {35.0 -90.0 40.0 -90.0 45.0 -90.0} {
    set Start_loc "$Start_lat $Start_lon $Start_hgt"
    set Output_file "$Output_base$Output_num"
    file delete Control
    set f [open Control w]
    puts $f "$Start_time"
    puts $f "1"
    puts $f "$Start_loc"
    puts $f "$Run_hours"
    puts $f "$Vert_coord"
    puts $f "$Top_model"
    puts $f "1"
    puts $f "$Meteo_path"
    puts $f "$Meteo_file"
    puts $f "$Output_path"
    puts $f "$Output_file"
    close $f
    exec "$Traj_path/hymodelt.exe"
    incr Output_num
}
```

In this particular example the test trajectory case is run for three different starting locations, each simulation writing a new endpoints file with a unique file name. The *CONTROL* file is recreated for each simulation. It would be trivial to rewrite the script to set the latitude-longitude and loop through a different number of starting days and hours. This script can be run under Windows or Unix. For instance, to compute new forecast trajectories each day, the process can be automated by including a data FTP at the beginning of the script to get the most recent meteorological forecast file, setting the starting time as "00 00 00 00" so that the trajectories will start at the beginning of the file, and finally calling the script once-a-day through the Unix crontable or the Windows' scheduler commands.

One problem with automated operations is that it is possible to generate simultaneous multiple jobs which may interfere with each other. Unix versions of Hymodelc and Hymodelt have a command line option of adding the process ID (PID): e.g. hymodelt [PID]. In this situation all standard named input and output files [those not defined in the Control file] have the PID added as a suffix to the file name: e.g. Control.[PID], Setup.[PID], Message.[PID].

An example of another type of operational configuration is the extended simulation of a pollutant emission using archive data to bring the simulation to the current time and then using forecast meteorological data to provide a daily projection. Each day the archive simulation must be updated with the new archive data and a new forecast product generated. This process can also be automated through a script, but for illustration purposes one can use the advanced features of the GUI to configure such a case. Assume a one-hour duration accidental pollutant release that occurred 48 hours prior to the current time. The following sequence applies:

- 1) From the "*Meteorology*" menu tab download the appropriate archive meteorological data and the most recent forecast meteorological data (assume it is available to +48h).
- 2) Setup the concentration simulation to run 96 hours using two meteorological files starting with the archive data and then switching to the forecast data.
- 3) Under the *Advanced* menu tab and *Configuration Setup* write the initialization file at 72 hrs.
- 4) Run the model.

At the completion of the simulation you will have the plume projection from release (-48 h) through the current forecast (+48 h). The *PARDUMP* file will contain all the endpoint positions at +24 hours, corresponding to the initialization time of when the next forecast will be available (assume there is one forecast per day).

The next day, when the new forecast data are available, reconfigure the model to run only with the forecast meteorological data for a duration of 48 hours and write the initialization file after 24 hours, then run the model to obtain the new projection. In this second part we assume that the first 24 hours of the forecast are not much different than the analysis. In practice, this procedure can be run at the same frequency that the new forecast data are available and typically at 4 times per day, data at the initial forecast hour are identical to the analysis data.

## **4.8 Backward Dispersion Simulations**

A common application of atmospheric trajectory and dispersion models is to try to determine the source of a pollution measurement. If a high value has been collected at a particular receptor, from which pollutant source region did the air originate? One common approach is to calculate the trajectory "backwards" from the receptor site. In the trajectory calculation this is accomplished by setting the integration time step to a negative value. However the trajectory only represents the upwind path of a single point, while the pollutant measurement may require of hundreds or thousands of trajectories to represent the dispersion of the pollutant in time and space.

Another approach is to run the entire dispersion-trajectory model "backwards" which is computationally attractive because in a 3D particle model the dispersion process is represented by a turbulent component added to the trajectory calculation and the advection process is fully reversible. The trajectory equation can be correctly integrated in either direction. However, the interpretation of the output is a bit more complex because dispersion is an irreversible process and although the equivalent numerical calculation will yield a result because the integration of the dispersion equation is still in the normal downstream mode on top of the backward upstream integration of the advection. The meaning of the upwind dispersion result is less clear.

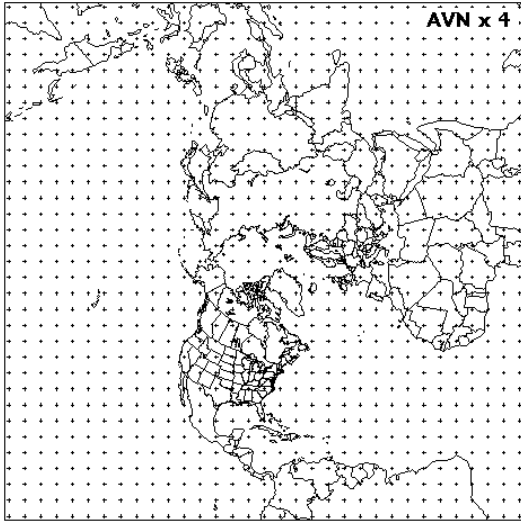
In any event as noted in the earlier instructions it is possible to run the dispersion model "backwards" by setting the run duration and emission hours to their equivalent negative value. The stop time of the sampling should be set prior to the start time. All start and stop times should be set to their exact values – relative start-stop times are not supported in the backward mode. To simplify interpretation of the results, horizontal dispersion is "turned off" in the backward calculation, resulting in a more reversible calculation.

## **4.9 Time Variation of the Emission Rate**

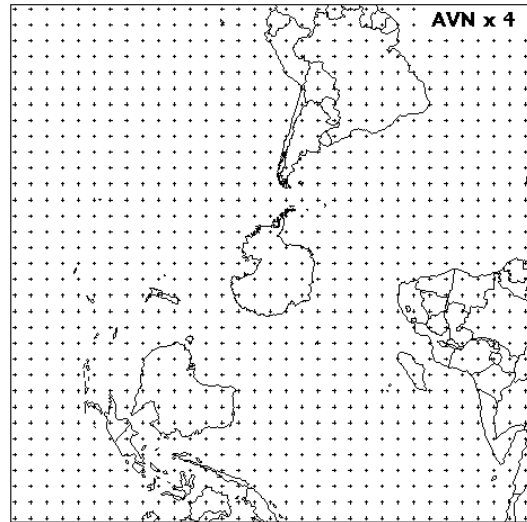
The most efficient way to incorporate a time varying emission rate into the existing model structure is still under evaluation. Currently the only way to accomplish this is to use the particle dump feature to restart the model each time with a new emission rate.

## 5. METEOROLOGICAL INPUT DATA

Meteorological data fields to run the model are usually already available from routine archives or from forecast model outputs. Archive meteorological data to run the model may be ordered from NOAA's National Climatic Data Center<sup>1</sup> or more recent data files can be copied from <ftp://gus.arlhq.ssmc.noaa.gov/pub/archives>. There are more complete descriptions of the different data sets available on-line. Figs. 3 and 4 illustrate the domain of the global Aviation model data, applicable to both archives and forecasts, available at a 181 km resolution but shown only at every fourth grid point.



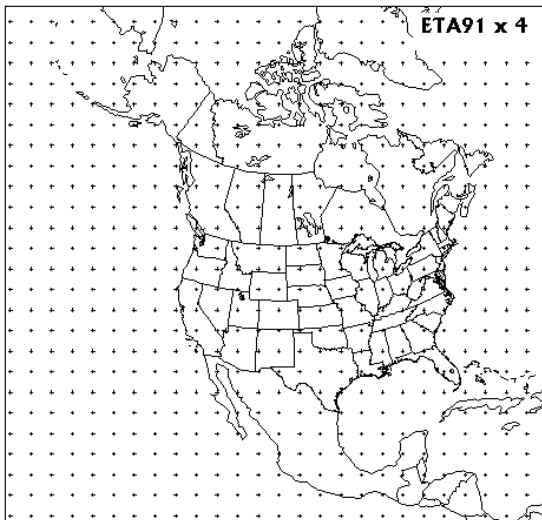
**Figure 3.** Global northern hemisphere grid.



**Figure 4.** Global southern hemisphere grid.

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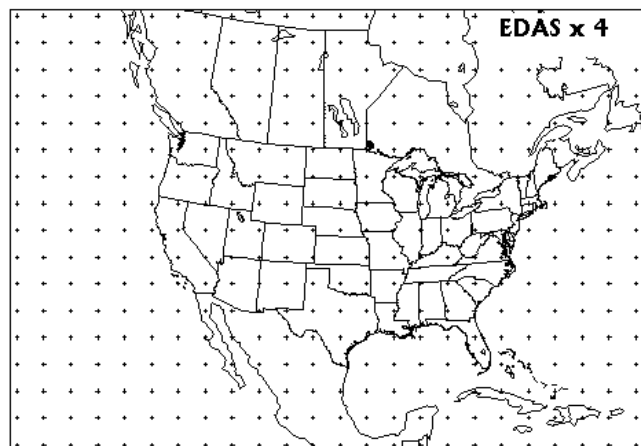
<sup>1</sup>Climate Services Branch, National Climatic Data Center, 151 Patton Avenue, Asheville, NC 28801, <http://www.ncdc.noaa.gov/archives>: TD-6140 and TD-6141 .



**Figure 5.** ETA forecast model grid.

The following sections describe the data in a little more detail so that customized applications can be developed. Additional subroutines are provided to simplify the task of creating a model compatible data file. This data file is composed of one or more time periods of data. Each time period begins with one or more ASCII index records that summarize the time period, the grid definition, variable and level information. Each subsequent record contains one horizontal data field, consisting of 50 ASCII bytes of time, variable, and level information for that record, followed by X times Y bytes of data, where X and Y are the number of data points in each direction. Floating point or integer data are packed as one byte per variable. Precision is maintained by packing the differences between adjacent grid points rather than packing the absolute values. In any one time period, although not required, the surface data precede the upper-level data fields. All records are the same length because the model needs to read the file in "direct access" mode. Data files can be read on most computing platforms without any transformation and appended to each other using routine operating system commands. Only binary transfers or copies are permitted. All of the routines discussed in this section can be found in the `\metdata\source` directory.

Note that the NCEP models are run at much higher spatial resolution than what is archived in a Hysplit compatible format. If calculations at higher resolution are required, then the original GRIB encoded files must be obtained and converted as described later in this section. Although sample software is provided, no additional guidance except for the information contained in this report is available. Higher resolution data over North America using the ETA model forecasts on a 91 km grid is shown in Fig. 5. The ETA data analysis and assimilation (EDAS) archive grid at 80 km resolution is shown in Fig. 6. Both illustrations only show every fourth grid point for clarity.



**Figure 6.** EDAS archive data grid.



## 5.1 Valid Meteorological Data Types

Meteorological variables are identified to the model by a unique 4 character identification that is written to the first 50 byte header portion of each data record. Some of the variables that can be decoded by the model and their units are identified below.

<u>Surface-Level Parameters Description</u>	<u>Units</u>	<u>Identification</u>
Pressure at surface	hPa	PRSS
Pressure reduced to mean sea level	hPa	MSLP
Temperature at surface	K	TMPS
Total precipitation (6-h accumulation)	m	TPP6
Momentum flux, u-component at surface	N/m <sup>2</sup>	UMOF
Momentum flux, v-component at surface	N/m <sup>2</sup>	VMOF
Sensible heat net flux at surface	W/m <sup>2</sup>	SHTF
Latent heat net flux at surface	W/m <sup>2</sup>	LHTF
Sfc downward short wave radiation flux	W/m <sup>2</sup>	DSWF
Temperature at 2 m AGL	K	T02M
Relative humidity at 2 m AGL	%	RH2M
U-component of wind at 10 m AGL	m/s	U10M
V-component of wind at 10 m AGL	m/s	V10M
Volumetric soil moisture content (<10cm)	fraction	SOLW
Total cloud cover, entire atmosphere	%	TCLD
<u>Upper-level Parameters Description</u>	<u>Units</u>	<u>Identification</u>
U-component of wind with respect to grid	m/s	UWND
V-component of wind with respect to grid	m/s	VWND
Geopotential height	gpm*	HGTS
Temperature	K	TEMP
Pressure vertical velocity	hPa/s	WWND
Relative humidity	%	RELH

Data may be obtained from any source, however there are certain minimum data requirements to run the model: surface pressure or terrain height, u-v wind components, temperature, and moisture. In addition, precipitation is required for wet removal calculations. Not required, but certainly necessary to improve vertical mixing calculations would be some measure of low-level stability. This may take the form of a near-surface wind and temperature or the fluxes of heat and momentum.

It is also important to have sufficient vertical resolution in the meteorological data files. Some of the NOAA higher resolution data files have five or more levels in the boundary layer (<850 hPa) in addition to wind and temperatures near the surface, usually 2 and 10 m agl. These surface values are especially important when the data are only available on absolute pressure surfaces, rather than terrain following sigma surfaces, to avoid interpolation to the surface between data levels when the local terrain is well above sea-level.

## 5.2 Creation of a Meteorological Input Data File

One may prepare data from other sources to be in a suitable format for the model through a series of routines described in this section. In general it is assumed you have access to some meteorological data source, either the data fields are already on a grid, such as output from a meteorological model, or perhaps observations that have been interpolated to a grid. Some example conversion programs are provided to convert from either NOAA, NCAR (re-analysis), or ECMWF GRIB format data files to Hysplit format (see *avn2arl*, *ncr2arl*, *ecm2arl* in directory *\data2arl*). If these programs are to be used directly, it will be necessary to obtain special platform dependent GRIB decoding subroutines. These are usually provided in conjunction with the GRIB data.

The meteorological data are processed in time-sequence, calling the subroutines provided, to create a Hysplit compatible output file. These subroutines will pack the data and write the index record. The index record, which precedes the data records for each time period, can only be written after the data records are processed. The routine must first be initialized by setting all the appropriate grid parameters and defining the meteorological variables that will be written to the file. Multiple output grids may be defined and written simultaneously by invoking the *PAKSET* routine with a different unit number for each grid. The grid parameters are all defined in a configuration file which should be in the directory from which the procedure is invoked:

```
CALL PAKSET(kunit,fname,krec,nx,ny,nz).
```

*Kunit* is the Fortran unit number to which the data records will be written. *Fname* is the character string of the name of the configuration file. It can be any name, but it will default to *metdata.cfg*. The file is opened internally on *kunit* to read the configuration file. This routine needs to be called once for each grid. *Krec* is the starting record number (of the index record) to which output will be written. It is normally set to 1 unless you are writing to the middle of a file. The remaining parameters (*nx*, *ny*, *nz*) are returned by the subroutine and define the horizontal and vertical grid dimensions. These values can be used to set variable dimensions. It is your responsibility to open *kunit* for output after having completed the *pakset* calls:

```
OPEN(file=, unit=kunit, form='unformatted', access='direct', recl={50+nx*ny})
```

The individual data records are packed and written by a call to *PAKREC*, once for each variable at each level. The routine calculates the record offset from the index record according to the variable and level information provided in the arguments and writes the record according to the order specified in *metdata.cfg*. The data can be supplied in any order. Note that although the level indicator (*ll*) goes from 1 to the number of levels, one is subtracted before it is written to the 50 byte header to be consistent with the definition of surface data to be at level "0". All the records in a time period may be initialized according to the value of the *kimi* flag. Initialization fills in the time variable for all records and assigns the variable identification field as *null*.

*CALL PAKREC(kunit,rvar,cvar,nx,ny,nxy,kvar,iy,im,id,ih,mn,ic,ll,kini)*

kunit - integer unit number of the defined file  
 rvar - input array of real\*4 data values  
 cvar - character\*1 array of packed data values  
 nx,ny - integer horizontal grid dimensions  
 nxy - integer product nx\*ny and length of cvar  
 kvar - character\*4 descriptor of variable being written  
 iy,im,id,ih - integer time identification  
 mn - integer minutes associated with the time (usually 0)  
 ic - integer forecast hour (hours from initialization)  
 ll - integer level indicator (1 to NZ)  
 kini - integer time period initialization flag (0-no; 1-yes)

When all the data records for a time period have been written, it is necessary to close that time period by writing its index record:

*CALL PAKNDX(kunit).*

At this point your program can return to *PAKREC* if data records for additional time periods are to be added to the file.

The key to the process is creating the proper configuration file for the data set you want to create. A sample *metdata.cfg* file for NOAA's NGM is shown below. The format is such that the first 20 characters are a dummy identification field followed by the data.

<u>20X,Format</u>	<u>Column_1 ID</u>	<u>Column_21 DATA</u>
A4	Data Source:	NGM
I4	Grid Number:	6
I4	Z-Coordinate:	1
F10	Pole Latitude:	90.0
...	Pole Longitude:	0.0
...	Reference Latitude:	60.0
...	Reference Longitude:	-105.0
...	Reference grid size:	182.9
...	Orientation:	0.0
...	Tangent Latitude:	90.0
...	Synch point X:	13.25
...	Synch point Y:	42.75
...	Synch Latitude:	90.0
...	Synch Longitude:	0.0
...	Reserved:	0.0
I4	Number X points:	33
I4	Number Y points:	28
I4	Number of Levels:	11
F6.,I3,(1X,A4)	Level 1:	1.0000 04 TPPT EXCO HFLX PRSS
...	Level 2:	0.9823 05 UWND VWND WWND SPHU TEMP
...	Level 3:	0.9432 05 UWND VWND WWND SPHU TEMP
...	Level 4:	0.8967 05 UWND VWND WWND SPHU TEMP
...	Level 5:	0.8437 05 UWND VWND WWND SPHU TEMP
...	Level 6:	0.7848 05 UWND VWND WWND SPHU TEMP
...	Level 7:	0.7210 05 UWND VWND WWND SPHU TEMP
...	Level 8:	0.6531 05 UWND VWND WWND SPHU TEMP

...	Level 9:	0.5820 05 UWND VWND WWND SPHU TEMP
...	Level 10:	0.5086 05 UWND VWND WWND SPHU TEMP
...	Level 11:	0.4341 05 UWND VWND WWND SPHU TEMP

It is important that the information contained in this file is correct as it not only controls the writing of the file, but much of the information is written into the index record of each time period. The model decodes this information to set up the internal processing of the meteorological data. A complete description of *metadata.cfg* follows:

Record 1 consists of a four character string that identifies the source of the meteorological data. This string will be passed through to many of the output graphics.

Record 2 is an optional integer identification of the meteorological data grid. It was used extensively in previous meteorological data file formats. It is not used in Hysplit\_4 applications.

Record 3 is an integer number that identifies the vertical coordinate system. Only four coordinate types are recognized: 1-pressure sigma; 2-pressure absolute; 3-terrain sigma; 4-hybrid sigma.

Records 4 & 5 identifies the pole position of the grid projection. Most projections will either be defined at +90 or -90 depending upon the hemisphere. The longitude would be the point 180 degrees from which the projection is cut.

Records 6 & 7 is the reference position at which the grid spacing is defined.

Record 8 is the grid spacing in km at the reference position.

Record 9 is the grid orientation or the longitude of the meridian which is parallel to the up-down direction of the grid.

Record 10 is the angle between the axis and the surface of the cone. For regular projections it is equal to the latitude at which the grid is tangent to the earth's surface. A polar stereographic grid would be tangent at either 90 or -90, while a Mercator projection is tangent at 0 latitude. A Lambert Conformal projection would be in between the two limits. An oblique stereographic projection would have a cone angle of 90.

Records 11 & 12 are used to equate a position on the grid with a position on the earth as given in Records 13 & 14. In this example, the position indicated is the center of the grid located over the North Pole. Any position is acceptable. It need not even be on the grid.

Record 15 is not currently used.

Records 16 & 17 identify the number of grid points in each direction.

Record 18 is the number of levels in the vertical, including the surface level.

Record 19, through the number of levels, identifies the height of each level in appropriate units according the definition of the vertical coordinate, the number of variables at that level, and the four character identification string for each variable. The height units are as follows for each coordinate: 1-sigma (fraction); 2-pressure (mb); 3-terrain (fraction); 4-hybrid (mb-offset.fraction)

### 5.3 Decoding Meteorological Data Files

One may want to develop other applications for Hysplit compatible meteorological data files. For these situations, some lower-level routines are provided in the source code library. The key to reading the meteorological files is decoding the index record. The format for this record is given below. Complete descriptions are similar to the variables in the discussion above.

<u>FORMAT</u>	<u>DESCRIPTION</u>
A4	Data Source
I3	Forecast hour (>99 the header forecast hr = 99)
I2	Minutes associated with data time
12F7.	1) Pole Lat, 2) Pole Long, 3) Tangent Lat, 4) Tangent Long, 5) Grid Size, 6) Orientation, 7) Cone Angle, 8) X-Synch point, 9) Y-Synch point, 10) Synch point lat, 11) Synch point long, 12) Reserved
3I3	1) Number x points, 2) Number y points, 3) Number levels
I2	Vertical coordinate system flag
I4	Length in bytes of the index record, excluding the first 50 bytes
<u>LOOP through the number of data levels</u>	
F6.	height of the first level
I2	number of variables at that level
<u>LOOP through the number of variables</u>	
A4	variable identification
I3	rotating checksum of the packed data
1X	Reserved space for future use

Once the index record has been read and decoded you have sufficient information to read and decode the data records. An un-packer is provided to convert the packed character\*1 array to a real\*4 array. It can also be used to extract a sub-grid from the full domain through specification of the sub-grid lower left corner:

*CALL PAKINP(rvar,cvar,nx,ny,nx0,ny0,lx,ly,prec,nexp,var1,ksum)*

rvar - real output array of integer dimensions lx,ly  
 cvar - character\*1 packed input array of length nx\*ny  
 nx,ny - integer dimensions of the full grid  
 nx0 - integer sub-grid position of left edge in nx units  
 ny0 - integer sub-grid position of lower edge in ny units  
 lx - integer first dimension of sub-grid length  
 ly - integer second dimension of sub-grid length  
 prec - real precision of packed data array  
 nexp - integer scaling exponent of packed data array  
 var1 - real value of array at position (1,1)  
 ksum - integer rotating checksum of packed data array

If the entire grid is to be unpacked then  $nx0=ny0=1$  and  $nx=lx$ ,  $ny=ly$ . The checksum (*ksum*) that is returned should be compared with the corresponding value in a table generated

from reading the index record. If you are not going to compare the checksum, set  $ksum = -1$ , this will save a little computer time. Due to the sub-grid option the checksum cannot be computed in the regular unpacking loop, but requires a second pass through the data. The checksum pass is enabled when  $ksum=0$ . It will then return a non-zero value. If you don't reset it to zero, no further checksums will be computed.

If you want to create your own packed data by converting a  $real*4$  array to the  $character*1$  packed data array use the following:

```
CALL PAKOUT(rvar,cvar,nx,ny,nxy,prec,nexp,var1,ksum)
```

Although the structure of the packed data may seem complex, unpacking is a very simple process, the basic elements of which are summarized in the Fortran code shown below. The value of each element is the sum of the initial value and the difference stored in that element location.

```
SUBROUTINE UNPACK (CPACK, RVAR, NX, NY, NXY, NEXP, VAR1)
CHARACTER*1 CPACK (NXY)
REAL*4 RVAR (NX, NY)
SCALE=2.0** (7-NEXP)
VOLD=VAR1
INDX=0
DO J=1, NY
  DO I=1, NX
    INDX=INDX+1
    RVAR (I, J) = ( ICHAR (CPACK (INDX)) - 127. ) /SCALE+VOLD
    VOLD=RVAR (I, J)
  END DO
  VOLD=RVAR (1, J)
END DO
RETURN
```

## 5.4 Sample Meteorological Programs

The source code for many different meteorological data applications can be found in */metdata* and */data2arl*. Most of these programs require the routines found in */metdata/source*. In addition, utility programs that convert GRIB formatted meteorological data files require GRIB decoding routines specific to the meteorological center that created the GRIB files; ECMWF decoders are provided with the data, NOAA decoders are available from the NCEP web site. In addition many of the GRIB decoders use some platform specific subroutines to read variable length records in Fortran in a direct-access mode (see the *\pcsubs* directory).

Programs found in *\metdata*:

*chk\_rec* - program to dump the first 50 bytes of each meteorological data record. Those bytes contain ASCII data describing the packing.

*chk\_file* - program to examine header and data records of an ARL packed meteorological data file. The program uses the same I/O subroutines common to Hysplit code. If this program does not work with your data file, neither will Hysplit.

*profile* - creates text file of the profile of meteorological variables at a specified location and time. The output written to the screen and to the file - profile.txt.

*display* - creates postscript file of meteorological data contoured and color filled for a single variable at a specified time. Output written to display.ps

*showgrid* - creates a postscript file that shows the extent of the domain of the meteorological data grid with marks at each grid point. Output written to showgrid.ps

Programs found in *\data2arl*:

*rsm2arl* - decodes the Regional Spectral Model (RSM) GRIB output fields on sigma surfaces to ARL packed format.

*avn2arl* - decodes NOAA Aviation model GRIB fields and converts them to ARL packed format.

*ncr2arl* - decodes NCEP/NCAR re-analysis GRIB fields and converts them to ARL packed format.

*ecm2arl* - decodes ECMWF model GRIB fields and converts them to ARL packed format. Requires ECMWF supplied GRIBEX subroutines.

*dat2arl* - creates a packed meteorological file for Hysplit using dummy fields hardwired into the program. The input meteorological data subroutines should be replaced by routines reading user supplied meteorological data files.

*content* - decodes NOAA grib sections without using the W3LIB routines. This program does not unpack the data but only lists the contents of the GRIB file.

*inventory* - decodes all the records within a NOAA grib file (without unpacking) providing content information. The program does not use the W3LIB libraries.

*unpacker* - decodes NOAA grib sections to a real data array without using the W3LIB routines.

## 5.5 Meteorological GUI Menu Tab

Some of the programs noted in Section 5.4 are also available through the *Meteorology* tab of the Hysplit GUI - in particular *display*, *profile*, *showgrid*, and *avn2arl*. These routines have not been fully integrated into an interactive environment and require entries through standard input. Some additional guidance in using these applications follows where the left side indicates the prompt the program writes to standard output and the right side in red indicates the user response to standard input. All examples use the sample meteorological data file.

### *showgrid*

```
Enter meteorological directory ...  
Enter meteorological file name ...  
Enter integer plotting increment:  
Enter lat/lon interval on background:  
Completed showgrid, enter to Continue ...
```

..\metdata\  
Oct1618.BIN  
1  
10  
Enter

### *profile*

```
Enter meteorological directory ...  
Enter meteorological file name ...  
File start time : 95 10 16 0 0  
File ending time: 95 10 18 22 0  
Enter Day & Hour: 16 0  
Enter Lat & Lon: 42.6532 -97.4386  
Updated file profile.txt ... 0,0 to exit  
Enter Day & Hour: 17 0
```

..\metdata\  
Oct1618.BIN  
17 0  
40.0 -90.0  
0 0



*display*

Enter meteorological directory ...

..\metdata\

Enter meteorological file name ...

Oct1618.BIN

File start time : 95 10 16 0 0

File ending time: 95 10 18 22 0

Enter Day & Hour: 16 0

17 0

Enter Center Lat & Lon: 42.6532 -97.4386

40.0 -100.0

Enter Radius in deg Lat: 10.0000

20.0

Enter 0 (B&W) or 1 (Color): 1

1

Level Height - Variable Character IDs

- 1 1.000 - TPPT EXCO HFLX PRSS ICWT SNOW SHGT MXLR WFLX CPPT MSLP
- 2 .982 - UWND VWND WWND SPHU TEMP
- 3 .943 - UWND VWND WWND SPHU TEMP
- 4 .897 - UWND VWND WWND SPHU TEMP
- 5 .844 - UWND VWND WWND SPHU TEMP
- 6 .785 - UWND VWND WWND SPHU TEMP
- 7 .721 - UWND VWND WWND SPHU TEMP
- 8 .653 - UWND VWND WWND SPHU TEMP
- 9 .582 - UWND VWND WWND SPHU TEMP
- 10 .509 - UWND VWND WWND SPHU TEMP
- 11 .434 - UWND VWND WWND SPHU TEMP

Enter Level Number:

2

Enter 4-Character Variable ID:

TEMP

Enter # of contours (max=8): 8

8

Enter contours interval (-1=auto): -1.000000000

-1.0

Data Load: TEMP at Level: 3 at Time: 17 0

Updated output file (display.ps) ... 0,0 to exit

Enter Day & Hour: 17 0

0 0

*avn2arl*

The NOAA Global Aviation Model (AVN) grib decoder can be run through the *Anonymous FTP / NCEP Server* tab of the Meteorology menu. The grib decoder is compiled to generate a Hysplit compatible input file by interpolating the AVN model output files from NCEP's global one degree latitude-longitude grid to 100x100 point conformal map at 100 km resolution centered about the point selected by the slider bars on the lower portion of the menu. AVN data consist of one grib file per forecast hour (0, +3, ... , out to +72 hours). Each grib file is 24Mb. The AVN decoder is run for each downloaded file, which creates an output file called *Data.avn*. This file is appended to *avnfcst.bin* to create one final output file that contains all forecast hours. It requires 970 Kb per forecast time.

The menu is divided into two parts. The upper portion is used to select the forecast initialization time and the duration of the forecast. The lower portion is used to set the output directory and center latitude-longitude of the extracted grid. If "Save all intermediate files" is checked, the global grib files are saved rather than deleted after each forecast hour is processed. In this way a different regional grid can be extracted from the global files without going through the FTP process again. To reprocess the saved grib files it is only necessary to set the FTP password to "skip".

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