

USER'S GUIDE FOR THE DEGADIS 2.1 DENSE GAS DISPERSION MODEL

By

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PREFACE

Version 2.1 of the elevated dense gas dispersion model, DEGADIS, has been developed by Dr. Jerry Havens and Dr. Tom Spicer of the University of Arkansas with the support of funding from the United States Environmental Protection Agency (EPA). While this model has not been extensively tested against field data and is subject to specific limitations and uncertainties, the EPA is making it publicly available through the National Technical Information Service (NTIS) as a research tool pending further model evaluation and development.

As developed, the DEGADIS 2.1 model is intended to completely replace the previous version of the model, DEGADIS 2.0. Version 2.1 incorporates a newly developed jet-plume model which will result in a plume which disperses in a fashion consistent with current Gaussian plume models if the plume becomes neutrally buoyant before it is predicted to fall to the ground. This represents a substantial improvement over the previous version and extends the applicability of the model into the not-denser-than-air regime. Specifically, the new jet-plume model provides for:

- (a) automatic adjustment of integration step size using the Runge-Kutta-Gill method;
- (b) elliptical plume cross-section with air entrainment specified consistent with the Pasquill-Gifford plume dispersion coefficient representation of atmospheric turbulent entrainment;
- (c) user specification of averaging time;
- (d) ground reflection of the plume when its lower boundary reaches ground level; and
- (e) application to scenarios where the plume remains aloft.

A technical description of the new jet-plume model is found in Section III of this User's Guide.

Source code for the model is provided in archived form with a dearchiving program. The code may be dearchived on an IBM-compatible PC using the instructions listed in the file named "README.NOW". The dearchived source code files should then be transferred to a VAX computer. At this point several files must then be renamed prior to compilation and execution. Specific information on the renaming process is contained in the file AAREADME.TXT. Print this file and the compilation batch file, BUILD.COM, prior to attempting compilation.

It is the concern of the EPA that this model be applied only within the framework of its intended use. To this end the user is referred to the specific recommendations in Section VI for model application. In addition the EPA is in the process of preparing a general guidance document to assist the typical regulatory user in the execution of denser-than-air cloud dispersion models.

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LIST OF SYMBOLS

a_e	empirical coefficient in Equation (III.2)
a_v	empirical constant (1.3) in Equations (V.6) and (V.27)
B_{EFF}	effective width of gas plume (m)
B'_i	local half width of source seen by observer i (m)
b	half width of horizontally homogeneous central section of gas plume (m)
b_e	empirical power in Equation (III.2)
b_j	characteristic jet width (m)
b_v	empirical constant (1.2) in Equations (V.8) and (V.28)
C_E	constant (1.15) in density intrusion (spreading) relation
c_p	heat capacity (J/kg K)
c_{p_a}	heat capacity of air (J/kg K)
c_{p_c}	heat capacity of contaminant (J/kg K)
c_{p_w}	heat capacity of water (liquid phase) (J/kg K)
c	local concentration (kg/m^3)
c_c	centerline concentration (kg/m^3)
$c_{c,L}$	vertically averaged layer concentration (kg/m^3)
c_f	friction coefficient
c'_c	centerline, ground-level concentration corrected for x-direction dispersion (kg/m^3)
D	source diameter (m)
D_h	added enthalpy (J/kg)
D'	diffusivity (m^2/s)
d_v	empirical constant (0.64) in Equations (V.4) and (V.12)
E	plume strength (kg/s)

$E(t)$	contaminant primary source rate (kg contaminant/s)
E_1	air entrainment associated with a free turbulent jet (kg/ms)
E_2	air entrainment associated with a bending plume (kg/ms)
E_3	jet/plume air entrainment associated with passive atmospheric dispersion (kg/ms)
$E_{3,a}$	air entrainment associated with passive atmospheric dispersion (kg/ms)
e_v	empirical constant (20.) in Equations (V.5) and (V.15)
F	overall mass transfer coefficient ($\text{kg}/\text{m}^2 \text{ s}$)
F_f	mass transfer coefficient due to forced convection ($\text{kg}/\text{m}^2 \text{ s}$)
F_n	mass transfer coefficient due to natural convection ($\text{kg}/\text{m}^2 \text{ s}$)
Fr	Froude number, $\rho_a u_a^2 / (g(\rho_e - \rho_a)D)$
Fr_h	"horizontal" Froude number, $u_a / [gD(\rho_e - \rho_a)/\rho_a]^{1/2}$
Fr_r	"release" Froude number, $V / [gD(\rho_e - \rho_a)/\rho_a]^{1/2}$
Gr	Grashoff number
g	acceleration of gravity (m/s^2)
H	characteristic release depth or depth of density intrusion or cloud (m)
H_a	ambient absolute humidity (kg water/kg dry air)
H_{EFF}	effective cloud depth (m)
H_h	height of head in density-driven flow (m)
H_L	total layer depth (m)
H_r	maximum plume rise above stack (m)
H_s	stack (exit) height (m)
H_t	height of tail in density-driven flow (m)
H_1	average depth of gravity current head (m)
H_4	depth of inward internal flow in a gravity current head (m)

h	enthalpy of source blanket (J/kg)
h_a	enthalpy of ambient humid air (J/kg)
h_E	enthalpy associated with primary source mass rate (J/kg)
h_f	heat transfer coefficient due to forced convection (J/m ² s K)
h_L	enthalpy of vertically averaged layer (J/kg)
h_n	heat transfer coefficient due to natural convection (J/m ² s K)
h_0	overall heat transfer coefficient (J/m ² s K)
h_p	enthalpy of primary source (J/kg)
h_w	enthalpy associated with mass flux of water from surface (J/kg)
J	release momentum ratio
K_0	constant in Equation (V.96) (m ^{1-γ₁})
K_y	horizontal turbulent diffusivity (m ² /s)
K_z	vertical turbulent diffusivity (m ² /s)
k	von Karman's constant, 0.35
$k_1 - k_6$	constants defined by Equations (III.14)-(III.19)
L	source length (m)
L_B	buoyancy length scale (m)
M	total cloud mass (kg)
M_a	total mass of air in the cloud (kg)
M_c	total mass of contaminant in the cloud (kg)
M_i	initial cloud mass (kg)
MW	molecular weight
$\cdot M_a$	mass rate of air entrainment into the cloud (kg/s)
$\cdot M_{w,s}$	mass rate of water transfer to the cloud from the water surface under the source (kg/s)

N	number of observers
Nu	Nusselt number
P	cloud momentum (kg m/s)
P _e	perimeter of R' (m)
P _h	momentum of head in density-driven flow (kg m/s)
P _t	momentum of tail in density-driven flow (kg m/s)
P _v	virtual momentum due to acceleration reaction (kg m/s)
Pr	Prandtl number
p	atmospheric pressure (atm)
p _{w,c}	partial pressure of water in the cloud (atm)
p _{w,s} [*]	vapor pressure of water at the surface temperature (atm)
Q	volumetric release rate (m ³ /s)
Q _E	source mass flux (kg/m ² s)
Q _e	volumetric entrainment flux (m/s)
Q ₁	flux of ambient fluid into front of gravity current head (m/s)
· Q _s	rate of heat transfer from the surface (J/s)
Q _*	atmospheric takeup flux (kg/m ² s)
Q _{*max}	maximum potential atmospheric takeup flux of contaminant (kg/m ² s)
q _s	surface heat flux (J/m ² s)
R	gas source radius (m)
R _h	inner radius of head in density-driven flow (m)
R _m	value of R when ($\pi R^2 Q_*$) is a maximum (m)
R _{max}	maximum radius of the cloud (m)
R _p	primary source radius (m)
R'	jet/plume region perpendicular to the axis

Ri_c	release Richardson number
Ri_f	Richardson number associated with the front velocity, Equation (V.11)
Ri_T	Richardson number associated with temperature differences, Equation (V.88)
Ri'_*	Richardson number associated with density differences corrected for convective scale velocity
Ri_*	Richardson number associated with density differences, Equation (V.78)
r	radial distance to jet/plume axis (m)
S	length of zone of flow establishment (m)
Sc	Schmidt number
Sh	Sherwood number
St_H	Stanton number for heat transfer
St_M	Stanton number for mass transfer
S_y	horizontal concentration scaling parameter (m)
S_z	vertical concentration scaling parameter (m)
S_{z0}	S_z at the downwind edge of the source ($x = L/2$) (m)
S_{z0_m}	value of S_{z0} when $(\pi R^2 Q_*)$ is a maximum (m)
s	distance along plume axis (m)
T	temperature associated with source blanket enthalpy (K)
$T_{c,L}$	temperature associated with layer-averaged enthalpy (K)
T_s	surface temperature (K)
t	time (s)
t_a	averaging time (s)
$t_{a,i}$	"instantaneous" averaging time associated with puff dispersion coefficients
t_p	averaging time associated with peak measured concentrations (s)

t_s	specified time (s)
t_{dn_i}	time when observer i encounters downwind edge (s)
t_{up_i}	time when observer i encounters upwind edge (s)
u	ambient (wind) velocity (m/s)
u_a	ambient average velocity (m/s)
u_c	jet/plume excess velocity at axis (m)
u_e	horizontal or frontal entrainment velocity (m/s)
u_{EFF}	effective cloud advection velocity (m/s)
u_f	cloud front velocity (m/s)
u_i	velocity of observer i (m/s)
u_L	average transport velocity associated with H_L (m/s)
u_x	wind velocity, along x-direction (m/s)
u_0	wind velocity measured at $z = z_0$ (m/s)
u_3	internal flow out of gravity current head (m/s)
u_4	internal flow into gravity current head (m/s)
u_*	friction velocity (m/s)
\bar{u}	characteristic average velocity (m/s)
v	jet velocity (m/s)
v_H	heat transfer velocity (0.0125 m/s) in Equation (V.38) (m/s)
w_a	mass fraction of air
w_c	mass fraction of contaminant
$w_{c,p}$	mass fraction of contaminant in primary source
w_e	vertical entrainment velocity associated with H_L (m/s)
w_*	convective scale velocity (m/s)
w'_e	entrainment velocity associated with H_{EFF} (m/s)
x_c	downwind distance to centerline ground contact (m)

$x_i(t)$	x position of observer i at time t (m)
x_{pi}	position of puff center due to observer i (m)
x_t	downwind distance where gravity spreading terminates (m)
x_v	virtual point source distance (m)
x_{dn_i}	x position of downwind edge of source for observer i
x_{up_i}	x position of upwind edge of source for observer i
\bar{x}	downwind distance to maximum rise (m)
x, y, z	Cartesian coordinates (m)
x_0	downwind edge of the gas source (m)
y'	local lateral dimension (m)
z_j	elevation of jet/plume centerline (m)
z_R	surface roughness (m)
z_0	reference height in wind velocity profile specification (m)
z'	local vertical dimension perpendicular to the jet/plume axis (m)
α	constant in power law wind profile
α_1	jet entrainment coefficient
α_2	line thermal entrainment coefficient
β_y	constant in σ_y correlation in Equation (V.97)
β_z	power in σ_z correlation
Γ	gamma function
γ	ratio of $(\rho - \rho_a)/c$
γ_1	constant in Equation (V.96)
γ_z	constant in σ_z correlation
Δ	ratio of $(\rho - \rho_a)/\rho$
ΔT	temperature driving force (K) ($T_s - T_{c,L}$) or ($T_s - T$)

Δ'	ratio of $(\rho - \rho_a)/\rho_a$
δ	empirical constant (2.15)
δ_L	empirical constant (2.15) in Equation (V.53)
δ_v	constant (0.20) in Equation (V.25)
δ_y	constant in σ_y correlation in Equation (V.97)
δ_z	constant in σ_z correlation
ϵ	frontal entrainment coefficient (0.59) in Equation (V.33)
ς	collection of terms defined by Equation (V.62) ($m^{-1/(1+\alpha)}$)
λ	Monin-Obukhov length (m)
μ	viscosity (kg/m s)
ρ	density of gas-air mixture (kg/m ³)
ρ_a	air density (kg/m ³)
ρ_c	jet/plume excess density at axis (kg/m ³)
ρ_e	density of released gas (kg/m ³)
ρ_L	vertically averaged layer density (kg/m ³)
ρ_0	density of contaminant's saturated vapor at T_0 (kg/m ³)
θ	angle between plume axis and horizontal (radians)
λ^2	turbulent Schmidt number, 1.42
σ_r	concentration profile parameter for an axisymmetrical jet/plume (m)
$\sigma_{x,a}$	Pasquill-Gifford x-direction (ambient) dispersion coefficient (m)
$\sigma_{y,a}$	Pasquill-Gifford y-direction (ambient) dispersion coefficient (m)
$\sigma_{y'}$	concentration profile parameter in the y' direction (m)
$\sigma_{z,a}$	Pasquill-Gifford z-direction (ambient) dispersion coefficient (m)
$\sigma_{z'}$	concentration profile parameter in the z' direction (m)
σ_1^2	variance associated with E_1 (m ²)
σ_2^2	variance associated with E_2 (m ²)

ϕ function describing influence of stable density stratification on vertical diffusion, Equation (V.76)
 $\hat{\phi}$ integrated source entrainment function
 Ψ logarithmic velocity profile correction function
 x_c jet centerline concentration at centerline touchdown

I. INTRODUCTION

Episodic releases of hazardous chemical gases from chemical process pressure relief operations can pose significant hazards to public health. Conventional air pollutant dispersion models may not be applicable for assessing the consequences of such releases, particularly when the gases released are denser than air. The DEGADIS model (Havens and Spicer, 1985) was designed to model the atmospheric dispersion of ground-level, area-source dense gas (or aerosol) clouds released with zero (initial) momentum into an atmospheric boundary layer over flat, level terrain. DEGADIS describes the dispersion processes which accompany the ensuing gravity-driven flow and entrainment of the gas into the atmospheric boundary layer, and it has been verified by comparison with a wide range of laboratory and field-scale heavy gas release/dispersion data. However, DEGADIS made no provision for processes which occur in high velocity releases, as from pressure relief valves.

Ooms, Mahieu, and Zelis (1974) reported a mathematical model for estimating the trajectory and dilution of dense gas jet plumes. The model comprised simplified balance equations for mass, momentum, and energy, with Gaussian similarity profiles for velocity, density, and concentration in the jet. Havens (1988) interfaced Ooms' jet-plume model with DEGADIS to provide for prediction of the trajectory and dilution of an elevated dense gas jet to ground contact, with (DEGADIS) prediction of the ensuing ground-level plume dispersion.

The purpose of this work was to improve the jet-plume/DEGADIS model as follows:

- The jet-plume part of the model was modified to provide for elliptical plume shape (cross-section) with air entrainment specified to be consistent with the Pasquill-Gifford plume dispersion coefficient representation of atmospheric turbulent entrainment. This modification allows application of the model to scenarios where the plume remains aloft.
- The jet-plume part of the model was modified to incorporate ground reflection when the plume (lower) boundary reaches ground level.
- The jet-plume part of the model was modified to provide for automatic adjustment of integration step-size (using the Runge-Kutta-Gill method as in DEGADIS). This modification improves the computational efficiency of the model.

II. CHARACTERIZING GAS DENSITY EFFECTS ON ATMOSPHERIC DISPERSION

Atmospheric dispersion of gas releases may involve the following fluid flow regimes:

jet • buoyancy-dominated • stably-stratified • passive dispersion

All four regimes, which may be present in different degrees depending on the rate and (characteristic) dimensions of the release, gas density, and characteristics of the atmospheric flow, should be accounted for in a general application dispersion model. The dispersion (and trajectory) of vertical jets (perpendicular to the wind flow) can be modeled with the jet-plume model described here, and the buoyancy-dominated, stably-stratified, and passive dispersion regimes (on level, unobstructed terrain) can be modeled with DEGADIS (Havens and Spicer, 1985).

The jet-plume model currently does not provide for simulation of horizontal jets (such as might occur from ruptured gas transfer lines). Consequently, we consider here other methods for estimating the relative importance of the four dispersion regimes for ground level releases with horizontal momentum. In rapid releases of large quantities of dense gas (with low initial momentum) a cloud having similar vertical and horizontal dimensions may form. In this "buoyancy-dominated regime", (gravity-induced) slumping and lateral spreading motion ensues until the kinetic energy of the buoyancy-driven flow is dissipated. The gravity-induced flow, which may effect mixing (primarily at the advancing vapor cloud front), can be an important determinant of the shape and extent of the gas cloud. After the kinetic energy of the buoyancy-driven flow is dissipated, the dispersion process following can be described as a "stably stratified" plume (or cloud) embedded in the mean wind flow. The density stratification present in this regime, which can be much stronger than that occurring naturally in the atmospheric boundary layer, tends to damp turbulence and reduce vertical mixing. As the dispersion proceeds, the stable stratification due to the dense gas decreases until the dispersion process can be represented as a neutrally buoyant plume (or cloud) in the mean wind flow. This (final) dispersion regime can be predicted with passive contaminant dispersion theory.

Havens and Spicer (1985) suggested criteria (based on water tunnel experiments reported by Britter (1980)) for determining the importance of each of the low-momentum flow regimes described above. In Britter's experiments brine was released (at floor level) into a water tunnel flow, and the lateral and upwind extent of the brine/water plume was measured as a function of the buoyancy length scale

$$L_B = Qg(\rho_e - \rho_a)/(\rho_e u^3)$$

where Q was the volumetric (brine) emission rate and u was the water-tunnel flow velocity. Britter's data indicated releases were passive

from the source when $L_B/D \lesssim 0.005$ and were dominated by the negative buoyancy dispersion regime when $L_B/D \gtrsim 0.1$. The following Release Richardson number criteria, based on these observations, were suggested:

If $Ric \gtrsim 30$	negative buoyancy-dominated
If $1 \lesssim Ric \lesssim 30$	stably stratified shear flow
If $Ric \lesssim 1$	passive dispersion

(II.1)

where $Ric = g(\rho_e - \rho_a)H/(\rho_a u_*^2)$. The reported values reflect the ratio $(u/u_*) = 16$ for Britter's water-tunnel flow, and the length scale corresponding to the depth of the layer was approximated by $H = Q/uD$.

High (initial) momentum releases require additional consideration if jet entrainment dominates other air entrainment mechanisms. In the negative buoyancy-dominated regime, the ambient flow does not strongly affect the rate of air entrainment. Consequently, the relative importance of buoyancy-driven and (horizontal, near-ground-level) jet flow effects can be evaluated using the criterion based on Britter's data (with the release velocity V used in place of the ambient velocity in the buoyancy length scale). Jet entrainment dominates the negative buoyancy regime entrainment when

$$\left(\frac{V}{u_*} \right)^2 \gtrsim 10 Ric \quad (\text{II.2})$$

This criterion suggests that dominance of jet effects decreases with increased jet density (since buoyancy-driven flow frontal air entrainment increases).

The air entrainment into a turbulent free jet can be estimated as (Wheatley, 1986):

$$\frac{dM_a}{dx} = \left(\frac{0.159}{2} \right) \left(\frac{2\pi R}{2} \right) V \rho_a \quad (\text{II.3})$$

The rate of (vertical) air entrainment (per unit width) for the stably-stratified shear flow and passive dispersion regimes predicted by DEGADIS is

$$\frac{d}{dx} (\rho_L u_L H_L) = \frac{\rho_a \delta_L k u_* (1 + \alpha)}{\phi(Ri_*)} \quad (\text{II.4})$$

where $\phi(Ri_*) = 0.88 + 0.099 Ri_*$. Using an effective width of $2\pi R$, a typical value of $\alpha = 0.2$, $\delta_L = 2.1$, $k = 0.35$, and $(u/u_*) = 30$ (typical of atmospheric boundary layers), the above equations can be combined to show that jet entrainment dominates the stably-stratified flow regime entrainment when

$$\frac{V}{u} \geq 16/(19 + Ri_c) \quad (II.5)$$

This criterion suggests that dominance of jet effects increases with increased jet density (since stably-stratified flow entrainment decreases). Further, when $Ri_c \lesssim 1$ the criterion suggests that jet effects dominate the passive dispersion regime when $(V/u) \geq 0.8$, which is consistent with the criterion suggested by Cude (1974) and Wheatley (1986).

Summarizing, the following procedure can be used to determine the (starting) dominant dispersion regime:

- (1) Calculate $Ri_c = g(\rho_e - \rho_a)H/(\rho_a u_*^2)$.
- (2) Determine the dominant (nonjet) dispersion regime using Equation (II.1).
- (3) Determine if horizontal jet effects dominate the regime determined in (2), using the relationships summarized in Table II.1.

Table II.1. Criteria for Determining Dominance of Jet Effects for Ground-Level Releases

Ground-level jet effects dominate:

- negative buoyancy-dominated regimes when $\left[\frac{V}{u_*} \right]^2 \geq 10 Ri_c$,
- stably stratified shear flow regimes when $V/u \geq 16/(19 + Ri_c)$, and
- passive dispersion regimes when $V/u \geq 0.8$

III. DESCRIPTION OF THE JET-PLUME MODEL

Gas (or aerosol) jet releases, such as might occur in chemical process pressure relief operations, are modeled as vertically directed releases of pure material into the atmospheric flow field. The jet is assumed to be discharged from a circular vent with a (constant) uniform velocity profile. At some distance downwind of the release, the velocity and concentration profiles are assumed Gaussian.

Zone of Flow Establishment

The region before Gaussian similarity velocity and concentration profiles are established is the so-called zone of flow establishment. Pratte and Baines (1967) reported that the length of the potential core (S_0) of a jet released from a tube perpendicular to a cross flow approaches the potential core length for a turbulent free jet with no cross flow as the ratio of the jet velocity to the cross-flow velocity (V/u_a) becomes large. Although S_0 is also a function of the Reynolds number (in the tube), Reynolds number dependence is less important for large release momentum. Pratte and Baines' (largest Reynolds number) data were correlated here as

$$(S_0/D) = 6.4 (1 - \exp(-0.48(V/u_a))) \quad (\text{III-1})$$

where D is the diameter of the release. The form of Equation (III-1) was chosen so that (S_0/D) approaches 0 as (V/u_a) approaches 0 and (S_0/D) approaches 6.4 as (V/u_a) becomes large. Chassaing et al. (1974) point out that similarity profiles are not established until some distance after the potential core has vanished. Consequently, the length of the zone of flow establishment (S) is modeled as

$$(S/D) = 7.7 \left\{ 1 - \exp \left[-0.48 \frac{\rho_e V}{\rho_a u_a} \right] \right\} \quad (\text{III.2})$$

where (V/u_a) has been replaced with $(\rho_e V / \rho_a u_a)$ to allow extended application for non-ambient-density gases.

Kamotani and Greber (1972) parameterized the trajectory of the jet-plume velocity profile in the zone of flow establishment as

$$x/D = a_e (z/D)^{b_e} \quad (\text{III.3})$$

where a_e and b_e are empirical constants determined as a function of the momentum ratio of the release ($J = \rho_e V^2 / (\rho_a u_a^2)$). Astleford et al. (1983) determined equations for a_e and b_e as functions of J from Kamotani and Greber's data. Morrow (1985) revised the equations to extend the range for smaller values of J and to include the effect of a release Froude number ($Fr = \rho_a u_a^2 / (g(\rho_e - \rho_a)D)$); these equations are shown in Table III.1. Correlations were also determined here for a_e and

b_e for $50 < J < 600$ from Kamotani and Greber's original work, based on a least squares fit; these equations are also assumed to apply for $J > 600$.

Table III.1. Equations for a_e and b_e .

Momentum Ratio (J)	Froude Number (Fr) Range	a_e	b_e	Ref.
$J \leq 0.036$	all	$18.519 J$	0.4	1
$0.036 < J \leq 10$	$Fr > 1.688$ or $Fr \leq 0$	$\exp[0.405465 + 0.24386 \ln J]$	0.4	1
	$0 < Fr \leq 1.688$	$\exp[0.2476 + 0.3016 \ln Fr + 0.24386 \ln J]$	0.4	1
$10 < J \leq 50$	all	$\exp[0.405465 + 0.131386 \ln J + 0.054931 (\ln J)^2]$	$\exp[-0.744691 - 0.074525 \ln J]$	2
$50 < J \leq 600$	all	$\exp[-2.58012 + 1.49202 \ln J - 0.097623 (\ln J)^2]$	$\exp[-0.446718 - 0.1950694 \ln J]$	3

References:

- 1 Morrow (1985)
- 2 Astleford, Morrow, and Buckingham (1983)
- 3 this work

At the end of the zone of flow establishment, the maximum jet-plume velocity is (approximately) the release velocity (V), and the centerline concentration is (approximately) the pure contaminant concentration. The jet angle (with the horizontal) is determined by x , z , and S from Equations (III.2) and (III.3).

Zone of Established Flow

For the zone of established flow, Ooms (1972) assumed Gaussian similarity profiles (single length scale) for velocity, density, and concentration. The concentration profile was

$$c(r,s) = c_c(s) \exp \left\{ - \left(\frac{r}{\lambda b_j(s)} \right)^2 \right\} \quad (\text{III.4})$$

where c_c represents the centerline concentration and λ^2 is a turbulent Schmidt number which represents the square of the ratio of the characteristic length of the concentration profile to the characteristic length of the velocity profile (b_j). By contrast, the Pasquill-Gifford representation of atmospheric turbulent dispersion assumes the following Gaussian profile:

$$c(x,y,z) = c_c(x) \exp \left\{ - \frac{1}{2} \left(\frac{y}{\sigma_y(x)} \right)^2 - \frac{1}{2} \left(\frac{z}{\sigma_z(x)} \right)^2 \right\} \quad (\text{III.5})$$

For a symmetric plume, $\sigma_y = \sigma_z$ ($= \sigma_r$), and $r^2 = y^2 + z^2$; Equation (III.5) becomes

$$c(x,r) = c_c(x) \exp \left\{ - \frac{1}{2} \left(\frac{r}{\sigma_r(x)} \right)^2 \right\}, \quad (\text{III.6})$$

and the profile,

$$c(s,y',z') = c_c(s) \exp \left\{ - \frac{1}{2} \left(\frac{y'}{\sigma_{y'}(s)} \right)^2 - \frac{1}{2} \left(\frac{z'}{\sigma_{z'}(s)} \right)^2 \right\} \quad (\text{III.7})$$

where y' and z' denote the local lateral and vertical dimensions in the jet-plume (Figure III.1), incorporates Equations (III.4) and (III.5) and provides for a circular cross section initially ($\sigma_{y'} = \sigma_{z'}$) with transition to an elliptical cross section.

Associated profiles for velocity and density are also assumed:

$$\bar{u}(s,y',z') = u_a \cos \theta + u_c(s) \exp \left\{ - \frac{1}{2} \left(\frac{\lambda y'}{\sigma_{y'}(s)} \right)^2 - \frac{1}{2} \left(\frac{\lambda z'}{\sigma_{z'}(s)} \right)^2 \right\} \quad (\text{III.8})$$

$$\rho(s,y',z') = \rho_a + \rho_c(s) \exp \left\{ - \frac{1}{2} \left(\frac{y'}{\sigma_{y'}(s)} \right)^2 - \frac{1}{2} \left(\frac{z'}{\sigma_{z'}(s)} \right)^2 \right\} \quad (\text{III.9})$$

Note that the turbulent Schmidt number now appears in the velocity profile. $\sigma_{y'}$ and $\sigma_{z'}$ should approach $\sigma_{y,a}$ and $\sigma_{z,a}$ for passive atmospheric

dispersion as the density and momentum of the jet/plume approach ambient values.

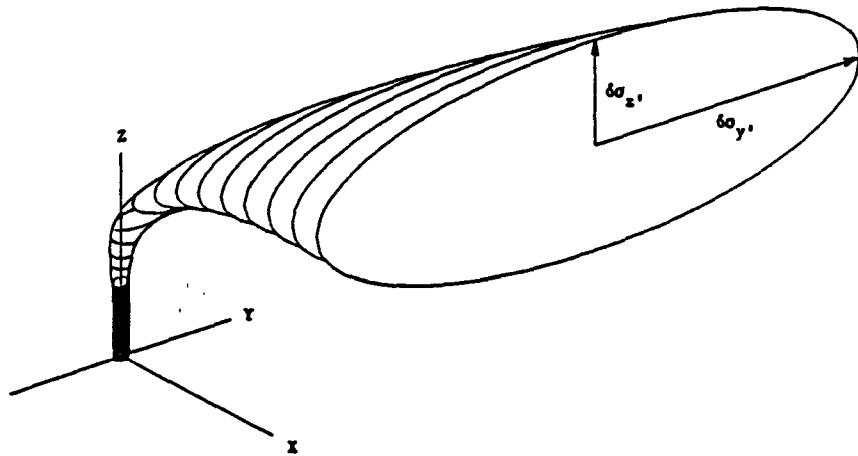


Figure III.1. Sketch of a Developing Jet in a Cross Flow.

The fundamental balance equations are:

Contaminant Mass Balance

$$\frac{d}{ds} \int_{R'} cu dR' = 0 \quad (\text{III.10})$$

Total Mass Balance

$$\frac{d}{ds} \int_{R'} \rho u dR' = E_1 + E_2 + E_3 \quad (\text{III.11})$$

X-Direction Momentum Balance

$$\frac{d}{ds} \int_{R'} \rho u (u \cos\theta) dR' = u_a (E_1 + E_2 + E_3) + c_d P_e \rho u_a^2 |\sin^3 \theta| / 2 \quad (\text{III.12})$$

Z-Direction Momentum Balance

$$\begin{aligned} \frac{d}{ds} \int_{R'} \rho u (u \sin\theta) dR' &= \int_{R'} g(\rho_a - \rho) dR' \\ &\quad - \text{sign}(\theta) c_d P_e \rho u_a^2 \sin^2 \theta \cos \theta / 2 \quad (\text{III.13}) \end{aligned}$$

The jet/plume boundary is assumed located where the concentration is one-tenth of the centerline value and encloses the region R' . (R' is an

ellipse or circle with major axes $\delta\sigma_y'$ and $\delta\sigma_z'$ where δ is a constant equal to 2.15; P_e is the perimeter of R' .) The term $(E_1 + E_2 + E_3)$ represents air entrainment into the jet plume by the three (independent) mechanisms discussed below. A drag force term assumed to act at right angles to the jet-plume trajectory is included on the right hand sides of Equations (III.12) and (III.13). The value used for c_d is 0.2; although this value of c_d is different from the value used by Ooms (1972) (since different length scales were used), its use in Equations (III.12) and (III.13) provides drag forces consistent with Ooms' predictions.

A (thermal) energy balance is not included in the balance equations given above. In the original formulation and in later work (Ooms, Mahieu, and Zelis, 1974), the energy balance was used to determine the developing jet temperature. However, since no provision was made for thermal energy transfer between the jet and its surroundings (e.g., the ground), the contaminant was (implicitly) assumed to mix adiabatically with ambient air, and an energy balance is not required if the adiabatic mixing properties are calculated beforehand.

For adiabatic mixing of two ideal gases with equal constant molal heat capacities, the ratio $\gamma = (\rho - \rho_a)/c$ is constant. Assumption of the density profile in Equation (III.9) and the concentration profile in Equation (III.7) is equivalent to assuming γ constant at any given distance, s . In the model development, $d\gamma/ds$ was assumed to be negligible. In the model implementation, γ is updated with distance based on the adiabatic mixing properties.

In Equations (III.10) - (III.13), u_a is considered constant for evaluation of the integrals. The value of u_a used is the averaged velocity over the top half of the jet plume, and u_a is updated with distance.

The integrals on the left-hand sides of Equations (III.10) - (III.13) do not have closed form solutions when R' represents an ellipse. If R' is approximated by a rectangle with an area equal to that of the original ellipse, the integrations are readily performed. (The approximating rectangle has sides with length $\sqrt{\pi}\delta\sigma_y'$ and $\sqrt{\pi}\delta\sigma_z'$.) The constants in Ooms' original formulation were compared to constants determined by approximating the circular cross section as square; the difference was less than 2%.

With the approximations stated above, six constants appear in the resulting equations:

$$k_1 = \frac{1}{\sigma_y' \sigma_z'} \int_{R'} \exp \left\{ -\frac{1}{2} \left(\left(\frac{y'}{\sigma_y'} \right)^2 + \left(\frac{z'}{\sigma_z'} \right)^2 \right) \right\} dR' \quad (\text{III.14})$$

$$k_2 = \frac{1}{\sigma_{y'} \sigma_{z'}} \int_{R'} \exp \left\{ -\frac{(1+\lambda^2)}{2} \left(\left(\frac{y'}{\sigma_{y'}} \right)^2 + \left(\frac{z'}{\sigma_{z'}} \right)^2 \right) \right\} dR' \quad (\text{III.15})$$

$$k_3 = \frac{1}{\sigma_{y'} \sigma_{z'}} \int_{R'} dR' = \pi \delta^2 \quad (\text{III.16})$$

$$k_4 = \frac{1}{\sigma_{y'} \sigma_{z'}} \int_{R'} \exp \left\{ -\frac{\lambda^2}{2} \left(\left(\frac{y'}{\sigma_{y'}} \right)^2 + \left(\frac{z'}{\sigma_{z'}} \right)^2 \right) \right\} dR' \quad (\text{III.17})$$

$$k_5 = \frac{1}{\sigma_{y'} \sigma_{z'}} \int_{R'} \exp \left\{ -\lambda^2 \left(\left(\frac{y'}{\sigma_{y'}} \right)^2 + \left(\frac{z'}{\sigma_{z'}} \right)^2 \right) \right\} dR' \quad (\text{III.18})$$

$$k_6 = \frac{1}{\sigma_{y'} \sigma_{z'}} \int_{R'} \exp \left\{ -\frac{(1+2\lambda^2)}{2} \left(\left(\frac{y'}{\sigma_{y'}} \right)^2 + \left(\frac{z'}{\sigma_{z'}} \right)^2 \right) \right\} dR' \quad (\text{III.19})$$

Air Entrainment

The balance equations for mass and x-direction momentum require specification of the air entrainment rate (E_1 , E_2 , and E_3 in Equations (III.11) and (III.13)).

E_1 represents the (near-field) air entrainment associated with jetting of the release. Although a cross flow is present, the momentum of the release dominates the ambient momentum when jetting is important, and the region near the jet release can be modeled as a free turbulent jet:

$$E_1 = \alpha_1 \rho_a P_e u_c \quad (\text{III.20})$$

where P_e is the perimeter of R' and α_1 is the entrainment coefficient associated with this area. Note that E_1 is taken to be zero if $u_c < 0$. A value of $\alpha_1 = 0.028$ is based on a summary of free turbulent jet experiments reported by List (1982); although this value of α_1 is different from the value used by Ooms (1972) (since different length scales were used), its use in Equation (III.20) provides rates of air entrainment consistent with Ooms' predictions.

E_2 represents the air entrainment associated with a cross-flow velocity component perpendicular to the jet-plume centerline. In the absence of jetting, large scale vortices (which entrain air) will form

in a plume due to the cross flow. Richards (1963) measured the air entrainment rate into line thermals released in a calm environment. Based on Richards' experiments, E_2 is modeled as follows:

$$E_2 = \alpha_2 \rho_a u_a | \sin\theta | \cos\theta \quad (\text{III.21})$$

where $u_a |\sin\theta|$ is the velocity component perpendicular to the jet-plume centerline and $\alpha_2 = 0.37$ is based on Richards' data. Ooms (1972) added the $\cos\theta$ term so that this contribution is present only in the far field. Again, although this value of α_2 is different from the value used by Ooms (1972), its use in Equation (III.21) provides rates of air entrainment consistent with Ooms' predictions.

E_3 represents the atmospheric air entrainment for a passive plume. For a passive plume, the atmospheric air entrainment rate ($E_{3,a}$) is given by

$$E_{3,a} = \frac{d}{dx} \int_{R'} \rho_a u_a dR' = \rho_a u_a \frac{d}{dx} \int_{R'} dR' = \rho_a u_a \frac{d}{dx} \left[k_3 \sigma_{y,a} \sigma_{z,a} \right] \quad (\text{III.22})$$

where $\sigma_{y,a}$ and $\sigma_{z,a}$ denote the ambient lateral and vertical Pasquill-Gifford dispersion coefficients, respectively. Since k_3 is constant, Equation (III.22) becomes

$$E_{3,a} = k_3 \rho_a u_a \left[\sigma_{z,a} \frac{d\sigma_{y,a}}{dx} + \sigma_{y,a} \frac{d\sigma_{z,a}}{dx} \right] \quad (\text{III.23})$$

where the derivatives can be calculated from accepted correlations for $\sigma_{y,a}$ and $\sigma_{z,a}$, such as

$$\sigma_{y,a} = \delta_y x^{\beta_y} \quad (\text{III.24})$$

$$\sigma_{z,a} = \delta_z x^{\beta_z} \exp(\gamma_z (\ln(x))^2) \quad (\text{III.25})$$

(Seinfeld, 1986). Typical values for δ_z , β_z , and γ_z are given in Table III.2.

Methods recommended in Section V for determining the effect of averaging time are applied for the jet-plume model, and values for δ_y and β_y as a function of stability and averaging time are given in Equation (V.101). For the jet plume, E_3 is modeled as

$$E_3 = k_3 \rho_a u_a \left[\sigma_{z,a} \frac{d\sigma_{y,a}}{dx} + \sigma_{z,a} \frac{d\sigma_{y,a}}{dx} \right] \cos\theta \quad (\text{III.26})$$

where the $\cos\theta$ term is included because E_3 is required as a function of s .

Table III.2. Constants for Determination of σ_z
from Seinfeld (1986) after Turner
(1970)

Stability Class	δ_z^1	β_z	γ_z
A	107.7	-1.7172	0.2770
B	0.1355	0.8752	0.0136
C	0.09623	0.9477	-0.0020
D	0.04134	1.1737	-0.0316
E	0.02275	1.3010	-0.0450
F	0.01122	1.4024	-0.0540

¹for use in $\sigma_z = \delta_z^1 x^z \exp(\gamma_z (\ln(x))^2)$ where
 σ_z and x are in meters.

Gravitational Force

The gravitational force term in Equation (III.13) is given by

$$\int_{R'} g(\rho_a - \rho) dR' = g\gamma \int_{R'} c dR' = k_1 g \gamma c \sigma_y' \sigma_z', \quad (\text{III.27})$$

As discussed earlier, γ is assumed constant in the equation development but is updated with distance in the model implementation. Note that Equation (III.27) applies for $\rho \leq \rho_a$ as well as $\rho > \rho_a$.

Additional Constraints

Along with an equation of state (such as the ideal gas law or an adiabatic mixing relationship between concentration and density), balance Equations (III.10) - (III.13) provide five constraints on the six unknowns c_c , ρ_c (or γ), u_c , θ , σ_y' , and σ_z' . Therefore, an additional constraint is required to solve the equations. The momentum balances provide the information needed to determine θ and u_c . The contaminant mass balance provides the information to determine c_c . The total mass balance provides the information to determine the product $\sigma_y' \sigma_z'$, but the balance equations provide no guidance for the individual values of σ_y' and σ_z' . (No additional constraint was needed in Ooms' model since a single length scale was used.) The last constraint is to

be specified so that the values of σ_y' and σ_z' are consistent with atmospheric processes.

The model treats three (assumed) independent mechanisms for air entrainment. For independent processes, the variances are additive:

$$\sigma_{y'}^2 = \sigma_1^2 + \sigma_2^2 + \sigma_{y,a}^2 \quad (\text{III.28})$$

$$\sigma_{z'}^2 = \sigma_1^2 + \sigma_2^2 + \sigma_{z,a}^2 \quad (\text{III.29})$$

where σ_1^2 and σ_2^2 represent the variances associated with entrainment processes E_1 and E_2 , respectively. Combining Equations (III.28) and (III.29):

$$\sigma_{y'}^2 - \sigma_{z'}^2 = \sigma_{y,a}^2 - \sigma_{z,a}^2 \quad (\text{III.30})$$

Equation (III.30) acts as the final constraint on the system. In the near field, $\sigma_{y,a}^2 - \sigma_{z,a}^2$ is small and $\sigma_{y'} \approx \sigma_{z'}$. In the far field, $\sigma_{y'}$ and $\sigma_{z'}$ will approach $\sigma_{y,a}$ and $\sigma_{z,a}$, respectively.

Closure

Initial conditions for the zone of established flow are determined from the zone of flow establishment; initial values of $\sigma_{y'}$ and $\sigma_{z'}$ are determined from the contaminant mass balance and the (known) release rate. With $\lambda^2 = 1.42$ (List, 1982), the assumed profiles, balance equations, and assumptions are combined to give a system of ordinary differential equations as follows:

$$\left| \begin{array}{cccc} A_{11} & A_{12} & A_{13} & A_{14} \\ A_{21} & A_{22} & A_{23} & A_{24} \\ A_{31} & A_{32} & A_{33} & A_{34} \\ A_{41} & A_{42} & A_{43} & A_{44} \end{array} \right| \left| \begin{array}{c} \frac{dc}{ds} \\ \frac{d}{ds} (\sigma_{y'}, \sigma_{z'}) \\ \frac{d\theta}{ds} \\ \frac{du_c}{ds} \end{array} \right| = \left| \begin{array}{c} B_1 \\ B_2 \\ - \\ B_3 \\ B_4 \end{array} \right|$$

where

(Contaminant Mass Balance)

$$A_{11} = (k_1 u_a \cos \theta + k_2 u_c) \sigma_{y'} \sigma_{z'}$$

$$A_{12} = (k_1 u_a \cos\theta + k_2 u_c) c_c$$

$$A_{13} = -k_1 u_a \sin\theta c_c \sigma_y, \sigma_z,$$

$$A_{14} = k_2 c_c \sigma_y, \sigma_z,$$

$$B_1 = 0$$

(Total Mass Balance)

$$A_{21} = 0$$

$$A_{22} = \rho_a (k_1 u_a \cos\theta + k_2 u_c)$$

$$A_{23} = -\rho_a k_3 u_a \sin\theta \sigma_y, \sigma_z,$$

$$A_{24} = \rho_a k_4 \sigma_y, \sigma_z,$$

$$B_2 = E_1 + E_2 + E_3$$

(z-Direction Momentum Balance)

$$A_{31} = (k_1 u_a^2 \cos^2\theta \sin\theta + 2k_2 u_a u_c \cos\theta \sin\theta + k_6 u_c^2 \sin\theta) \gamma \sigma_y, \alpha_z,$$

$$A_{32} = \rho_a (k_3 u_a^2 \cos^2\theta \sin\theta + 2k_4 u_a u_c \cos\theta \sin\theta + k_5 u_c^2 \sin\theta)$$

$$+ (k_1 u_a^2 \cos^2\theta \sin\theta + 2k_2 u_a u_c \cos\theta \sin\theta + k_6 u_c^2 \sin\theta) \gamma c_c$$

$$\begin{aligned}
A_{33} = & \rho_a (-2k_3 u_a^2 \cos\theta \sin^2\theta + k_3 u_a^2 \cos^3\theta \\
& - 2k_4 u_a u_c \sin^2\theta + 2k_4 u_a u_c \cos^2\theta \\
& + k_5 u_c^2 \cos\theta) \sigma_y, \sigma_z, \\
& + (-2k_1 u_a^2 \cos\theta \sin^2\theta + k_1 u_a^2 \cos^3\theta \\
& - 2k_2 u_a u_c \sin^2\theta + 2k_2 u_a u_c \cos^2\theta \\
& + k_6 u_c^2 \cos\theta) \gamma c_c \sigma_y, \sigma_z, \\
A_{34} = & \rho_a (2k_4 u_a \cos\theta \sin\theta + 2k_5 u_c \sin\theta) \sigma_y, \sigma_z, \\
& + (2k_2 u_a \cos\theta \sin\theta + 2k_6 u_c \sin\theta) c_c \sigma_y, \sigma_z, \\
B_3 = & g k_1 \gamma c_c \sigma_y, \sigma_z,
\end{aligned}$$

(x-Direction Momentum Balance)

$$\begin{aligned}
A_{41} = & (k_1 u_a^2 \cos^3\theta + 2k_2 u_a u_c \cos^2\theta + k_6 u_c^2 \cos\theta) \gamma \sigma_y, \sigma_z, \\
A_{42} = & \rho_a (k_3 u_a^2 \cos^3\theta + 2k_4 u_a u_c \cos^2\theta + k_5 u_c^2 \cos\theta) \\
& + (k_1 u_a^2 \cos^3\theta + 2k_2 u_a u_c \cos^2\theta + k_6 u_c^2 \cos\theta) \gamma c_c \\
A_{43} = & \rho_a (-3u_a^2 \cos^2\theta \sin\theta - 4k_4 u_a u_c \cos\theta \sin\theta \\
& - k_5 u_c^2 \sin\theta) \sigma_y, \sigma_z, \\
& + (-3k_1 u_a^2 \cos^2\theta \sin\theta - 4k_2 u_a u_c \cos\theta \sin\theta \\
& - k_6 u_c^2 \sin\theta) \gamma c_c \sigma_y, \sigma_z, \\
A_{44} = & \rho_a (2k_4 u_a \cos^2\theta + 2k_5 u_c \cos\theta) \sigma_y, \sigma_z, \\
& + (2k_2 u_a \cos^2\theta + 2k_6 u_c \cos\theta) \gamma c_c \sigma_y, \sigma_z, \\
B_4 = & u_a (E_1 + E_2 + E_3)
\end{aligned}$$

and

$$\begin{aligned}
 k_1 &= 2\pi \left[\operatorname{erf} \left(\frac{\delta}{2} \left(\frac{\pi}{2} \right)^{\frac{1}{2}} \right) \right]^2 \\
 k_2 &= \left(\frac{2\pi}{1 + \lambda^2} \right) \left(\operatorname{erf} \left(\frac{\delta}{2} \left(\frac{\pi(1 + \lambda^2)}{2} \right)^{\frac{1}{2}} \right) \right)^2 \\
 k_3 &= \pi \delta^2 \\
 k_4 &= \frac{2\pi}{\lambda^2} \left(\operatorname{erf} \left(\frac{\delta \lambda}{2} \left(\frac{\pi}{2} \right)^{\frac{1}{2}} \right) \right)^2 \\
 k_5 &= \frac{\pi}{\lambda^2} \left(\operatorname{erf} \left(\frac{\delta \lambda \sqrt{\pi}}{2} \right) \right)^2 \\
 k_6 &= \left(\frac{2\pi}{1 + 2\lambda^2} \right) \left(\operatorname{erf} \left(\frac{\delta}{2} \left(\frac{\pi(1 + 2\lambda^2)}{2} \right)^{\frac{1}{2}} \right) \right)^2
 \end{aligned}$$

After c_c is determined from the system of equations above, the contribution from the reflected image is approximated by

$$c_c \exp \left\{ -\frac{1}{2} \left(\frac{2 z_j}{\sigma_{z'}} \right)^2 \right\} \quad (\text{III.31})$$

so that the reported concentration is

$$c_c'' = c_c \left\{ 1 + \exp \left\{ -\frac{1}{2} \left(\frac{2 z_j}{\sigma_{z'}} \right)^2 \right\} \right\} \quad (\text{III.32})$$

The reported mole fraction and density are calculated based on c_c'' . Although the method of images is a common approach for accounting for ground interactions with the plume, application of the method of images to this problem may cause c_c'' to increase with downwind distance. Note that at the centerline touchdown, $c_c'' = 2c_c$.

For Ooms model predictions, DEGADIS is used to model the subsequent dispersion when the jet-plume reaches ground level ($z_j = 0$). The ground-level source input to DEGADIS is modeled as a circular area source with radius $\delta \sigma_y'$, and concentration equal to the centerline value from the Ooms model. Note that the downwind distance reported in DEGADIS includes the distance to touchdown calculated in the Ooms model. A description of the DEGADIS model is included in Section V.

IV. EVALUATION OF THE JET-PLUME MODEL

A series of simulations were made with the jet-plume model to:

- compare the model predictions with wind tunnel dense gas jet trajectory and dilution data
- characterize the sensitivity of the model to the specification of entrainment coefficients α_1 and α_2 .

Table IV.1 shows the "typical" jet releases simulated.

Table IV.1. Specification of Gas Release Rates for Modeling.

Jet Diameter m	Jet Velocity m/s	Gas Release Rate kg/s
0.05 (~2 in)	30.6 (~100 ft/s)	0.24
0.2 (~8 in)	91.7 (~300 ft/s)	11.52
0.5 (~20 in)	213.9 (~700 ft/s)	168.0

Comparison with Wind Tunnel Test Data

Hoot, Meroney, and Peterka (1973) reported plume rise, downwind distance to plume centerline touchdown, and dilution at touchdown for wind tunnel jet releases of Freon-12/air mixtures. The ranges of experimental variables studied were:

gas specific gravity (air = 1)	1.1 - 4.6
gas exit diameter, cm	0.32 and 0.64
gas exit height, cm	7.6 and 15.2
gas exit velocity/wind velocity ratio	2.5 - 25
wind (tunnel) velocity, m/s	0.23 and 0.46

Correlations (of the wind tunnel data) were presented for plume rise, distance to plume touchdown, and plume centerline concentration at (centerline) ground contact, in a laminar crosswind:

Plume Rise

$$H_r = 1.32 D [(V/u_a)(\rho_e/\rho_a)]^{1/3} Fr_r^{2/3} \quad (\text{IV.1})$$

Downwind Distance to Maximum Rise

$$\bar{X} = (D u_a/V) Fr_r^2 \quad (\text{IV.2})$$

Downwind Distance to Centerline Ground Contact

$$x_c = 0.56 D \left\{ \left(H_r/D \right)^3 \left[\left(2 + H_s/H_r \right)^3 - 1 \right] u_a/V \right\}^{1/2} Fr_h + \bar{x} \quad (\text{IV.3})$$

Centerline Concentration at Ground Contact

$$x_c = \frac{6.0 Q}{u_a (2H_r + H_s)^2} \quad (\text{IV.4})$$

where D = initial jet diameter (m)

Fr = "release" Froude number, $V/[g D (\rho_e - \rho_a)/\rho_a]^{1/2}$

Fr_h = "horizontal" Froude number, $u_a/[g D (\rho_e - \rho_a)/\rho_a]^{1/2}$

H_r = maximum height of plume rise above stack (m)

H_s = exit height (m)

Q = steady-state jet rate (kg/s)

u_a = wind velocity (m/s)

V = initial jet velocity (m/s)

x_c = downwind distance to ground contact (centerline) (m)

\bar{x} = downwind distance to maximum rise (m)

ρ_a = ambient air density (kg/m^3)

ρ_e = initial jet density (kg/m^3)

x_c = jet centerline concentration at centerline touchdown (kg/m^3)

Note that Equation (IV.4) is based on analysis by Briggs presented in Guinnup (1989).

Table IV.2 compares the jet-plume model predictions for plume rise, downwind distance to ground contact, and concentration at ground contact with the results of Hoot et al.'s wind-tunnel data correlations for the "low diameter/low velocity", "typical diameter/typical velocity", and "high diameter/high velocity" cases (Table IV.1) in 3 and 6 m/s winds. Based on a scaled release height of 10 m, a surface roughness of 2.2 mm was assumed with the 3 and 6 m/s velocities because the jet-plume model implementation assumes a logarithmic velocity profile. This assumption causes overprediction of the wind tunnel velocities by greater than 20% for scaled heights greater than ~40 m. Since the wind tunnel correlations were for jets in a laminar crosswind, the simulations shown in Table IV.2 were made with the entrainment due to atmospheric turbulence (E_3) set to zero.

Table IV.2. Comparison of Jet-Plume Model Prediction with Hoot et al.'s Wind Tunnel Data Correlation

Gas Jet Density: 4.0 kg/m ³		Gas Jet Elevation: 10 m		
(Hoot et al. correlation / Jet-Plume model prediction)				
Maximum Rise, m (3, 6 m/s)	Distance to Contact, m (3, 6 m/s)	Centerline Concentration at Ground Contact kg/m ³ x 10 ³ (3, 6 m/s)		
<u>Low Diameter (0.05 m) - Low Velocity (30.6 m/s)</u>				
3.0/2.6	2.4/1.7	150/160	375/370	1.9/4.2 1.1/2.6
<u>Typical Diameter (0.2 m) - Typical Velocity (91.7 m/s)</u>				
22.6/20.9	17.9/15.2	165/170	350/320	7.6/17.4 5.4/14.3
<u>High Diameter (0.5 m) - High Velocity (213.9 m/s)</u>				
97.0/87.4	76.9/66.5	320/380	650/680	8.1/18.3 6.3/15.5

The model predictions of maximum rise and distance to centerline ground contact are in good agreement with the wind tunnel correlations. The agreement between model predictions of distance to centerline ground contact and the wind tunnel correlations tends to degrade as the maximum rise increases because the logarithmic velocity profile assumed in the model overpredicts the (laminar) wind tunnel velocity. The model prediction of the maximum concentration at centerline ground contact differs by a factor of about two from the observed concentration. Note that if interaction with the ground were not included in the calculations, the predicted model concentrations would be reduced by exactly a factor of 2. Therefore, the agreement between model predictions of the maximum concentration at centerline ground contact and the wind tunnel correlations is considered good.

Sensitivity to Entrainment Coefficient Specification

The model sensitivity to the coefficients α_1 and α_2 was determined by systematic variation around the values given above. All simulations were of denser-than-air gas jets (initial density 4.0 kg/m³) exiting vertically upward at 10 meters elevation. The predictions assumed a

logarithmic velocity profile with 3 and 6 m/s velocities at 10 m elevation, a surface roughness of 2.2 mm, and D stability. Table IV.3 shows the effect of individually varying the entrainment coefficients α_1 and α_2 by factors of two below and above the values given above, for the "low diameter/low velocity", "typical diameter/typical velocity", and "high diameter/high velocity" cases (Table IV.1) in 3 and 6 m/s winds.

The predictions of maximum rise and distance to centerline ground contact are relatively insensitive to factor-of-four variations in α_1 and α_2 over the range of release conditions in Table IV.1. The predictions of maximum concentration at centerline ground contact are more sensitive to variations in α_1 and α_2 ; factor-of-four variations in α_1 and α_2 resulted in concentration changes of less than a factor of ~2 when concentrations were compared at the same distances.

Table IV.3. Sensitivity of Jet-Plume Model Prediction to Variation of Entrainment Coefficients α_1 and α_2

Gas Jet Density: 4.0 kg/m ³				Gas Jet Elevation: 10 m		
Entrainment Coefficients α_1 α_2	Maximum Rise, m (3, 6 m/s)	Distance to Ground Contact, m (3, 6 m/s)	Centerline Concentration at Ground Contact kg/m ³ x 10 ³ (3, 6 m/s)			
<u>Low Diameter (0.05 m) - Low Velocity (30.6 m/s)</u>						
0.014 0.37	2.9 1.6	740* 420*	--	--		
0.028 0.37	2.5 1.5	750* 420*	--	--		
0.056 0.37	2.1 1.4	750* 420*	--	--		
0.028 0.185	2.9 1.7	750* 420*	--	--		
0.028 0.37	2.5 1.5	750* 420*	--	--		
0.028 0.74	2.1 1.2	740* 420*	--	--		
<u>Typical Diameter (0.2 m) - Typical Velocity (91.7 m/s)</u>						
0.014 0.37	25.7 16.9	300 1770	4.7	0.22		
0.028 0.37	20.6 14.4	240 1440	6.2	0.31		
0.056 0.37	16.4 12.2	210 1190	7.7	0.42		
0.028 0.185	23.0 16.4	220 1530	9.1	0.30		
0.028 0.37	20.6 14.4	240 1440	6.2	0.31		
0.028 0.74	17.3 11.9	280 1400	3.6	0.30		
<u>High Diameter (0.5 m) - High Velocity (213.9 m/s)</u>						
0.014 0.37	112 79.4	510 1280	10.2	3.2		
0.028 0.37	86.6 64.7	420 1030	13.1	4.4		
0.056 0.37	66.9 52.6	350 860	15.4	5.7		
0.028 0.185	95.0 73.4	380 1010	21.4	5.6		
0.028 0.37	86.6 64.7	420 1030	13.1	4.4		
0.028 0.74	74.7 53.7	480 1100	6.8	3.0		

*The plume remained aloft. The reported distance is the estimated distance to 10 ppm.

V. DESCRIPTION OF THE DEGADIS DENSE GAS DISPERSION MODEL

DEGADIS (DEnse GAs DISpersion) model development was sponsored by the U.S. Coast Guard and the Gas Research Institute (Havens and Spicer, 1985). DEGADIS is an adaptation of the Shell HEGADAS model (Colenbrander, 1980 and Colenbrander and Puttock, 1983) and incorporates some techniques proposed by van Ulden (1983). It is designed to model the atmospheric dispersion of ground-level, area-source dense gas (or aerosol) clouds released with zero (initial) momentum into an atmospheric boundary layer flow over flat, level terrain.

If a gas release rate does not exceed the potential atmospheric takeup rate, the gas is taken up directly by the atmospheric flow and dispersed downwind. If a gas release rate exceeds the potential atmospheric takeup rate, a denser-than-air "secondary source" blanket is formed over the primary source, and this near-field, buoyancy-dominated regime is modeled using a lumped parameter model (spatially averaged properties) which incorporates air entrainment at the gravity-spreading front using a frontal entrainment velocity (Figure V.1). Downwind dispersion is modeled with a power law concentration distribution in the vertical direction and a modified Gaussian profile in the horizontal direction with a power law specification for the wind profile. The downwind dispersion models the ensemble average of the gas concentrations.

Secondary Source Blanket Formation

A denser-than-air gas "secondary source" cloud or blanket may be formed over an evaporating liquid pool (or otherwise specified ground-level emission source) or by the instantaneous release of a volume of gas. A lumped parameter model of such a blanket is illustrated in Figure V.1. The blanket is represented as a cylindrical gas volume which spreads laterally as a density-driven flow with entrainment from the top of the source blanket by wind shear and air entrainment into the advancing front edge. The blanket spreads laterally until the atmospheric takeup rate from the top is matched by the air entrainment rate from the side and, if applicable, by the rate of gas addition from under the blanket. The blanket center is assumed stationary over the source.

Secondary Source Blanket Extent for Ground-Level Emission Sources

The downwind emission rate from the blanket is equal to the potential atmospheric takeup rate, Q_{\max} . For $E(t)/\pi R_p^2(t) > Q_{\max}$, a blanket is formed over the primary source. The blanket frontal (spreading) velocity is modeled as

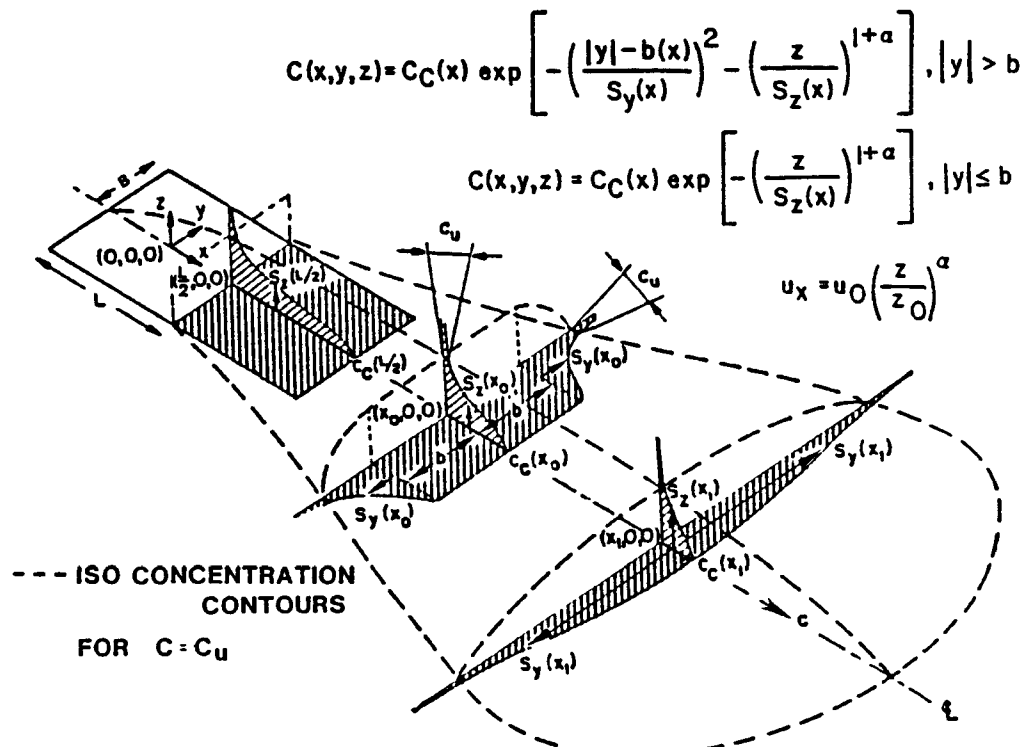
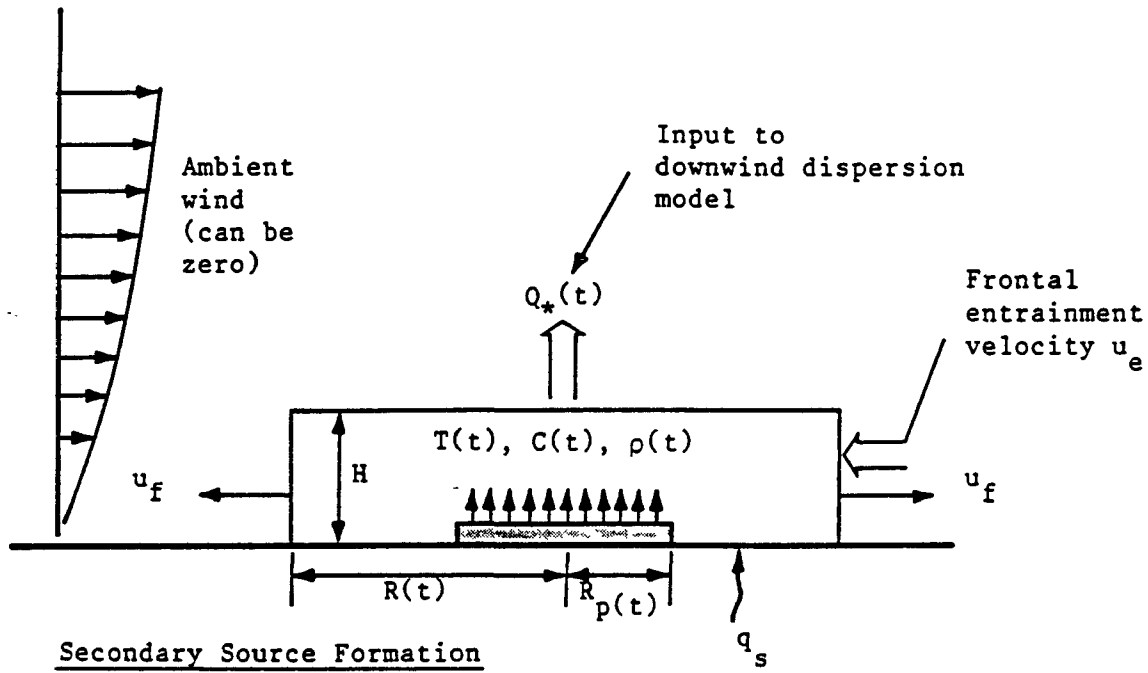


Figure V.1. Schematic Diagram of DEGADIS Dense Gas Dispersion Model.

$$u_f = C_E \left[g \left(\frac{\rho - \rho_a}{\rho_a} \right) H^2 \right]^{1/2} \quad (V.1)$$

where ρ is the average density of the blanket for ($\rho > \rho_a$). $C_E = 1.15$, based on laboratory measurements of cloud spreading velocity (Havens and Spicer, 1985).

The blanket radius R is determined by integrating $dR/dt = u_f$. When the total mass of the blanket is decreasing with time (i.e., the atmospheric takeup exceeds the input rate from the primary source plus the air entrainment rate), the radius is modeled as $(dR/dt)/R = (dH/dt)/H$, with the radius of the blanket constrained to be greater than or equal to the primary source radius R_p .

Secondary Source Blanket Extent for Instantaneous Gas Releases

The gravity intrusion model (Equation (V.1)) overpredicts initial velocities for instantaneous, above-ground releases of a denser-than-air gas since the initial acceleration phase is not modeled. Consequently, the following procedure adapted from van Ulden (1983) is incorporated in DEGADIS.

For instantaneous gas releases, the radially symmetric cloud is considered to be composed of a tail section with height H_t and radius R_h and a head section with height H_h (Figure V.2). A momentum balance is used to account for the acceleration of the cloud from rest; the effect of ambient (wind) momentum is ignored. Although the following equations are derived assuming the absence of a primary source, the resulting equations are assumed to model the secondary source cloud development when the primary source rate is nonzero. When the frontal velocity from the momentum balance is the same as Equation (V.1), the momentum balance is no longer applied and the frontal velocity is given by Equation (V.1).

There are three main forces acting on the cloud: a static pressure force (F_p), a dynamic drag force (F_d), and a force which accounts for the acceleration reaction of the ambient fluid, represented as a rate of virtual momentum change with respect to time ($-dP_v/dt$). Denoting the momentum of the head and tail as P_h and P_t , respectively, the momentum balance is

$$\frac{dP}{dt} = \frac{d}{dt} (P_h + P_t) = F_p + F_d - \frac{dP_v}{dt} \quad (V.2)$$

or

$$\frac{d}{dt} (P_h + P_t + P_v) = F_p + F_d \quad (V.3)$$

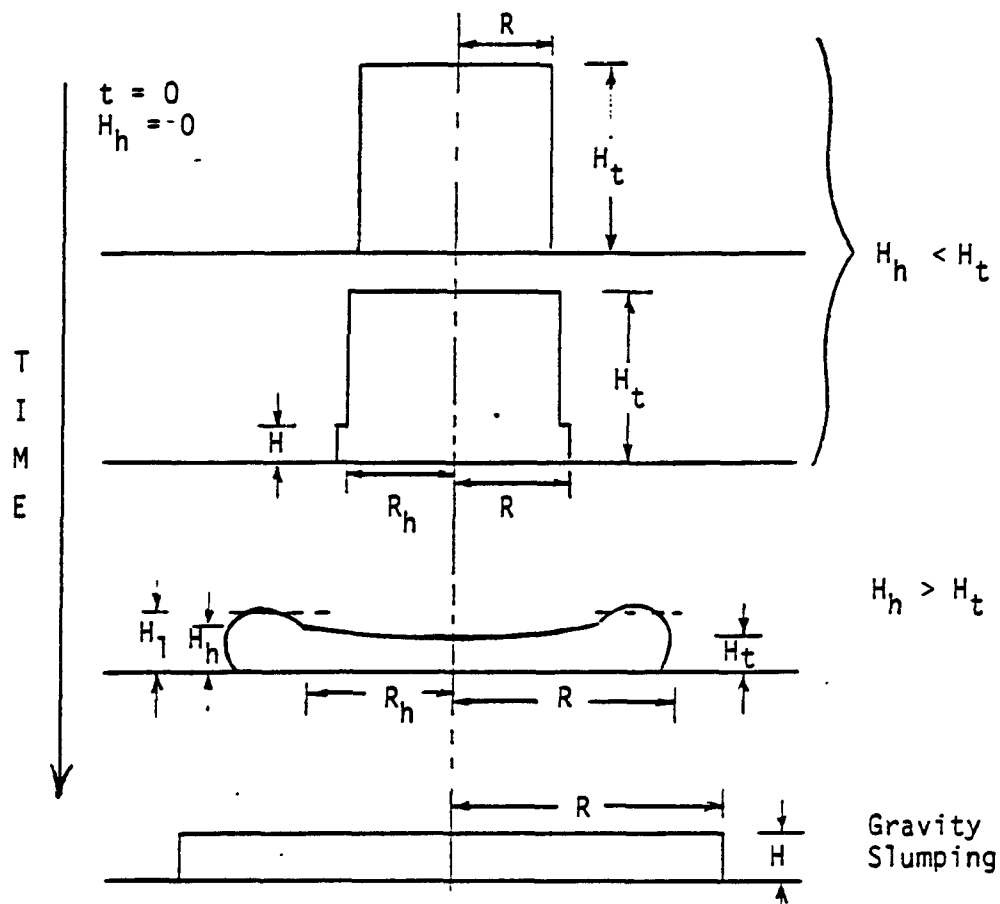


Figure V.2. Schematic Diagram of a Radially Spreading Cloud.

The terms in the momentum balance are evaluated differently for the time periods before a gravity current head has developed ($H_h < H_t$) and after the head has developed but the cloud is still accelerating (Figure V.2). Model equations describing the time period after the gravity current head forms ($H_h \geq H_t$) are derived first. The model equations describing the earlier time period ($H_h < H_t$) use simplifications of the equations applicable for $H_h \geq H_t$.

Unsteady Gravity Current ($H_h \leq H_t$)

When $H_h \geq H_t$ (Figures V.2, V.3), the frontal velocity is determined from the momentum balance (Equation (V.2)) as follows.

The static pressure force, obtained by integrating the static pressure over the boundary of the current, is

$$F_p = \left(\frac{1}{2} g \Delta \rho H_t \right) \left[2\pi R H_t \right] = \pi g \Delta \rho R H_t^2 \quad (V.4)$$

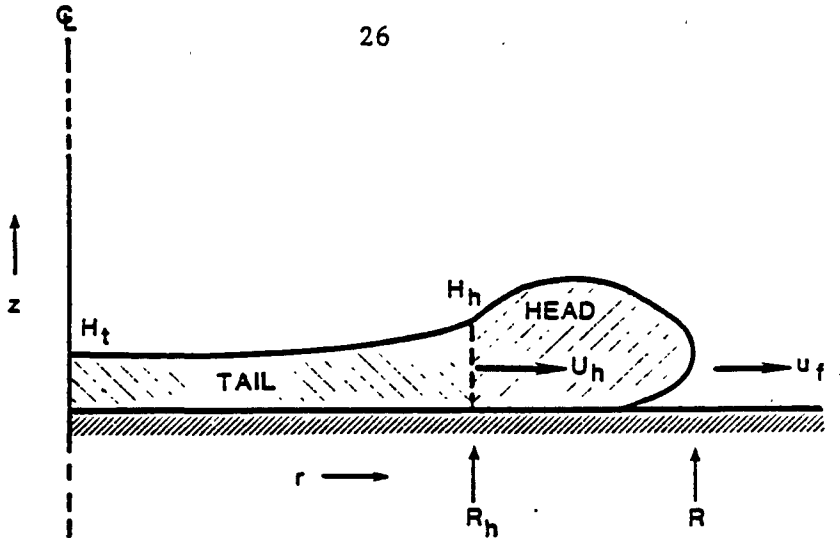


Figure V.3. The Unsteady Gravity Current (van Ulden, 1983).

Neglecting the shear stress at the bottom, the dynamic force on the current is the sum of the drag force on the head of the current and the lift force that arises due to asymmetry of the ambient flow around the head. The drag force is represented by

$$F_D = - \frac{d_v}{2} \rho_a u_f^2 \left(2\pi R_h a_v H_h \right) = - a_v d_v \pi R_h \rho_a u_f^2 \quad (V.5)$$

where d_v is an effective drag coefficient and the constant a_v is an empirical ratio of the average head depth H_1 to H_h ($a_v = H_1/H_h$).

The horizontal acceleration reaction ($-dP_v/dt$) is approximated by the reaction to an accelerating elliptical cylinder with an aspect ratio H/R (Batchelor, 1967):

$$- \left[\frac{dP_v}{dt} \right] = - \frac{d}{dt} \left(k_1 \rho_a \pi R H^2 u_f \right) \quad (V.6)$$

and the vertical acceleration reaction is represented as

$$- \left[\frac{dP_v}{dt} \right]_z = - \frac{d}{dt} \left(k_2 \rho_a \pi R H^2 u_f \right) \quad (V.7)$$

where k_1 and k_2 are coefficients of order one. Using a single constant, Equations (V.6) and (V.7) give

$$- \frac{dP_v}{dt} = - e_v \pi \rho_a \frac{d(RH^2 u_f)}{dt} \quad (V.8)$$

Using Equations (V.4), (V.5), and (V.8), the momentum balance (Equation (V.2)) becomes

$$\frac{dP}{dt} = \pi g \Delta \rho R H_t^2 - a_v d \frac{\pi \rho_a R H_h u_f^2}{v} - e_v \pi \rho_a \frac{d(RH^2 u_f)}{dt} \quad (V.9)$$

Following van Ulden (1979, 1983), it is assumed that the potential energy decrease due to slumping of the cloud is offset by the production of kinetic energy, which through the action of shear is partly transformed to turbulent kinetic energy. Part of the turbulent kinetic energy is transformed back into potential energy due to entrainment of air by the cloud. This "buoyant destruction" of kinetic energy is assumed to be proportional to the rate of production of turbulent kinetic energy, and following Simpson and Britter (1979) it is assumed that the turbulent kinetic energy production rate scales as $\pi \rho_a H R u_f^3$. Then,

$$\frac{1}{2} g \Delta \rho H \frac{dV}{dt} = \epsilon \pi \rho_a H R u_f^3 \quad (V.10)$$

which can be written

$$\frac{dV}{dt} = \frac{\epsilon (2\pi RH) u_f}{\left[\frac{g \Delta \rho H}{\rho_a u_f^2} \right]} = \frac{\epsilon (2\pi RH) u_f}{R i_f} \quad (V.11)$$

where ϵ is an empirical coefficient. Noting that dV/dt represents the air entrainment rate,

$$\frac{\dot{M}_a}{\rho_a} = \epsilon (2\pi RH) u_f \sqrt{\left[\frac{g \Delta \rho H}{\rho_a u_f^2} \right]} \quad (V.12)$$

where \dot{M}_a represents the air entrainment mass rate.

The volume integral

$$V = 2\pi \int_0^R h(r, t) r dr \quad (V.13)$$

where $h(r, t)$ is to be expressed in terms of H_h and H_t , and the momentum integral

$$P = 2\pi \int_0^R \rho u(r, t) h(r, t) r dr = P_t + P_h \quad (V.14)$$

are then approximated with separate analyses of the head and tail of the current.

In the tail of the current, the shallow layer approximation is applied. It is assumed that the shape of the current is quasi-stationary in time, and the layer-averaged density difference is assumed horizontally uniform. It follows that the volume and momentum of the tail are given by

$$v_t = \pi R_h^2 \left(H_t + H_h \right) / 2. \quad (V.15)$$

$$P_t = \frac{2}{5} \rho \left(\frac{2}{3} H_t + H_h \right) \pi R_h^3 \frac{u_f}{R} \quad (V.16)$$

A momentum balance for the head region, Figure V.4, assuming quasi-steady state, indicates that the static and dynamic pressure forces on the head should be balanced by the net flux of momentum due to flow into and out of the head. The static pressure and drag are, respectively

$$F_p = \left(\frac{1}{2} g \Delta \rho H_h \right) \left(2 \pi R_h H_h \right) = \pi g \Delta \rho R_h H_h^2 \quad (V.17)$$

and

$$\begin{aligned} F_D &= - d_v \left(\frac{1}{2} \rho_a u_f^2 \right) \left[2 \pi R_h (a_v H_h) \right] \\ &= - a_v d_v \rho_a u_f^2 \pi R_h H_h \end{aligned} \quad (V.18)$$

Near the surface, the inward flow (u_4 in Figure V.4) carries momentum into the head, while the return flow (u_3 in Figure V.4) carries momentum out of the head. Assuming $u_3 = u_4$, $H_4 = 1/2 H_h$, and $u_4 = \delta_v u_f$, the momentum flux into the head is approximately

$$Q_h = \delta_v^2 \rho_a u_f^2 \left[2 \pi R_h H_h \right] \quad (V.19)$$



Figure V.4. The Head of a Steady Gravity Current (Simpson and Britter, 1979; van Ulden, 1983).

Upon rearranging, the momentum balance on the head gives

$$\frac{\rho a_f^2}{g\Delta\rho H_h} = 1./ \left[a_v d_v - 2\delta_v^2 \right] - c_E^2 \quad (\text{V.20})$$

when $\delta_v = 0.2$ and $d_v = 0.64$; Equation (V.20) then specifies the head velocity boundary condition. The volume of the head is determined by assuming that the head length scales with H_1 . It follows that

$$R - R_h = b_v H_1 \quad (\text{V.21})$$

where b_v is an empirical constant, and the volume of the head becomes

$$V_h = \pi a_v^2 b_v (R + R_h) H_h^2 \quad (\text{V.22})$$

If the layer-averaged velocity is assumed to increase linearly with r , it follows that

$$u_h = u_f \left(\frac{R_h}{r} \right) \quad (\text{V.23})$$

and

$$P_h = \frac{2\pi}{3} \rho a_v \frac{u_f H_h}{R} \left[R^3 - R_h^3 \right] \quad (\text{V.24})$$

Along with the definition of u_f ,

$$\frac{dR}{dt} = u_f, \quad (\text{V.25})$$

Equations (V.9), (V.11), (V.20), (V.21), (V.23), and (V.25) are solved to determine ρ , H_t , H_h , V , P_h , and P_t when $H_h > H_t$.

The constants a_v , b_v , d_v , e_v , and c are assigned values 1.3, 1.2, 0.64, 20., and 0.59, respectively, based on analysis of the still-air denser-than-air gas release experiments of Havens and Spicer (1985).

Initial Gravity Current Development ($H_h < H_t$)

In order to model the initial cloud shape, the tail and head height are considered constant with respect to radius. The momentum balance on the cloud is then given by

$$\begin{aligned} \frac{d}{dt} \left[P_h + P_t \right] &= \pi g \Delta \rho \left[R_h H_t^2 + a_v b_v H_h^3 \right] \\ &\quad - \pi a_v d_v \rho_a R H_h u_f^2 - \frac{dP_v}{dt} \end{aligned} \quad (\text{V.26})$$

where the first term on the right-hand side represents the static pressure force on the head and the second term represents the drag force on the bottom surface of the cloud. The third force is the acceleration reaction by the ambient fluid, represented by Equation (V.8).

The dimensions of the head are again given by

$$R_h = R - a_v b_v H_h \quad (V.27)$$

and

$$H_h = \left(\frac{u_f}{C_E} \right)^2 [g\Delta\rho/\rho_a] \quad (V.28)$$

When the height of the tail H_t is assumed uniform with respect to radius, it follows that

$$H_t = \left[\frac{M}{\rho} - \pi a_v^2 b_v (R + R_h) H_h^2 \right]^{1/2} (\pi R_h^2) \quad (V.29)$$

where M is the total mass of the cloud. The momentum of the head P_h and tail P_t are then

$$P_h = \frac{2}{3} \pi a_v \frac{\rho H_h (R^3 - R_h^3)}{R} u_f \quad (V.30)$$

and

$$P_t = \frac{2}{3} \pi \frac{\rho H_t R_h^3}{R} u_f \quad (V.31)$$

Equations (V.26) through (V.31) determine the momentum of the blanket as a function of time, and thus the frontal velocity u_f . The cloud accelerates from rest because $H_h = 0$ initially.

Material and Energy Balances

The balance on the total mass of gas in the source blanket ($M = \pi R^2 H \rho$) is

$$\frac{dM}{dt} = \frac{d}{dt} \left[\pi R^2 H \rho \right] = \frac{E(t)}{w_{c,p}(t)} + \dot{M}_a + \dot{M}_{w,s} - \left[\frac{Q_{\max}}{w_c} \right] [\pi R^2] \quad (V.32)$$

where $E(t)$ is the contaminant evolution rate from the primary (liquid) source and $w_{c,p}(t)$ is the contaminant mass fraction in the primary source. For spills onto water, the water entrainment term ($\dot{M}_{w,s}$) is included in the source blanket description and is calculated from

Equation (V.46), and the (humid) air entrainment rate (Equation (V.12)) is

$$\dot{M}_a = 2\pi RH(\epsilon u_f) \rho_a \check{\left(g\Delta\rho H / (\rho_a u_f^2) \right)} \quad (V.33)$$

The balance on the mass of contaminant in the source blanket

$$(M_c = w_c \pi R^2 H \rho) \text{ is}$$

$$\frac{dM_c}{dt} = \frac{d}{dt} \left[w_c \pi R^2 H \rho \right] = E(t) - Q_{\max} (\pi R^2) \quad (V.34)$$

and the mass balance on the air in the source blanket

$$(M_a = w_a \pi R^2 H \rho) \text{ is}$$

$$\begin{aligned} \frac{dM_a}{dt} = \frac{d}{dt} \left[w_a \pi R^2 H \rho \right] &= \frac{E(t)}{w_{c,p}(t)} \left(\frac{1 - w_{c,p}(t)}{1 + H_a} \right) \\ &+ \frac{\dot{M}_a}{1 + H_a} - \left(\frac{Q_{\max}}{w_c} \right) w_a (\pi R^2) \end{aligned} \quad (V.35)$$

where the ambient humidity is H_a and the mass fraction of contaminant and air are $w_c = M_c/M$ and $w_a = M_a/M$, respectively. Note that any dilution of the primary source with air is assumed to be at ambient humidity.

The energy balance on the source blanket ($h\pi R^2 H \rho$) gives

$$\begin{aligned} \frac{d}{dt} \left[h\pi R^2 H \rho \right] &= \frac{h_p E(t)}{w_{c,p}(t)} + h_a \dot{M}_a + h_w \dot{M}_{w,s} \\ &- h \left(\frac{Q_{\max}}{w_c} \right) (\pi R^2) + \dot{Q}_s \end{aligned} \quad (V.36)$$

where h_p is the enthalpy of the primary source gas, h_a is the enthalpy of the ambient humid air, and h_w is the enthalpy of any water vapor entrained by the blanket (if on water). There are three alternate sub-models included for heat transfer (\dot{Q}_s) from the surface to the cloud.

The simplest method for calculating the heat transfer between the substrate and the gas cloud is to specify a constant heat transfer coefficient for the heat transfer relation

$$\dot{Q}_s = q_s \left[\pi \left(R^2 - R_p^2 \right) \right] = h_0 \Delta T \left[\pi \left(R^2 - R_p^2 \right) \right] \quad (V.37)$$

where \dot{Q}_s is the rate of heat transfer to the cloud, q_s is the heat flux, and ΔT is the temperature difference between the cloud and the surface. For the calculation of heat transfer over the source, the temperature difference is based on the average temperature of the blanket.

In the evaluation of the Burro and Coyote series of experiments, Koopman et al. (1981) proposed the following empirical heat transfer coefficient relationship for heat transfer between a cold LNG vapor cloud and the ground

$$h_0 = V_H \rho C_p \quad (V.38)$$

where the value of V_H was estimated to be 0.0125 m/s. (The constant V_H can be specified in this model.)

From the heat transfer coefficient descriptions for heat transfer from a flat plate, the following relationships can be applied. For natural convection, the heat transfer coefficient is estimated using the Nusselt (Nu), Grashoff (Gr), and Schmidt (Sc) numbers (McAdams, 1954) from

$$Nu = 0.14 (Gr Sc)^{1/3} \quad (V.39)$$

or

$$h_n = 0.14 \left[\frac{g \rho^2 C_p^3 \mu}{T Pr^2} \Delta T \right]^{1/3} \quad (V.40)$$

where h_n is the natural convection heat transfer coefficient and Pr is the Prandtl number. In order to simplify the calculations, the parameter group

$$\left[Pr^{-2} \left(C_p MW \right)^3 \left(\frac{\mu}{MW T} \right) \right]^{1/3} \quad (V.41)$$

is estimated to be 60 in mks units (actual values are 47.25, 58.5, and 73.4 for air, methane, and propane, respectively). Equation (V.40) becomes

$$h_n = 18 \left[\left(\frac{\rho}{MW} \right)^2 \Delta T \right]^{1/3} \quad (V.42)$$

where the density ρ , molecular weight MW , and temperature difference ΔT are based on the average composition of the gas blanket.

For forced convection, the Colburn analogy (Treybal, 1980) is applied to a flat plate using the Stanton number for heat transfer St_H and the Prandtl number as

$$St_H \Pr^{2/3} = \frac{c_f}{2} = \left(\frac{u_*}{\bar{u}} \right)^2 \quad (V.43)$$

or

$$h_f = (\bar{u} \rho C_p) \Pr^{-2/3} \left(\frac{u_*}{\bar{u}} \right)^2 \quad (V.44)$$

where h_f is the forced convection heat transfer coefficient. If the velocity is evaluated at $z = H/2$ and \Pr is specified 0.741,

$$h_f = \left[1.22 \frac{u_*^2}{u_0} \left(\frac{2z_0}{H} \right)^\alpha \right] \rho C_p \quad (V.45)$$

If $H/2 < z_R$, then the velocity is evaluated at $z = z_R$.

The overall heat transfer coefficient is then the maximum of the forced and natural convection coefficients, i.e. $h_0 = \max(h_f, h_n)$. The heat flux and transfer rate are then estimated by Equation (V.37).

If the gas blanket is formed over water, water may be transferred from the water surface to the blanket. The rate of mass transfer of water is modeled as

$$\dot{M}_{w,s} = \frac{F_0}{p} \left(p_{w,s}^* - p_{w,c} \right) \left[\pi \left(R^2 - R_p^2 \right) \right] \quad (V.46)$$

where F_0 is an overall mass transfer coefficient. The partial pressure driving force is the difference of the vapor pressure of water at the surface temperature $p_{w,s}^*$ and the partial pressure of water in the cloud, $p_{w,c}$. (The water partial pressure in the cloud is the minimum of: (a) the water mole fraction times the ambient pressure; or (b) the water vapor pressure at the cloud temperature ($p_{w,c}^*$).) The natural convection coefficient is based on the heat transfer coefficient and the analogy between the Sherwood number (Sh) and the Nusselt number (Nu) suggested by Bird et al. (1960)

$$Sh = Nu = 0.14 (Gr Sc)^{1/3} = \frac{F_n L}{D} \left(\frac{MW}{\rho} \right) \quad (V.47)$$

If the Schmidt number is taken as 0.6, and $\left(\frac{\mu}{T \cdot MW} \right)$ is estimated to be 2.2×10^{-9} in mks units,

$$F_n = 9.9 \times 10^{-3} \left[\left(\frac{\rho}{MW} \right)^2 \Delta T \right]^{1/3} \quad (V.48)$$

For forced convection, Treybal (1980) suggests that the Stanton number for mass transfer St_M and the Stanton number for heat transfer St_H are related by

$$St_M = St_H \left(\frac{Pr}{Sc} \right)^{2/3} = 1.15 St_H \quad (V.49)$$

or,

$$F_f = \frac{20.7 h_0}{\frac{MW C_p}{p}} \quad (V.50)$$

The overall mass transfer coefficient F_0 is the larger of the natural and forced convection coefficients.

For the case when the primary (liquid) source emission rate $E(t)$ is larger than the atmospheric takeup rate $Q_* \max \pi R_p^2$, Equations (V.32), (V.34), (V.35), and (V.36) are integrated for the mass, concentration, and enthalpy of the gas blanket along with an appropriate equation of state (i.e. relationship between enthalpy and temperature and between temperature and density).

When the emission rate is not sufficient to form a gas blanket, the flux of contaminant is not determined by the maximum atmospheric takeup rate. Consider the boundary layer formed by the emission of gas into the atmosphere above the primary source. If the source is modeled to have a uniform width $2b$ and entrain no air along the sides of the layer, the balance on the total material ($\rho_L u_L H_L$) in a differential slice of the layer is

$$\frac{d}{dx} \left[\rho_L u_L H_L \right] = \rho_a w_e + \left(\frac{Q_*}{w_c} \right)_p \quad (V.51)$$

where w_e is the vertical rate of air entrainment into the layer given by Equation (V.83), ρ_L is the average density of the slice, and $(Q_*/w_c)_p$ is the total flux of gas from the primary source. The balance on the mass flow rate of contaminant ($w_c \rho_L u_L H_L$) at any $(x - x_{up})$ is

$$c_{c,L} u_L H_L = Q_*(x - x_{up}) \quad (V.52)$$

With an equation of state to relate $c_{c,L}$ and ρ_L , Equation (V.51) is integrated from the upwind edge of the source ($x = x_{up}$) to the downwind edge ($x = L + x_{up}$).

In order to generate the initial conditions for the downwind dispersion calculations, the maximum concentration c_c and the vertical dispersion parameter S_z are needed. Since Equations (V.51) and (V.52) are written for a vertically averaged layer, consider the vertical average of the power law distribution. The height of the layer H_L is the height to some concentration level, say 10% of the maximum. Although strictly a function of α , this value is modeled as

$$H_L = \delta_L H_{EFF} \quad (V.53)$$

where H_{EFF} is the effective height defined by Equation (V.79) and $\delta_L = 2.15$. The vertically averaged concentration $c_{c,L}$ is defined by

$$c_{c,L} H_L = \int_0^{\infty} c dz \quad (V.54)$$

Similarly, the effective transport velocity u_L is defined by

$$c_{c,L} u_L H_L = \int_0^{\infty} c u_x dz \quad (V.55)$$

With Equation (V.53) and defining relations for H_{EFF} and u_{EFF} (Equations (V.79) and (V.93), respectively), it follows that

$$c_c = \delta_L c_{c,L} \quad (V.56)$$

$$u_L H_L = \delta_L \left(\frac{u_0 z_0}{1 + \alpha} \right) \left(\frac{s_z}{z_0} \right)^{1+\alpha} \quad (V.57)$$

and

$$\delta_L w'_e = w_e \quad (V.58)$$

where w'_e is given by Equation (V.83).

Maximum Atmospheric Takeup Rate

The maximum atmospheric takeup rate is determined as the largest takeup rate which satisfies Equations (V.51) and (V.52), and the maximum concentration of contaminant at the downwind edge of the source is the source contaminant concentration (c_c)_s. Combining Equations (V.51) and (V.52) and assuming adiabatic mixing of ideal gases with equal molal heat capacities (i.e. $(\rho - \rho_a)/c_c = \gamma = \text{constant}$), the maximum takeup flux is modeled as

$$Q_{max} = (c_c)_s \frac{k u_* (1 + \alpha)}{\hat{\phi}} \left(\frac{\delta_L}{\delta_L - 1} \right) \quad (V.59)$$

where

$$\hat{\phi} = \frac{1}{L} \int_0^L \frac{dx}{\phi(Ri_*)} \quad (V.60)$$

and $\phi(Ri_*)$ is defined (for $\rho \geq \rho_a$) by Equation (V.76).

An upper bound for the atmospheric takeup flux can be associated with the condition where the source begins to spread as a gravity intrusion against the approach flow. In water flume experiments, Britter (1980) measured the upstream and lateral extent of a steady-state plume from a circular source as a function of Ri_* . A significant upstream spread was obtained for $Ri_* > 32$, and lateral spreading at the center of the source was insignificant for $Ri_* < 8$. The presence of any significant lateral spreading represents a lower bound on the conditions of the maximum takeup flux.

The integral of Equation (V.60) is evaluated using a local Richardson number

$$Ri_*(x) = \xi(x - x_{up})^{\frac{1}{1+\alpha}} \quad (V.61)$$

where

$$\xi = g \left(\frac{\rho - \rho_a}{\rho_a} \right) \frac{z_0}{u_*^2} \frac{\Gamma\left(\frac{1}{1+\alpha}\right)}{1+\alpha} \left[\frac{ku_*(1+\alpha)}{\phi_c} \left(\frac{1+\alpha}{u_0 z_0} \right) \left(\frac{\delta_L}{\delta_{L-1}} \right) \right]^{\frac{1}{1+\alpha}} \quad (V.62)$$

and ϕ_c is 3.1 (corresponding to $Ri_* = 20$ ($8 < Ri_* < 32$)). Equation (V.60) then becomes

$$\hat{\phi} = \frac{1}{L} \int_0^L \frac{dx}{\frac{1.04}{0.88 + 0.099 \xi^{1.04} x}} \quad (V.63)$$

Equation (V.63) can be solved analytically (Gradshteyn and Ryzhik, 1980). With $\hat{\phi}$ determined, Q_{\max} can be determined with Equation (V.59).

Simulation of Transient Gas Releases

For simulating steady-state releases, the transient source calculation is carried out until the source characteristics no longer change (significantly) with time. The maximum centerline concentration c_c , the horizontal and vertical dispersion parameters S_y and S_z , the half width b , and if necessary, the enthalpy h are used as initial conditions for the downwind calculation specified in a transient spill.

Transient releases are modeled as a series of pseudo-steady-state releases. Consider a series of observers traveling with the wind over

the transient gas source described above; each observer originates from the point which corresponds with the maximum upwind extent of the gas blanket ($x = -R_{max}$). The observer velocity is u_{EFF} (Equation (V.93)), the average transport velocity of the gas. Since, in general, the value of u_{EFF} may differ from observer to observer, observer separation is not guaranteed. For a neutrally buoyant cloud, u_{EFF} is a function of downwind distance alone, and observer separation is guaranteed. Following Colenbrander (1980), the observer velocity is modeled as:

$$u_i(x) = \frac{u_0}{\Gamma\left(\frac{1}{1+\alpha}\right)} \left(\frac{S_{z_{0m}}}{z_0} \right)^\alpha \left[\frac{x + R_{max}}{\frac{\sqrt{\pi}}{2} R_m + R_{max}} \right]^{\alpha/(1+\alpha)} \quad (V.64)$$

where $S_{z_{0m}}$ is the value of S_{z_0} when the averaged source rate $(\pi R^2 Q_*)$ is a maximum, and the subscript i denotes observer i . Noting that $u_i(x) = dx_i/dt$, observer position and velocity are determined.

A pseudo-steady-state approximation of the transient source is determined as each observer passes over the source. If t_{up_i} and t_{dn_i} denote the times when observer i encounters the upwind and downwind edges of the source respectively, then the source fetch seen by observer i is:

$$L_i = x_{up_i} - x_{dn_i} \quad (V.65)$$

The width of the source $2B'_i(t)$ is defined by

$$B'^2_i(t) = R^2(t) - x_i^2(t) \quad (V.66)$$

Then the gas source area seen by observer i is

$$2L_i b_i = 2 \int_{t_{up_i}}^{t_{dn_i}} B'_i u_i dt \quad (V.67)$$

where $2b_i$ is the average width.

The takeup rate of contaminant $2(Q_* Lb)_i$ is calculated as

$$2(Q_* Lb)_i = 2 \int_{t_{up_i}}^{t_{dn_i}} Q_* B'_i u_i dt \quad (V.68)$$

The total mass flux rate from the source is

$$2(\rho_L u_L H_L b)_i = 2 \int_{t_{up_i}}^{t_{dn_i}} \left[\rho_a w'_e + \left(\frac{Q_*}{w_c} \right) \right] B'_i u_i dt \quad (V.69)$$

The average composition of the layer can now be determined at each $x - x_{up}$ over the source. The enthalpy of the layer is given by

$$2(h_L \rho_L u_L H_L b)_i = 2 \int_{t_{up_i}}^{t_{dn_i}} h \left(\frac{Q_*}{w_c} \right) B'_i u_i dt \quad (V.70)$$

(due to the choice of the reference temperature as the ambient temperature). With a suitable equation of state relating enthalpy, temperature, and density, the source can be averaged for each observer. After the average composition of the layer is determined at the downwind edge, an adiabatic mixing calculation is performed for gas of the layer average concentration and the ambient air. This calculation determines the function between density and concentration relationship for the remainder of the simulation if the calculation is adiabatic; it represents the adiabatic mixing condition if heat transfer is included in the downwind simulation.

For each of several observers released successively from $x = -R_{max}$, the observed dimensions L and b, the downwind edge of the source x_{dn} , the average vertical dispersion coefficient S_z , the average takeup flux Q_* , the centerline concentration c_c , and if applicable, the average enthalpy h_L is determined. A steady-state calculation is then made for each observer. The distribution parameters for any specified time t_s are determined by locating the position of the series of observers at time t_s , i.e. $x_i(t_s)$. The corresponding concentration distribution is then computed from the assumed profiles.

Steady-State Downwind Dispersion

The model treats dispersion of gas entrained into the wind field from an idealized, rectangularly shaped source of width 2b and length L. The circular source cloud is represented as an equivalent area rectangle ($\pi R^2 = 2bL$) with equivalent fetch ($L = 2R$). Similarity forms for the concentration profiles represent the plume as a horizontally homogeneous center section with Gaussian concentration profile edges:

$$c(x, y, z) = c_c(x) \exp \left[- \left(\frac{|y| - b(x)}{S_y(x)} \right)^2 - \left(\frac{z}{S_z(x)} \right)^{1+\alpha} \right]$$

for $|y| > b$

$$= c_c(x) \exp \left[- \left(\frac{z}{S_z(x)} \right)^{1+\alpha} \right] \quad \text{for } |y| \leq b \quad (V.71)$$

A power law wind velocity profile is assumed

$$u_x = u_0 \left(\frac{z}{z_0} \right)^\alpha \quad (V.72)$$

where the value of α is determined by a weighted least-squares fit of the logarithmic profile

$$u_x = \frac{u_*}{k} \left[\ln \left(\frac{z + z_R}{z_R} \right) - \Psi \left(\frac{z}{\lambda} \right) \right] \quad (V.73)$$

Functional forms for Ψ and typical values of α are given in Table V.1 for different Pasquill stability categories. With these profiles, the parameters of Equation (V.71) are constrained by ordinary differential equations.

TABLE V.1
TYPICAL ATMOSPHERIC BOUNDARY LAYER STABILITY
AND WIND PROFILE CORRELATIONS

Pasquill Stability Category	Monin-Obukhov Length (λ) as a Function of Surface Roughness $z_R(m)^1$	Typical Power Law Exponents α in Eqn. (V.72)	Corrections to Logarithmic Profiles as Given by Businger et al. (1971) Ψ in Eqn. (V.73)
A	$-11.4 z_R^{0.10}$	0.108	$\Psi = 2 \ln \left[\frac{1+a}{2} \right] + \ln \left[\frac{1+a^2}{2} \right]$
B	$-26.0 z_R^{0.17}$	0.112	$-2 \tan^{-1}(a) + \pi/2$ with
C	$-123 z_R^{0.30}$	0.120	$a = (1 - 15(z/\lambda))^{1/4}$
D	∞	0.142	$\Psi = 0$
E	$123 z_R^{0.30}$	0.203	$\Psi = -4.7 (z/\lambda)$
F	$26.0 z_R^{0.17}$	0.253	

¹Curve fit of data from Pasquill (1974).

Vertical Dispersion

The vertical dispersion parameter S_z is determined by requiring that it satisfy the diffusion equation

$$u_x \frac{\partial c}{\partial x} = \frac{\partial}{\partial z} K_z \frac{\partial c}{\partial z} \quad (\text{V.74})$$

with the vertical turbulent diffusivity given by

$$K_z = \frac{k u_* z}{\phi(Ri_*)} \quad (\text{V.75})$$

The function $\phi(Ri_*)$ is a curve fit of laboratory data for vertical mixing in stably density-stratified fluid flows ($Ri_* > 0$) reported by Kantha et al. (1977), Lofquist (1960), and McQuaid (1976). For $Ri_* < 0$, the function $\phi(Ri_*)$ was taken from Colenbrander and Puttock (1983) and modified so the passive limit of the two functions agree as follows:

$$\begin{aligned}\phi(Ri_*) &= 0.88 + 0.099 Ri_*^{1.04} & Ri_* \geq 0 \\ &- 0.88/(1 + 0.65 |Ri_*|^{0.6}) & Ri_* < 0\end{aligned}\quad (\text{V.76})$$

The friction velocity is calculated using Equation (V.73) from a known velocity u_0 at a specific height z_0 . Combining the assumed similarity forms for concentration and velocity, Equations (V.71), (V.72), (V.74), and (V.75) give

$$\frac{d}{dx} \left[\left(\frac{u_0 z_0}{1 + \alpha} \right) \left(\frac{s_z}{z_0} \right)^{1+\alpha} \right] = \frac{k u_* (1 + \alpha)}{\phi(Ri_*)} \quad (\text{V.77})$$

where the Richardson number Ri_* is computed as

$$Ri_* = g \left(\frac{\rho - \rho_a}{\rho_a} \right) \frac{H_{\text{EFF}}}{u_*^2} \quad (\text{V.78})$$

and the effective cloud depth is defined as

$$H_{\text{EFF}} = \frac{1}{c_c} \int_0^\infty c dz = \Gamma \left(\frac{1}{1 + \alpha} \right) \frac{s_z}{1 + \alpha} \quad (\text{V.79})$$

Equation (V.77) can be viewed as a volumetric balance on a differential slice downwind of the source. A mass balance over the same slice gives,

$$\frac{d}{dx} \left(\rho_L u_L H_L \right) = \rho_a w_e \quad (\text{V.80})$$

which is Equation (V.51) without the source term. With Equations (V.57) and (V.58), this becomes

$$\frac{d}{dx} \left(\rho_L u_{\text{EFF}} H_{\text{EFF}} \right) = \rho_a w'_e \quad (\text{V.81})$$

Assuming adiabatic mixing of ideal gases with equal constant molal heat capacities (i.e. $(\rho - \rho_a)/c_c = \text{constant}$) and using the contaminant mass balance, the mass balance becomes

$$\frac{d}{dx} \left(u_{\text{EFF}} H_{\text{EFF}} \right) = w'_e \quad (\text{V.82})$$

which leads to

$$w'_e = \frac{w_e}{\delta_L} = \frac{k u_* (1 + \alpha)}{\phi(Ri_*)} \quad (\text{V.83})$$

Equations (V.81) and (V.83) are combined to give

$$\frac{d}{dx} \left(\rho_L u_{\text{EFF}} H_{\text{EFF}} \right) = \frac{\rho_a k u_* (1 + \alpha)}{\phi(Ri_*)} \quad (\text{V.84})$$

Generalizing, Equation (V.84) is assumed to apply when $(\rho_a - \rho_c)/c$ is not constant.

When heat transfer from the surface is present, vertical mixing will be enhanced by the convection turbulence due to heat transfer. Zeman and Tennekes (1977) model the resulting vertical turbulent velocity as

$$\frac{w}{u_*} = \left[1 + \frac{1}{4} \left(\frac{w_*}{u_*} \right)^2 \right]^{1/2} \quad (\text{V.85})$$

where w_* is the convective scale velocity described as

$$\left(\frac{w_*}{u_*} \right)^2 = \left[\frac{gh}{u_* \bar{u}} \frac{\left(T_s - T_{c,L} \right)}{T_{c,L}} \right]^{2/3} \quad (\text{V.86})$$

If \bar{u} is evaluated at H_{EFF} ,

$$\frac{w}{u_*} = \left(1 + \frac{1}{4} \text{Ri}_T^{2/3} \right)^{1/2} \quad (\text{V.87})$$

where

$$\text{Ri}_T = g \left(\frac{T_s - T_{c,L}}{T_{c,L}} \right) \frac{H_{\text{EFF}}}{u_* u_0} \left(\frac{z_0}{H_{\text{EFF}}} \right)^\alpha \quad (\text{V.88})$$

and $T_{c,L}$ is the temperature obtained from the energy balance of Equations (V.103) and (V.104). Equation (V.84) is modified to account for this enhanced mixing by

$$\frac{d}{dx} \left(\rho_L u_{\text{EFF}}^H H_{\text{EFF}} \right) = \frac{\rho_a k w (1 + \alpha)}{\phi(Ri'_*)} \quad (\text{V.89})$$

where $Ri'_* = Ri_* \left(\frac{u_*}{w} \right)^2$.

Although derived for two-dimensional dispersion, Equation (V.89) is extended for application to a denser-than-air gas plume which spreads laterally as a density intrusion:

$$\frac{d}{dx} \left(\rho_L u_{\text{EFF}}^H H_{\text{EFF}} B_{\text{EFF}} \right) = \frac{\rho_a k w (1 + \alpha)}{\phi(Ri'_*)} B_{\text{EFF}} \quad (\text{V.90})$$

where the plume effective half width is defined by

$$B_{\text{EFF}} = b + \frac{\sqrt{\pi}}{2} S_y \quad (\text{V.91})$$

and determined using the gravity intrusion relation

$$\frac{dB_{\text{EFF}}}{dt} = C_E \left[g \left(\frac{\rho - \rho_a}{\rho_a} \right) H_{\text{EFF}} \right]^{1/2} \quad (\text{V.92})$$

The average transport velocity in the plume is defined by

$$u_{\text{EFF}} = \frac{\int_0^\infty c u_x dz}{\int_0^\infty cdz} = u_0 \left(\frac{S_z}{z_0} \right)^\alpha \Gamma \left(\frac{1}{1 + \alpha} \right) \quad (\text{V.93})$$

and the lateral spread of the cloud is modeled as

$$\frac{dB_{\text{EFF}}}{dx} = \frac{1}{u_{\text{EFF}}} \frac{dB_{\text{EFF}}}{dt}$$

$$= C_E \left[\frac{g z_0 \Gamma^3 \left(\frac{1}{1+\alpha} \right)}{u_0^2 (1 + \alpha)} \right]^{1/2} \left[\frac{\rho - \rho_a}{\rho_a} \right]^{1/2} \left[\frac{s_z}{z_0} \right]^{\left(\frac{1}{2} - \alpha \right)} \quad (V.94)$$

Horizontal Dispersion

The crosswind similarity parameter $S_y(x)$ is also determined by requiring that it satisfy the diffusion equation

$$u_x \frac{\partial c}{\partial x} = \frac{\partial}{\partial y} \left[K_y \frac{\partial c}{\partial y} \right] \quad (V.95)$$

with the horizontal turbulent diffusivity given by

$$K_y = K_0 u_x B_{EFF}^{\gamma_1} \quad (V.96)$$

For a passive plume ($b = 0$), $S_y = \sqrt{2} \sigma_{y,a}$ where $\sigma_{y,a}$ is the similarity parameter correlated by Pasquill (1974) in the form $\sigma_{y,a} = \delta_y x^{\beta_y}$ where δ_y and β_y are functions of the Pasquill stability category and the averaging time. Furthermore, Equations (V.95) and (V.96) require that

$$\sigma_{y,a} \frac{d\sigma_{y,a}}{dx} = K_0 B_{EFF}^{\gamma_1} \quad (V.97)$$

where $\gamma_1 = 2 - 1/\beta_y$ and $K_0 = \frac{2\beta_y}{\pi} (\delta_y \sqrt{\pi/2})^{1/\beta_y}$. Then,

$$S_y \frac{dS_y}{dx} = \frac{4\beta_y}{\pi} B_{EFF}^2 \left[\frac{\delta_y \sqrt{\pi/2}}{B_{EFF}} \right]^{1/\beta_y} \quad (V.98)$$

where Equation (V.98) is also assumed applicable for determining S_y when b is not zero.

At the downwind distance x_t where $b = 0$, the crosswind concentration profile is assumed Gaussian with S_y given by

$$S_y = \sqrt{2} \delta_y (x_t + x_v)^{\beta_y} \quad (V.99)$$

where x_v is a virtual source distance determined as

$$S_y(x_t) = \sqrt{2} \delta_y (x_t + x_v)^{\beta_y} \quad (V.100)$$

The gravity spreading calculation is terminated for $x > x_t$.

Averaging Time

The observed dispersion of a gas plume is a function of the observation (averaging) time. Herein we have assumed:

- The most important influence of averaging time on observed plume properties is associated with plume meander.
- For ground-level releases, vertical plume meander is much smaller than horizontal plume meander.

Gifford (1960) showed that the ratio of maximum measured concentrations based on different averaging times (t_a and t_p) was given by

$$\frac{c_c(t_p)}{c_c(t_a)} = \left(\frac{t_a}{t_p} \right)^{0.2}$$

for t_a/t_p up to about 200, when source and receptor were at the same level. Spicer and Havens (1988) proposed that

$$\frac{\sigma_{y,a}(t_a)}{\sigma_{y,a}(t_{a,i})} \approx \left(\frac{t_a}{t_{a,i}} \right)^{0.2}$$

where $t_{a,i}$ was suggested to be an "instantaneous" averaging time representing the smallest time scale associated with plume meander. (Practically, $t_{a,i}$ represents a nonzero averaging time associated with the puff coefficient $\sigma_{y,a}(t_{a,i})$.) Based on correlations from Gifford (1976) and Slade (1968), Spicer and Havens (1988) presented the following dispersion coefficient correlations for stabilities A through F:

$$\sigma_{y,a}(x; t_a) = 0.423 \left(\frac{t_a^*}{600 \text{ s}} \right)^{0.2} x^{0.9} \quad \text{A} \quad (\text{V.101a})$$

$$\sigma_{y,a}(x; t_a) = 0.313 \left(\frac{t_a^*}{600 \text{ s}} \right)^{0.2} x^{0.9} \quad \text{B} \quad (\text{V.101b})$$

$$\sigma_{y,a}(x; t_a) = 0.210 \left(\frac{t_a^*}{600 \text{ s}} \right)^{0.2} x^{0.9} \quad \text{C} \quad (\text{V.101c})$$

$$\sigma_{y,a}(x; t_a) = 0.136 \left(\frac{t_a^*}{600 \text{ s}} \right)^{0.2} x^{0.9} \quad \text{D} \quad (\text{V.101d})$$

$$\sigma_{y,a}(x; t_a) = 0.102 \left(\frac{t_a^*}{600 \text{ s}} \right)^{0.2} x^{0.9} \quad \text{E} \quad (\text{V.101e})$$

$$\sigma_{y,a}(x; t_a) = 0.0674 \left(\frac{t_a^*}{600 \text{ s}} \right)^{0.2} x^{0.9} \quad F \quad (\text{V.101f})$$

where $t_a^* = \max(t_a, t_{a,i})$ and t_a is the desired averaging time. Values of $t_{a,i}$ are shown in Table V.2.

Table V.2. Values of $t_{a,i}$ for Use in Eq. (V.101)

Pasquill Stability	$t_{a,i}(\text{s})$
A	18.4
B	18.4
C	18.4
D	18.3
E	11.4
F	4.6

Values for δ_y and β_y are then determined by Equation (V.101). (For example, for an averaging time of 60 s for F stability, $t_a^* = 60 \text{ s}$, $\delta_y = 0.0425$, and $\beta_y = 0.9$.)

(Thermal) Energy Balance

For some simulations of cryogenic gas releases, heat transfer to the plume in the downwind dispersion calculation may be important, particularly in low wind conditions. The source calculation determines a gas/air mixture initial condition for the downwind dispersion problem. Air entrained into the plume is assumed to mix adiabatically. Heat transfer to the plume downwind of the source adds additional heat. This added heat per unit mass D_h is determined by an energy balance on a uniform cross-section as

$$\frac{d}{dx} \left[D_h \rho_L u_{EFF}^H B_{EFF}^H \right] = q_s / \delta_L \quad (\text{V.102})$$

where q_s is determined by Equation (V.37) along with the desired method of calculating h_0 . Equation (V.102) is applied when $b = 0$ and is generalized to

$$\frac{d}{dx} \left[D_h \rho_L u_{EFF}^H B_{EFF}^H B_{EFF}^B \right] = q_s B_{EFF} / \delta_L \quad (\text{V.103})$$

when $b > 0$. Since the average density of the layer ρ_L cannot be determined until the temperature (i.e. D_h) is known, a trial and error procedure is required.

Closure

For a steady plume, the centerline concentration c_c is determined from the material balance

$$E = \int_0^{\infty} \int_{-0}^{\infty} c u_x dy dz = 2c_c \left(\frac{u_0 z_0}{1 + \alpha} \right) \left(\frac{s_z}{z_0} \right)^{1+\alpha} B_{EFF} \quad (V.104)$$

where E is the plume source strength.

Equations (V.76), (V.78), (V.79), (V.85)-(V.91), (V.93), (V.94), (V.98)-(V.100), and (V.102)-(V.104) are combined with an equation of state relating cloud density to gas concentration and temperature and are solved simultaneously to predict s_z , s_y , c_c , and b as functions of downwind distance beginning at the downwind edge of the gas source.

Correction for Along-Wind Dispersion

Following Colenbrander (1980), an adjustment to c_c is applied to account for dispersion parallel to the wind direction. The calculated centerline concentration $c_c(x)$ is considered to have resulted from the release of successive planar puffs of gas ($c_c(x)\Delta x$) without any dispersion in the x -direction. If it is assumed that each puff diffuses in the x -direction as the puff moves downwind independently of any other puff and that the dispersion is one-dimensional and Gaussian, the x -direction concentration dependence is given by

$$c'_c(x; x_{p_i}) = \frac{c_c(x_{p_i})\Delta x_i}{\sqrt{2\pi} \sigma_{x,a}} \exp \left[-\frac{1}{2} \left(\frac{x - x_{p_i}}{\sigma_{x,a}} \right)^2 \right] \quad (V.105)$$

where x_{p_i} denotes the position of the puff center due to observer i .

After Beals (1971), the x -direction dispersion coefficient $\sigma_{x,a}$ is assumed to be a function of distance from the downwind edge of the gas source ($X = x - x_0$) and atmospheric stability given by

$$\begin{aligned} \sigma_{x,a}(X) &= 0.02 X^{1.22} && \text{unstable, } x \geq 130 \text{ m} \\ &= 0.04 X^{1.14} && \text{neutral, } x \geq 100 \text{ m} \\ &= 0.17 X^{0.97} && \text{stable, } x \geq 50 \text{ m} \end{aligned} \quad (V.106)$$

where ($X = x - x_0$) and $\sigma_{x,a}$ are in meters. The concentration at x is then determined by superposition, i.e., the contribution to c_c at a given x from neighboring puffs is added to give an x -direction corrected value of c'_c . For N observers,

$$c'_c(x) = \sum_{i=1}^N \frac{c_c(x_{p_i})}{\sqrt{2\pi} \sigma_{x,a}} \exp \left[-\frac{1}{2} \left(\frac{x - x_{p_i}}{\sigma_{x,a}} \right)^2 \right] \Delta x_i \quad (V.107)$$

and for large N,

$$c'_c(x) = \frac{1}{\sqrt{2\pi}} \int_0^\infty \frac{c_c(\xi)}{\sigma_{x,a}(\xi - \xi_0)} \exp \left[-\frac{1}{2} \left[\frac{x - \xi}{\sigma_{x,a}(\xi - \xi_0)} \right]^2 \right] d\xi \quad (V.108)$$

The corrected centerline concentration c'_c is used in the assumed profiles in place of c_c , along with the distribution parameters S_y , S_z , and b .

VI. CONCLUSIONS AND RECOMMENDATIONS

DEGADIS was designed to model the atmospheric dispersion of ground-level, area-source dense gas (or aerosol) clouds released with zero (initial) momentum into an atmospheric boundary layer over flat, level terrain. DEGADIS describes the dispersion processes which accompany the ensuing gravity-driven flow and entrainment of the gas into the atmospheric boundary layer.

A jet-plume model has been interfaced with DEGADIS to provide for prediction of the trajectory and dilution of vertically oriented gas or aerosol jets. When the jet plume returns to ground level, DEGADIS predicts the ensuing ground-level plume dispersion. The jet-plume model provides for

- automatic adjustment of integration step-size (using the Runge-Kutta-Gill method as in DEGADIS),
- elliptical plume shape (cross-section), with air entrainment specified consistent with the Pasquill-Gifford plume dispersion coefficient representation of atmospheric turbulent entrainment,
- application to scenarios where the plume returns to ground level or remains aloft, and
- ground reflection when the plume (lower) boundary reaches ground level.

It is recommended that the following limitations and cautions be observed when using the jet-plume and DEGADIS models.

- The jet-plume model presently provides only for vertically oriented releases.
- Both models assume an unobstructed atmospheric flow field. The jet-plume model assumes a logarithmic wind profile, and the DEGADIS model assumes a power law wind profile which is consistent with the logarithmic wind profile. Application of the models should be limited to releases where the depth of the dispersing layer is much greater than the height of the surface roughness elements.
- Averaging time can be user-specified. There are three important time scales for the typical dispersion application: the averaging time input to the model (t_{av}), the time scale assumed for specification of the concentration/hazard relation (t_{haz}), and the time required for the contaminant material to be taken up by

the atmosphere (t_{rel}). The averaging time t_{av} influences the rate of lateral spread (dispersion) due to atmospheric turbulence (δ_y). For an elevated release which does not return to the ground, t_{rel} is the duration of the release. For a transient ground-level release or for an elevated release which returns to the ground, t_{rel} is the duration of the secondary source blanket in DEGADIS. For a steady-state release (large t_{rel}), t_{av} should be set to t_{haz} . (For example, if t_{haz} is based on toxicological data for one hour exposure, then $t_{haz} = t_{av} = 3600$ s.) For a transient release, t_{av} should be set to the minimum of t_{haz} or t_{rel} . If $t_{haz} \leq t_{rel}$, $t_{av} = t_{haz}$. If $t_{rel} < t_{haz}$, $t_{av} = t_{rel}$, and the concentration time history at a given position should be averaged over t_{haz} to determine the hazard.

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APPENDIX A
MODEL APPLICATION ON VAX/VMS

DEGADIS Version 2.1 (including the jet/plume front-end model) was developed under VAX/VMS V3.5 and VAX-11 Fortran V3.5; there should be no installation difficulty for VAX/VMS V4.0 or later.

Installation

The directory which contains the Fortran source code for DEGADIS must be equivalenced with the logical name SYS\$DEGADIS:. If the full directory specification is DQA0:[DEGADIS_V21], issue the VAX/VMS command:

```
$ ASSIGN DQA0:[DEGADIS_V21] SYS$DEGADIS:  
with either the /PROCESS, /GROUP, or /SYSTEM qualifier (/SYSTEM is recommended). Once this assignment is made, the files must be compiled and linked to form DEGADISIN, DEGADIS1, DEGADIS2, DEGADIS3, DEGADIS4, and SDEGADIS2; the VMS command procedure BUILD.COM performs these tasks. The process which compiles and links DEGADIS must have READ, WRITE, and EXECUTE access privileges to SYS$DEGADIS while only READ and EXECUTE access privileges are needed to execute the existing models.
```

Considerations for Installation other than VAX/VMS

There are two types of problems which may occur when attempting to install DEGADIS on a different computer or operating system. The first is due to the use of non-standard ANSI Fortran 77 language elements. The second is due to the use of external VAX/VMS routines in DEGADIS.

The following VAX-11 Fortran extensions have been used in DEGADIS:

- (*) In-line comments--An exclamation mark (!) is used to include comments at the end of a valid statement.
- (*) Special characters--The underscore (_) is used in variable names.
- (*) DO loops--DO loops are used with the structure:

```
DO v = e1,e2[,e3]
```

```
.
```

```
.
```

```
.
```

```
END DO
```

where v is a variable name and e1, e2, and e3 are numeric expressions. The numeric expressions have the standard Fortran 77 meaning.

- (*) INCLUDE statements--INCLUDE statements simply allow other source files to be inserted in the routine being compiled at this point in the source. The system table '(\$SSDEF)' is used to check the status or returning system routines.
- .
- (*) OPEN keyword NAME--The OPEN keyword NAME specifies the file name to be opened.
- (*) Fortran descriptors--The Q descriptor obtains the integer number of characters remaining in the input record during a READ operation. The dollar sign (\$) descriptor suppresses the carriage return at the end of a line on output.
- (*) Continuation lines--Continuation lines have been expressed by using either a nonblank character in column 6 or by beginning the line with a tab and a number in the next column.
- (*) Concatenation of character strings--Character strings are concatenated using two slashes (//).

The following VAX/VMS subroutines have been used in DEGADIS:

- (*) SECNDS
TIME = SECNDS(TIME0)
SECNDS returns to TIME the difference between the number of seconds after midnight on the system clock and the value of TIME0.
- (*) LIB\$DATE_TIME
ISTAT = LIB\$DATE_TIME (STRING)
LIB\$DATE_TIME returns a 24-character ASCII string with the system date and time. ISTAT is an integer variable which accepts the return status.
- (*) LIB\$DO_COMMAND
ISTAT = LIB\$DO_COMMAND (STRING)
LIB\$DO_COMMAND issues the command STRING (a character string) to VAX/VMS. If the command is not issued, ISTAT contains the failure code. If the command is issued, the calling process is terminated.

Model Implementation

The (combined) jet/plume (JETPLU) and DEGADIS models consist of nine separate programs:

- JETPLU_IN is the file-based input module which defines the simulation.
- JETPLU describes the trajectory and dilution of the vertically oriented jet release.
- DEGBRIDGE is an interface program which takes the output from JETPLU (if where the jet/plume returns to ground level) and provides appropriate input to DEGADIS.

- DEGADISIN is the interactive input module which defines DEGADIS-only simulation.
- DEGADIS1 determines α and describes the gas source for transient and steady-state releases.
- DEGADIS2 describes the pseudosteady-state downwind dispersion of the released gas.
- DEGADIS3 sorts the results of DEGADIS2 for a transient release at given times.
- DEGADIS4 sorts the results of DEGADIS2 for a transient release at given positions.
- SDEGADIS2 describes the steady-state downwind dispersion of the released gas.

Steady-state releases are simulated by executing JETPLU, DEGBRIDGE, DEGADIS1, and SDEGADIS2; time-limited (transient) releases are simulated by executing JETPLU, DEGBRIDGE, DEGADIS1, DEGADIS2, and DEGADIS3.

The JETPLU DEGADIS model uses three types of input information

- VAX/VMS command procedure for execution
- simulation definition
- numerical parameters.

The VAX/VMS command procedure used to execute JETPLU DEGADIS is generated by JETPLU_IN which also reads the simulation definition. DEGADIS can be run without JETPLU (for ground-level, zero momentum releases). Input to DEGADIS is made to run interactively using DEGADISIN. Example input sessions are included in Appendix B. The numerical parameters (convergence criteria, initial increments, etc.) are supplied to DEGADIS through a series of input files. Although these numerical parameters are easily changed, the user should need to change these only rarely (with the exception of the time-sort parameters which are explained later in this appendix).

VAX/VMS Command Procedures

The VAX/VMS command procedure generated by JETPLU_IN controls the execution of programs for the simulation. Program execution follows one of two paths: (a) a transient (time-limited) release or (b) a steady-state release. JETPLU_IN, which automatically generates the appropriate command procedure, requires a simulation name be specified. The simulation name must be a valid VAX/VMS file name without a file extension and is designated RUN_NAME. JETPLU and DEGADIS will use this file name with standard extensions for input, interprocess communication, and output. Figures A.1 and A.2 show example VAX/VMS command procedures created by JETPLU_IN for the run names TEST_S and TEST for steady-state and transient releases, respectively. The directory which contains the executable images has been assigned the system logical name SYS\$DEGADIS. The COPY/LOG command copies a file

from the first argument to the second argument, and the RUN command executes the specified image. These steps may also be carried out by issuing the commands at a terminal.

```
$ assign test_s.ino for001
$ assign test_s.out for003
$ assign test_s.ind for002
$ run sys$degadis:jetplu
$ deassign for001
$ deassign for002
$ deassign for003
$ run sys$degadis:degbridge
test_s
$ copy/log SYS$DEGADIS:example.er1 test_s.er1
$ copy/log SYS$DEGADIS:example.er2 test_s.er2
$ run SYS$DEGADIS:DEGADIS1
test_s
$ run SYS$DEGADIS:SDEGADIS2
test_s
$ copy/log test_s.out + test_s.scl + test_s.sr3 -
  test_s.lis
```

Figure A.1. Example DEGADIS Command Procedure on VAX/VMS for a Steady-State Simulation Named TEST_S.

```

$ assign test.ino for001
$ assign test.out for003
$ assign test.ind for002
$ run sys$degadis:jetplu
$ deassign for001
$ deassign for002
$ deassign for003
$ run sys$degadis:degbridge
test
$ copy/log SYS$DEGADIS:example.erl test.erl
$ copy/log SYS$DEGADIS:example.er2 test.er2
$ copy/log SYS$DEGADIS:example.er3 test.er3
$ run SYS$DEGADIS:DEGADIS1
test
$ run SYS$DEGADIS:DEGADIS2
test
$ run SYS$DEGADIS:DEGADIS3
TEST
$ copy/log test.out + test.scl + test.sr3 -
test.lis

```

Figure A.2. Example DEGADIS Command Procedure on VAX/VMS for a Transient Simulation Named Test.

Simulation Definition

JETPLU_IN specifies information about the ambient wind field, the properties of the released material, and details of the release.

The input uses free-formatted files; a value for each parameter must be specified in the input file even if it has no meaning for the simulation at hand. This form of input allows the user to ignore spacing in the input file. Numbers on a particular line can be separated by commas, spaces, or tabs. Comments may also be included at the end of any line in the input file as long as all required values are given before any comments are included. Lines which are only comments are allowed only after all required values have been specified. (Sample input files are included in Appendix B.)

The ambient wind field is characterized by a known velocity u_0 at a given height z_0 , a surface roughness z_R , and the Pasquill stability class or Monin-Obukhov length. The Pasquill stability class is required to estimate values of the lateral, vertical, and along-wind similarity coefficients. If specified, the Monin-Obukhov length λ is required to calculate the friction velocity u_* . In addition to these specifications, the ambient temperature, pressure, and humidity must be specified.

The contaminant properties are used by the model to fix the mixture density from the contaminant concentration. In the model, there are three possible ways to simulate the mixture density:

- Case 1: Treat the contaminant as an ideal gas and allow for condensation of ambient humidity. This is accomplished by setting $\langle NDEN \rangle = 0$. In DEGADIS, ground-to-cloud heat transfer can either be included ($\langle INDHT \rangle = 1$) or precluded ($\langle INDHT \rangle = 0$). The contaminant heat capacity is determined by $\langle CPK \rangle$ and $\langle CPP \rangle$ using

$$C_{p_c}(T) = (MW_c)^{-1} \left[3.33 \times 10^4 + q_1 \left[\frac{\frac{p_1}{T} - \frac{p_1}{T_0}}{\frac{T}{T_0} - 1} \right] \right] \quad (A.1)$$

where $\langle CPK \rangle$ is denoted by q_1 and $\langle CPP \rangle$ is denoted by p_1 . If a constant heat capacity is desired, set $\langle CPP \rangle$ to 0. and $\langle CPK \rangle$ to the desired heat capacity (in J/kg K).

- Case 2: Treat the contaminant as an ideal gas and do not allow for condensation of ambient humidity. In this case, ground-to-cloud heat transfer in DEGADIS is not allowed, and $(\rho - \rho_a)/c$ is assumed constant. This is accomplished by setting $\langle NDEN \rangle = -1$. Values input for $\langle INDHT \rangle$, $\langle CPK \rangle$, and $\langle CPP \rangle$ are ignored.
- Case 3: Estimate the contaminant/air mixture density as a function of contaminant concentration based on a user-supplied estimate. Ordered triples of the contaminant mole fraction, the contaminant concentration (kg/m^3), and the contaminant/air mixture density (kg/m^3) are input to the model. This is accomplished by setting $\langle NDEN \rangle$ equal to the number of triples. In the input file, the ordered triples (starting with pure air) are input on the next $NDEN$ lines immediately following the line for $\langle NDEN \rangle$. Values input for $\langle INDHT \rangle$, $\langle CPK \rangle$, $\langle CPP \rangle$, and $\langle TEMJET \rangle$ are ignored.

The user must choose to simulate the release as time-limited (transient) or steady-state (by the specification of $\langle TEND \rangle$), and the jet elevation and diameter must be specified.

Figure A.3 summarizes the structure of the input file RUN_NAME.IN required by JETPLU_IN. The description following is written for each of the variables in Figure A.3. After each line of input, an explanation is included. All units are SI (meters, kilograms, second) except as specified.

```

<TITLE1>
<TITLE2>
<TITLE3>
<TITLE4>
<U0> <Z0>
<ZR>
<INDVEL> <ISTAB> <RML>
<TAMB> <PAMB> <RELHUM>
<TSURF>
<GASNAM>
<GASMW>
<AVTIME>
<TEMJET>
<GASUL> <GASLL> <ZLL>
<INDHT> <CPK> <CPP>
<NDEN>
<ERATE>
<ELEJET> <DIAJET>
<TEND>
<DISTMX>

```

Figure A.3. RUN_NAME.IN Structure Required by JETPLU_IN

```

<TITLE1>
<TITLE2>
<TITLE3>
<TITLE4>
    <TITLE1>, <TITLE2>, <TITLE3>, and <TITLE4> are four lines of
    up to 80 characters each of a title for this simulation.

<U0> <Z0>
    <U0> (m/s) is the ambient wind velocity at <Z0> (m).

<ZR>
    <ZR> is the surface roughness (m).

<INDVEL> <ISTAB> <RML>
    <INDVEL> is an indicator which determines the method of
    calculation for the ambient velocity profile in the
    jet/plume model as follows:

    For <INDVEL>=1, the Pasquill-Gifford stability category (in
    <ISTAB> using 1 for A, 2 for B, etc.) is used along with

```

<ZR> to determine the Monin-Obukhov length <RML>; the log velocity profile is then determined using <RML>.

For <INDVEL>=2, the Monin-Obukhov length <RML> is supplied by the user; the log velocity profile is then determined using <RML>. Note that <ISTAB> must still be specified.

<TAMB> <PAMB> <RELHUM>

<TAMB>, <PAMB>, and <RELHUM> are the ambient temperature (K), the ambient pressure (atm or N/m²), and the relative humidity (%), respectively.

<TSURF>

<TSURF> is the surface temperature (K); if <TSURF> < 250 K, <TSURF> is set to <TAMB>.

<GASNAM>

<GASNAM> is a three-letter designation for the contaminant name. Any character string of three letters or less is valid; this is for user run identification and does not access property data.

<GASMW>

<GASMW> is the contaminant molecular weight (kg/kmole).

<AVTIME>

<AVTIME> is the averaging time (s). This parameter is used to estimate the value of <DELTAY>.

<TEMJET>

<TEMJET> is the temperature of the released contaminant (K).

<GASUL> <GASLL> <ZLL>

<GASUL> and <GASLL> are the concentrations to be used for estimating contours for an upper and lower concentration level in DEGADIS. The calculations are made for the elevation <ZLL>. Note that the JETPLU/DEGADIS computations will be carried out to <GASLL>/2.

<INDHT> <CPK> <CPP>

<INDHT> is used to include heat transfer in the DEGADIS computations. Heat transfer is not included for <INDHT>=0. For <INDHT>=1, heat transfer is included, and the heat transfer coefficient is calculated by DEGADIS. <CPP> and <CPK> are used to calculate the heat capacity as a function of temperature according to the correlation included in DEGADIS. If a constant heat capacity is desired, set <CPP> to 0. and <CPK> to the desired heat capacity (J/kg K).

<NDEN>

<NDEN> is used to specify the contaminant density profile. There are three cases for <NDEN>:

<NDEN> = -1; The simulation treats the contaminant as if it were an ideal gas with a molal heat capacity equal to that of air. Water condensation effects are ignored (equivalent to an "isothermal" release in DEGADIS).

<NDEN> = 0; The simulation treats the contaminant as if it were an ideal gas with the heat capacity indicated by <CPK> and <CPP>. Water condensation effects are taken into account as appropriate (equivalent to a nonisothermal release in DEGADIS).

<NDEN> > 0; <NDEN> specifies the number of triples which follow in the next <NDEN> lines. The triples are used to specify the contaminant concentration as a function of density based on adiabatic mixing with ambient air. The ordered triples represent (in order):

- (1) the contaminant mole fraction
- (2) the contaminant concentration (kg contam/m³ mix)
- (3) the mixture density (kg mixture/m³ mixture).

The ordered triples must go from pure air to pure contaminant (equivalent to an "isothermal" release in DEGADIS).

<ERATE>

<ERATE> is the mass evolution (release) rate (kg/s).

<ELEJET> <DIAJET>

<ELEJET> is the initial jet elevation (m), and <DIAJET> is the initial jet diameter (m).

<TEND>

<TEND> is the duration of the primary release (s). For steady-state releases, set <TEND> to 0.; to run the jet/plume model only, set <TEND> to a negative number.

<DISTMX>

<DISTMX> is the maximum distance between output points in the JETPLU output (m).

DEGBRIDGE takes the output from the jet/plume model and creates the file necessary for DEGADIS to complete the simulation.

Input Module--DEGADISIN

DEGADISIN is the optional interactive input module which defines a DEGADIS simulation; DEGADISIN is composed of two subroutines (Figure A.4):

(*) DEGADISIN contains the program overhead and generates the command file RUN_NAME.COM which can be used to control simulation execution (D-38).

(*) IOT contains the interactive question-and-answer sequence which defines the simulation; IOT also creates the file RUN_NAME.INP (D-76).

An example of a DEGADISIN query sequence is included in Appendix B. As this information is gathered, it is written to the file RUN_NAME.INP (Figure A.5). Once DEGADISIN is completed, RUN_NAME.INP may be edited to correct minor input mistakes. If major revisions are necessary, the recommended practice is to execute DEGADISIN again.

Once the information required by DEGADISIN has been entered properly, DEGADIS may be executed using the command procedure generated by DEGADISIN under the file name RUN_NAME.COM. If DEGADIS is not to be run using this command file, the user must enter the simulation name (RUN_NAME) after each of the programs are begun. (In this case, the user must also provide copies of the numerical parameter files.)



Figure A.4. DEGADISIN Flowchart.

```

TITLE(1)
TITLE(2)
TITLE(3)
TITLE(4)
U0, Z0, ZR
ISTAB
OODIST, AVTIME
DELTA, BETA, RML
SIGX_COEFF, SIGX_POW, SIGX_MIN_DIST
TAMB, PAMB, HUMID
ISOFL, TSURF
IHTFL, HTCO
IWTFL, WTCO
GAS_NAME
GAS_MW, GAS_TEMP, GAS_RHOE
GAS_CPK, GAS_CPP
GAS_UFL, GAS_LFL, GAS_ZSP
NP
If <ISOFL>0) then {  

  (for  

  external density  

  calculations)  

  .  

  .  

  .  

  . DEN((J,1),J=1,5)  

  . DEN((J,2),J=1,5)  

  .  

  .  

  . DEN((J,NP),J=1,5)  

  . CCLOW  

  . GMASSO  

  . NT  

  . PTIME(1),ET(1),R1T(1),PWC(1),PTEMP(1),PFRACV(1)  

  . PTIME(2),ET(2),R1T(2),PWC(2),PTEMP(2),PFRACV(2)  

  .  

  .  

  . PTIME(NT),ET(NT),R1T(NT),PWC(NT),PTEMP(NT),PFRACV(NT)  

  . CHECK1, CHECK2, AGAIN, CHECK3, CHECK4, CHECK5  

  . TINP  

  for steady-state  

  only      ESS, SRCLEN, SRCB
}
NT of these {  

}

```

Figure A.5. Structure for Free-Formatted RUN_NAME.INP File.

Source Module--DEGADIS1

DEGADIS1 estimates values for the friction velocity and ambient wind profile power α and characterizes the primary gas source for the remainder of the model; DEGADIS1 is composed of the following subroutines (Figure A.6):

(*) AFGEN	linearly interpolates between a pair of points based on a list of supplied values (D-3).
-----------	--

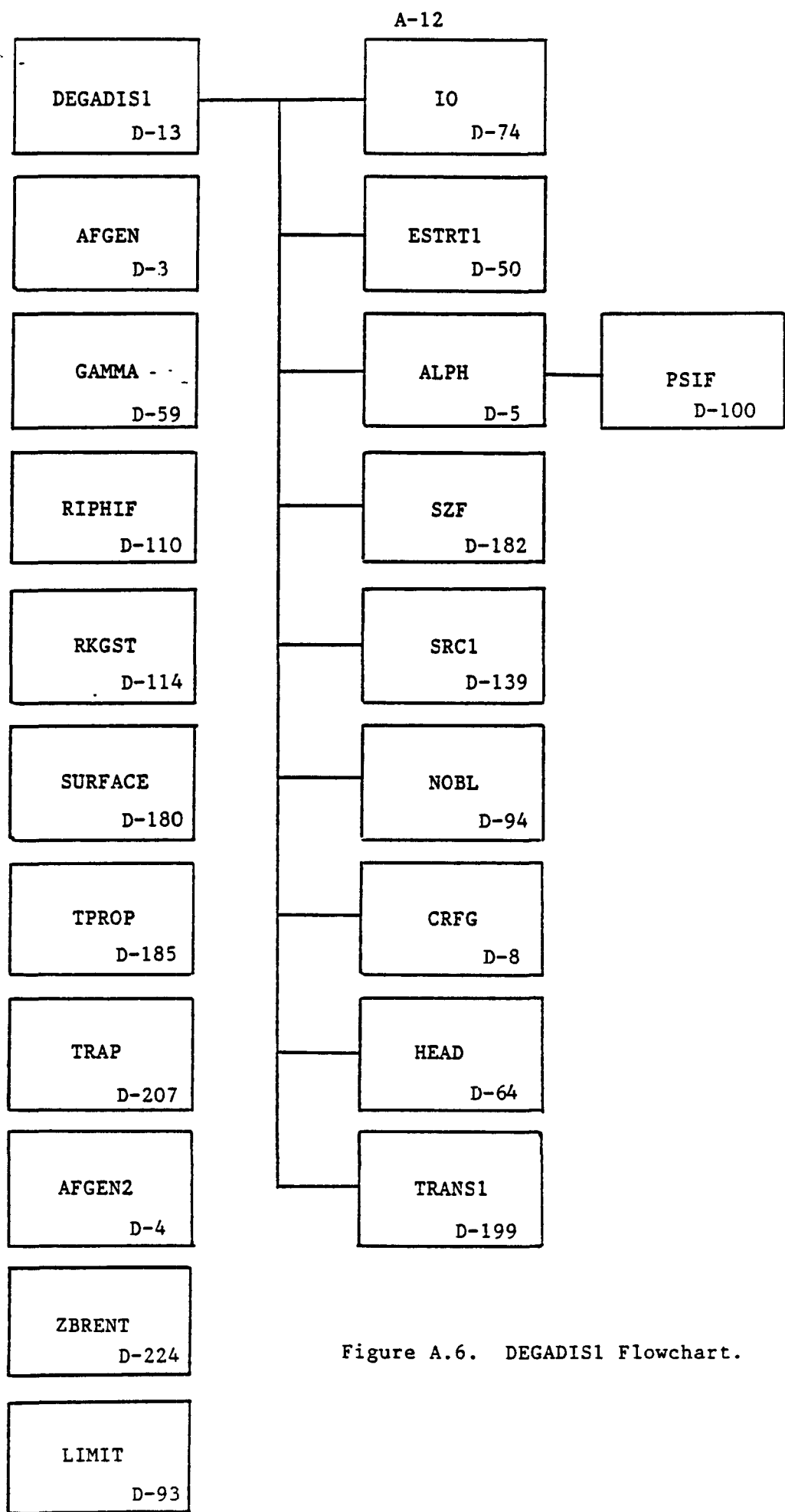


Figure A.6. DEGADIS1 Flowchart.

- (*) AFGEN2 linearly interpolates between a pair of points based on a list of supplied values (D-4).
- (*) ALPH estimates the ambient wind profile power α by minimizing the integral of the difference between an ambient logarithmic velocity profile and the assumed power law velocity profile (D-5).
- (*) CRFG creates a table of calculated values which will describe the secondary gas source for the downwind dispersion calculations (D-8).
- (*) DEGADIS1 contains the program overhead and sequentially calls the routines required to estimate the ambient wind profile power α and to characterize the primary gas source (D-14).
- (*) ESTRT1 recovers the numerical parameters contained in the file RUN_NAME.ER1 (D-50).
- (*) GAMMA calculates the gamma function of the argument (I.e. $\Gamma(x)$) (D-59).
- (*) HEAD writes a formatted output heading to the file RUN_NAME.SCL (D-64).
- (*) IO recovers the simulation definition contained in RUN_NAME.INP (D-74).
- (*) LIMIT establishes the limit for ZBRENT (D-93).
- (*) NOBL estimates gas source behavior when no gas blanket is present (D-94).
- (*) PSIF calculates the ψ function in the logarithmic velocity profile (D-100).
- (*) RIPHIF calculates the Richardson number and the values of $\phi(R_{i*})$ and $\hat{\phi}(R_{i*})$ (D-110).
- (*) RKGST performs numerical integration of a specified system of equations using a variable-step, modified fourth-order Runge-Kutta method (D-114).
- (*) SCR1 contains the ordinary differential equations which describe the gas blanket formed as a result of the primary gas source (D-139).

- (*) SURFACE estimates heat and water transfer rates across the bottom surface of the gas layer (D-180).
- (*) SZF estimates the value of S_z if the primary source can just form a gas blanket over the source (D-182).
- (*) TPROP is a series of utility routines which estimate the thermodynamic properties of a given gas mixture (D-185).
- (*) TRANS1 writes the information to continue the next simulation step to the file RUN_NAME.TR2 (D-199).
- (*) TRAP is a utility included for program diagnostics (D-207).
- (*) ZBRENT finds the bracketed root of an equation using Brent's method (D-224).

As input, DEGADIS1 requires two files:

- (*) RUN_NAME.ER1 contains various numerical parameters. For most simulations, a copy of the SYS\$DEGADIS:EXAMPLE.ER1 file will be adequate. A copy of SYS\$DEGADIS:EXAMPLE.ER1 is included in Figure A.7.
- (*) RUN_NAME.INP contains the simulation definition as discussed in Appendix B (Figure A.5).

As output, DEGADIS1 generates the following files:

- (*) RUN_NAME.SCD contains the calculated values which describe the secondary gas source. It is generated by SRC1 and NOBL and is then read by CRFG; it is a temporary file.
- (*) RUN_NAME.SCL is the listed output which describes the input information for the simulation and the calculated secondary gas source. It is written by HEAD and CRFG.
- (*) RUN_NAME.TR2 contains the information to continue the next simulation step.

```

!This is an example of how to set up and use the run parameter
! input files. Comment lines start with an exclamation mark(!)
! in the first column. The only restrictions for data input are
! as follows:
!      1) The data must be entered in the same order
!          all of the time.
!      2) Only the number must be between columns 10 and 20.
!      3) Always include the decimal point in the number
!
! Column layout:
!23456789012345678901234567890
!-----1-----2-----3
!      I           I
STPIN    0.01      MAIN - RKGST - INITIAL STEP SIZE
ERBND    0.0025    MAIN - RKGST - ERROR BOUND
STPMX    5.12      MAIN - RKGST - MAXIMUM STEP SIZE
WTRG     1.         MAIN - RKGST - WEIGHT FOR RG
WTTM     1.         MAIN - RKGST - WEIGHT FOR Total Mass
WTYA     1.         MAIN - RKGST - WEIGHT FOR ya
WTYC     1.         MAIN - RKGST - WEIGHT FOR yc
WTEB     1.         MAIN - RKGST - WEIGHT FOR Energy Balance
WTmB     1.         MAIN - RKGST - WEIGHT FOR Momentum Balance
WTuh     1.         MAIN - RKGST - WEIGHT FOR Ueff*Heff
XLI      0.05      ALPH - LOWER LIMIT OF SEARCH FOR ALPHA
XRI      0.50      ALPH - UPPER LIMIT OF SEARCH FOR ALPHA
EPS      0.001     ALPH - ERROR BOUND USED BY "RTMI"
ZLOW     0.01      ALPHI - maximum BOTTOM HEIGHT FOR FIT OF ALPHA
!
STPINZ   -0.02     ALPHI - INITIAL RKGST STEP <0.
!
ERBNDZ   0.005     ALPHI - ERROR BOUND FOR RKGST
!
STPMXZ   -2.00     ALPHI - MAXIMUM STEP FOR RKGST <0.
!
! Note that comment lines can be mixed with the numbers.
!
SRCOER   0.007     SRC10 - OUTPUT Error criterion
SRCSS    5.2        SRC10 - min time for Steady; STPMX
SRCcut   .00001    SRC10 - min height for blanket
htcut    .0         SRC1 - min height for blanket heat transfer
ERNOBL   1.0005    NOBL - CONVERGENCE ratio
NOBLPT   100.       NOBL - NUMBER OF POINTS
!
!                         USED ON THE LAST PORTION OF THE SOURCE
!
crfger   0.008     error criterion in building GEN3 vectors
!
```

Figure A.7. SYS\$DEGADIS:EXAMPLE.ERL Listing.

```

epsilon    0.59      epsilon USED IN AIR ENTRAINMENT SPECIFICATION
!
! /SPRD_CON/
!
ce        1.15      constant in gravity slumping equation
delrhomin 0.0       stop cloud spread if delrho<delrhomin
!
!
! /SZFC/
!
szstp0   0.01      SZF - Initial step size
szerr    0.001     SZF - Error criterion
szstpmx   5.0       SZF - Maximum step size [-] m
szsz0    0.01      SZF - Initial Value of dellay*Ueff*Heff
!
!
! /ALPHcom/
!
ialpf1    1.         ALPHI - calculation flag; 0) alpha=alpco; 1)1/(1+z); 2)1
alpco    0.2         ALPHI - Value for alpha if IALPFL = 0
!
!
! /PHIcom/
!
iphifl    3.         PHIF - calc flag
dellay   2.15       Ratio of H1/Heff
!
!
!
! /VUcom/
!
vua      1.3         Constant Av in source model
vub      1.2         Constant Bv in source model
vuc      20.0        Constant Ev in source model
vud      .64         Constant Dv in source model
vudelta  0.20       Constant DELTA v in source model
!
! End-of-File

```

Figure A.7. (concluded)

Pseudosteady-State Module--DEGADIS2

DEGADIS2 performs the downwind dispersion portion of the calculation for each of several observers released successively over the transient source described by DEGADIS1. (Note that the routines INCGAMMA, GAMMA, and SERIES are linked in DEGADIS2 but never are called in DEGADIS2.) DEGADIS2 is composed of the following subroutines (Figure A.8):

- (*) AFGEN linearly interpolates between a pair of points based on a list of supplied values (D-3).
- (*) AFGEN2 linearly interpolates between a pair of points based on a list of supplied values (D-4).
- (*) DEGADIS2 contains the program overhead and sequentially calls the routines to recover the information generated in DEGADIS1, recover the numerical parameter file RUN_NAME.ER2, and perform the simulation (D-23).
- (*) ESTRT2 recovers the numerical parameters contained in the file RUN_NAME.ER2, particularly the number of observers NOBS (D-54).
- (*) LIMIT establishes the limit for ZBRENT (D-93).
- (*) OB contains the ordinary differential equations which average the gas source for each observer (D-97).
- (*) PSS contains the ordinary differential equations which describe the portion of the downwind dispersion calculation when $b > 0$ (D-100).
- (*) PSSOUT governs the output of calculated points to the file RUN_NAME.PSD when PSS is active (D-104).
- (*) RIPHIF is a series of utilities which calculates the Richardson number and the value of $\phi(R_i)$ (D-110).
- (*) RKGST performs numerical integration of a specified system of equations using a modified fourth-order Runge-Kutta method (D-114).
- (*) SSG contains the ordinary differential equations which describe the portion of the downwind dispersion calculation when $b = 0$ (D-152).
- (*) SSGOUT governs the output of calculated points to the file RUN_NAME.PSD when SSG is active (D-155).

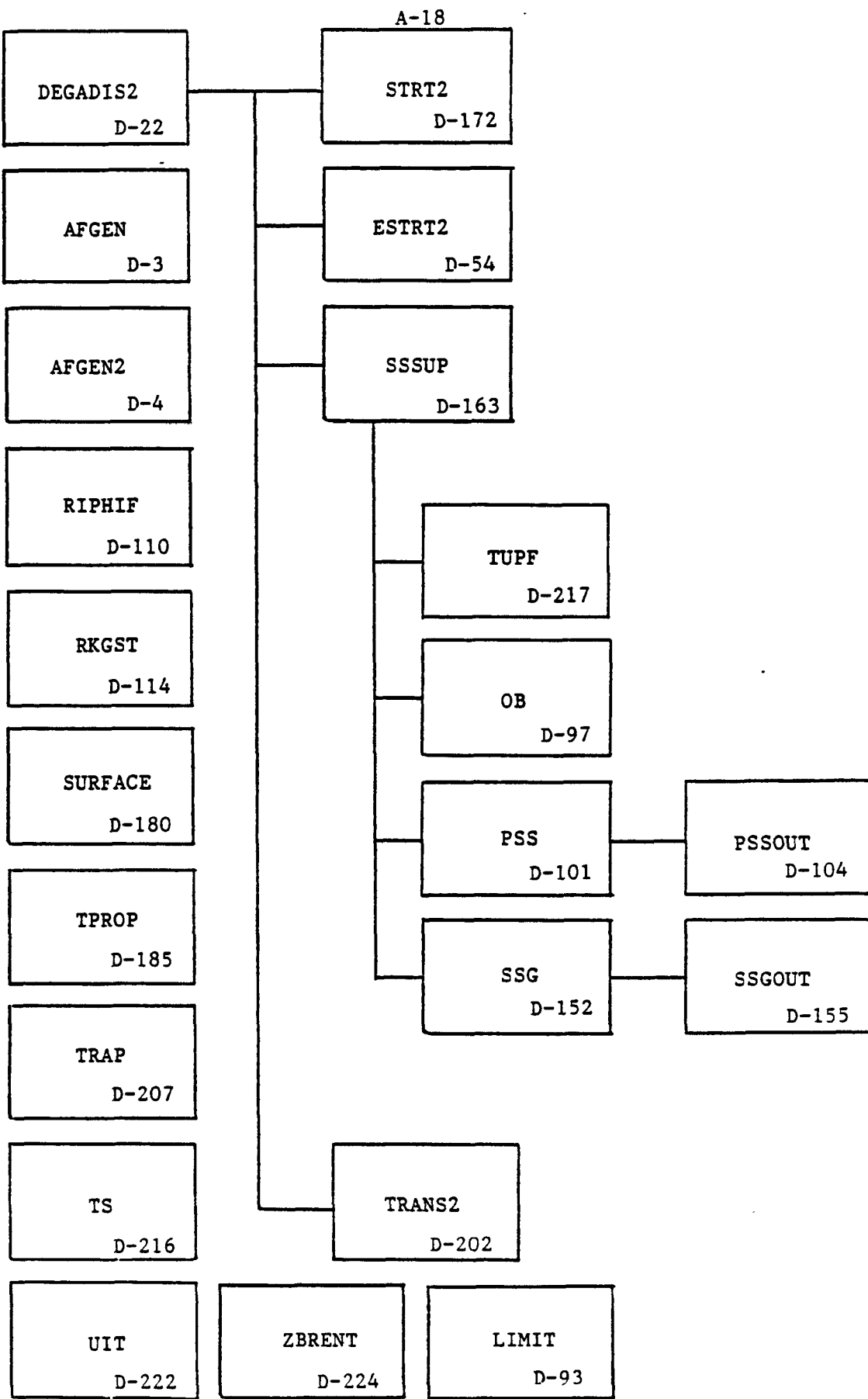


Figure A.8. DEGADIS2 Flowchart.

- (*) SSSUP is a supervisor routine which controls the averaging of the source for each observer, the portion of the downwind dispersion calculation when $b > 0$, and the portion of the downwind dispersion calculation when $b = 0$ (D-163).
- (*) STRT2 recovers the information generated in DEGADIS1 contained in the file RUN_NAME.TR2 (D-172).
- (*) SURFACE estimates heat and water transfer rates across the bottom surface of the gas layer (D-180).
- (*) TPROP is a series of utility routines which estimates the thermodynamic properties of a given gas mixture (D-185).
- (*) TRANS2 writes the information necessary for DEGADIS3 to the file (RUN_NAME.TR3) (D-202).
- (*) TRAP is a utility included for program diagnostics (D-207).
- (*) TS calculates the time when a given observer will be at a given downwind distance (D-216).
- (*) TUPF contains the routines which determine the intersection of the upwind/downwind edge of the secondary gas source with a given observer (D-217).
- (*) UIT is a series of routines to calculate observer position and velocity as a function of time (D-222).
- (*) ZBRENT finds the bracketed root of an equation using Brent's method (D-224).

As input, DEGADIS2 requires two files:

- (*) RUN_NAME.ER2 contains the various numerical parameters, particularly the number of observers NOBS. For most simulations, a copy of the SYS\$DEGADIS:EXAMPLE.ER2 file will be adequate. A copy of SYS\$DEGADIS:EXAMPLE.ER2 is included in Figure A.9.
- (*) RUN_NAME.TR2 contains the basic simulation definition as well as calculated secondary source parameters.

DEGADIS2 generates the following output files:

- (*) RUN_NAME.OBS contains a summary of the source parameters for each observer.

```

! This is an example for an "ER2" run parameter file.
! The same rules apply as for the "ER1" files.
!
!23456789012345678901234567890
!-----1-----2-----3
!
! These values are in common area /ERROR/
!
SYOER      0.0      SSSUP - RKGST - INITIAL SY
ERRO       0.005    SSSUP - RKGST(OBS) - ERROR BOUND
SZOER      0.01     SSSUP - RKGST(OBS) - INITIAL SZ
WTAIO       1.0      SSSUP - RKGST(OBS) - WEIGHT FOR AI
WTQOO      1.0      SSSUP - RKGST(OBS) - WEIGHT FOR Q
WTSZO      1.0      SSSUP - RKGST(OBS) - WEIGHT FOR SZ
ERRP       0.003    SSSUP - RKGST(PSS) - ERROR BOUND
SMXP        10.     SSSUP - RKGST(PSS) - MAXIMUM STEP
WTSZP      1.0      SSSUP - RKGST(PSS) - WEIGHT FOR SZ
WTSYP      1.0      SSSUP - RKGST(PSS) - WEIGHT FOR SY
WTBEP      1.0      SSSUP - RKGST(PSS) - WEIGHT FOR BEFF
WTDH       1.0      SSSUP - RKGST(PSS) - WEIGHT FOR DH
ERRG       0.003    SSSUP - RKGST(SSG) - ERROR BOUND
SMXG        10.     SSSUP - RKGST(SSG) - MAXIMUM STEP SIZE
ERTDNF     0.0005   TDNF - CONVERGENCE CRITERION
ERTUPF     0.0005   TUPF - CONVERGENCE CRITERION
WTRUH       1.0      SSSUP - RKGST(SSG) - WEIGHT FOR RUH
WTDHG      1.0      SSSUP - RKGST(SSG) - WEIGHT FOR DH
!
! These values are in common area /STP/
!
STPO        0.05    SSSUP - RKGST(OBS) - INITIAL STEP
STPP        0.05    SSSUP - RKGST(PSS) - INITIAL STEP
ODLP        0.06    SSSUP - RKGST(PSS) - RELATIVE OUTPUT DELTA
ODLLP      80.      SSSUP-RKGST(PSS)-MAXIMUM DISTANCE BETWEEN
OUTPUTS(m)
STPG        0.05    SSSUP - RKGST(SSG) - INITIAL STEP
ODLG        0.045   SSSUP - RKGST(SSG) - RELATIVE OUTPUT DELTA
ODLLG      80.      SSSUP-RKGST(SSG)-MAXIMUM DISTANCE BETWEEN
OUTPUTS(m)
!
! The last variable NOBS is in /CNOBS/
!
! Note: it is read in as a real value even though it is integer type
!       in the program.
!
NOBS       30.
!
!
!
! End-of-File

```

Figure A.9. SYS\$DEGADIS:EXAMPLE.ER2 Listing.

- (*) RUN_NAME.PSD contains the calculated downwind dispersion parameters for each observer. DEGADIS3 and DEGADIS4 sort this information to determine the downwind concentration profiles as a function of position and time.
- (*) RUN_NAME.TR3 contains the simulation definition and the number of each record type written to RUN_NAME.PSD.

Time Sort Modules--DEGADIS3 and DEGADIS4

DEGADIS3 sorts the downwind dispersion calculation for each of several observers for concentration information at several given times; the along-wind dispersion correction is then applied as desired. DEGADIS3 uses the following subroutines (Figure A.10).

- (*) DEGADIS3 contains the program overhead and sequentially calls the routines to recover the information generated in DEGADIS2, recover the numerical parameter file RUN_NAME.ER3, sort and apply the along-wind dispersion correction to the results of DEGADIS2, and output the results (D-28).
- (*) ESTRT3 recovers the numerical parameters contained in the file RUN_NAME.ER3, particularly the time sort parameters (D-58).
- (*) GAMMA calculates the gamma function of the argument (i.e. $\Gamma(x)$) D-59).
- (*) GETTIM sets the default time sort parameters as needed (D-61).
- (*) INCGAMMA calculates the incomplete gamma function of the two arguments (D-70).
- (*) LIMIT establishes the limit for ZBRENT (D-93).
- (*) SERIES evaluates a series needed to estimate the mass of gas above a given concentration level (D-130).
- (*) SORTS recovers the information in RUN_NAME.PSD and arranges the information according to the time sort parameters in the file RUN_NAME.ER3 (D-131).
- (*) SORTS1 applies the along-wind dispersion correction to the time-sorted information (D-134).
- (*) SRTOUT generates the formatted output file RUN_NAME.SR3 (D-148).
- (*) STRT3 recovers the information generated in DEGADIS2 contained in the file RUN_NAME.TR3 (D-178).

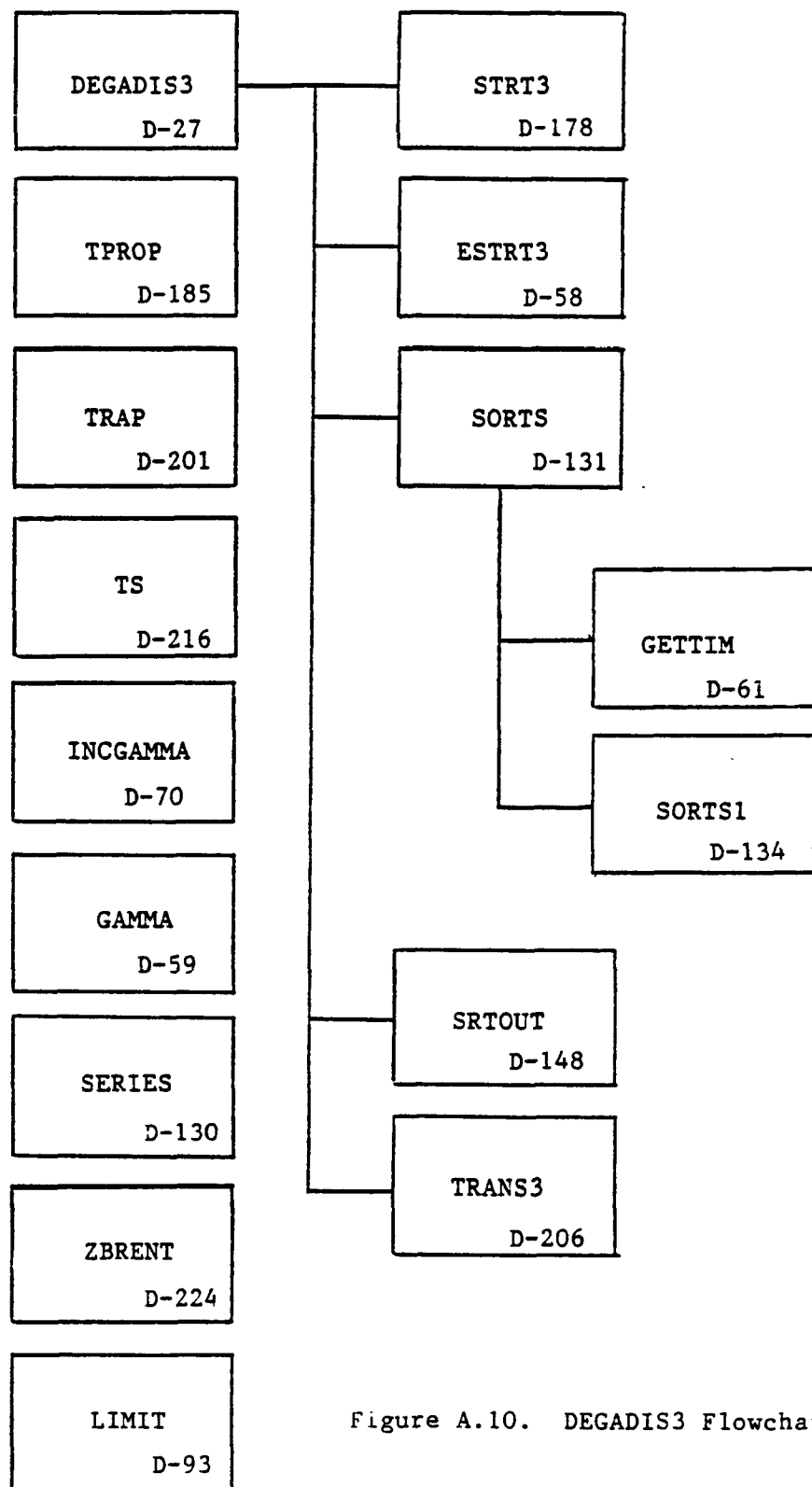


Figure A.10. DEGADIS3 Flowchart.

- (*) TPROP is a series of utility routines which estimate the thermodynamic properties of a given gas mixture (D-894).
- (*) TRANS3 writes RUN_NAME.TR4 which contains the necessary information to recover the other output files for this simulation (D-206).
- (*) TRAP is a utility included for program diagnostics (D-207).
- (*) TS calculates the time when a given observer will be at a given downwind distance (D-216).
- (*) ZBRENT finds the bracketed root of an equation using Brent's method (D-224).

As input, DEGADIS3 requires three files:

- (*) RUN_NAME.ER3 contains various numerical parameters including the time sort parameters and the flag which dictates whether the along-wind dispersion correction is applied. A copy of SYS\$DEGADIS:EXAMPLE.ER3 file uses the default time sort parameters and includes the along-wind dispersion correction which should apply for most simulations. A copy of SYS\$DEGADIS:EXAMPLE.ER3 is included in Figure A.11.
- (*) RUN_NAME.PSD contains the calculated downwind dispersion parameters for each observer. DEGADIS3 sorts this information to determine the downwind concentration profiles as a function of position at a given time.
- (*) RUN_NAME.TR3 contains the number of each record type written to RUN_NAME.PSD as well as the simulation definition.

As output, DEGADIS3 generates two new files:

- (*) RUN_NAME.SR3 is the formatted output list of the time-sorted concentration parameters. Concentration contours are generated for the specified upper and lower levels of concern at the specified height entered in DEGADISIN or JETPLU_IN.
- (*) RUN_NAME.TR4 contains the necessary information to recover the other output files to facilitate further processing.

```
! This is an example for an "ER3" run parameter file.  
! The same rules apply as for the "ER1" files.  
!  
!23456789012345678901234567890  
!-----1-----2-----3  
!  
! These values are in common area /ERROR/  
!  
ERT1      20.      FIRST SORT TIME  
ERDT      5.       SORT TIME DELTA  
ERNTIM    20.      NUMBER OF TIMES FOR THE SORT  
!  
! Note: ERNTIM is entered as a real variable even though  
! it is an integer type variable in the program.  
!  
! The value of CHECK5 determines whether the above sort parameters  
! are used. CHECK5 is initialized through the passed transfer  
! files to .FALSE. CHECK5 is set to .TRUE. if a real value of 1.  
! is passed in this file.  
!  
CHECK5     0.      USE THE DEFAULT TIME PARAMETERS  
!CHECK5     1.      USE THE TIME PARAMETERS GIVEN ABOVE  
!  
!  
sigx_flag   1.      correction for x-direction dispersion is to be made  
!sigx_flag   0.      no correction for x-direction dispersion  
!  
!  
!  
! End-of-File
```

Figure A.11. SYS\$DEGADIS:EXAMPLE.ER3 Listing.

DEAGDIS4 sorts the downwind dispersion calculation for each of several observers for concentration information at several given positions; the along-wind dispersion correction is then applied as desired. DEGADIS4 uses the following subroutines (Figure A.12):

- (*) DEGADIS4 contains the program overhead and sequentially calls the routines to recover the information generated in DEGADIS2, recover the numerical parameter file RUN_NAME.ER3, sort and apply the along-wind dispersion correction to the results of DEGADIS2, and output the results (D-33).
- (*) DOSOUT generates the formatted output file RUN_NAME.SR4 (D-45).
- (*) ESTRT3 recovers the numerical parameters contained in the file RUN_NAME.ER3, particularly whether the x-direction dispersion correction is to be applied (D-58).
- (*) GAMMA calculates the gamma function of the argument (i.e. $\Gamma(x)$) (D-59).
- (*) GETTIMDOS sets the time sort parameters as required to output the concentration time history at the desired positions (D-63).
- (*) INCGAMMA calculates the incomplete gamma function of the two arguments (D-70).
- (*) LIMIT establishes the limit for ZBRENT (D-93).
- (*) SERIES evaluates a series needed to estimate the mass of gas above a given concentration level (D-130).
- (*) SORTS recovers the information in RUN_NAME.PSD and arranges the information according to the time sort parameters (D-134).
- (*) SORTS1 applies the x-direction dispersion correction to the time-sorted information (D-131).
- (*) STRT3 recovers the information generated in DEGADIS2 contained in the file RUN_NAME.TR3 (D-178).
- (*) TPROP is a series of routines which estimate the thermodynamic properties of a given gas mixture (D-185).
- (*) TRANS3 writes RUN_NAME.TR4 which contains the necessary information to recover the other output files for this simulation (D-206).

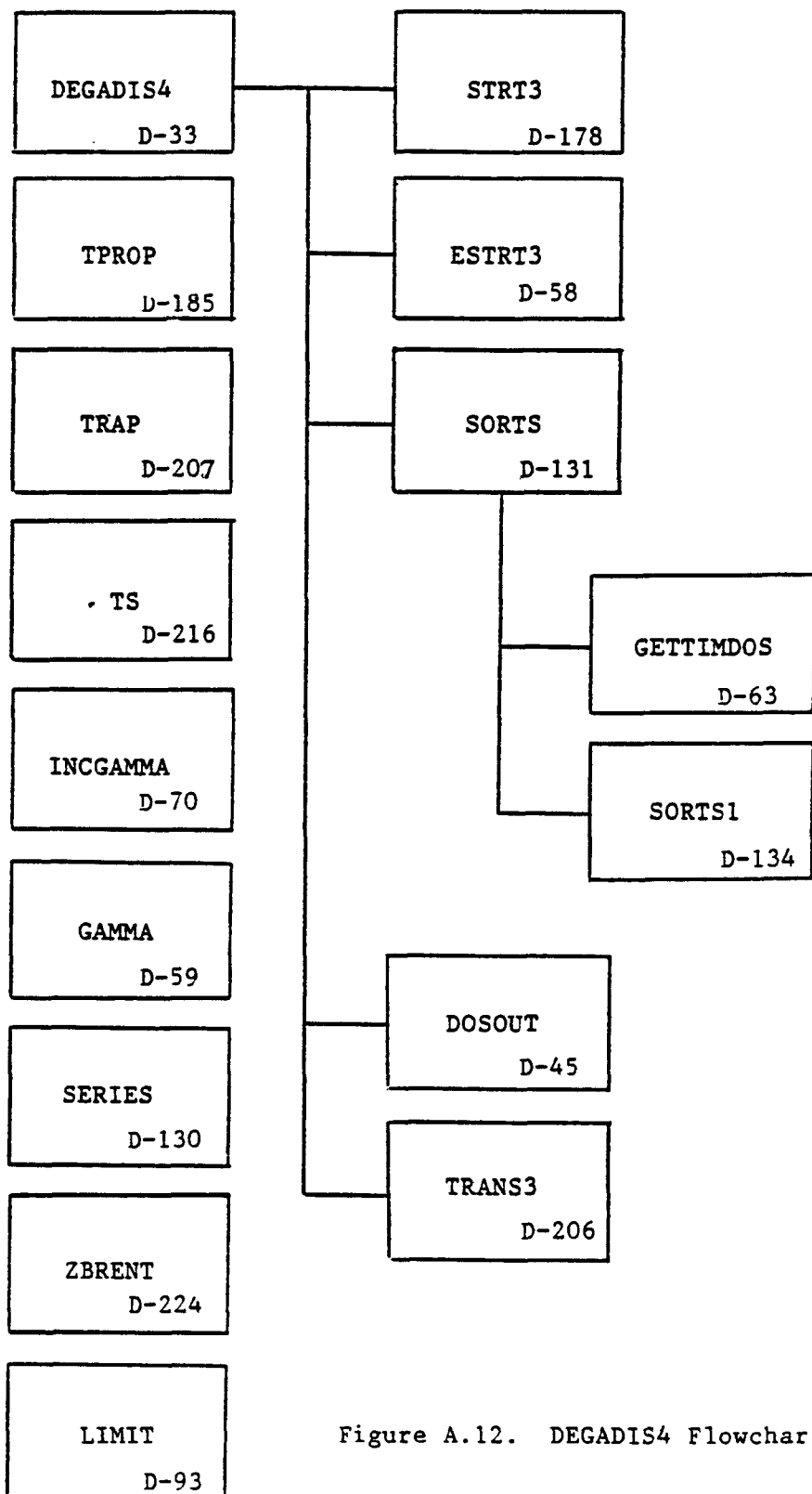


Figure A.12. DEGADIS4 Flowchart

- (*) TRAP is a utility included for program diagnostics (D-207).
- (*) TS calculates the time when a given observer will be at a given downwind distance (D-216).
- (*) ZBRENT finds the bracketed root of an equation using Brent's method (D-224).

As input, DEGADIS4 requires three files and input from the terminal:

- (*) RUN_NAME.ER3 contains the flag which dictates whether the x-direction dispersion correction is applied. A copy of SYS\$DEGADIS:EXAMPLE.ER3 file includes the x-direction dispersion correction which should apply for most simulations.
- (*) RUN_NAME.PSD contains the calculated downwind dispersion parameters for each observer. DEGADIS4 sorts this information to determine the downwind concentration time histories at the desired positions.
- (*) RUN_NAME.TR3 contains the number of each record type written to RUN_NAME.PSD as well as the simulation definition.
- (*) terminal input DEGADIS4 prompts the user for the file name to be used for this run. In addition, DEGADIS4 requests the number of downwind distances (JDOS). For each downwind distance, DEGADIS4 asks for the x-position (DOSDISX(I)). Four positions are allowed for each downwind distance (DOSDISY(IJ,I)) and DOSDISZ(IJ,I) for IJ=1 to 4). If fewer than four positions are desired, negative or zero values are entered for the first position which is not desired. A summary of this input information is included in Figure A.13. Note that the same information can be put in a command file for batch processing. The same information can be put in a file which can be associated with FOR005.DAT so that a file can be used for input.

FOR I=1 to JDOS	$\left\{ \begin{array}{l} \text{RUN_NAME} \\ \text{JDOS} \\ \text{DOSDISX}(I) \\ * \left\{ \begin{array}{l} \text{DOSDISY}(1,I), \text{DOSDISZ}(1,I) \\ \text{DOSDISY}(2,I), \text{DOSDISZ}(2,I) \\ \text{DOSDISY}(3,I), \text{DOSDISZ}(3,I) \\ \text{DOSDISY}(4,I), \text{DOSDISZ}(4,I) \end{array} \right. \end{array} \right.$
-----------------	--

Figure A.13. Structure of Input for DEGADIS4.

As output, DEGADIS4 generates two new files:

- (*) RUN_NAME.SR4 is the formatted output list of the sorted concentration time histories.
- (*) RUN_NAME.TR4 contains the necessary information to recover the other output files to facilitate further processing.

Steady-State Module--SDEGADIS2

SDEGADIS2 is a simplification of DEGADIS2 which uses many of the same subroutines. SDEGADIS2 performs the downwind dispersion portion of the calculation for a steady-state source described by DEGADIS1, SDEGADIS2 is composed of the following subroutines (Figure A.14):

- (*) AFGEN linearly interpolates between a pair of points based on a list of supplied values (D-3).
- (*) GAMMA calculates the gamma function of the argument (i.e. $\Gamma(x)$) (D-59).
- (*) ESTRT2SS recovers a subset of the numerical parameters contained in the file RUN_NAME.ER2 (D-56).
- (*) INCGAMMA calculates the incomplete gamma function of the two arguments (D-70).
- (*) LIMIT establishes the limit for ZBRENT (D-93).
- (*) PSS is the same subroutine used in DEGADIS2; it contains the ordinary differential equations which describe the downwind dispersion calculation when $b > 0$ (D-100).
- (*) PSSOUTSS governs the output of calculated points to the file RUN_NAME.SR3 when PSS is active (D-107).
- (*) RIPHIF calculates the Richardson number and the value of $\phi(R_i)$ (D-108).
- (*) RKGST performs numerical integration of a specified system of equations using a modified fourth-order Runge-Kutta methods (D-114).
- (*) SDEGADIS2 contains the program overhead and sequentially calls the routines to recover the information generated in DEGADIS1, recover the numerical parameters file RUN_NAME.ER2, and perform the steady-state simulation (D-122).
- (*) SERIES evaluates a series needed to estimate the mass of gas above a given concentration level (D-130).

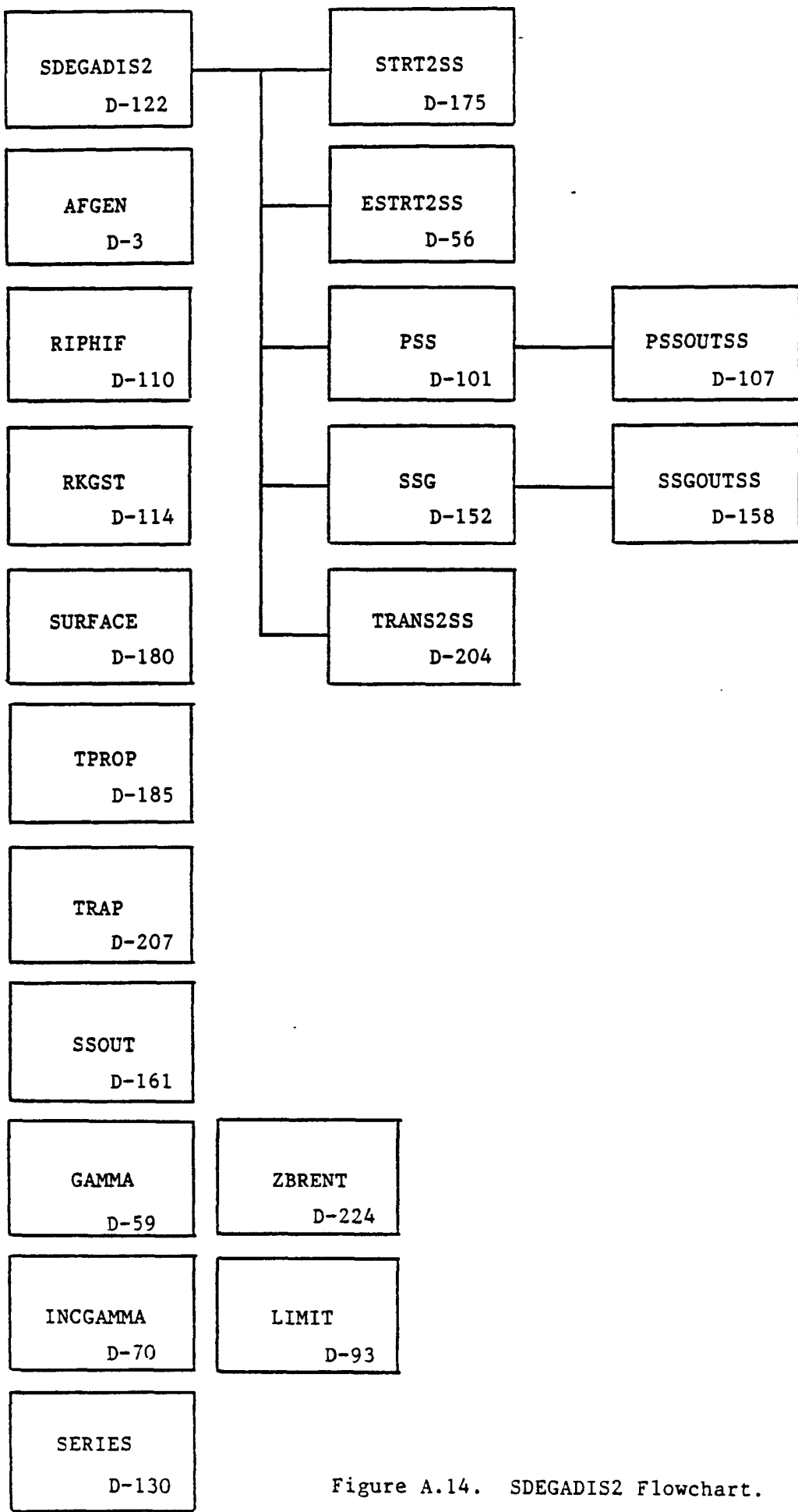


Figure A.14. SDEGADIS2 Flowchart.

- (*) SSG is the same subroutine used in DEGADIS2; it contains the ordinary differential equations which describe the downwind dispersion calculation when b = 0 (D-152).
- (*) SSGOUTSS governs the output of calculated points to the file RUN_NAME.SR3 when SSG is active (D-158).
- (*) SSOUT writes RUN_NAME.SR3 and calculates the concentration contours (D-161).
- (*) STRT2SS recovers a subset of the information generated in DEAGDIS1 contained in the file RUN_NAME.TR2 (D-175).
- (*) SURFACE estimates heat and water transfer rates across the bottom surface of the gas layer (D-180).
- (*) TPROP is a series of routines which estimate the thermodynamic properties of a given gas mixture (D-185).
- (*) TRANS2SS writes RUN_NAME.TR3 (D-204).
- (*) TRAP is a utility included for program diagnostics (D-207).
- (*) ZBRENT finds the bracketed root of an equation using Brent's method (D-224).

As input, SDEAGDIS2 requires two files:

- (*) RUN_NAME.ER2 contains various numerical parameters, the steady-state simulation typically requires only a copy of SYS\$DEGADIS:EXAMPLE.ER2.
- (*) RUN_NAME.TR2 contains the basic simulation definition as well as calculated secondary source parameters; the steady-state simulation requires only part of these.

As output, SDEAGDIS2 generates the following files:

- (*) RUN_NAME.SR3 is the formatted output list of the downwind concentration parameters. Concentration contours are generated for the specified upper and lower levels of concern at the specified height entered in DEAGDISIN.
- (*) RUN_NAME.TR3 contains the necessary information to recover the other output files to facilitate further processing.

APPENDIX B

EXAMPLE MODEL INPUT AND OUTPUT

Bhopal, India, MIC Release

The example conditions shown in Table B.1 for the accidental release of methylisocyanate (MIC) on December 3, 1984, in Bhopal, India, were reported by Singh (1986). MIC was assumed released as a pure, ambient temperature gas at a steady rate of 6.72 kg/s.

Figure B.1. MIC Example Simulation Release Conditions

Mass of MIC released	40 tons
Duration of release	90 min
Vent elevation	33 m
Vent diameter	0.2 m
Wind velocity at 10 m	1.9 m/s
Atmospheric stability	E/F

The input file for a steady-state release is shown in Figure B.1. A surface roughness of 0.1 m and an averaging time of 3600 s have been specified. For the lowest concentration of interest, the lethal concentrations to 50% of laboratory animals exposed to MIC (LC₅₀) for one and two hours are about 30 and 20 ppm, respectively. The concentrations of 30 ppm (GASUL=0.00003 mole fraction) and 20 ppm (GASLL=0.00002 mole fraction) are used to determine concentration contours in DEGADIS. The steady-state condition is indicated by TEND=0.

A batch command file is shown in Figure B.2 for the simulation called EX1. (The logical symbol SY\$DEGADIS: is assigned to the directory which contains the executable image of the model.) The output of the model is in the file with the original name and extension LIS for (for this example, EX1.LIS). To run the model interactively, the lines shown in Figure B.2 can be entered at a terminal.

The program JETPLU_IN reads the input file and generates the input file to the JETPLU program; when JETPLU_IN finishes, JETPLU_IN writes the line "JETPLU_IN - beginning command file". At this point, the JETPLU program begins; several lines of output are generated showing the values of various parameters calculated by the program. When JETPLU finishes, DEGBRIDGE begins; DEGBRIDGE takes the output of JETPLU and

Methylisocyanate (MIC) release simulation

```
2.9 10.          UO, ZO
0.1
1 6 0.          ZR
298. 1. 50.      INDVEL, ISTAB, RML
298.          TAMB, PAMB, RELHUM
MIC          TSURF
57.           GASNAM
3600.          GASMW
298.          AVTIME - based on 1 hr toxic levels
0.00003 0.00002 0.5  JETTEM
0 2000. 0.      GASUL, GASLL, ZLL
-1           INDHT, CPK, CPP
6.72          NDEN
33. 0.2        ERATE
0.0           ELEJET, DIAJET
100.          TEND
                      DISTMX
```

This is the end of the file. Any comments can be included here since the file is not read after the line for DISTMX above.

Figure B.1. Listing of EX1.IN.

```
$ RUN SYS$DEGADIS:JETPLU_IN
EX1
```

Figure B.2. Example Command File Used to Simulate the EX1 Simulation.

generates the necessary input file for DEGADIS. When DEGBRIDGE finishes, the standard DEGADIS procedure begins by copying the ER1 and ER2 parameter files. DEGADIS generates several lines of output showing the values of various numerical parameters calculated by the model. Last, the model generates the LIS file from the output files of JETPLU and DEGADIS (Figure B.3).

In the JETPLU model output, columns 3-5 correspond to plume centerline position. For the specified height, columns 9 through 11 correspond to the maximum concentration and horizontal distances to the specified mole fractions, respectively. For the MIC example case, the JETPLU calculations are continued until the centerline mole fraction drops below 5 ppm (GASLL/4). The model predicts the maximum extent of the 30 and 20 ppm concentration levels to be 2500 and 4200 m, respectively.

***** JETPLU/DEGADIS v2.1 *****

6-SEP-1989 14:40:54.50

Methylisocyanate (MIC) release simulation

Ambient Meteorological Conditions...

Ambient windspeed at reference height:	2.9000	m/s
Reference height:	10.000	m
Surface roughness:	0.10000	m
Pasquill stability class:	F	
Monin-Obukhov length:	17.524	m
Friction velocity:	0.13910	m/s
Ambient temperature:	298.00	K
Ambient pressure:	1.0000	atm
Ambient humidity:	1.00882E-02	
Relative humidity:	50.000	%
Specified averaging time:	3600.0	s
DELTAy:	9.64473E-02	
BETAy:	0.90000	
DELTAz:	1.12200E-02	
BETAz:	1.4024	
GAMMAz:	-5.40000E-02	

Contaminant Properties...

Contaminant molecular weight:	57.000	
Initial temperature:	298.00	
Upper level of interest:	3.00000E-05	
Lower level of interest:	2.00000E-05	
Heat capacity constant:	80700.	
Heat capacity power:	1.0000	

NDEN flag: -1

ISOFL flag: 1

Release Properties...

Release rate:	6.7200	kg/s
Discharge elevation:	33.000	m
Discharge diameter:	0.20000	m

Model Parameters...

ALFA1:	2.80000E-02	
ALFA2:	0.37000	
DISTMX:	100.00	m

Figure B.3.. LIS File from the Output Files of JETPLU and DEGADIS.

Downwind Distance (m)	Elevation (m)	Mole Fraction	Centerline Concentration (kg/m³)	Density (kg/m³)	Temperature (K)	Sigma y (m)	Sigma z (m)	At z= 0.500 m	Width to: 2.00E-03molz 3.00E-03m
4.122E+03	34.4	1.00	2.33	2.33	298.	0.110	0.110	0.000E+00	
56.9	45.8	8.888E-03	2.072E-02	1.19	298.	4.03	2.14	0.000E+00	
110.	45.8	3.728E-03	8.590E-03	1.18	298.	6.87	3.02	0.000E+00	
178.	45.0	1.736E-03	4.047E-03	1.18	298.	10.4	4.28	0.000E+00	
257.	44.1	9.506E-04	2.216E-03	1.18	298.	14.4	5.66	0.000E+00	
327.	43.2	6.391E-04	1.490E-03	1.18	298.	17.9	6.79	0.000E+00	
417.	42.3	4.304E-04	1.003E-03	1.18	298.	22.2	8.12	0.000E+00	
517.	41.3	3.039E-04	7.086E-04	1.18	298.	27.0	9.49	0.000E+00	
617.	40.3	2.289E-04	5.337E-04	1.18	298.	31.6	10.8	0.000E+00	
717.	39.4	1.803E-04	4.205E-04	1.18	298.	36.1	11.9	0.000E+00	
817.	38.6	1.468E-04	3.423E-04	1.18	298.	40.6	13.0	0.000E+00	
917.	37.7	1.225E-04	2.857E-04	1.18	298.	45.0	14.1	0.000E+00	
1.017E+03	36.9	1.043E-04	2.432E-04	1.18	298.	49.4	15.1	0.000E+00	
1.117E+03	36.2	9.023E-05	2.104E-04	1.18	298.	53.8	16.0	0.000E+00	
1.217E+03	35.4	7.909E-05	1.844E-04	1.18	298.	58.1	16.9	1.788E-05	
1.317E+03	34.6	7.010E-05	1.634E-04	1.18	298.	62.3	17.8	2.119E-05	21.2
1.417E+03	33.9	6.275E-05	1.463E-04	1.18	298.	66.5	18.7	2.405E-05	40.4
1.517E+03	33.2	5.667E-05	1.321E-04	1.18	298.	70.7	19.5	2.642E-05	52.8
1.617E+03	32.5	5.161E-05	1.203E-04	1.18	298.	74.9	20.2	2.831E-05	62.4
1.717E+03	31.8	4.736E-05	1.104E-04	1.18	298.	79.1	21.0	2.976E-05	70.5
1.817E+03	31.1	4.378E-05	1.021E-04	1.18	298.	83.2	21.7	3.083E-05	77.4
1.917E+03	30.5	4.075E-05	9.500E-05	1.18	298.	87.3	22.4	3.156E-05	83.4
2.017E+03	29.8	3.818E-05	8.901E-05	1.18	298.	91.3	23.1	3.201E-05	88.6
2.117E+03	29.2	3.599E-05	8.391E-05	1.18	298.	95.4	23.8	3.223E-05	93.2
2.217E+03	28.6	3.411E-05	7.953E-05	1.18	298.	99.4	24.4	3.225E-05	97.2
2.317E+03	28.0	3.250E-05	7.578E-05	1.18	298.	103.	25.0	3.212E-05	101.
2.417E+03	27.4	3.111E-05	7.252E-05	1.18	298.	107.	25.6	3.186E-05	104.
2.517E+03	26.9	2.989E-05	6.968E-05	1.18	298.	111.	26.2	3.151E-05	106.
2.617E+03	26.3	2.882E-05	6.719E-05	1.18	298.	115.	26.8	3.108E-05	108.
2.717E+03	25.8	2.788E-05	6.499E-05	1.18	298.	119.	27.4	3.060E-05	110.
2.817E+03	25.2	2.703E-05	6.302E-05	1.18	298.	123.	27.9	3.007E-05	111.
2.917E+03	24.7	2.627E-05	6.124E-05	1.18	298.	127.	28.4	2.951E-05	112.
3.017E+03	24.2	2.557E-05	5.962E-05	1.18	298.	131.	29.0	2.892E-05	113.
3.117E+03	23.7	2.494E-05	5.814E-05	1.18	298.	135.	29.5	2.833E-05	113.
3.217E+03	23.2	2.435E-05	5.676E-05	1.18	298.	139.	30.0	2.773E-05	112.
3.317E+03	22.7	2.380E-05	5.548E-05	1.18	298.	143.	30.5	2.712E-05	111.
3.417E+03	22.2	2.328E-05	5.427E-05	1.18	298.	147.	31.0	2.652E-05	110.
3.517E+03	21.7	2.279E-05	5.313E-05	1.18	298.	150.	31.4	2.592E-05	108.
3.617E+03	21.2	2.232E-05	5.204E-05	1.18	298.	154.	31.9	2.533E-05	106.
3.717E+03	20.8	2.188E-05	5.100E-05	1.18	298.	158.	32.3	2.475E-05	103.
3.817E+03	20.3	2.145E-05	5.000E-05	1.18	298.	162.	32.8	2.418E-05	99.8
3.917E+03	19.9	2.103E-05	4.904E-05	1.18	298.	166.	33.2	2.362E-05	95.6
4.017E+03	19.4	2.063E-05	4.810E-05	1.18	298.	169.	33.7	2.308E-05	90.7
4.117E+03	19.0	2.025E-05	4.720E-05	1.18	298.	173.	34.1	2.255E-05	84.9
4.217E+03	18.6	1.987E-05	4.632E-05	1.18	298.	177.	34.5	2.204E-05	78.0

Figure B.3. (continued)

4.317E+03	18.2	1.850E-05	4.547E-05	1.18	298.	181.	34.9	2.153E-05	69.5
4.417E+03	17.7	1.915E-05	4.463E-05	1.18	298.	185.	35.3	2.105E-05	58.9
4.517E+03	17.3	1.880E-05	4.382E-05	1.18	298.	188.	35.7	2.057E-05	44.7
4.617E+03	16.9	1.846E-05	4.303E-05	1.18	298.	192.	36.1	2.011E-05	20.4
4.717E+03	16.5	1.812E-05	4.226E-05	1.18	298.	196.	36.5	1.967E-05	
4.817E+03	16.1	1.780E-05	4.150E-05	1.18	298.	200.	36.8	1.923E-05	
4.917E+03	15.7	1.748E-05	4.076E-05	1.18	298.	203.	37.2	1.882E-05	
5.017E+03	15.4	1.717E-05	4.004E-05	1.18	298.	207.	37.6	1.841E-05	
5.117E+03	15.0	1.687E-05	3.933E-05	1.18	298.	211.	37.9	1.802E-05	
5.217E+03	14.6	1.657E-05	3.864E-05	1.18	298.	214.	38.3	1.763E-05	
5.317E+03	14.2	1.628E-05	3.796E-05	1.18	298.	218.	38.6	1.726E-05	
5.417E+03	13.8	1.600E-05	3.730E-05	1.18	298.	222.	39.0	1.690E-05	
5.517E+03	13.5	1.572E-05	3.665E-05	1.18	298.	225.	39.3	1.656E-05	
5.617E+03	13.1	1.545E-05	3.602E-05	1.18	298.	229.	39.6	1.622E-05	
5.717E+03	12.8	1.518E-05	3.540E-05	1.18	298.	233.	40.0	1.589E-05	
5.817E+03	12.4	1.492E-05	3.479E-05	1.18	298.	236.	40.3	1.558E-05	
5.917E+03	12.1	1.467E-05	3.420E-05	1.18	298.	240.	40.6	1.527E-05	
6.017E+03	11.7	1.442E-05	3.362E-05	1.18	298.	244.	40.9	1.497E-05	
6.117E+03	11.4	1.418E-05	3.306E-05	1.18	298.	247.	41.3	1.468E-05	
6.217E+03	11.0	1.394E-05	3.250E-05	1.18	298.	251.	41.6	1.440E-05	
6.317E+03	10.7	1.371E-05	3.196E-05	1.18	298.	254.	41.9	1.413E-05	
6.417E+03	10.4	1.348E-05	3.143E-05	1.18	298.	258.	42.2	1.387E-05	
6.517E+03	10.0	1.326E-05	3.091E-05	1.18	298.	262.	42.5	1.361E-05	
6.617E+03	9.71	1.304E-05	3.041E-05	1.18	298.	265.	42.8	1.336E-05	
6.717E+03	9.38	1.283E-05	2.991E-05	1.18	298.	269.	43.1	1.312E-05	
6.817E+03	9.06	1.262E-05	2.943E-05	1.18	298.	273.	43.3	1.289E-05	
6.917E+03	8.74	1.242E-05	2.895E-05	1.18	298.	276.	43.6	1.266E-05	
7.017E+03	8.43	1.222E-05	2.849E-05	1.18	298.	280.	43.9	1.244E-05	
7.117E+03	8.12	1.203E-05	2.804E-05	1.18	298.	283.	44.2	1.222E-05	
7.217E+03	7.81	1.184E-05	2.759E-05	1.18	298.	287.	44.5	1.201E-05	
7.317E+03	7.50	1.165E-05	2.716E-05	1.18	298.	290.	44.7	1.181E-05	
7.417E+03	7.19	1.147E-05	2.674E-05	1.18	298.	294.	45.0	1.161E-05	
7.517E+03	6.89	1.129E-05	2.632E-05	1.18	298.	298.	45.3	1.142E-05	
7.617E+03	6.59	1.112E-05	2.592E-05	1.18	298.	301.	45.5	1.123E-05	
7.717E+03	6.30	1.095E-05	2.552E-05	1.18	298.	305.	45.8	1.105E-05	
7.817E+03	6.00	1.078E-05	2.514E-05	1.18	298.	308.	46.0	1.087E-05	
7.917E+03	5.71	1.062E-05	2.476E-05	1.18	298.	312.	46.3	1.070E-05	
8.017E+03	5.42	1.046E-05	2.439E-05	1.18	298.	315.	46.5	1.053E-05	
8.117E+03	5.13	1.031E-05	2.403E-05	1.18	298.	319.	46.8	1.037E-05	
8.217E+03	4.84	1.015E-05	2.367E-05	1.18	298.	322.	47.0	1.021E-05	
8.317E+03	4.56	1.000E-05	2.332E-05	1.18	298.	326.	47.3	1.005E-05	
8.417E+03	4.28	9.859E-06	2.298E-05	1.18	298.	329.	47.5	9.898E-06	
8.517E+03	4.00	9.716E-06	2.265E-05	1.18	298.	333.	47.8	9.750E-06	
8.617E+03	3.72	9.577E-06	2.233E-05	1.18	298.	336.	48.0	9.605E-06	
8.717E+03	3.45	9.440E-06	2.201E-05	1.18	298.	340.	48.2	9.464E-06	
8.817E+03	3.17	9.307E-06	2.170E-05	1.18	298.	343.	48.5	9.326E-06	
8.917E+03	2.90	9.176E-06	2.139E-05	1.18	298.	347.	48.7	9.192E-06	
9.017E+03	2.63	9.048E-06	2.109E-05	1.18	298.	350.	48.9	9.061E-06	
9.117E+03	2.37	8.923E-06	2.080E-05	1.18	298.	354.	49.2	8.933E-06	
9.217E+03	2.10	8.800E-06	2.052E-05	1.18	298.	357.	49.4	8.808E-06	
9.317E+03	1.84	8.680E-06	2.024E-05	1.18	298.	361.	49.6	8.685E-06	
9.417E+03	1.58	8.562E-06	1.996E-05	1.18	298.	364.	49.8	8.566E-06	
9.517E+03	1.32	8.447E-06	1.969E-05	1.18	298.	368.	50.0	8.449E-06	
9.617E+03	1.06	8.334E-06	1.943E-05	1.18	298.	371.	50.3	8.335E-06	
9.717E+03	0.803	8.223E-06	1.917E-05	1.18	298.	375.	50.5	8.224E-06	
9.817E+03	0.549	8.115E-06	1.892E-05	1.18	298.	378.	50.7	8.115E-06	
9.917E+03	0.296	8.009E-06	1.867E-05	1.18	298.	382.	50.9	8.008E-06	
1.002E+04	4.518E-02	7.905E-06	1.843E-05	1.18	298.	385.	51.1	7.904E-06	
1.003E+04	0.000E+00	7.886E-06	1.839E-05	1.18	298.	386.	51.1	7.886E-06	

Figure B.3. (continued)

U O A _ D E G A D I S M O D E L O U T P U T -- V E R S I O N 2 . 1

***** 6-SEP-1989 14:44:19.11 *****

Data input on 6-SEP-1989 14:44:11.16
 Source program run on 6-SEP-1989 14:44:19.11

TITLE BLOCK

Methylisocyanate (MIC) release simulation

Wind velocity at reference height	2.90	m/s	
Reference height	10.00	m	
Surface roughness length	0.100	m	
Pasquill Stability class	F		
Monin-Obukhov length	17.5	m	
Gaussian distribution constants			
Specified averaging time	3600.00	s	
Delta	0.09645		
Beta	0.90000		
Wind velocity power law constant	Alpha	0.44904	
Friction velocity		0.13910 m/s	
Ambient Temperature		298.00 K	
Ambient Pressure		1.000 atm	
Ambient Absolute Humidity		1.009E-02 kg/kg BDA	
Ambient Relative Humidity		50.00 %	
Input:	Mole fraction	CONCENTRATION OF C	GAS DENSITY
		kg/m***3	kg/m***3
0.00000	0.00000		1.17737
1.00000	2.33112		2.33112

Specified Gas Properties:

Molecular weight:	57.000
Storage temperature:	298.00 K
Density at storage temperature and ambient pressure:	2.3311 kg/m***3
Mean heat capacity constant:	80700.
Mean heat capacity power:	1.0000
Upper mole fraction contour:	3.00000E-05
Lower mole fraction contour:	2.00000E-05
Height for isolopleths:	0.50000 m

Source input data points

Initial mass in cloud: 0.00000E+00

Time	Contaminant Mass Rate	Source Radius	Contaminant Mass Fraction	Temperature	Enthalpy
s	kg/s	m	kg contam/kg mix	K	J/kg
0.00000E+00	6.7200	829.28	1.56157E-05	298.00	0.00000E+00
60230.	6.7200	829.28	1.56157E-05	298.00	0.00000E+00
60231.	0.00000E+00	0.00000E+00	1.56157E-05	298.00	0.00000E+00
60232.	0.00000E+00	0.00000E+00	1.56157E-05	298.00	0.00000E+00

Calculation procedure for ALPHA: 1

Entrainment prescription for PHI: 3

Layer thickness ratio used for average depth: 2.1500

Air entrainment coefficient used: 0.590

Gravity slumping velocity coefficient used: 1.150

Isothermal calculation

Heat transfer not included

Figure B.3. (continued)

***** CALCULATED SOURCE PARAMETERS *****

Time sec	Gas Radius m	Height m	Qstar kg/m**2/s	SZ(x=L/2.) m	Mole frac C	Density kg/m**3	Rich No.
0.00000E+00 5.75485	829.280 829.293	1.10000E-05 0.115404	2.741708E-06 2.737199E-06	56.6987 61.9269	7.886120E-06 7.873082E-06	1.17738 1.17738	0.756144 0.756144

Source strength [kg/s] : 6.7200 Equivalent Primary source radius [m] : 829.28
 Equivalent Primary source length [m] : 1658.6 Equivalent Primary source half-width [m] : 651.32

Secondary source concentration [kg/m**3] : 1.83552E-05 Secondary source SZ [m] : 61.927

Contaminant flux rate: 3.11030E-06

Secondary source mass fractions... contaminant: 1.558989E-05 air: 0.99000
 Enthalpy: 0.00000E+00 Density: 1.1774

Secondary source length [m]	1658.6	Secondary source half-width [m]	651.33					
Distance (m)	Mole Fraction	Concentration Density (kg/m**3)	Density (kg/m**3)	Temperature (K)	Half Width (m)	Sz (m)	Sy (m)	Width at z= 0.50 m to: 2.000E-03 mole% 3.000E-03 mole%

1.086E+04	7.873E-06	1.836E-05	1.18	298.	651.	61.9	0.000E+00
1.086E+04	7.873E-06	1.836E-05	1.18	298.	650.	61.9	1.86
1.086E+04	7.873E-06	1.835E-05	1.18	298.	649.	61.9	2.64
1.087E+04	7.872E-06	1.835E-05	1.18	298.	635.	62.0	18.8
1.087E+04	7.861E-06	1.833E-05	1.18	298.	628.	62.0	26.5

For the UFL of 3.00000E-03 mole percent, and the LFL of 2.00000E-03 mole percent:

The mass of contaminant between the UFL and LFL is: 0.00000E+00 kg.
 The mass of contaminant above the LFL is: 0.00000E+00 kg.

Figure B.3. (concluded)

Enid, Oklahoma, Ammonia Release

The model was also used to simulate an ammonia pipeline failure which occurred near Enid, Oklahoma, on May 7, 1976 (Atwood, 1976). The release occurred during early morning hours (~ 8 AM) when a bulldozer struck and ruptured an eight-inch ammonia pipeline operating at approximately 700 psi. Approximately 500 tons of ammonia were released in 4.5 hours from the ten-mile section isolated between check valves. The windspeed at the time of the accident was reported to be approximately 10 mph. The dense aerosol plume reportedly etched a parabolic-shaped scar about 6 miles long and one-half mile wide. The conditions used to simulate the release are shown in Table B.2.

Table B.2. Ammonia Example Simulation
Release Conditions

Ammonia release rate	56 kg/s (twice 500 tons per 4.5 hours)
Release elevation	zero (ground level)
Vent diameter	0.2 m
Wind speed (at 10 m)	4.5 m/s
Surface roughness	1 cm
Atmospheric stability	D

The input file is shown in Figure B.4. The release elevation has been set to 0.02 m ($2 z_R$) to avoid numerical problems. The ammonia aerosol/(humid) air mixture density shown in Figure B.5 was determined by assuming the ambient air and release ammonia aerosol mixes adiabatically (Spicer and Havens, 1988). The relationship between mixture density and ammonia concentration is input to the model with 20 ordered triples of ammonia mole fraction, ammonia concentration (kg/m^3), and ammonia/air mixture density (kg/m^3).

Figure B.6 shows the predicted 1000 ppm and 100 ppm ground-level concentration isopleths which assume one-hour averaging time for the estimation of lateral dispersion by meander. The predicted plume rise was approximately 70 meters, and the plume was predicted to return to ground level about 525 meters downwind of the release. The LIS file from the output files of JETPLU and DEGADIS is shown in Figure B.7.

Enid, Oklahoma, ammonia pipeline release simulation

4.5	10.	U0, Z0
0.01		ZR
1	4	INDVEL, ISTAB, RML
298.	0.98	TAMB, PAMB, RELHUM
298.		TSURF
NH3		GASNAM
17.		GASMW
3600.		AVTIME
298.		TEMJET
0.001	0.0001	GASUL, GASLL, ZLL
0	3845.000	INDHT, CPK, CPP
22		NDEN
0.0000000E+00	0.0000000E+00	1.153908
1.4000000E-02	9.8185645E-03	1.185840
2.0000000E-02	1.4134786E-02	1.192022
4.2000000E-02	3.0499677E-02	1.213623
6.2000000E-02	4.6361663E-02	1.239219
8.1000000E-02	6.2597632E-02	1.270430
0.1050000	8.5226245E-02	1.320677
0.1420000	0.1255372	1.415537
0.1610000	0.1486259	1.465818
0.1760000	0.1675721	1.501815
0.1890000	0.1841250	1.527787
0.2040000	0.2030910	1.550787
0.2230000	0.2269812	1.571988
0.4430000	0.5594106	1.755602
0.5560000	0.8019109	1.890973
0.6540000	1.081982	2.055484
0.7390000	1.410169	2.257174
0.8120000	1.797091	2.504694
0.8740000	2.255920	2.809017
0.9250000	2.790716	3.175532
0.9670000	3.424175	3.622905
1.0000000	4.149494	4.149494
56.		ERATE
0.02	0.2	ELEJET, DIAJET
0.0		TEND
20.		DISTMX

This is the end of the file. Any comments can be included here since the file is not read after the line for DISTMX above.

Figure B.4. Listing of EX2.IN.

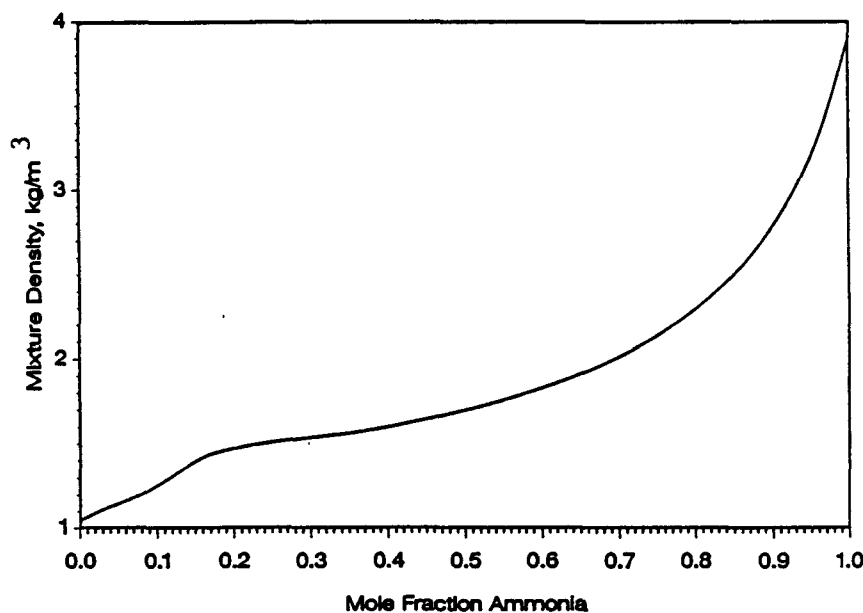


Figure B.5. Ammonia Aerosol/Air Adiabatic Mixture Density.

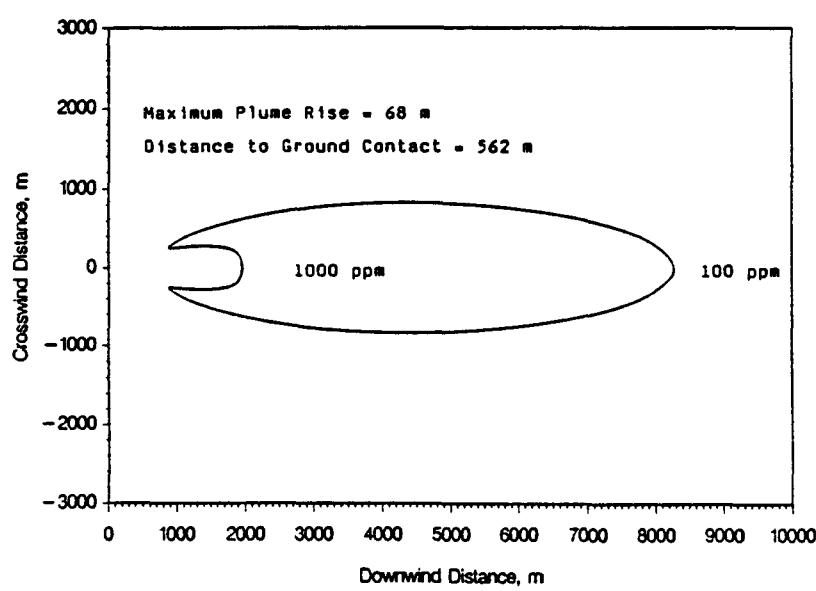


Figure B.6. Ground-Level Isocontours--Oklahoma Ammonia Pipeline Break

***** JETPLU/DEGADIS v2.1 *****

6-SEP-1989 15:48:31.08

Enid, Oklahoma, ammonia pipeline release simulation

Ambient Meteorological Conditions...

Ambient windspeed at reference height: 4.5000 m/s
 Reference height: 10.000 m
 Surface roughness: 1.00000E-02 m

Pasquill stability class: D

Monin-Obukhov length: 0.00000E+00 m
 Friction velocity: 0.22797 m/s
 Ambient temperature: 298.00 K
 Ambient pressure: 0.98000 atm
 Ambient humidity: 1.03009E-02
 Relative humidity: 50.000 %

Specified averaging time: 3600.0 s
 DELTAy: 0.19461
 BETAY: 0.90000
 DELTAz: 4.13400E-02
 BETAZ: 1.1737
 GAMMAz: -3.16000E-02

Contaminant Properties...

Contaminant molecular weight: 17.000
 Initial temperature: 298.00
 Upper level of interest: 1.00000E-03
 Lower level of interest: 1.00000E-04
 Heat capacity constant: 3845.0
 Heat capacity power: 1.0000

NDEN flag: 22

Mole fraction	Concentration (kg/m ³)	Density (kg/m ³)
0.00000E+00	0.00000E+00	1.1539
1.40000E-02	9.81856E-03	1.1858
2.00000E-02	1.41348E-02	1.1920
4.20000E-02	3.04997E-02	1.2136
6.20000E-02	4.63617E-02	1.2392
8.10000E-02	6.25976E-02	1.2704
0.10500	8.52262E-02	1.3207
0.14200	0.12554	1.4155
0.16100	0.14863	1.4658
0.17600	0.16757	1.5018
0.18900	0.18413	1.5278
0.20400	0.20309	1.5508
0.22300	0.22698	1.5720
0.44300	0.55941	1.7556
0.55600	0.80191	1.8910
0.65400	1.0820	2.0555
0.73900	1.4102	2.2572
0.81200	1.7971	2.5047
0.87400	2.2559	2.8090
0.92500	2.7907	3.1755
0.96700	3.4242	3.6229
1.00000	4.1495	4.1495

ISOFL flag: 1

Figure B.7. LIS File from Output Files of JETPLU and DEGADIS.

Release Properties...

Release rate: 56.000 kg/s
 Discharge elevation: 2.00000E-02 m
 Discharge diameter: 0.20000 m

Model Parameters...

ALFA1: 2.80000E-02

Downwind Distance (m)	Elevation (m)	Mole Fraction	Centerline Concentration (kg/m³)	Density (kg/m³)	Temperature (K)	Sigma y (m)	Sigma z (m)	At z= 0.000E+00 m -----		
								Mole Fraction	Width to: 1.00E-02molz	(m)
2.151E-17	1.56	1.00	4.15	4.15	298.	0.110	0.110	0.000E+00		
13.2	45.9	6.181E-02	4.642E-02	1.24	298.	5.00	4.64	0.000E+00		
27.5	56.0	4.044E-02	2.944E-02	1.21	298.	7.36	6.44	0.000E+00		
41.3	61.2	3.127E-02	2.251E-02	1.20	298.	9.14	7.56	0.000E+00		
60.9	65.3	2.423E-02	1.729E-02	1.20	298.	11.2	8.57	0.000E+00		
80.8	67.2	2.004E-02	1.423E-02	1.19	298.	13.1	9.18	0.000E+00		
101.	67.5	1.724E-02	1.220E-02	1.19	298.	14.9	9.54	0.000E+00		
121.	66.6	1.460E-02	1.029E-02	1.19	298.	16.8	10.1	0.000E+00		
141.	64.7	1.219E-02	8.553E-03	1.18	298.	18.8	10.8	0.000E+00		
160.	62.2	1.019E-02	7.112E-03	1.18	298.	20.9	11.7	0.000E+00		
180.	59.4	8.577E-03	5.965E-03	1.17	298.	23.1	12.6	0.000E+00		
200.	56.3	7.297E-03	5.060E-03	1.17	298.	25.3	13.6	0.000E+00		
220.	53.2	6.278E-03	4.343E-03	1.17	298.	27.4	14.6	0.000E+00		
240.	50.0	5.459E-03	3.769E-03	1.17	298.	29.6	15.7	0.000E+00		
259.	46.7	4.794E-03	3.304E-03	1.16	298.	31.7	16.7	0.000E+00		
279.	43.5	4.246E-03	2.923E-03	1.16	298.	33.8	17.7	0.000E+00		
299.	40.2	3.791E-03	2.607E-03	1.16	298.	35.9	18.7	7.628E-04	72.5	
318.	37.0	3.411E-03	2.343E-03	1.16	298.	38.0	19.8	1.183E-03	81.8	22.1
338.	33.8	3.098E-03	2.127E-03	1.16	298.	40.1	20.7	1.638E-03	86.2	39.9
358.	30.7	2.857E-03	1.960E-03	1.16	298.	42.2	21.7	2.077E-03	90.7	51.1
378.	27.5	2.700E-03	1.852E-03	1.16	298.	44.2	22.7	2.461E-03	95.1	59.5
397.	24.4	2.635E-03	1.807E-03	1.16	298.	46.3	23.7	2.764E-03	99.5	66.1
417.	21.4	2.653E-03	1.819E-03	1.16	298.	48.3	24.6	2.978E-03	104.	71.5
437.	18.3	2.728E-03	1.871E-03	1.16	298.	50.4	25.5	3.105E-03	108.	76.0
457.	15.3	2.821E-03	1.935E-03	1.16	298.	52.4	26.5	3.155E-03	113.	79.6
477.	12.4	2.896E-03	1.987E-03	1.16	298.	54.4	27.4	3.141E-03	117.	82.4
496.	9.48	2.928E-03	2.009E-03	1.16	298.	56.4	28.3	3.077E-03	121.	84.7
516.	6.60	2.904E-03	1.993E-03	1.16	298.	58.3	29.1	2.975E-03	125.	86.3
536.	3.75	2.826E-03	1.939E-03	1.16	298.	60.3	30.0	2.848E-03	130.	87.4
556.	0.937	2.703E-03	1.854E-03	1.16	298.	62.3	30.9	2.704E-03	134.	88.0
562.	0.000E+00	2.654E-03	1.820E-03	1.16	298.	62.9	31.2	2.653E-03	135.	88.1

(END OF JETPLU SIMULATION)

Figure B.7. (continued)

***** U O A _ D E G A D I S M O D E L O U T P U T -- V E R S I O N 2.1 *****
 ***** 6-SEP-1989 15:49:46.45 *****

Data input on 6-SEP-1989 15:49:38.76
 Source program run on 6-SEP-1989 15:49:46.45

TITLE BLOCK

Enid, Oklahoma ammonia pipeline release simulation

Wind velocity at reference height	4.50	m/s	
Reference height	10.00	m	
Surface roughness length	1.000E-02	m	
Pasquill Stability class	D		
Monin-Obukhov length	infinite		
Gaussian distribution constants			
Specified averaging time	3600.00	s	
Delta	0.19461		
Beta	0.90000		
Wind velocity power law constant	Alpha	0.18680	
Friction velocity		0.22797 m/s	
Ambient Temperature	298.00 K		
Ambient Pressure	0.980 atm		
Ambient Absolute Humidity	1.030E-02 kg/kg BDA		
Ambient Relative Humidity	50.00 %		
Input:	Mole fraction	CONCENTRATION OF C	GAS DENSITY
		kg/m**3	kg/m**3
0.00000	0.00000	1.15391	
0.01400	0.00982	1.18584	
0.02000	0.01413	1.19202	
0.04200	0.03050	1.21362	
0.06200	0.04636	1.23922	
0.08100	0.06260	1.27043	
0.10500	0.08523	1.32068	
0.14200	0.12554	1.41554	
0.16100	0.14863	1.46582	
0.17600	0.16757	1.50182	
0.18900	0.18413	1.52779	
0.20400	0.20309	1.55079	
0.22300	0.22698	1.57199	
0.44300	0.55941	1.75560	
0.55600	0.80191	1.89097	
0.65400	1.08198	2.05548	
0.73900	1.41017	2.25717	
0.81200	1.79709	2.50469	
0.87400	2.25592	2.80902	
0.92500	2.79072	3.17553	
0.96700	3.42418	3.62290	
1.00000	4.14949	4.14949	

Specified Gas Properties:

Molecular weight:	17.000
Storage temperature:	298.00 K
Density at storage temperature and ambient pressure:	4.1495 kg/m**3
Mean heat capacity constant:	3845.0
Mean heat capacity power:	1.0000
Upper mole fraction contour:	1.00000E-03
Lower mole fraction contour:	1.00000E-04
Height for isopleths:	0.00000E+00m

Figure B.7. (continued)

Source input data points

Initial mass in cloud: 0.00000E+00

Time s	Contaminant Mass Rate kg/s	Source Radius m	Contaminant Mass Fraction kg contam/kg mix	Temperature K	Enthalpy J/kg
0.00000E+00	56.000	135.30	1.56929E-03	298.00	0.00000E+00
60230.	56.000	135.30	1.56929E-03	298.00	0.00000E+00
60231.	0.00000E+00	0.00000E+00	1.56929E-03	298.00	0.00000E+00
60232.	0.00000E+00	0.00000E+00	1.56929E-03	298.00	0.00000E+00

Calculation procedure for ALPHA: 1

Entrainment prescription for PHI: 3

Layer thickness ratio used for average depth: 2.1500

Air entrainment coefficient used: 0.590

Gravity slumping velocity coefficient used: 1.150

Isothermal calculation

Heat transfer not included

Water transfer not included

***** CALCULATED SOURCE PARAMETERS *****

Time sec	Gas Radius m	Height m	Qstar kg/m**2/s	SZ(x=L/2.) m	Mole frac C	Density kg/m**3	Rich No.
0.00000E+00	135.300	1.100000E-05	2.902896E-04	11.1516	2.654004E-03	1.15996	0.756144
4.79323	136.413	1.77907	2.883740E-04	11.2280	2.636814E-03	1.15992	0.756144
23.4441	146.849	7.82487	2.719845E-04	11.9374	2.490347E-03	1.15959	0.756144
64.4041	181.434	15.3666	2.327196E-04	14.1936	2.145477E-03	1.15880	0.756144
105.364	219.626	17.5763	2.059423E-04	16.5237	1.919935E-03	1.15829	0.756144
146.324	256.937	17.0954	1.896346E-04	18.6339	1.793855E-03	1.15800	0.756144
197.524	300.368	14.6524	1.789255E-04	20.8536	1.731507E-03	1.15786	0.756144
218.004	315.190	13.4747	1.770343E-04	21.5390	1.730059E-03	1.15785	0.756144
228.244	317.744	13.2504	1.769059E-04	21.6129	1.732304E-03	1.15786	0.756144

Source strength [kg/s] : 56.000 Equivalent Primary source radius [m] : 135.30

Equivalent Primary source length [m] : 270.60 Equivalent Primary source half-width [m] : 106.26

Secondary source concentration [kg/m**3] : 1.18555E-03 Secondary source SZ [m] : 21.613

Contaminant flux rate: 1.76556E-04

Secondary source mass fractions... contaminant: 1.023911E-03 air: 0.98879
Enthalpy: 0.00000E+00 Density: 1.1579

Secondary source length [m] : 635.49 Secondary source half-width [m] : 249.56

Distance (m)	Mole Fraction	Concentration (kg/m**3)	Density (kg/m**3)	Temperature (K)	Half Width (m)	Sz (m)	Sy (m)	Width at z= 0.00 m to: 1.000E-02moleZ	0.100 moleZ (m)
880.	1.732E-03	1.186E-03	1.16	298.	250.	21.8	0.000E+00	250.	250.
884.	1.729E-03	1.183E-03	1.16	298.	237.	21.6	15.4	263.	248.
894.	1.719E-03	1.176E-03	1.16	298.	226.	21.5	29.8	276.	248.
914.	1.704E-03	1.166E-03	1.16	298.	215.	21.4	46.9	294.	249.
949.	1.678E-03	1.148E-03	1.16	298.	204.	21.2	67.4	317.	252.
1.009E+03	1.631E-03	1.116E-03	1.16	298.	192.	20.9	93.2	348.	258.
1.089E+03	1.569E-03	1.073E-03	1.16	298.	184.	20.7	120.	384.	265.
1.169E+03	1.507E-03	1.030E-03	1.16	298.	179.	20.5	143.	415.	270.
1.249E+03	1.444E-03	9.878E-04	1.16	298.	175.	20.4	164.	443.	274.

Figure B.7. (continued)

1.329E+03	1.383E-03	9.457E-04	1.16	298.	173.	20.4	183.	469.	277.
1.409E+03	1.323E-03	9.045E-04	1.16	298.	171.	20.5	201.	493.	277.
1.489E+03	1.265E-03	8.644E-04	1.16	298.	170.	20.6	218.	516.	275.
1.569E+03	1.208E-03	8.255E-04	1.16	298.	169.	20.8	234.	538.	270.
1.644E+03	1.157E-03	7.904E-04	1.16	298.	168.	21.0	248.	557.	263.
1.724E+03	1.104E-03	7.543E-04	1.16	298.	168.	21.2	264.	576.	251.
1.804E+03	1.054E-03	7.198E-04	1.16	298.	167.	21.5	278.	595.	231.
1.884E+03	1.005E-03	6.868E-04	1.16	298.	167.	21.8	293.	612.	189.
1.964E+03	9.595E-04	6.555E-04	1.16	298.	167.	22.2	307.	628.	
2.044E+03	9.160E-04	6.256E-04	1.16	298.	167.	22.5	321.	644.	
2.124E+03	8.746E-04	5.973E-04	1.16	298.	167.	22.9	334.	659.	
2.204E+03	8.353E-04	5.704E-04	1.16	298.	166.	23.3	347.	673.	
2.284E+03	7.981E-04	5.450E-04	1.16	298.	166.	23.8	360.	686.	
2.364E+03	7.629E-04	5.209E-04	1.16	298.	166.	24.2	373.	698.	
2.444E+03	7.296E-04	4.981E-04	1.16	298.	166.	24.7	386.	710.	
2.524E+03	6.982E-04	4.766E-04	1.16	298.	166.	25.2	398.	721.	
2.604E+03	6.684E-04	4.563E-04	1.16	298.	165.	25.7	411.	732.	
2.684E+03	6.403E-04	4.370E-04	1.16	298.	165.	26.2	423.	742.	
2.764E+03	6.137E-04	4.188E-04	1.16	298.	165.	26.7	435.	751.	
2.844E+03	5.885E-04	4.017E-04	1.16	298.	164.	27.2	447.	760.	
2.924E+03	5.647E-04	3.854E-04	1.16	298.	164.	27.8	459.	768.	
3.004E+03	5.422E-04	3.700E-04	1.16	298.	164.	28.3	470.	776.	
3.084E+03	5.210E-04	3.555E-04	1.16	298.	163.	28.9	482.	783.	
3.164E+03	5.008E-04	3.417E-04	1.16	298.	163.	29.5	493.	789.	
3.244E+03	4.817E-04	3.287E-04	1.15	298.	162.	30.1	505.	795.	
3.324E+03	4.637E-04	3.164E-04	1.15	298.	162.	30.6	516.	801.	
3.404E+03	4.465E-04	3.047E-04	1.15	298.	161.	31.2	527.	806.	
3.484E+03	4.303E-04	2.936E-04	1.15	298.	160.	31.8	538.	811.	
3.564E+03	4.149E-04	2.830E-04	1.15	298.	160.	32.4	549.	815.	
3.644E+03	4.002E-04	2.730E-04	1.15	298.	159.	33.1	560.	819.	
3.724E+03	3.863E-04	2.636E-04	1.15	298.	158.	33.7	571.	822.	
3.804E+03	3.731E-04	2.545E-04	1.15	298.	157.	34.3	582.	825.	
3.884E+03	3.606E-04	2.460E-04	1.15	298.	156.	34.9	593.	828.	
3.964E+03	3.486E-04	2.378E-04	1.15	298.	155.	35.6	603.	830.	
4.044E+03	3.372E-04	2.300E-04	1.15	298.	155.	36.2	614.	831.	
4.124E+03	3.264E-04	2.226E-04	1.15	298.	154.	36.8	624.	833.	
4.204E+03	3.161E-04	2.156E-04	1.15	298.	153.	37.5	635.	834.	
4.284E+03	3.062E-04	2.089E-04	1.15	298.	152.	38.1	645.	834.	
4.364E+03	2.968E-04	2.024E-04	1.15	298.	151.	38.8	655.	834.	
4.444E+03	2.878E-04	1.963E-04	1.15	298.	150.	39.4	665.	834.	
4.524E+03	2.792E-04	1.904E-04	1.15	298.	148.	40.1	676.	833.	
4.604E+03	2.710E-04	1.849E-04	1.15	298.	147.	40.7	686.	832.	
4.684E+03	2.632E-04	1.795E-04	1.15	298.	146.	41.4	696.	831.	
4.764E+03	2.557E-04	1.744E-04	1.15	298.	145.	42.1	706.	829.	
4.844E+03	2.485E-04	1.695E-04	1.15	298.	144.	42.7	716.	827.	
4.924E+03	2.416E-04	1.648E-04	1.15	298.	143.	43.4	726.	824.	
5.004E+03	2.350E-04	1.603E-04	1.15	298.	141.	44.1	735.	821.	
5.084E+03	2.287E-04	1.559E-04	1.15	298.	140.	44.8	745.	818.	
5.164E+03	2.226E-04	1.518E-04	1.15	298.	139.	45.4	755.	814.	
5.244E+03	2.168E-04	1.478E-04	1.15	298.	138.	46.1	765.	810.	
5.324E+03	2.112E-04	1.440E-04	1.15	298.	136.	46.8	774.	806.	
5.404E+03	2.058E-04	1.403E-04	1.15	298.	135.	47.5	784.	801.	
5.484E+03	2.006E-04	1.368E-04	1.15	298.	133.	48.1	794.	796.	
5.564E+03	1.957E-04	1.334E-04	1.15	298.	132.	48.8	803.	790.	
5.644E+03	1.909E-04	1.301E-04	1.15	298.	131.	49.5	813.	784.	
5.724E+03	1.863E-04	1.270E-04	1.15	298.	129.	50.2	822.	778.	
5.804E+03	1.818E-04	1.240E-04	1.15	298.	128.	50.9	831.	771.	
5.884E+03	1.775E-04	1.210E-04	1.15	298.	126.	51.6	841.	763.	
5.964E+03	1.734E-04	1.182E-04	1.15	298.	125.	52.3	850.	756.	
6.044E+03	1.694E-04	1.155E-04	1.15	298.	123.	53.0	859.	747.	

Figure B.7. (continued)

6.124E+03	1.656E-04	1.129E-04	1.15	298.	122.	53.7	869.	739.
6.204E+03	1.619E-04	1.104E-04	1.15	298.	120.	54.3	878.	730.
6.284E+03	1.583E-04	1.079E-04	1.15	298.	119.	55.0	887.	720.
6.364E+03	1.548E-04	1.056E-04	1.15	298.	117.	55.7	896.	710.
6.444E+03	1.515E-04	1.033E-04	1.15	298.	116.	56.4	905.	699.
6.524E+03	1.483E-04	1.011E-04	1.15	298.	114.	57.1	914.	688.
6.604E+03	1.452E-04	9.896E-05	1.15	298.	112.	57.8	923.	676.
6.684E+03	1.421E-04	9.690E-05	1.15	298.	111.	58.5	932.	664.
6.764E+03	1.392E-04	9.491E-05	1.15	298.	109.	59.2	941.	651.
6.844E+03	1.364E-04	9.298E-05	1.15	298.	107.	59.9	950.	637.
6.924E+03	1.336E-04	9.111E-05	1.15	298.	106.	60.6	959.	622.
7.004E+03	1.310E-04	8.929E-05	1.15	298.	104.	61.3	968.	607.
7.084E+03	1.284E-04	8.754E-05	1.15	298.	102.	62.0	977.	591.
7.164E+03	1.259E-04	8.584E-05	1.15	298.	101.	62.7	986.	574.
7.244E+03	1.235E-04	8.419E-05	1.15	298.	99.0	63.4	994.	556.
7.324E+03	1.211E-04	8.259E-05	1.15	298.	97.3	64.1	1.003E+03	537.
7.404E+03	1.189E-04	8.103E-05	1.15	298.	95.6	64.8	1.012E+03	516.
7.484E+03	1.166E-04	7.952E-05	1.15	298.	93.8	65.5	1.021E+03	494.
7.564E+03	1.145E-04	7.806E-05	1.15	298.	92.1	66.2	1.029E+03	471.
7.644E+03	1.124E-04	7.664E-05	1.15	298.	90.3	66.9	1.038E+03	445.
7.724E+03	1.104E-04	7.526E-05	1.15	298.	88.5	67.6	1.046E+03	418.
7.804E+03	1.084E-04	7.391E-05	1.15	298.	86.7	68.3	1.055E+03	387.
7.884E+03	1.065E-04	7.261E-05	1.15	298.	84.9	69.0	1.064E+03	352.
7.964E+03	1.046E-04	7.134E-05	1.15	298.	83.1	69.7	1.072E+03	312.
8.044E+03	1.028E-04	7.010E-05	1.15	298.	81.3	70.5	1.081E+03	262.
8.124E+03	1.011E-04	6.890E-05	1.15	298.	79.5.	71.2	1.089E+03	192.
8.204E+03	9.936E-05	6.773E-05	1.15	298.	77.6	71.9	1.098E+03	
8.284E+03	9.769E-05	6.660E-05	1.15	298.	75.8	72.6	1.106E+03	
8.364E+03	9.607E-05	6.549E-05	1.15	298.	74.0	73.3	1.114E+03	
8.444E+03	9.448E-05	6.441E-05	1.15	298.	72.1	74.0	1.123E+03	
8.524E+03	9.294E-05	6.336E-05	1.15	298.	70.2	74.7	1.131E+03	
8.604E+03	9.144E-05	6.234E-05	1.15	298.	68.3	75.4	1.140E+03	
8.684E+03	8.998E-05	6.134E-05	1.15	298.	66.5	76.1	1.148E+03	
8.759E+03	8.864E-05	6.043E-05	1.15	298.	64.7	76.8	1.156E+03	
8.839E+03	8.725E-05	5.948E-05	1.15	298.	62.8	77.5	1.164E+03	
8.919E+03	8.589E-05	5.855E-05	1.15	298.	60.9	78.2	1.172E+03	
8.999E+03	8.457E-05	5.765E-05	1.15	298.	58.9	78.9	1.180E+03	
9.079E+03	8.327E-05	5.677E-05	1.15	298.	57.0	79.6	1.189E+03	
9.159E+03	8.201E-05	5.591E-05	1.15	298.	55.1	80.3	1.197E+03	
9.239E+03	8.078E-05	5.507E-05	1.15	298.	53.2	81.0	1.205E+03	
9.319E+03	7.958E-05	5.425E-05	1.15	298.	51.2	81.7	1.213E+03	
9.399E+03	7.841E-05	5.345E-05	1.15	298.	49.3	82.4	1.221E+03	
9.479E+03	7.726E-05	5.267E-05	1.15	298.	47.3	83.1	1.229E+03	
9.559E+03	7.614E-05	5.190E-05	1.15	298.	45.3	83.8	1.237E+03	
9.639E+03	7.505E-05	5.116E-05	1.15	298.	43.4	84.5	1.245E+03	
9.719E+03	7.398E-05	5.043E-05	1.15	298.	41.4	85.2	1.253E+03	
9.799E+03	7.293E-05	4.972E-05	1.15	298.	39.4	85.9	1.261E+03	
9.879E+03	7.191E-05	4.902E-05	1.15	298.	37.4	86.6	1.269E+03	
9.959E+03	7.091E-05	4.834E-05	1.15	298.	35.4	87.3	1.277E+03	
1.004E+04	6.994E-05	4.767E-05	1.15	298.	33.4	88.0	1.285E+03	
1.011E+04	6.904E-05	4.706E-05	1.15	298.	31.5	88.7	1.293E+03	
1.018E+04	6.822E-05	4.650E-05	1.15	298.	29.8	89.3	1.300E+03	
1.025E+04	6.747E-05	4.600E-05	1.15	298.	28.2	89.9	1.306E+03	
1.031E+04	6.680E-05	4.553E-05	1.15	298.	26.6	90.4	1.312E+03	
1.036E+04	6.618E-05	4.512E-05	1.15	298.	25.3	90.9	1.318E+03	
1.042E+04	6.558E-05	4.470E-05	1.15	298.	23.9	91.4	1.323E+03	
1.047E+04	6.504E-05	4.434E-05	1.15	298.	22.6	91.8	1.328E+03	
1.052E+04	6.451E-05	4.397E-05	1.15	298.	21.3	92.3	1.333E+03	
1.056E+04	6.403E-05	4.365E-05	1.15	298.	20.2	92.7	1.337E+03	
1.060E+04	6.362E-05	4.336E-05	1.15	298.	19.2	93.0	1.341E+03	

Figure B.7. (continued)

1.064E+04	6.320E-05	4.308E-05	1.15	298.	18.1	93.4	1.345E+03
1.068E+04	6.279E-05	4.280E-05	1.15	298.	17.1	93.7	1.349E+03
1.072E+04	6.244E-05	4.256E-05	1.15	298.	16.2	94.0	1.352E+03
1.075E+04	6.209E-05	4.232E-05	1.15	298.	15.3	94.3	1.356E+03
1.078E+04	6.179E-05	4.212E-05	1.15	298.	14.6	94.6	1.359E+03
1.081E+04	6.150E-05	4.192E-05	1.15	298.	13.8	94.9	1.362E+03
1.084E+04	6.120E-05	4.172E-05	1.15	298.	13.0	95.1	1.365E+03
1.087E+04	6.096E-05	4.155E-05	1.15	298.	12.4	95.3	1.367E+03
1.089E+04	6.072E-05	4.139E-05	1.15	298.	11.7	95.6	1.369E+03
1.092E+04	6.048E-05	4.123E-05	1.15	298.	11.1	95.8	1.372E+03
1.094E+04	6.029E-05	4.110E-05	1.15	298.	10.6	96.0	1.374E+03
1.096E+04	6.010E-05	4.097E-05	1.15	298.	10.1	96.1	1.376E+03
1.098E+04	5.991E-05	4.084E-05	1.15	298.	9.55	96.3	1.378E+03
1.100E+04	5.972E-05	4.071E-05	1.15	298.	9.03	96.5	1.380E+03
1.101E+04	5.958E-05	4.061E-05	1.15	298.	8.64	96.6	1.381E+03
1.103E+04	5.944E-05	4.052E-05	1.15	298.	8.26	96.8	1.382E+03
1.104E+04	5.930E-05	4.042E-05	1.15	298.	7.87	96.9	1.384E+03
1.106E+04	5.916E-05	4.033E-05	1.15	298.	7.48	97.0	1.385E+03
1.107E+04	5.903E-05	4.023E-05	1.15	298.	7.10	97.2	1.387E+03
1.109E+04	5.889E-05	4.014E-05	1.15	298.	6.71	97.3	1.388E+03
1.110E+04	5.880E-05	4.008E-05	1.15	298.	6.45	97.4	1.389E+03
1.111E+04	5.871E-05	4.002E-05	1.15	298.	6.19	97.5	1.390E+03
1.112E+04	5.861E-05	3.995E-05	1.15	298.	5.93	97.5	1.391E+03
1.113E+04	5.852E-05	3.989E-05	1.15	298.	5.68	97.6	1.392E+03
1.114E+04	5.843E-05	3.983E-05	1.15	298.	5.42	97.7	1.393E+03
1.115E+04	5.834E-05	3.977E-05	1.15	298.	5.16	97.8	1.394E+03
1.116E+04	5.825E-05	3.971E-05	1.15	298.	4.90	97.9	1.395E+03
1.117E+04	5.816E-05	3.965E-05	1.15	298.	4.64	98.0	1.396E+03
1.118E+04	5.807E-05	3.959E-05	1.15	298.	4.38	98.1	1.397E+03
1.119E+04	5.798E-05	3.952E-05	1.15	298.	4.12	98.2	1.398E+03
1.119E+04	5.794E-05	3.949E-05	1.15	298.	3.99	98.2	1.398E+03
1.120E+04	5.785E-05	3.943E-05	1.15	298.	3.73	98.3	1.399E+03
1.121E+04	5.781E-05	3.940E-05	1.15	298.	3.60	98.3	1.400E+03
1.122E+04	5.772E-05	3.934E-05	1.15	298.	3.35	98.4	1.401E+03
1.122E+04	5.767E-05	3.931E-05	1.15	298.	3.22	98.5	1.401E+03
1.123E+04	5.758E-05	3.925E-05	1.15	298.	2.96	98.6	1.402E+03
1.124E+04	5.754E-05	3.922E-05	1.15	298.	2.83	98.6	1.403E+03
1.125E+04	5.745E-05	3.916E-05	1.15	298.	2.57	98.7	1.404E+03
1.125E+04	5.741E-05	3.913E-05	1.15	298.	2.44	98.7	1.404E+03
1.126E+04	5.732E-05	3.907E-05	1.15	298.	2.18	98.8	1.405E+03
1.127E+04	5.728E-05	3.904E-05	1.15	298.	2.05	98.9	1.406E+03
1.127E+04	5.723E-05	3.901E-05	1.15	298.	1.92	98.9	1.406E+03
1.128E+04	5.719E-05	3.898E-05	1.15	298.	1.79	99.0	1.407E+03
1.128E+04	5.714E-05	3.895E-05	1.15	298.	1.66	99.0	1.407E+03
1.129E+04	5.710E-05	3.892E-05	1.15	298.	1.53	99.0	1.407E+03
1.129E+04	5.706E-05	3.889E-05	1.15	298.	1.40	99.1	1.408E+03
1.130E+04	5.701E-05	3.886E-05	1.15	298.	1.27	99.1	1.408E+03
1.130E+04	5.697E-05	3.883E-05	1.15	298.	1.14	99.2	1.409E+03
1.131E+04	5.693E-05	3.880E-05	1.15	298.	1.01	99.2	1.409E+03
1.131E+04	5.688E-05	3.877E-05	1.15	298.	0.880	99.3	1.410E+03
1.132E+04	5.684E-05	3.875E-05	1.15	298.	0.750	99.3	1.410E+03
1.132E+04	5.680E-05	3.872E-05	1.15	298.	0.620	99.3	1.411E+03
1.133E+04	5.675E-05	3.869E-05	1.15	298.	0.490	99.4	1.411E+03
1.133E+04	5.671E-05	3.866E-05	1.15	298.	0.360	99.4	1.412E+03
1.134E+04	5.667E-05	3.863E-05	1.15	298.	0.230	99.5	1.412E+03
1.134E+04	5.662E-05	3.860E-05	1.15	298.	0.100	99.5	1.413E+03
1.135E+04	5.658E-05	3.857E-05	1.15	298.	0.000E+00	99.6	1.412E+03
1.142E+04	5.570E-05	3.797E-05	1.15	298.	0.000E+00	101.	1.420E+03
1.150E+04	5.474E-05	3.732E-05	1.15	298.	0.000E+00	102.	1.427E+03
1.158E+04	5.381E-05	3.668E-05	1.15	298.	0.000E+00	103.	1.435E+03

Figure B.7. (continued)

1.166E+04	5.290E-05	3.606E-05	1.15	298.	0.000E+00	104.	1.443E+03
1.174E+04	5.202E-05	3.546E-05	1.15	298.	0.000E+00	105.	1.450E+03
1.182E+04	5.115E-05	3.487E-05	1.15	298.	0.000E+00	106.	1.458E+03
1.190E+04	5.031E-05	3.430E-05	1.15	298.	0.000E+00	107.	1.466E+03
1.194E+04	4.995E-05	3.405E-05	1.15	298.	0.000E+00	107.	1.469E+03

For the UFL of 0.10000 mole percent, and the LFL of 1.00000E-02 mole percent:

The mass of contaminant between the UFL and LFL is: 40143. kg.
The mass of contaminant above the LFL is: 42885. kg.

Figure B.7. (concluded)

Burro Test LNG Releases

In 1980, the U.S. Department of Energy sponsored at China Lake, California, the BURRO series of LNG releases (Koopman, et al., 1982). The Burro tests were ground-level releases; their simulation does not require use of the JETPLU model. Burro 9 (Table B.3) was modeled here, both as a steady-state and transient (time-limited) release, using the DEGADIS model with interactive input via DEGADISIN.

Table B.3. Burro 9 Test Conditions

Source rate	130.0 kg/s
Source radius	22.06 m
Wind speed	6.5 m/s at 8.0 m
Atmospheric stability	C (Pasquill)
Monin-Obukhov length	-140. m
Surface roughness	2.05×10^{-4} m
Air temperature	33.4 C
Atmospheric humidity	12.5 %
Surface temperature	310 K

The source radius assumes LNG release onto water, with a constant evaporative flux of 0.085 kg/m² s. The input procedures for simulation of the transient release (RUN_NAME=BURRO9) and the steady-state release (RUN_NAME=BURRO9S) are very similar. Therefore, only the specification of the surface rate and extent have been included for the transient release. In the following line-by-line description of the input procedure:

- (*) A line terminator (normally a carriage return) must end every line entered by the user.
- (*) The file name specification RUN_NAME must satisfy system restrictions.
- (*) When DEGADISIN requests the user to choose an option, responses are a single character (capital or lower case). The default responses are denoted by a capital letter inside angle brackets (e.g. <N>). When applicable, a menu of responses is included inside parentheses.
- (*) For numerical responses, a comma, space, tab, or line terminator (carriage return) may separate the numbers.
- (*) When a file is used as input (i.e. for the density or transient source input), DEGADISIN reads the same information from the file which would be entered at the terminal in the same order and in the same format.

Notes on Steady-State Simulation of BURRO9

- (1) Begin the input procedure by execution of DEGADISIN.
- (2) The file name specification must follow system restrictions. The DEGADIS model uses this file name along with various file extensions for input and output.
- (3) The Title Block is used to carry any desired comments such as information on the specification of certain parameters.
- (4) The wind field parameters include the wind velocity (m/s) at a specified height (m) and the surface roughness (m).
- (5) The Pasquill stability class is used to generate estimates of other atmospheric parameters which follow.
- (6) The averaging time is used to determine the value of δ_y (DELTAY) in the lateral dispersion coefficient specification. Changes to the values of δ_y are calculated assuming that the effect of averaging time only influences the lateral plume meander of a steady-state release.
- (7) The current settings of pertinent atmospheric parameters are displayed in this list. If any of these are to be changed, the first letter of the parameter to be changed is entered. Note that the default--indicated by <ND>--is No for no changes.
- (8) The Monin-Obukhov length (Length in the list) is to be changed, so L is entered in response to the prompt.
- (9) The list is redisplayed to verify the change and to request any further changes. The (default) response of No causes the program to go to the next question.
- (10) The ambient temperature and pressure are entered.
- (11) DEGADISIN calculates the ambient air density for the given input parameters.
- (12) If the release is isothermal, respond "Y". A positive response causes DEGADISIN to ask for a list of concentration, density, and mole fraction points for the gas mixture. The default response is negative.
- (13) If the release is simulated as adiabatic, the default negative response is chosen. For inclusion of heat transfer effects, the surface temperature and the method of calculating the heat transfer coefficient must be specified.
- (14) Water transfer to the source blanket (if present) can be included in the calculation.

- (1) RUN SYSSDEGADIS:DEGADISIN
DEnse GAs DISpersion Model input module.
- (2) Enter the simulation name : [DIR]RUNNAME BURRO9S
INPUT MODULE -- DEGADIS MODEL

Enter Title Block -- up to 4 lines of 80 characters
To stop, type "/"
Steady-state simulation of BURRO 9
//
- (3) ENTER WIND PARAMETERS -- U0 (m/s), Z0 (m), and ZR(m)
U0 -- Wind velocity at reference height Z0
ZR -- Surface Roughness
6.5,8.,2.05E-4
- (4) Enter the Pasquill stability class: (A,B,C,D,E,F) <D> C
Enter the averaging time (s) for estimating DELTAY: 0.
- (5)
- (6)
- (7) The values for the atmospheric parameters are set as follows:
DELTAY: 0.1046
BETAY: 0.9000
Monin-Obukhov length: -9.3344 m
Sigma X Coefficient: 0.0200
Sigma X Power: 1.2200
Sigma X Minimum Distance: 130.0000 m
Do you wish to change any of these?
(No,Delтай,Bетай,Length,Coefficient,Power,Minimum) <N> L
Note: For infinity, RML = 0.0
Enter the desired Monin-Obukhov length: (m) -140.
- (8)
- (9) The values for the atmospheric parameters are set as follows:
DELTAY: 0.1046
BETAY: 0.9000
Monin-Obukhov length: -140.0000 m
Sigma X Coefficient: 0.0200
Sigma X Power: 1.2200
Sigma X Minimum Distance: 130.0000 m
Do you wish to change any of these?
(No,Delтай,Bетай,Length,Coefficient,Power,Minimum) <N>
- (10) Enter the ambient temperature(K) and pressure(atm): 305.88,0.94
The ambient humidity can be entered as Relative or Absolute.
Enter either R or A <R or a>
Enter the relative humidity (%): 12.5
- (11) Ambient Air density is 1.081886 kg/m**3
- (12) Is this an Isothermal spill? <y or N>
- (13) Is heat transfer to be included in the calculations <y or N> Y
Enter the surface temperature [=] K : 310.
Do you want to use the built in correlation, the LLNL correlation, or
enter a particular value?
(Corr,LLNLcorr,Value) <C>
- (14) Is water transfer to be included in the source <y or N>

- (15) Enter the three-letter designation of the diffusing gas. The properties of LNG as methane, LPG as propane, and NH₃ (ammonia) are included among others.
- (16) A list of the properties for the specified gas (if available) is given. If any of the parameters are to be changed, the first letter of the parameter to be changed in the list is given to the prompt. Here, the level at which the flammability contours are calculated is changed from 0.5 m to 1.0 m.
- (17) The gas property list is displayed again. The default response is no change.
- (18) The lowest concentration of interest is the concentration at which the calculations are stopped.
- (19) The BURRO9 case is for a release of "pure" (undiluted) LNG. For diluted releases, DEGADIS requires the mass fraction of the contaminant and the mixture temperature.
- (20) If a steady-state release is to be simulated, type "Y" to the prompt. For a steady simulation, the steady-state mass evolution rate (kg/s) and primary source extent (m) are required.
- (21) A note about the numerical parameter files is included. These files contain various constant values used in the programs to which the user has access without recompiling the programs. Access is granted as a convenience.
- (22) DEGADISIN will generate a command procedure suitable for running the model under VMS.
- (23) If so desired, DEGADISIN will initiate the command procedure under VMS. If not, the program returns to the operating system.

- (15) Enter the code name of the diffusing species: LNG
- (16) The characteristics for the gas are set as follows:
 Molecular weight: 16.04
 Storage temperature [K]: 111.70
 Density at storage temperature, PAMB [kg/m**3]: 1.6845
 Mean Heat capacity constant 5.60000E-08
 Mean Heat capacity power 5.0000
 Upper Flammability Limit [mole frac] 0.15000
 Lower Flammability Limit [mole frac] 5.00000E-02
 Height of Flammability Limit [m] 0.50000
 Do you wish to change any of these? (No,Mole,Temp,Den,Heat,Power,Upper,Lower,Z)
 <N> Z
 Enter the desired Height for the flammable limit calculations: 1.
- (17) The characteristics for the gas are set as follows:
 Molecular weight: 16.04
 Storage temperature [K]: 111.70
 Density at storage temperature, PAMB [kg/m**3]: 1.6845
 Mean Heat capacity constant 5.60000E-08
 Mean Heat capacity power 5.0000
 Upper Flammability Limit [mole frac] 0.15000
 Lower Flammability Limit [mole frac] 5.00000E-02
 Height of Flammability Limit [m] 1.00000
 Do you wish to change any of these? (No,Mole,Temp,Den,Heat,Power,Upper,Lower,Z)
 <N>
- (18) The suggested LOWEST CONCENTRATION OF INTEREST (gas_lfl/2.)
 is 1.50191E-02 kg/m**3. Enter the desired value: 0.015
- Specification of source parameters.
- (19) Is this a release of pure (P) or diluted (d) material specified above? <P or d>
- (20) Is this a Steady state simulation? <y or N> Y
 Enter the desired evolution rate [=] kg/sec : 130.
 Enter the desired source radius [=] m : 22.06
- (21) In addition to the information just obtained, DEGADIS
 requires a series of numerical parameter files which use
 the same name as [DIR]RUNNAME given above.
 For convenience, example parameter files are included for
 each step. They are:
 EXAMPLE.ER1 and
 EXAMPLE.ER2
 Note that each of these files can be edited during the course of the
 simulation if a parameter proves to be out of specification.
- (22) Do you want a command file to be generated to execute the procedure? <Y or n>
 The command file will be generated under the file name:
 BURROSS.com
- (23) Do you wish to initiate this procedure? <y or N>
 \$

Notes on Transient Simulation of BURRO9

Beginning with the specification of the source rate and extent, the responses to all of the previous questions except the simulation name (RUN_NAME) are the same for the steady-state case and are not repeated.

- (19) The BURRO9 case is for a release of "pure" (undiluted) LNG. For diluted releases, DEGADIS requires the mass fraction of the contaminant and the mixture temperature.
- (20) The default response is for a transient release.
- (21) An initial mass of gas can be specified over the source. This can be used to model aboveground releases such as the Thorney Island Trials.
- (22) The transient source description consists of ordered triples of time, evolution rate, and source radius for pure sources. For diluted sources, values of evolution rate, source radius, contaminant mass fraction, and source temperature must be specified as functions of time.
- (23) An input file can be used to enter the data triples to avoid typing errors or to use as output from another model such as a liquid spreading model. The file format is the same as the terminal entry format.
- (24) The first item is the number of triples used in the description followed by the triples with the last two values showing no gas present.
- (25) A note about the numerical parameter files is included. These files contain various constant values used in the programs to which the user has access without recompiling the programs. Access is granted as a convenience.
- (26) DEGADISIN will generate a command procedure suitable for running the model under VMS.
- (27) If so desired, DEGADISIN will initiate the procedure under VMS. If not, the program returns to the operating system.

- (19) Specification of source parameters.
- (20) Is this a release of pure (P) or diluted (d) material specified above? <P or d>
- (21) Is this a Steady state simulation? <y or N>
- Enter the initial mass of pure gas over the source. (kg)
(Positive or zero): 0.
- (22) Source Description
- The description of the primary source mass evolution rate E and radius R1 for a transient release is input by ordered triples as follows:
- ```

first point -- t=0, E(t=0), R1(t=0) (initial, nonzero values)
second point -- t=t1, E(t=t1), R1(t=t1)
.
.
.
last nonzero point -- t=TEND, E(t=TEND), R1(t=TEND)
next to last point -- t=TEND+1., E=0., R1=0.
last point -- t=TEND+2., E=0., R1=0.

```
- Note: the final time (TEND) is the last time when E and R1 are non-zero.
- (23) Do you have an input file for the Source Description? [y or N]
- Enter the number of triples (max= 30) starting with t=0. and ending with t=TEND+2. for the source description: 4
- (24) Enter TIME (sec), EVOLUTION RATE (kg/s), and POOL RADIUS (m)
- ```

0.,130.,22.06
80.,130.,22.06
81.,0.,0.
82.,0.,0.

```
- (25) In addition to the information just obtained, DEGADIS requires a series of numerical parameter files which use the same name as [DIR]RUNNAME given above.
- For convenience, example parameter files are included for each step. They are:
EXAMPLE.ER1,
EXAMPLE.ER2, and
EXAMPLE.ER3
- Note that each of these files can be edited during the course of the simulation if a parameter proves to be out of specification.
- (26) Do you want a command file to be generated to execute the procedure? <Y or n>
The command file will be generated under the file name:
BURRO9.com
- (27) Do you wish to initiate this procedure? <y or N>
\$

The INP files for BURR09S and BURR09 are shown in Figures B.8 and B.9. If necessary, the user may edit the INP file before beginning the simulation.

Example Simulation Output

BURR09.LIS and BURR09S.LIS contain the output listing for the transient and steady-state releases, respectively. A discussion of the steady-state and transient simulation listings follows. Because of the similarities between the steady-state and transient simulation listings, the first portion of the transient simulation is not included.

Steady-state simulation of BURRO 9

```

6.500000      8.000000      2.0500000E-04
3
0.0000000E+00 0.0000000E+00
0.1046050     0.9000000     -140.0000
2.0000000E-02  1.220000     130.0000
305.8800      0.9400000    4.3809577E-03   12.50000
0   310.0000
1   0.0000000E+00
0   0.0000000E+00
LNG
16.04000     111.7000     1.684480
5.6000000E-08 5.000000
0.1500000    5.0000000E-02  1.000000
1.5000000E-02
0.0000000E+00
4
0.0000000E+00 130.0000    22.06000    1.000000    111.7000    1.000000
6023.000     130.0000    22.06000    1.000000    111.7000    1.000000
6024.000     0.0000000E+00 0.0000000E+00    1.000000    111.7000    1.000000
6025.000     0.0000000E+00 0.0000000E+00    1.000000    111.7000    1.000000
F F F T F
9-JUN-1989 15:55:42.38
130.0000     44.12000    17.32588

```

Figure B.8. BURRO9S.INP Listing.

Time-limited simulation of Burro 9

```

6.500000      8.000000      2.0500000E-04
3
0.0000000E+00 0.0000000E+00
0.1046050     0.9000000     -140.0000
2.0000000E-02  1.220000     130.0000
305.8800      0.9400000    4.3809577E-03   12.50000
0   310.0000
1   0.0000000E+00
0   0.0000000E+00
LNG
16.04000     111.7000     1.684480
5.6000000E-08 5.000000
0.1500000    5.0000000E-02  1.000000
1.5000000E-02
0.0000000E+00
4
0.0000000E+00 130.0000    22.06000    1.000000    111.7000    1.000000
80.00000     130.0000    22.06000    1.000000    111.7000    1.000000
81.00000     0.0000000E+00 0.0000000E+00    1.000000    111.7000    1.000000
82.00000     0.0000000E+00 0.0000000E+00    1.000000    111.7000    1.000000
F F F F F
9-JUN-1989 16:02:24.47

```

Figure B.9. BURRO9.INP Listing.

Notes on Steady-State Simulation of BURRO9

- (1) The date and time DEGADISIN was run are included.
- (2) The input information gathered by DEGADISIN is repeated to assist in documentation of the simulations. Included here are the Title Block and the atmospheric conditions.

***** U O A _ D E G A D I S M O D E L O U T P U T -- V E R S I O N 2.1 *****
 ***** 6-SEP-1989 16:05:07.54 *****

1 Data input on 9-JUN-1989 15:55:42.38
 Source program run on 6-SEP-1989 16:05:07.54

2) TITLE BLOCK

Steady-state simulation of BURRO 9

Wind velocity at reference height	6.50	m/s
Reference height	8.00	m
Surface roughness length	2.050E-04	m
Pasquill Stability class	C	
Monin-Obukhov length	-140.	m
Gaussian distribution constants		
Specified averaging time	0.00	s
Delta	0.10461	
Beta	0.90000	
Wind velocity power law constant	Alpha	0.10638
Friction velocity		0.21878 m/s
Ambient Temperature		305.88 K
Surface Temperature		310.00 K
Ambient Pressure		0.940 atm
Ambient Absolute Humidity		4.381E-03 kg/kg BDA
Ambient Relative Humidity		12.50 %

Adiabatic Mixing:	Mole fraction	CONCENTRATION OF C kg/m**3	GAS DENSITY kg/m**3	Enthalpy J/kg	Temperature K
	0.00000	0.00000	1.08189	0.00000E+00	305.88
	0.00897	0.00542	1.08476	-2018.3	303.89
	0.01786	0.01088	1.08771	-4036.6	301.93
	0.02669	0.01636	1.09065	-6054.8	299.98
	0.04413	0.02741	1.09652	-10091.	296.16
	0.06131	0.03858	1.10235	-14128.	292.41
	0.08657	0.05555	1.11110	-20183.	286.93
	0.11126	0.07278	1.11976	-26238.	281.62
	0.14331	0.09616	1.13126	-34311.	274.78
			*		
			*		
			*		
	0.74203	0.86864	1.41242	-2.48249E+05	160.04
	0.79281	0.98925	1.45478	-2.74487E+05	150.27
	0.84028	1.11626	1.49833	-3.00724E+05	141.26
	0.88805	1.26067	1.54684	-3.28980E+05	132.29
	0.93269	1.41351	1.59719	-3.57236E+05	124.01
	0.97738	1.58729	1.65343	-3.87511E+05	115.81
	1.00000	1.68448	1.68448	-4.03657E+05	111.70

- (3) Continuing with the input information, the contaminant gas properties are output.
- (4) The specification of the mass evolution rate, source radius, contaminant mass fraction, and source temperature are output. For a steady-state release, there is no initial mass in the cloud, and the source parameters are held constant for an arbitrarily large period of time.
- (5) Finally, certain numerical parameters and calculation flags are displayed. Some of these are set in DEGADISIN while others are set in the numerical parameter files.
- (6) A summary of the calculated secondary source parameters is included. The secondary source gas radius and height are output as functions of time along with other secondary parameters including the source mass flux (Qstar), the vertical concentration distribution parameter at the downwind edge ($SZ(x - L/2.)$), the contaminant mole fraction (Mole frac C), the gas mixture density (Density), and the Richardson number based on the cloud spreading velocity (Rich No.).
- (7) For a steady-state release, the source calculations are terminated after the calculated parameters are no longer changing as a function of time. A summary of the steady-state secondary source is included.
- (8) The downwind portion of the calculations is included. The distance downwind of the source is given in the first column. Columns 2 through 6 contain the mole fraction, contaminant concentration, mixture density, ratio of $(\rho - \rho_a)/c_c$ (Gamma), and mixture temperature on the centerline of the gas cloud at ground level. Columns 7 through 9 contain the contour shape parameters b (Half Width), S_z , and S_y . Finally, columns 10 and 11 contain the width from the centerline to the indicated concentration levels at the indicated height. Note that the output is prematurely terminated. Output actually continues until the centerline, ground-level concentration drops below the lowest concentration of interest.

3

Specified Gas Properties:

Molecular weight:	16.040
Storage temperature:	111.70 K
Density at storage temperature and ambient pressure:	1.6845 kg/m**3
Mean heat capacity constant:	5.60000E-08
Mean heat capacity power:	5.0000
Upper mole fraction contour:	0.15000
Lower mole fraction contour:	5.00000E-02
Height for isopleths:	1.0000 m

4

Source input data points

Initial mass in cloud: 0.00000E+00

Time s	Contaminant Mass Rate kg/s	Source Radius m	Contaminant Mass Fraction kg contam/kg mix	Temperature K	Enthalpy J/kg
0.00000E+00	130.00	22.060	1.0000	111.70	-4.03657E+05
6023.0	130.00	22.060	1.0000	111.70	-4.03657E+05
6024.0	0.00000E+00	0.00000E+00	1.0000	111.70	-4.03657E+05
6025.0	0.00000E+00	0.00000E+00	1.0000	111.70	-4.03657E+05

5

Calculation procedure for ALPHA: 1

Entrainment prescription for PHI: 3

Layer thickness ratio used for average depth: 2.1500

Air entrainment coefficient used: 0.590

Gravity slumping velocity coefficient used: 1.150

NON Isothermal calculation

Heat transfer calculated with correlation: 1

Water transfer not included

CALCULATED SOURCE PARAMETERS

6

Time sec	Gas Radius m	Height m	Qstar kg/m**2/s	SZ(x=L/2.) m	Mole frac C	Density kg/m**3	Temperature K	Rich No.
0.000000E+00	22.0600	1.100000E-05	8.329550E-02	0.498870	1.00000	1.68448	111.700	0.756144
1.93242	22.1998	1.304959E-03	8.367377E-02	0.508692	0.999937	1.66670	112.897	0.756144
7.25031	22.2263	1.331373E-03	8.375806E-02	0.510725	0.999935	1.66315	113.139	0.756144

7

Source strength [kg/s] : 130.00 Equivalent Primary source radius [m] : 22.060

Equivalent Primary source length [m] : 44.120 Equivalent Primary source half-width [m] : 17.326

Secondary source concentration [kg/m**3] : 1.6630 Secondary source SZ [m] : 0.51072

Contaminant flux rate: 8.37642E-02

Secondary source mass fractions... contaminant: 0.999884 air: 1.15782E-04
Enthalpy: -4.00663E+05 Density: 1.6631

Secondary source length [m] : 44.453 Secondary source half-width [m] : 17.457

8

Distance (m)	Mole Fraction	Concentration (kg/m**3)	Density (kg/m**3)	Gamma	Temperature (K)	Half Width (m)	Sz (m)	Sy (m)	Width at z= 5.00 mole%	1.00 m to 15.0 m
22.2	1.00	1.66	1.6631	0.350	113.	17.5	0.511	0.000E+00	17.5	17.5
22.5	0.998	1.65	1.6570	0.348	114.	16.7	0.510	0.948	18.0	17.5
25.2	0.980	1.58	1.6061	0.332	119.	15.9	0.504	3.05	20.0	18.3

•

•

•

Notes on Transient Simulation Output of BURRO9

- (6) A summary of the calculated secondary source parameters is included. The secondary source gas radius and height are output as functions of time along with other secondary parameters including the source mass flux (Qstar), the vertical concentration distribution parameter at the downwind edge ($SZ(x = L/2.)$), the contaminant mole fraction (Mole frac C), the gas mixture density (Density), and the Richardson number based on the cloud spreading velocity (Rich No.). Note that only a portion of the output is shown. The source calculation ends when all the primary and secondary source gas has been taken up in the atmospheric flow.

6

CALCULATED SOURCE PARAMETERS								
Time sec	Gas Radius m	Height m	Qstar kg/m**2/s	SZ(x=L/2.) m	Mole frac C	Density kg/m**3	Temperature K	Rich No.
0.000000E+00	22.0600	1.100000E-05	8.329550E-02	0.498870	1.00000	1.68448	111.700	0.756144
7.812500E-05	22.0600	1.108054E-05	8.329550E-02	0.498870	1.00000	1.68448	111.700	0.756144
1.562500E-04	22.0600	1.116108E-05	8.329550E-02	0.498870	1.00000	1.68448	111.700	0.756144
2.995998E-04	22.0600	1.130885E-05	8.329549E-02	0.498870	1.00000	1.68448	111.700	0.756144
4.429495E-04	22.0600	1.145663E-05	8.329549E-02	0.498870	1.00000	1.68448	111.700	0.756144
6.200244E-04	22.0600	1.163917E-05	8.329548E-02	0.498870	1.00000	1.68448	111.700	0.756144
7.970994E-04	22.0600	1.182171E-05	8.329548E-02	0.498870	1.00000	1.68448	111.700	0.756144
1.302401E-03	22.0600	1.234260E-05	8.329547E-02	0.498870	1.00000	1.68448	111.700	0.756144
5.584851E-03	22.0601	1.675684E-05	8.329537E-02	0.498870	1.00000	1.68448	111.700	0.756144
•								
•								
•								
9.775356E-02	22.0620	1.112523E-04	8.330184E-02	0.499023	0.999998	1.68419	111.719	0.756144
0.134816	22.0631	1.488579E-04	8.330547E-02	0.499110	0.999998	1.68403	111.730	0.756144
0.184839	22.0649	1.991522E-04	8.331114E-02	0.499247	0.999995	1.68377	111.747	0.756144
0.251490	22.0676	2.652104E-04	8.331988E-02	0.499457	0.999994	1.68338	111.774	0.756144
0.355980	22.0726	3.662174E-04	8.333587E-02	0.499841	0.999991	1.68266	111.822	0.756144
0.523614	22.0821	5.206466E-04	8.336356E-02	0.500537	0.999984	1.68137	111.908	0.756144
1.15444	22.1268	9.974017E-04	8.348547E-02	0.503776	0.999959	1.67548	112.304	0.756144
1.47042	22.1565	1.162954E-03	8.355603E-02	0.505664	0.999948	1.67209	112.532	0.756144
1.79110	22.1863	1.274689E-03	8.363808E-02	0.507764	0.999940	1.66834	112.786	0.756144
2.07986	22.2139	1.324084E-03	8.371218E-02	0.509677	0.999936	1.66496	113.015	0.756144
36.8751	22.2250	1.345084E-03	8.369677E-02	0.509883	0.999934	1.66475	113.030	0.756144
79.9765	22.2283	1.370607E-03	8.375954E-02	0.510787	0.999933	1.66299	113.149	0.756144
80.0499	21.8594	1.392848E-03	8.688047E-02	0.535701	0.999932	1.60926	116.927	0.756144
80.0918	21.1203	1.375235E-03	9.024113E-02	0.548069	0.999933	1.57472	119.492	0.756144
80.3716	15.7908	1.086961E-03	0.110060	0.530173	0.999954	1.48958	126.319	0.756144
80.6057	11.3499	8.196970E-04	0.132618	0.486207	0.999969	1.41975	132.531	0.756144
80.7910	7.57905	5.692027E-04	0.157312	0.408026	0.999979	1.36527	137.818	0.756144
80.8842	5.36474	4.112889E-04	0.174600	0.334241	0.999984	1.33743	140.686	0.756144
80.9319	3.99387	3.084351E-04	0.186437	0.273428	0.999987	1.32770	141.717	0.756144
80.9590	3.04157	2.356705E-04	0.195637	0.223938	0.999990	1.32331	142.187	0.756144
80.9750	2.34519	1.820257E-04	0.203107	0.183421	0.999991	1.32103	142.432	0.756144
80.9846	1.82960	1.421910E-04	0.209143	0.150660	0.999992	1.31932	142.617	0.756144
80.9904	1.44346	1.123626E-04	0.213940	0.124304	0.999993	1.31719	142.848	0.756144
80.9939	1.15560	9.016800E-05	0.217575	0.103447	0.999994	1.31406	143.188	0.756144
80.9964	0.900469	7.055920E-05	0.220619	8.393002E-02	0.999994	1.30850	143.796	0.756144
80.9978	0.714082	5.630183E-05	0.222380	6.893466E-02	0.999995	1.30040	144.691	0.756144
80.9986	0.582421	4.629137E-05	0.222961	5.789226E-02	0.999995	1.28998	145.861	0.756144
80.9992	0.462480	3.725033E-05	0.222315	4.744542E-02	0.999995	1.27292	147.816	0.756144
80.9995	0.384735	3.146655E-05	0.220537	4.044054E-02	0.999995	1.25354	150.101	0.756144
80.9998	0.308874	2.594198E-05	0.216437	3.339515E-02	0.999995	1.22064	154.147	0.756144
80.9999	0.251972	2.196174E-05	0.209942	2.794875E-02	0.999995	1.17617	159.974	0.756144
81.0000	0.217901	1.971995E-05	0.203099	2.460883E-02	0.999995	1.13276	166.105	0.756144
81.0000	0.199541	1.859155E-05	0.197788	2.278038E-02	0.999995	1.10028	171.009	0.756144

•

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•

- (7) An indication of the constants used in the x-direction dispersion correction is included. The output will also indicate if the x-direction dispersion correction was not applied.
- (8) The concentration field is shown for different times after the beginning of the spill. Unless otherwise specified, DEGADIS will choose default values for the time to output the concentration field. If other times are desired, DEGADIS3 can be executed again after the appropriate changes to RUN_NAME.ER3 have been made.
- (9) The downwind portion of the calculations is included. The distance downwind of the source is given in the first column. Columns 2 through 6 contain the mole fraction, contaminant concentration, mixture density, ratio of $(\rho - \rho_a)/c_c$ (Gamma), and mixture temperature on the centerline of the gas cloud at ground level. Columns 7 through 9 contain the contour shape parameters b (Half Width), S_z, and S_y. Finally, columns 10 and 11 contain the width from the centerline to the indicated concentration levels at the indicated height. Note that the output is prematurely terminated.

Sorted values for each specified time.

7

X-Direction correction was applied.

Coefficient: 2.00000E-02
 Power: 1.2200
 Minimum Distance: 130.00 m

8

Time after beginning of spill 12.00000 sec

9

Distance (m)	Mole Fraction	Concentration (kg/m**3)	Density (kg/m**3)	Gamma	Temperature (K)	Half Width (m)	Sz	Sy	Width at z=		1.00 m to: 15.0 mole%
									5.00 mole%	(m)	
31.2	1.03	1.29	1.2269	0.113	166.	14.8	0.572	5.37	22.2	(m)	19.4
44.8	0.757	0.804	1.2680	0.232	182.	15.4	0.589	8.97	26.3	(m)	20.4
58.6	0.500	0.425	1.1880	0.250	222.	14.0	0.602	11.3	24.5	(m)	
72.7	0.152	0.100	1.1046	0.227	281.	6.89	0.777	11.3			

For the UFL of 15.000 mole percent, and the LFL of 5.0000 mole percent:

The mass of contaminant between the UFL and LFL is: 89.846 kg.
 The mass of contaminant above the LFL is: 641.91 kg.

Time after beginning of spill 23.00000 sec

Distance (m)	Mole Fraction	Concentration (kg/m**3)	Density (kg/m**3)	Gamma	Temperature (K)	Half Width (m)	Sz	Sy	Width at z=		1.00 m to: 15.0 mole%
									5.00 mole%	(m)	
29.0	1.06	1.38	1.2314	0.108	163.	14.8	0.568	4.57	21.1	(m)	18.8
42.6	0.888	1.05	1.2857	0.195	168.	14.7	0.630	8.28	26.0	(m)	21.6
56.4	0.759	0.842	1.3227	0.286	172.	16.0	0.662	11.2	30.6	(m)	24.4
70.4	0.684	0.686	1.2564	0.254	190.	17.5	0.706	13.7	35.0	(m)	26.8
84.7	0.596	0.543	1.2040	0.225	209.	18.7	0.777	16.2	38.9	(m)	28.8
99.1	0.429	0.337	1.1457	0.189	239.	18.6	0.896	18.6	39.5	(m)	24.6
114.	0.235	0.163	1.1145	0.201	268.	14.7	1.10	19.6	31.8		
129.	6.338E-02	3.939E-02	1.0875	0.143	296.	5.46	1.47	17.7			

For the UFL of 15.000 mole percent, and the LFL of 5.0000 mole percent:

The mass of contaminant between the UFL and LFL is: 359.47 kg.
 The mass of contaminant above the LFL is: 2105.8 kg.

Time after beginning of spill 34.00000 sec

Distance (m)	Mole Fraction	Concentration (kg/m**3)	Density (kg/m**3)	Gamma	Temperature (K)	Half Width (m)	Sz	Sy	Width at z=		1.00 m to: 15.0 mole%
									5.00 mole%	(m)	
26.9	1.10	1.45	1.2190	9.445E-02	162.	15.0	0.553	3.80	20.3	(m)	18.3
40.4	0.912	1.09	1.2781	0.180	167.	14.6	0.623	7.77	25.2	(m)	21.2
54.1	0.778	0.871	1.3185	0.272	171.	15.7	0.657	10.7	29.9	(m)	23.9
68.1	0.697	0.710	1.2658	0.259	187.	17.3	0.695	13.3	34.4	(m)	26.5
82.4	0.620	0.576	1.2130	0.228	205.	18.5	0.760	15.8	38.3	(m)	28.6
96.8	0.544	0.468	1.1756	0.200	220.	19.5	0.844	18.2	41.8	(m)	30.3

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In addition to the output of the concentration field at specified times (from DEGADIS3), DEGADIS allows for transient releases output of concentration time histories at specified positions using DEGADIS4. DEGADIS4 can be executed interactively or in batch mode. An example DEGADIS4 interactive run and output follow.

Notes on Using DEGADIS4

- (1) DEGADIS4 can be run interactively as demonstrated here or in batch mode by supplying the same responses demonstrated here. DEGADIS4 sends the output to RUN_NAME.SR4.
- (2) The RUN_NAME of the transient simulation is used here.
- (3) The number of downwind positions where output is desired is entered here. For each downwind distance, questions 4 and 5 are required.
- (4) The x-coordinate is the downwind distance from the source where output is desired.
- (5) At each downwind distance, DEGADIS4 asks for the y- and z-coordinates desired. DEGADIS4 automatically supplies information about the centerline concentration ($y = 0$ and $z = 0$). Note that negative numbers signal that no more positions are desired for this particular downwind distance.

1 Enter the file name used for this run: BURRO9
2 Enter the number of downwind distances desired:
3 max of 10 downwind distances; 4 positions at each distance
1
4 enter the x coordinate:
100.
5 enter the y and z coordinate pairs at this distance:
0.,1.
30.,1.
-1.,-1.
t1: 18.000 tf: 99.263 dt: 2.0000 dist: 100.00
FORTRAN STOP
\$

Notes on DEGADIS4 Output

- (1) DEGADIS4 indicates the constants used in the x-direction dispersion correction. The user has the option of not making the x-direction dispersion correction if desired by making the appropriate changes to RUN_NAME.ER3.
- (2) At each downwind distance specified, the concentration profile parameters are output as a function of time. Note that part of the output has been removed.
- (3) DEGADIS4 outputs the concentration time history at the off-centerline positions specified.

1

X-Direction correction was applied.
 Coefficient: 2.00000E-02
 Power: 1.2200
 Minimum Distance: 130.00 m

2

Centerline values for the position -->
 x: 100.00 m

Time (s)	Mole Fraction	Concentration (kg/m**3)	Density (kg/m**3)	Gamma	Temperature (K)	Half Width (m)	Sz (m)	Sy (m)	Width at z=	1.00 m to:
									5.00 mole%	15.0 mole%
20.0	0.264	0.192	1.1202	0.203	264.	13.4	0.984	17.2	28.8	
22.0	0.370	0.282	1.1350	0.185	248.	17.0	0.931	18.3	36.5	
24.0	0.455	0.367	1.1525	0.191	235.	18.9	0.899	18.7	40.7	27.2
26.0	0.506	0.426	1.1652	0.194	227.	19.4	0.880	18.7	42.1	29.9
28.0	0.529	0.451	1.1714	0.197	223.	19.6	0.868	18.6	42.5	30.7
30.0	0.529	0.448	1.1678	0.191	224.	19.6	0.867	18.6	42.5	30.6
32.0	0.529	0.450	1.1704	0.196	223.	19.6	0.868	18.6	42.5	30.6
34.0	0.530	0.452	1.1706	0.195	223.	19.6	0.868	18.6	42.5	30.7

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90.0	0.530	0.452	1.1694	0.193	223.	19.6	0.869	18.6	42.5	30.7
92.0	0.531	0.453	1.1702	0.194	223.	19.6	0.869	18.6	42.5	30.8
94.0	0.529	0.448	1.1678	0.191	224.	19.6	0.869	18.7	42.5	30.7
96.0	0.514	0.431	1.1637	0.189	226.	19.8	0.880	18.9	42.9	30.6

3

Time (s)	Mole fraction at: y= 0.00000E+00 m	Mole fraction at: y= 30.000 m	Mole fraction at: y= -1.0000 m	Mole fraction at: y= 0.00000E+00 m
	z= 1.0000 m	z= 1.0000 m	z= -1.0000 m	z= 0.00000E+00 m
18.00000	0.0000000E+00	0.0000000E+00	0.0000000E+00	0.0000000E+00
20.00000	0.1086350	4.4177441E-02		
22.00000	0.1464778	9.1616339E-02		
24.00000	0.1790154	0.1289436		
26.00000	0.1992250	0.1492197		
28.00000	0.2065024	0.1560519		
30.00000	0.2052136	0.1552692		
32.00000	0.2060279	0.1555801		
34.00000	0.2068415	0.1560748		
90.00000	0.2070588	0.1563020		
92.00000	0.2075295	0.1565798		
94.00000	0.2058755	0.1559159		
96.00000	0.2017301	0.1552051		

\$

```

Program jetplu_in
c
c
c   JETPLU_IN is designed to perform two tasks including:
c
c     a) Read the input file for the JETPLU/DEGADIS model input. From
c        this information, JETPLU_IN prepares the necessary file to run the
c        JETPLU/DEGADIS model.
c
c     b) Generate a command file to execute the JETPLU/DEGADIS model from
c        the input information. This portion is highly dependent on the
c        VAX/VMS environment because of the manipulation of character strings
c        and the instructions necessary to invoke the VAX/VMS commands.
c
c.....  

c
c   This program can be invoked interactively or in a command file
c   submitted to batch. For either case, the syntax is:
c
c   $ RUN SYS$DEGADIS:JETPLU_IN
c   RUN_NAME
c
c   where RUN_NAME is the file name given to the particular simulation.
c   The input file RUN_NAME.IN is to be created before this program
c   is invoked using free formats. The order of input is as follows:
c
c   <TITLE1>
c   <TITLE2>
c   <TITLE3>
c   <TITLE4>
c
c   <U0> <Z0>
c   <ZR>
c   <INDVEL> <ISTAB> <RML>
c   <TAMB> <PAMB> <RELHUM>
c   <TSURF>
c
c   <GASNAM>
c   <GASMW>
c   <AVTIME>
c   <TEMJET>
c   <GASUL> <GASLL> <ZLL>
c   <INDHT> <CPK> <CPP>
c   <NDEN>
c
c   <ERATE>
c   <ELEJET> <DIAJET>
c   <TEND>
c   <DISTMX>
c
c   Note that for readability, blank lines were inserted between the

```

c input sections specifying a simulation title, atmospheric conditions,
c gas properties, and the particular release conditions. Symbol
c definitions are as follows:
c
c <TITLE1>, <TITLE2>, <TITLE3>, and <TITLE4> are four lines of up
c to 80 characters each of a title for this simulation.
c
c <U0> (m/s) is the ambient wind velocity at <Z0> (m).
c
c <ZR> is the surface roughness (m).
c
c <INDVEL> is an indicator which determines the method of calculation
c for the ambient velocity profile in the jet/plume model as
c follows:
c
c For <INDVEL>=1, the Pasquill-Gifford stability category
c (in <ISTAB> using 1 for A, 2 for B, etc.) is used
c along with <ZR> to determine the Monin-Obukhov length
c <RML>; the log velocity profile is then determined
c using <RML>.
c
c For <INDVEL>=2, the Monin-Obukhov length <RML> is supplied
c by the user; the log velocity profile is then determined
c using <RML>. Note that <ISTAB> must still be specified.
c
c <TAMB>, <PAMB>, and <RELHUM> are the ambient temperature (K), the
c ambient pressure (atm or N/m**2), and the relative humidity (%),
c respectively.
c
c <TSURF> is the surface temperature (K); if <TSURF> < 250 K, <TSURF> is
c set to <TAMB>.
c
c <GASNAM> is a three-letter designation for the contaminant's name.
c Any character string of three letters or less is valid; this
c is for user run identification and does not access property
c data.
c
c <GASMW> is the contaminant's molecular weight (kg/kmole).
c
c <AVTIME> is the averaging time (s). This parameter is
c used to estimate the value of <DELTAY>.
c
c <TEMJET> is the temperature of the jet (K).
c
c <GASUL> and <GASLL> are the concentrations to be used for estimating
c contours for an upper and lower concentration level in
c DEGADIS. The calculations are made for the elevation
c <ZLL>. Note that the JETPLU/DEGADIS computations will be carried
c out to <GASLL>/2.
c
c <INDHT> is used to include heat transfer in the DEGADIS computations.

c Heat transfer is not included for <INDHT>=0. For <INDHT>=1,
c heat transfer is included, and the heat transfer coefficient
c is calculated by DEGADIS. <CPP> and <CPK> are used to
c calculate the heat capacity as a function of
c temperature according to the correlation included in
c DEGADIS. If a constant heat capacity is
c desired, set <CPP> to 0. and <CPK> to the desired heat
c capacity (J/kg K).

c <NDEN> is used to specify the contaminant density profile.
c There are three cases for <NDEN>:

c <NDEN> = -1; The simulation treats the contaminant as if
c it were an ideal gas with a molal heat capacity
c equal to that of air. Water condensation effects
c are ignored. (equivalent to ISOFL=1 in DEGADIS)

c <NDEN> = 0; The simulation treats the contaminant as if
c it were an ideal gas with the heat capacity indicated
c by <CPK> and <CPP>. Water condensation effects
c are taken into account as appropriate.
c (equivalent to ISOFL=0 in DEGADIS)

c <NDEN> > 0; <NDEN> specifies the number of triples which follow
c in the next <NDEN> lines. The triples are used to
c specify the contaminant concentration as a function of
c density based on adiabatic mixing with ambient air.
c The ordered triples represent (in order):
c (1) the contaminant mole fraction
c (2). the contaminant concentration (kg contam/m³ mix)
c (3) the mixture density (kg mixture/m³ mixture)
c The ordered triples must go from pure air to pure
c contaminant. (equivalent to ISOFL=1 in DEGADIS)

c <ERATE> is the mass evolution (release) rate (kg/s).

c <ELEJET> is the initial jet elevation (m); the minimum jet elevation
c is twice the surface roughness. <DIAJET> is the initial
c jet diameter (m).

c <TEND> is the duration of the primary release (s). For steady-state
c releases, set <TEND> to 0.; to run the jet/plume model only, set
c <TEND> to a negative number.

c <DISTMX> is the maximum distance between output points in the JETPLU
c output (m).

c-----

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c University of Arkansas

```

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c      Fayetteville, AR 72701
c
c      (501) 575-4951
c
c-----
c

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

include 'sys$degadis:DEGADISIN.dec'

dimension DEN(5,igen)

c
data vkc/0.35D0/
c
logical check4
c
character*80 TITLE(4)
c
character*3 gas_name
c
character*100 OPNRUP
character OPNRUP1(100)
equivalence (opnrup1(1),opnrup)
character*4 IN,er1,er2,er3,com,scl,sr3,lis
character*4 dummy
character*3 plus, gasnam
character*2 con
DATA POUND//  '/,POUNDN/-1.E-20/
c
DATA IN'/.IN '//,er1'/.er1'//,er2'/.er2'//,er3'/.er3'/
data scl'/.scl'//,sr3'/.sr3'//,lis'/.lis'/
data com'/.com'/
data plus' + '/,con' - '/

c
c... GET THE FILE NAME TO BE USED BY ALL OF THE ROUTINES.....
c
READ(5,820) NCHAR,opnrup
opnrup = opnrup(1:nchar) // in(1:4)
c
c... Now get the rest of the desired information from RUN_NAME.IN
c
open(unit=8,name=opnrup,type='old')

read(8,8000) title(1)
read(8,8000) title(2)
read(8,8000) title(3)
read(8,8000) title(4)
8000  format(a80)

```

```

8010  format(a3)
      read(8,*) u0, z0
      read(8,*) zr

      read(8,*) indvel, istab, rml
c
c... Based on INDVEL, set the Monin-Obukhov length as desired. Also,.....
c      calculate USTAR
c
      if( indvel.ne.1 .or. indvel.ne.2) indvel = 1
      if( istab.le.0 .or. istab.gt.6) istab = 4
      if(indvel .eq. 1) then
          if(istab.eq.1) then
              rml = -11.43D0 * zr**0.103D0
          else if(istab.eq.2) then
              rml = -25.98D0 * zr**0.171D0
          else if(istab.eq.3) then
              rml = -123.4D0 * zr**0.304D0
          else if(istab.eq.4) then
              rml = 0.0D0
          else if(istab.eq.5) then
              rml = 123.4D0 * zr**0.304D0
          else if(istab.eq.6) then
              rml = 25.98D0 * zr**0.171D0
          endif
      endif
      ustar = u0*vkc/(dlog((z0+zr)/zr) + psif(z0,rml))

      read(8,*) tamb, pamb, relhum

      if( tamb.le.0. ) tamb = tamb+273.15D0
      if( pamb.gt.1.1 ) pamb = pamb/101325.0D0
      if( relhum.lt.0. .or. relhum.gt.100. ) relhum = 50.
c
c... Calculate the absolute humidity HUMID
c
      vaporp = 6.0298D-3* exp(5407.0D*(1.0D/273.15D0 - 1.0D/tamb))
      sat = 0.622D0*vaporp/(pamb - vaporp)
      humid = relhum/100.0D * sat

      read(8,*)
      if( tsurf.lt.250. ) tsurf = tamb

      read(8,8010) gasnam
      read(8,*) gasmw
      read(8,*) avtime
      read(8,*) TEMJET
      read(8,*) gasul, gasll, zll
      if( gasll.le.0. ) gasll = 0.01
      if( gasul.le.gasll ) gasul = dmax1( 1.1D0*gasll, 1.0D0)

```

```

c
c... Now that AVTIME has been set, the value of DELTAY can be fixed. Also
c      set the values of BETAY, DELTAZ, BETAZ, and GAMMAZ
c
      goto(161,162,163,164,165,166) istab
161   timeav = dmax1( avtime, 18.4D0)           ! A
      deltay = 0.423D0*(timeav/600.D0)**0.2D0
      betay = 0.9D0
      deltz = 107.66D0
      betaz = -1.7172D0
      gammaz = 0.277D0
      goto 170
162   timeav = dmax1( avtime, 18.4D0)           ! B
      deltay = 0.313D0*(timeav/600.D0)**0.2D0
      betay = 0.9D0
      deltz = 0.1355D0
      betaz = 0.8752D0
      gammaz = 0.0136D0
      goto 170
163   timeav = dmax1( avtime, 18.4D0)           ! C
      deltay = 0.210D0*(timeav/600.D0)**0.2D0
      betay = 0.9D0
      deltz = 0.09623D0
      betaz = 0.9477D0
      gammaz = -0.0020D0
      goto 170
164   timeav = dmax1( avtime, 18.3D0)           ! D
      deltay = 0.136D0*(timeav/600.D0)**0.2D0
      betay = 0.9D0
      deltz = 0.04134D0
      betaz = 1.1737D0
      gammaz = -0.0316D0
      goto 170
165   timeav = dmax1( avtime, 11.4D0)           ! E
      deltay = 0.102D0*(timeav/600.D0)**0.2D0
      betay = 0.9D0
      deltz = 0.02275D0
      betaz = 1.3010D0
      gammaz = -0.0450D0
      goto 170
166   timeav = dmax1( avtime, 4.6D0)            ! F
      deltay = 0.0674D0*(timeav/600.D0)**0.2D0
      betay = 0.9D0
      deltz = 0.01122D0
      betaz = 1.4024D0
      gammaz = -0.0540D0
c
170   continue
c

```

```

c... Recover INDHT, CPK, and CPP. If CPP is set to 0, then CPK contains.....
c      the (constant) heat capacity.
c
read(8,*) indht, CPK, CPP
if(cpp .eq. 0.D0) then
    CPP = 1.D0
    CPK = CPK*gasmw - 3.33D4
endif

c
c... recover NDEN and set ISOFL and DEN accordingly.....
c
read(8,*) nden
if(nden .lt. -1) nden=-1

if(nden .eq. -1) then
    isofl = 1
    rhoe = pamb*101325.D0*gasmw/8314.D0/TEMJET
    rhoa = pamb*
        (1.D0+humid)/(0.002833D0 + 0.004553D0*humid)/tamb
    den(1,1) = 0.D0
    den(2,1) = 0.D0
    den(3,1) = rhoa
    den(4,1) = 0.D0
    den(5,1) = tamb
    den(1,2) = 1.D0
    den(2,2) = rhoe
    den(3,2) = rhoe
    den(4,2) = 0.D0
    den(5,2) = tamb
else if(nden .eq. 0) then
    isofl = 0
else
    isofl = 1

do iii = 1,nden
read(8,*) den(1,iii), den(2,iii), den(3,iii)
den(4,iii) = 0.D0
den(5,iii) = tamb
enddo
endif

nden0 = nden
if(nden .eq. -1) nden0 = 2

read(8,*) erate
read(8,*) elejet, DIAJET
if(elejet .lt. 2.D0*zr) then
    elejet = 2.D0*zr
    write(6,*) 'JETPLU_IN: ELEJET has been increased to: ',elejet

```

```

        endif
        read(8,*) tend
        check4 = .true.
        if(tend .gt. 0.) check4 = .false.

        read(8,*) distmx
c
c... It is time to prepare the input file for the JETPLU model.....
c
        opnrup = opnrup(1:nchar) // '.ino'
        open(unit=1,name=opnrup,type='new')

        write(1,603) title(1)
        write(1,603) title(2)
        write(1,603) title(3)
        write(1,603) title(4)

c
c... atmospheric parameters
c
        write(1,*) u0, z0
        write(1,*) zr
        write(1,*) istab, rml, ustarr
        write(1,*) tamb, pamb, humid, relhum, tsurf
        write(1,*) avtime, deltay, betay
        write(1,*) deltaz, betaz, gammaz

c
c... contaminant parameters
c
        write(1,*) gasmw
        write(1,*) temjet
        write(1,*) gasul, gasll, zll
        write(1,*) CPK, CPP
        write(1,*) nden, nden0
        if(nden0 .gt. 0) then

            do iii = 1,nden0
            write(1,*) (den(jjj,iii),jjj=1,5)
            enddo
        endif
        write(1,*) isofl

c
c... spill parameters
c
        write(1,*) erate
        write(1,*) elejet, diajet

c
c... coding parameters
c
        alfa1 = 0.028D0
        alfa2 = 0.37D0
        write(1,*) alfa1, alfa2

```

```

        write(1,*) distmx

603   format(a80)
      close(unit=1)

c
c... Now, prepare the command file..... .
c
c
c... FORMATS
c
820   FORMAT(Q,A40)
c
c
1210  format(a4)
      opnrup = opnrup(1:nchar) // com(1:4)
c
      open(unit=8,name=opnrup,type='new',
      $ carriagecontrol='list',recordtype='variable')
c
c*** Lines to start the jet/plume model and invoke DEGBRIDGE
c
      opnrup = opnrup(1:nchar) // '.ino for001'
      write(8,1100) (opnrup1(i),i=1,nchar+11)
1100  format('$ assign ',51a1)
      opnrup = opnrup(1:nchar) // '.out for003'
      write(8,1100) (opnrup1(i),i=1,nchar+11)
      opnrup = opnrup(1:nchar) // '.ind for002'
      write(8,1100) (opnrup1(i),i=1,nchar+11)
      write(8,1160)
1160  format('$ run sys$degadis:jetplu')
      write(8,1170)
1170  format('$ deassign for001',//,$ deassign for002',//,
      '$ deassign for003')

c
c... Bypass DEGBRIDGE if this is an "JETPLU only" run
c
      if(tend .lt. 0.) goto 3000
      write(8,1180)
1180  format('$ run sys$degadis:degbridge')
      write(8,1290) (opnrup1(i),i=1,nchar)

c
c
c
      opnrup = opnrup(1:nchar) // er1(1:4)
c
      write(8,1250) (opnrup1(i),i=1,nchar+4)
1250  format('$ copy/log SYS$DEGADIS:example.er1 ',40a1)

```

```

        IF(u0 .eq. 0.) then
            write(8,1280)
            write(8,1290) (opnrup1(i),i=1,nchar)
            goto 1340
            endif
        opnrup = opnrup(1:nchar) // er2(1:4)

c
        write(8,1260) (opnrup1(i),i=1,nchar+4)
1260    format('$ copy/log SYS$DEGADIS:example.er2 ',40a1)
        opnrup = opnrup(1:nchar) // er3(1:4)

c
        if(.not.check4) then           ! transient
c
            write(8,1270) (opnrup1(i),i=1,nchar+4)
1270    format('$ copy/log SYS$DEGADIS:example.er3 ',40a1)
c
            write(8,1280)
1280    format('$ run SYS$DEGADIS:DEGADIS1')
            write(8,1290) (opnrup1(i),i=1,nchar)
1290    format(40a1)
            write(8,1300)
1300    format('$ run SYS$DEGADIS:DEGADIS2')
            write(8,1290) (opnrup1(i),i=1,nchar)
            write(8,1320)
1320    format('$ run SYS$DEGADIS:DEGADIS3')
            write(8,1290) (opnrup1(i),i=1,nchar)

c
        else
            write(8,1280)
            write(8,1290) (opnrup1(i),i=1,nchar)
c
            write(8,1330)
1330    format('$ run SYS$DEGADIS:SDEGADIS2')
            write(8,1290) (opnrup1(i),i=1,nchar)

c
        endif

        opnrup = opnrup(1:nchar) // '.out' //
        plus(1:3) // opnrup(1:nchar) // scl(1:4) //
        plus(1:3) // opnrup(1:nchar) // sr3(1:4) // con(1:2)
        write(8,1370) (opnrup1(i),i=1,3*nchar+20)
1370    format('$ copy/log ',100a1)
        opnrup = opnrup(1:nchar) // lis(1:4)
        write(8,1390) (opnrup1(i),i=1,nchar+4)
1390    format(' ',40a1)
c
        1340  close(unit=8)

3000  continue
        write(6,2099)
2099  format(//,' JETPLU_IN - beginning command file.')

```

```
opnrup = '0' // opnrup(1:nchar) // ''
istat = lib$do_command(opnrup)
write(6,2100)
2100 format(,/ ?JETPLU_IN? command file failed to start.')
c
CALL EXIT
END
#####
.
```

```

PROGRAM JETPLU_MAIN
c
c-----
c
c      This program calculates the trajectory and dilution of a steady
c      gas jet released at right angles to the wind. The zone before Gaussian
c      profiles can be assumed for the velocity and concentration profiles
c      is termed the zone of flow establishment; calculations for this zone
c      are done in the routine SETJET. The zone after Gaussian profiles can
c      be assumed to apply is termed the zone of established flow. The
c      ordinary differential equations describing this zone are included in
c      MODEL; these are integrated with the routine RKGST. Interactions
c      between the jet/plume and the ground are accounted for using the method
c      of images.
c
c-----
c
c      T. Spicer
c      University of Arkansas
c      Department of Chemical Engineering
c      Fayetteville, AR 72701
c
c      501-575-6516
c
c
Implicit real*8(a-h,o-z), integer*4(i-n)
c
include 'sys$degadis:DEGADIS1.dec'
c
external model,modout

character*80 title(4)
character*24 tinp
character*1 stabil(6)

dimension prmt(6)
dimension aux(8,6)

DIMENSION YR(6),dYR(6)

common
./GEN2/ DEN(5,igen)
./parm/ u0,z0,zr,rml,ustar,vkc,gg,rhoe,rhoa,deltay,betay,gammaf,
        .          ccflow
./com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
        .  gas_ufl,gas_lfl,gas_zsp,gas_name
./comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
        .  humsrc
./PHYS/ UA, deltaz,betaz,gammaz,rho, temp,yc
./COEFF/ ALFA1,ALFA2, sc,cd,delta
./rks/ rk1,rk2,rk3,rk4,rk5,rk6, rate,totrte,xmo,zmo

```

```

./sys/ sysz,sy,sz,sya,sza
./lfl/ cufl, clfl

c
character*3 gas_name

data vkc/0.35D0/
data gg/9.81D0/

data sc/1.42D0/
data cd/0.2D0/
data delta/2.15D0/

data stabil/'A','B','C','D','E','F'/

c
c
c... READ IN DATA FROM FOR001.DAT .....
OPEN (UNIT=1,STATUS='OLD')
OPEN (UNIT=3,STATUS='NEW')

READ(1,600) (title(j),j=1,4)
READ(1,*) u0,z0
READ(1,*) ZR
read(1,*) istab, rml, ustarr
read(1,*) tamb, pamb, humid, relhum, tsurf
read(1,*) avtime, deltay, betay
read(1,*) deltaz, betaz, gammaz

read(1,*) gas_mw
read(1,*) gas_temp
read(1,*) gas_ufl, gas_lfl, gas_zsp
read(1,*) gas_cpk, gas_cpp
read(1,*) nden, nden0
if(nden0 .gt. 0) then
  do iii = 1,nden0
    read(1,*) (den(jjj,iii),jjj=1,5)
  enddo

  den(1,nden0+1) = 2.00
  rhoa = den(3,1)
  rhoe = den(3,nden0)
else
  rhoa = Pamb*
    (1.00+humid)/(0.002833D0 + 0.004553D0*humid)/Tamb
  rhoe = pamb*101325.D0*gas_mw/8314.D0/gas_temp
endif
gas_rhoe = rhoe

read(1,*) isofl

```

```

read(1,*) erate
QINIT = erate/rhoe
read(1,*) elejet, diajet

READ(1,*) ALFA1, ALFA2
read(1,*) distmx
c

gamma = (rhoe - rhoa)/rhoe
cclow = (gas_lfl/4.)*pamb*101325.D0*gas_mw/8314.D0/tamb
c
c... initialize DEN if necessary .....
c
wc = 1.00
wa = 0.00
enth = cpc(gas_temp)*(gas_temp - tamb)
call setden(wc, wa, enth)

call adiabat(2,twc,twa,gas_lfl,ya,clfl,r,w,t,p)
call adiabat(2,twc,twa,gas_ufl,ya,cufl,r,w,t,p)

c
c
c... PRINT VALUES OF PARAMETERS INITIAL CONDITIONS .....
OPEN(UNIT=3,STATUS='NEW')
iii = lib$date_time(tinp)
write(3,700) tinp
write(3,710) (title(j),j=1,4)
write(3,720)
write(3,730) u0,z0,zr
write(3,740) stabil(istab)
write(3,750) rml,ustar,tamb,pamb,humid,relhum
write(3,760) avtime, deltax, betay, deltaz, betaz, gammaz
write(3,770)
write(3,780) gas_mw,gas_temp,gas_ufl,gas_lfl,gas_cpk,gas_cpp
write(3,790) nden
if(nden .gt. 0) then
    write(3,800)
    do iii = 1,nden
        write(3,810) den(1,iii), den(2,iii), den(3,iii)
    enddo
endif
write(3,820) isofl
write(3,830)
write(3,840) erate,elejet,diajet
write(3,850)
write(3,860) alfa1,alfa2,distmx
write(3,870) gas_zsp,(100.D0*gas_lfl),(100.D0*gas_ufl)
c

```

```

C
C... CALCULATE WIND SPEED AT THE JET DISCHARGE HEIGHT .....
C

UA=USTar/vkc*(dLOG((elejet+ZR)/ZR)-PSIF(elejet,rml))

C
C... CALL SETJET TO CALCULATE THE INITIAL CONDITIONS .....
C      FOR JETPLU MODEL INTEGRATION.
C

DO 17 I=1,6
dyr(i) = 1.0D0
17 YR(I)=0.0D0

YR(1) = rhoe
CALL SETJET(YR,UA,QINIT,DIAJET,elejet)
rho  = gas_rhoe
temp = gas_temp

C
C
C... CALCULATED jet parameters:
c      Yr(1) ... CENTERLINE CONCENTRATION
c      Yr(2) ... sysz (product of local lateral and vertical dimension)
c      Yr(3) ... theta (ANGLE OF TRAJECTORY)
c      Yr(4) ... uc (VELOCITY DECREMENT)
c      Yr(5) ... x (downwind distance)
c      Yr(6) ... zj (elevation)
c

prmt(1) = 0.0D0          ! lower limit
prmt(2) = 100000.         ! upper limit
prmt(3) = max(Yr(5)/20.0D0, 1.D-30) ! initial step
prmt(4) = 0.0001          ! error criteria
prmt(5) = distmx         ! maximum step size
prmt(6) = distmx/2.0D0    ! approximate output step size

ndim = 6
call rkgst(prmt,yr,dyr,ndim,ihlf,model,modout,aux)

c
if(ihlf.ge.10) then
    write(3,*) '----ihlf.ge.10----'
    dist = 0.0D0
    write(2,*) dist
    stop 'ihlf.ge.10'
endif
c

c

600  FORMAT(a80)
C     .

```

```

700  format(1x,1x,'*****',35x,'JETPLU/DEGADIS v2.1 ',35x,
        '*****',//,1x,a24,/)
710  format(1x,a80)
720  format(//,' Ambient Meteorological Conditions...')
730  format( //,' Ambient windspeed at reference height: ',1pg13.5,' m/s',
        .,' Reference height: ',1pg13.5,' m',
        .,' Surface roughness: ',1pg13.5,' m')
740  format( //,' Pasquill stability class: ',12x,A1)
750  format( //,' Monin-Obukhov length: ',1pg13.5,' m',
        .,' Friction velocity: ',1pg13.5,' m/s',
        .,' Ambient temperature: ',1pg13.5,' K',
        .,' Ambient pressure: ',1pg13.5,' atm',
        .,' Ambient humidity: ',1pg13.5,
        .,' Relative humidity: ',1pg13.5,' %')
760  format( //,' Specified averaging time: ',1pg13.5,' s',
        .,' DELTAY: ',1pg13.5,
        .,' BETAY: ',1pg13.5,
        .,' DELTAZ: ',1pg13.5,
        .,' BETAZ: ',1pg13.5,
        .,' GAMMAZ: ',1pg13.5)
770  format(//,' Contaminant Properties...')
780  format( //,' Contaminant molecular weight: ',1pg13.5,
        .,' Initial temperature: ',1pg13.5,
        .,' Upper level of interest: ',1pg13.5,
        .,' Lower level of interest: ',1pg13.5,
        .,' Heat capacity constant: ',1pg13.5,
        .,' Heat capacity power: ',1pg13.5)
790  format( //,' NDEN flag: ',I3)
800  format(1x,1x,'Mole fraction',1x,1x,'Concentration',1x,4x,'Density',
        .,1x,15x,4x,'(kg/m3)',4x,4x,'(kg/m3)',4x)
810  format(1x,3(1x,1pg13.5))
820  format( //,' ISOFL flag: ',I3)
830  format(//,' Release Properties...')
840  format( //,' Release rate: ',1pg13.5,' kg/s',
        .,' Discharge elevation: ',1pg13.5,' m',
        .,' Discharge diameter: ',1pg13.5,' m')
850  format(//,' Model Parameters...')
860  format( //,' ALFA1: ',1pg13.5,
        .,' ALFA2: ',1pg13.5,
        .,' DISTMX: ',1pg13.5,' m',//)
870  format(1x,96x,'-----',1x,'At z= ',1pg11.3,' m',1x,'-----',//,
        .,1x,2x,'Downwind',2x,1x,'Elevation',2x,4x,'Mole',4x,
        .,1x,'Centerline',1x,2x,'Density',3x,'Temperature',1x,
        .,2x,'Sigma y',3x,2x,'Sigma z',3x,4x,'Mole',4x,7x,'Width to:',//,
        .,1x,2x,'Distance',2x,12x,2x,'Fraction',2x,'Concentration',
        .,47x,2x,'Fraction',1x,1pE8.2,'mol%',1pE8.2,'mol%',//,
        .,1x,4x,'(m)',5x,4x,'(m)',5x,12x,2x,'(kg/m3)',3x,
        .,2x,'(kg/m3)',3x,4x,'(K)',5x,
        .,4x,'(m)',5x,4x,'(m)',5x,12x,4x,'(m)',5x,4x,'(m)')/

```

C

```

      CALL EXIT
      END

c-----.
c
c
      SUBROUTINE model(sss,yr,dyr,prmt)

      IMPLICIT REAL*8(A-H,O-Z), INTEGER*4(I-N)
c
      INCLUDE 'SYS$DEGADIS:DEGADIS1.DEC'
      PARAMETER (RT2 = 1.414213562D0)           ! SQRT(2.)

      EXTERNAL SYFUN
c

      COMMON
      ./GEN2/ DEN(5,igen)
      ./PARM/ u0,z0,zr,rml,ustar,vkc,gg,rhoe,rhoa,deltay,betay,gammaf,
      .          ccflow
      ./COM_GPROP/ gas_mw,gas_temp,gas_rhoe,gas_cpK,gas_cpp,
      .          gas_ufl,gas_lfl,gas_zsp,gas_name
      ./COMATM/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
      .          humsrc
      ./PHYS/ UA, deltaz,betaz,gammaz,rho, temp,yc
      ./COEFF/ ALFA1,ALFA2, sc,cd,delta
      ./RKS/ rk1,rk2,rk3,rk4,rk5,rk6, rate,totrte,xmo,zmo
      ./SYS/ sysz,sy,sz,sya,sza

c
      CHARACTER*3 gas_name

      DIMENSION yr(6),dyr(6),prmt(6)
      DIMENSION A(4,5)

c
c
c...integrated variables..... .
c
      CC     = yr(1)
      SYSZ  = ABS(yr(2))
      THETA = yr(3)
      UC    = yr(4)
      DIST  = yr(5)
      ZJ    = yr(6)

c
c... Calculate UENTR so that E1 is set to zero when UC<0
c
      UENTR = MAX(UC,0.D0)

```

```

c
c...and some hybrid variables
c
      ST    = SIN(theta)
      ST2   = ST*ST
      CT    = COS(theta)
      CT2   = CT*CT
c
c... Estimate the density
c
      call adiabat(0,wc,wa,yc,ya,cc,rho,wm,enth,temp)
      gamma = (rho-rhoa)/cc
c
c...calculate the ambient sigmas.....
c
      sya = deltay*dist**betay
      sza = deltz*dist**betaz * exp(gammaz*log(dist)**2)
      dsya = 0.
      dsza = 0.
      if(dist .gt. 1.D0) then
          dsya = sya/dist** betay * ct
          dsza = sza/dist*(betaz + 2.D0*gammaz*log(dist)) * ct
      endif
c
c...calculate the "constants"/properties for the ellipse including:.....
c      - the values for SY and SZ
c      - the curved perimeter and area of the ellipse
c      - the ambient windspeed averaged over z in the ellipse
c      - RK1 through RK6
c
c... Values for SY and SZ
c
      sstart = sqrt(sysz)
      sinc   = sstart/20.D0
      sylow  = 1.D-10
      syhigh = max(sysz, 10.D0)
      call limit(syfun, sstart, sinc, syhigh, sylow)

      tol = 0.0001D0
      call zbrent(sy, syfun, sylow, syhigh, tol, ierr)
      if(ierr .ne. 0) stop 'ZBRENT failed.'
      sz = sysz/sy
c
c
c...the curved perimeter, area of the ellipse
c
      call ellips(sy*delta, sz*delta, sz*delta, pe, area)
c

```

```

c...the ambient windspeed averaged over z in the ellipse (UA).
c      Calculate ALPHA from two velocity points calculated using the log
c      wind profile. The integrated profile based on ALPHA gives UA.
c      Don't let ZJ be less than ZR. The 0.01 offset for ZTOP gets
c      around a divide by zero when calculating ALPHA.
c
c      zj = max(zj, zr)

ztop = zj + delta*sz*ct + 0.01D0
utop = ustar/vkc*(dlog((ztop+zr)/zr) - psif(ztop,rml))
umid = ustar/vkc*(dlog((zj +zr)/zr) - psif(zj ,rml))
alpha = dlog(utop/umid) / dlog(ztop/zj)
alpha1 = 1.0D0+alpha

zbot = zj - delta*sz*ct
zbot = max(zbot, zr)
ua = umid/(alpha1*(ztop-zbot)*zj**alpha)
        *(ztop**alpha1 - zbot**alpha1)

vj = uc+ua*ct
vj = max(vj, 0.0D0)

c
c...some additional hybrid variables
c
UA2    = UA*UA
UACT   = UA*CT
UAST   = UA*ST
uauc   = ua*uc
uc2    = uc*uc
ucct   = uc*ct
ucst   = uc*st
ccsy   = cc*sy
ccsz   = cc*sz
ccsysz= cc*sy*sz

c
c... Gas Component Mass Balance Equation .....
c
qqq = rk1*uact + rk2*uc
rate = ccsysz*qqq

A(1,1) = qqq * sysz
A(1,2) = qqq * cc
A(1,3) = - rk1 * ccsysz * uast
A(1,4) = rk2 * ccsysz
A(1,5) = 0.0D0

c
c... Overall Mass Balance Equation .....
c
qqq = rk3*uact + rk4*uc
totrte = rhoa*qqq*sysz + gamma*rate

```

```

A(2,1) = 0.00
A(2,2) = rhoa * qqq
A(2,3) = - rhoa * rk3 * sysz * uast
A(2,4) = rhoa * rk4 * sysz

e3 = rk3*ua*(sza*dsya + sya*dsza)
e2 = alfa2*uact*abs(st)*pe
A(2,5) = rhoa*(ALFA1*uentr*pe + e2 + e3)

C
C... Z-direction Momentum Balance Equation .....
C
drag = cd * pe * rhoa/2.00*uast**2

rrr = 2.00*rk2*uauc*ct + rk1*ua2*ct2 + rk6*uc2
qqq = 2.00*rk4*uauc*ct + rk3*ua2*ct2 + rk5*uc2
xmo = rhoa*st*qqq*sysz + gamma*st*rrr*ccsysz

A(3,1) = gamma*sysz*rrr*st
A(3,2) = rhoa*qqq*st + gamma*cc*rrr*st
A(3,3) =
rhoa*sysz*(ct*qqq + st*
- 2.00*rk4*uauc*st - 2.00*rk3*ua2*ct*st))
+ gamma*ccsysz*(ct*rrr + st*
- 2.00*rk2*uauc*st - 2.00*rk1*ua2*ct*st))
A(3,4) =
rhoa*sysz*st*(2.00*rk4*uact + 2.00*rk5*uc)
+ gamma*ccsysz*st*(2.00*rk2*uact + 2.00*rk6*uc)
A(3,5) = -RK1*gg*gamma*ccsysz - sign(1.00,theta)*drag*CT

C
C... X-Direction Momentum Balance Equation .....
C
rrr = rk1*ua2*ct2*ct + 2.00*rk2*uauc*ct2 + rk6*uc2*ct
qqq = rk3*ua2*ct2*ct + 2.00*rk4*uauc*ct2 + rk5*uc2*ct
xmo = rhoa*qqq*sysz + gamma*rrr*ccsysz

A(4,1) = gamma*sysz*rrr
A(4,2) = rhoa*qqq + gamma*cc*rrr
A(4,3) =
rhoa*sysz*(-3.00*rk3*ua2*ct2*st
- 4.00*rk4*uauc*ct*st - rk5*uc2*st)
+ gamma*ccsysz*(-3.00*rk1*ua2*ct2*st
- 4.00*rk2*uauc*ct*st - rk6*uc2*st)
A(4,4) =
rhoa*sysz*(2.00*rk4*ua*ct2 + 2.00*rk5*ucct)
+ gamma*ccsysz*(2.00*rk2*ua*ct2 + 2.00*rk6*ucct)
A(4,5) = UA*A(2,5) + drag*ABS(ST)

C
C... SIMUL USED FOR MATRIX INVERSION
C
nnn = 4
call SIMUL(nnn,A,dyr)
c

```

```

c... Force any stray derivatives to be physically realistic.
c
c      if(dyr(1) .gt. 0.00) dyr(1) = 0.00
c      if(dyr(2) .lt. 0.00) dyr(2) = 0.00
c
c...d(x) and d(z) are calculated directly from cos(theta) and sin(theta),
c      respectively.
c
c      dyr(5) = ct
c      dyr(6) = st
c
c      RETURN
c      END

c-----
c
c----- subroutine modout(sss,yr,dyr,ihlf,ndim,prmt)
c
c      Implicit real*8(a-h,o-z), integer*4(i-n)
c
c      include 'sys$degadis:DEGADIS1.dec'
c
c      common
c./parm/ u0,z0,zr,rml,ustar,vkc,gg,rhoe,rhoa,deltay,betay,gammaf,
c          .       cclow
c./com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
c          .       gas_ufl,gas_lfl,gas_zsp,gas_name
c./PHYS/ UA, deltz,gammatz,rho, temp,yc
c./COEFF/ ALFA1,ALFA2, sc,cd,delta
c./rks/ rk1,rk2,rk3,rk4,rk5,rk6, rate,totrte,xmo,zmo
c./sys/ sysz,sy,sz,sya,sza
c./lfl/ cufl,clfl

      dimension yr(6),dyr(6),prmt(6)

      character*3 gas_name

      data ooo1/0.00/, ooo2/0.00/
      data iip/0/

c
c
c      cc     = yr(1)
c      sysz  = yr(2)
c      theta = yr(3)
c      uc    = yr(4)
c      dist  = yr(5)
c      zj    = yr(6)
c
c... Estimate CZSP and YZSP at y'=0 and z=GAS_ZSP.....
c      Estimate as though z' and z represent the same distances. Note that
c      no contribution from either exponential is included if the location

```

```

c      is outside the plume (outside DELTA*SZ which means ARG<2.31125).
c
      arg1 = 0.5D0*((zj - gas_zsp)/sz)**2
      arg2 = 0.5D0*((zj + gas_zsp)/sz)**2
      if(arg1 .lt. 2.31125D0) then
          arg1 = exp(-arg1)
      else
          arg1 = 0.0D0
      endif
      if(arg2 .lt. 2.31125D0) then
          arg2 = exp(-arg2)
      else
          arg2 = 0.0D0
      endif
      czsp = cc*(arg1 + arg2)

      call adiabat(0,wc,wa,yzsp,ya,czsp,rho,wm,enth,temp)

      if(czsp .gt. cufl) then
          wufl = sy*sqrt(2.0D0*log(czsp/cufl))
          wlfl = sy*sqrt(2.0D0*log(czsp/clfl))
      else if(czsp .gt. clfl) then
          wufl = 0.0D0
          wlfl = sy*sqrt(2.0D0*log(czsp/clfl))
      else
          wufl = 0.0D0
          wlfl = 0.0D0
      endif
      wufl = min(wufl, delta*sy)
      wlfl = min(wlfl, delta*sy)

c
c... IF THE JET REACHES THE GROUND, STOP THE COMPUTATION..... .
c
      IF (zj.LE.0.0D0) then
          cc = cc - zj*(occ - cc)/(ozj - zj)
          sy = sy - zj*(osy - sy)/(ozj - zj)
          sz = sz - zj*(osz - sz)/(ozj - zj)
          theta = theta - zj*(otheta - theta)/(ozj - zj)
          uc = uc - zj*(ouc - uc)/(ozj - zj)
          dist = dist - zj*(odist - dist)/(ozj - zj)
          ua = ua - zj*(oua - ua)/(ozj - zj)
          yzsp = yzsp - zj*(oyzsp - yzsp)/(ozj - zj)
          wlfl = wlfl - zj*(owlfl - wlfl)/(ozj - zj)
          wufl = wufl - zj*(owufl - wufl)/(ozj - zj)

          zj = 0.0D0
          cc = 2.0D0*cc

          call adiabat(0,wc,wa,yc,ya,cc,rho,wm,enth,temp)

          WRITE(2,*) dist,cc,delta*sy

```

```

if(wufl.eq.0.00 .and. wlfl.eq.0.00) then
    WRITE(3,734) dist,zj,yc,cc,rho,temp,sy,sz,yzsp
else if(wufl.eq.0.00 .and. wlfl.ne.0.00) then
    WRITE(3,734) dist,zj,yc,cc,rho,temp,sy,sz,yzsp,wlfl
else
    WRITE(3,734) dist,zj,yc,cc,rho,temp,sy,sz,yzsp,wlfl,wufl
endif
    WRITE(6,9999) dist,zj,cc,sy,sz,uc,ua,theta
    prmt(5) = 1.
    return
endif
c
c... store the present values for interpolation when ZJ=0
c
    occ    = cc
    osy    = sy
    osz    = sz
    otheta = theta
    ouc    = uc
    odist  = dist
    ozj    = zj
    oua    = ua
    oyzsp  = yzsp
    owlfl  = wlfl
    owufl  = wufl
c
c... Calculate the image contribution .....
c
    cimage = cc*exp(-0.500*(2.00*zj/sz)**2)
    cc = cc + cimage

    call adiabat(0,wc,wa,yc,ya,cc,rho,wm,enth,temp)
c
c... IF THE JET cc drops below cclow, stop the calculation.....
c
    IF (cc.lt.cclow) then
        if(wufl.eq.0.00 .and. wlfl.eq.0.00) then
            WRITE(3,734) dist,zj,yc,cc,rho,temp,sy,sz,yzsp
        else if(wufl.eq.0.00 .and. wlfl.ne.0.00) then
            WRITE(3,734) dist,zj,yc,cc,rho,temp,sy,sz,yzsp,wlfl
        else
            WRITE(3,734) dist,zj,yc,cc,rho,temp,sy,sz,yzsp,wlfl,wufl
        endif
        WRITE(6,9999) dist,zj,cc,sy,sz,uc,ua,theta
    c
    c... set DIST to 0 to preclude DEGADIS from continuing
        dist = 0.00
        WRITE(2,*) dist,cc,delta*sy
        prmt(5) = 1.
        return
    endif

```

```

c
c... printed output
c
      if(dist .ge. ooo1+ooo2) then
          ooo1 = dist
          ooo2 = prmt(6)

          if(wufl.eq.0.D0 .and. wlfl.eq.0.D0) then
              WRITE(3,734) dist,zj,yc,cc,rho,temp,sy,sz,yzsp
          else if(wufl.eq.0.D0 .and. wlfl.ne.0.D0) then
              WRITE(3,734) dist,zj,yc,cc,rho,temp,sy,sz,yzsp,wlfl
          else
              WRITE(3,734) dist,zj,yc,cc,rho,temp,sy,sz,yzsp,wlfl,wufl
          endif
          WRITE(6,9999) dist,zj,cc,sy,sz,uc,ua,theta

          iip = iip + 1
          if(iip .eq. 3) then
              write(3,735)
              write(6,735)
              iip = 0
          endif
      endif

c
      734  FORMAT(1X,10(1pg11.3,1x),1pg11.3)
      735  format(1x)
      9999 FORMAT (1x,5(1pg9.3,1x),1pg10.3,1pg9.3,1pg10.3,3(1pg9.3,1x),
                   1pg10.3,1x,1pg9.3)
      .
      return
      end

c
c
c-----



      SUBROUTINE SETJET(YR,ua,QINIT,DIAJET,elejet)
c
c      THIS SUBROUTINE TRANSFORMS THE EXIT 'TOPHAT' VELOCITY PROFILE OF
c      A CONTINUOUS JET TO THE SIMILARITY (GAUSSIAN) FORM REQUIRED FOR
c      INPUT TO THE JETPLU MODEL - USES WIND TUNNEL DATA CORRELATIONS BY
c      Y. KAMOTANI AND I. GREBER, NASA CONTRACTOR REPORT CR-2392, 3/74.
c
c      This has been simplified for vertical releases only.
c

      implicit real*8(a-h,o-z), integer*4(i-n)
c
      include 'sys$degadis:DEGADIS1.dec'
      parameter (rt2 = 1.414213562D0)           ! sqrt(2.)

```

```

c
      common
./parm/ u0,z0,zr,rml,ustar,vkc,gg,rhoe,rhoa,deltay,betay,gammaf,
.          cclow
./COEFF/ ALFA1,ALFA2, sc,cd,delta
./rks/ rk1,rk2,rk3,rk4,rk5,rk6, rate,totrte,xmo,zmo
./sys/ sysz,sy,sz,sya,sza

DIMENSION YR(6)

c
c... initial values for CC and UJ .....
c
cc = yr(1)
uj = qinit/pi/diajet**2*4.00
c
c... Calculate the length of the zone of flow development based on analysis
c      of Pratte and Baines work in the ASCE Journal of the Hydraulics
c      Division.
c
c.....SOD represents (S/D) or length of development zone (S) nondimensionalized
c      using the jet diameter (D).
c
sod = 7.7D0*(1.00 - exp(-0.48D0*sqrt(rhoe*uj/rhoa/ua)))

c
c...Based on RJ ("J" in Kamotani and Grebber) and FROUDE (Froude number), .....
c      calculate "ae" and "be" (represented herein as AJ and BJ).

rj = (rhoe/rhoa)*(Uj/ua)**2
delrho = rhoe - rhoa
froude = 1.688D0
if(DELRHO .GT. 0.) froude = rhoa*ua**2/gg/delrho/diajet
froude = min(froude, 1.688D0)

if(rj .le. 0.036D0) then
    aj = 18.519D0*rj
    bj = 0.400
else if(rj .gt. 0.036D0 .and. rj .le. 10.00) then
    aj = EXP(0.2476D0 +0.3016D0*log(froude) +0.24386D0*LOG(rj))
    bj = 0.400
else if(rj .gt. 10.00 .and. rj .le. 50.00) then
    aj = EXP(0.405465D0 +0.131386D0*LOG(rj) +0.054931D0*log(rj)**2)
    bj = exp(-0.744691D0 - 0.074525D0*log(rj))
else if(rj .gt. 50.00 .and. rj .le. 600.00) then
    aj = EXP(-2.55104D0 +1.49202D0*log(rj) -0.097623D0*log(rj)**2)
    bj = exp(-0.446718D0 - 0.150694D0*log(rj))
else
    write(6,*) 'J exceeded in SETJET; Aj and Bj are extrapolated.'
    aj = EXP(1.44099D0 + 0.243045D0*log(rj))

```

```

bj = exp(-0.446718D0 - 0.150694D0*log(rj))
endif

bji = 1.00/bj
c
c...Estimate (z/D) (as ZOD) from AJ, BJ, and SOD using a Newton-Raphson .....
c      procedure.
c
zod = sod

iii = 0
100  continue
iii = iii+1
if(iii .gt. 100) then
    stop 'SETJET Loop failed'
endif

xod = (zod/aj)**bji
fff = xod**2 + zod**2 - sod**2
ffffp= xod**2*2.00/bj/zod + 2.00*zod
zodn = zod - fff/ffffp
check = abs((zodn-zod)/zod)
if(check .gt. 0.00001) then
    zod = zodn
    goto 100
endif
c
c... Calculate DIST, THETA, ZJ, and UC from ZOD and XOD. Note that XOD is .....
c      subject to underflows which is cured on a VAX by (XOD = ZOD).
c
xod = xod
dist = xod*diajet
if(xod .gt. 0.00) then
    slope = bj*zod/xod
    theta = atan(slope)
else if(xod .eq. 0.00) then
    theta = pi/2.00
else
    stop 'XOD < 0. in SETJET.'
endif
zj = zod*diajet + elejet
uc = uj - ua*cos(theta)
c
c...Determine SY and SZ to close the material balance .....
c
erate = qinit*rhoe
c
c... Calculate the constants RK1 to RK6; PPP "corrects" the area of
c      the assumed rectangle to the ellipse.
c

```

```

      ppp = sqrt(pi)/2.00

      qqq = erf(delta /rt2 *ppp)
      rk1 = pi * qqq * (2.00*qqq)

      qqq = erf(delta /rt2*sqrt(1.00+sc) *ppp)
      rk2 = pi/(1.00+sc) * qqq * (2.00*qqq)

      rk3 = pi*delta**2

      qqq = erf(delta /rt2*sqrt(sc) *ppp)
      rk4 = pi/sc * qqq * (2.00*qqq)

      qqq = erf(delta *sqrt(sc) *ppp)
      rk5 = pi/(2.00*sc) * qqq * (2.00*qqq)

      qqq = erf(delta /rt2*sqrt(1.00+2.00*sc) *ppp)
      rk6 = pi/(1.00+2.00*sc) * qqq * (2.00*qqq)

c
c... Calculate SYSZ.
c
      sysz = erate/cc/(rk1*ua*cos(theta) + rk2*uc)

      sy = sqrt(sysz)
      sz = sy

      yr(1) = cc
      yr(2) = sysz
      yr(3) = theta
      yr(4) = uc
      yr(5) = max(dist, 1.0-30)
      yr(6) = zj

      write(6,*) 'JETPLU/SETJET initial conditions...'
      write(6,*) ''
      write(6,*) '      u0: ',u0    , '      z0: ',z0
      write(6,*) '      erate: ',erate , '      uj: ',uj
      write(6,*) '      rhoe: ',rhoe , '      rhoa: ',rhoa
      write(6,*) '      rj: ',rj    , '      froude: ',froude
      write(6,*) ''
      write(6,*) '      dist: ',dist , '      zj: ',zj
      write(6,*) '      diajet: ',diajet,'      uc: ',uc
      write(6,*) '      sy: ',sy    , '      sz: ',sz
      write(6,*) '      theta: ',theta,'      sod: ',sod
      write(6,*) ''
      write(6,*) '      rk1: ',rk1   , '      rk2: ',rk2
      write(6,*) '      rk3: ',rk3   , '      rk4: ',rk4
      write(6,*) '      rk5: ',rk5   , '      rk6: ',rk6
      write(6,*) ''

```

C-30

```
      write(6,*) ' DIST      ZJ      ',  
      'CC      SY      SZ      UC      UA      THETA'  
      write(6,*) ''  
      RETURN  
      END  
  
c-----  
c  
c      function syfun(sytry)  
c  
c      implicit real*8(a-h,o-z), integer*4(i-n)  
  
c      common  
.rks/ rk1,rk2,rk3,rk4,rk5,rk6, rate,tot rte,xmo,zmo  
.sys/ sysz,sy,sz,sya,sza  
  
c      sztry = sysz/sytry  
c  
c      syfun = sya**2 - sza**2 - sytry**2 + sztry**2  
c... to improve the accuracy of the calculation...  
c  
c      syfun = (sya-sza)*(sya+sza) + (sztry-sytry)*(sztry+sytry)  
  
c      return  
c      end  
#####
```

APPENDIX D

DEGADIS MODEL SOURCE CODE

AFGEN.FOR	D-3	PSSOUT.FOR	D-104
AFGEN2.FOR	D-4	PSSOUTSS.FOR	D-107
ALPH.FOR	D-5	RIPHIF.FOR	D-110
CRFG.FOR	D-8	RKGST.FOR	D-114
DEGADIS1.DEC	D-13	SDEGADIS2.FOR	D-122
DEGADIS1.FOR	D-14	SERIES.FOR	D-130
DEGADIS2.DEC	D-22	SORTS.FOR	D-131
DEGADIS2.FOR	D-23	SORTS1.FOR	D-134
DEGADIS3.DEC	D-27	SRC1.FOR	D-139
DEGADIS3.FOR	D-28	SRTOUT.FOR	D-148
DEGADIS4.DEC	D-32	SSG.FOR	D-152
DEGADIS4.FOR	D-33	SSGOUT.FOR	D-155
DEGADISIN.DEC	D-38	SSGOUTSS.FOR	D-158
DEGADISIN.FOR	D-39	SSOUT.FOR	D-161
DOSOUT.FOR	D-45	SSSUP.FOR	D-163
ESTRT1.FOR	D-50	STR2.FOR	D-172
ESTRT2.FOR	D-54	STR2SS.FOR	D-175
ESTRT2SS.FOR	D-56	STR3.FOR	D-178
ESTRT3.FOR	D-58	SURFACE.FOR	D-180
GAMMA.FOR	D-59	SZF.FOR	D-182
GETTIM.FOR	D-61	TPROP.FOR	D-185
GETTIMDOS.FOR	D-63	TRANS1.FOR	D-199
HEAD.FOR	D-64	TRANS2.FOR	D-202
INCGAMMA.FOR	D-70	TRANS2SS.FOR	D-204
IO.FOR	D-74	TRANS3.FOR	D-206
IOT.FOR	D-76	TRAP.FOR	D-207
LIMIT.FOR	D-93	TS.FOR	D-216
NOBL.FOR	D-94	TUPF.FOR	D-217
OB.FOR	D-97	UIT.FOR	D-222
PSIF.FOR	D-100	ZRENT.FOR	D-224
PSS.FOR	D-101		

```

C.....  

C  

C      THIS FUNCTION LINEARLY INTERPOLATES FROM THE GIVEN  

C          PAIR OF DATA POINTS.  

C  

C      FUNCTION AFGEN(TAB,X,SPEC)  

C  

Implicit Real*8 ( A-H, D-Z ), Integer*4 ( I-N )  

C  

C      include 'sys$degadis:DEGADIS1.dec'  

C      common/nend/poundn,pound  

C  

C      character*4 pound  

C      character*(*) SPEC  

C      DIMENSION TAB(1)  

C  

C      IF(X .GE. TAB(1)) GO TO 95  

C      WRITE(lunlog,1100) X,spec  

C      AFGEN = TAB(2)  

C      RETURN  

C  

95    continue  

      ix = 1  

100   IX = ix + 2  

C  

      IY = IX + 1  

      IF( TAB(IX).EQ.POUNDN .AND. TAB(IY).EQ.POUNDN ) GO TO 500  

      IF(X .GE. TAB(IX)) GO TO 100  

C  

      IXP = IX-2  

      IYP = IXP + 1  

C  

      SL = (TAB(IY) - TAB(IYP))/(TAB(IX) - TAB(IXP))  

      AFGEN = SL*(X - TAB(IXP)) + TAB(IYP)  

      RETURN  

C  

500   CONTINUE  

      IX = IX-2  

      IY = IY-2  

      IXP = IX-2  

      IYP = IY-2  

C  

      SL = (TAB(IY) - TAB(IYP))/(TAB(IX) - TAB(IXP))  

      AFGEN = SL*(X - TAB(IXP)) + TAB(IYP)  

C  

1100  FORMAT(2X,'?AFGEN? UNDERFLOW; argument: ',1pg13.5,5X,A20)  

      RETURN  

      END  

#####

```

```

C.....  

C  

C      THIS FUNCTION LINEARLY INTERPOLATES FROM THE GIVEN  

C          PAIR OF DATA POINTS.  

C  

C      FUNCTION AFGEN2(XTAB,TAB,X,SPEC)  

C  

C      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      include 'sys$degadis:DEGADIS1.dec'  

C      common/nend/poundn,pound  

C  

C      character*4 pound  

C      character*(*) SPEC  

C      DIMENSION XTAB(1),TAB(1)  

C  

C      IF(X .GE. XTAB(1)) GO TO 95  

C      WRITE(lunlog,1100) X,spec  

C      AFGEN2 = TAB(2)  

C      RETURN  

C  

C      95    continue  

C      ix = 1  

C      100   IX = ix + 1  

C  

C      IF( XTAB(IX).EQ.POUNDN ) GO TO 500  

C      IF(X .GE. XTAB(IX)) GO TO 100  

C  

C      IXP = IX-1  

C  

C      SL = (TAB(IX) - TAB(IXP))/(XTAB(IX) - XTAB(IXP))  

C      AFGEN2 = SL*(X - XTAB(IXP)) + TAB(IXP)  

C      RETURN  

C  

C      500  CONTINUE  

C      IX = IX-1  

C      IXP = IX-1  

C  

C      SL = (TAB(IX) - TAB(IXP))/(XTAB(IX) - XTAB(IXP))  

C      AFGEN2 = SL*(X - XTAB(IXP)) + TAB(IXP)  

C  

C      1100 FORMAT(2X,'?AFGEN2? UNDERFLOW; argument: ',1pg13.5,5X,A20)  

C      RETURN  

C      END  

#####

```

```

C.....  

C  

C      SUBROUTINES TO CALCULATE THE VALUE OF ALPHA  

C  

C      SUBROUTINE ALPH  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      include 'sys$degadis:DEGADIS1.dec'  

C  

C      COMMON  

$/ERROR/STPIN,ERBND,STPMX,WTRG,WTtm,WTya,WTyc,WTeb,WTmb,WTuh,XLI,  

$ XRI, EPS, ZLOW, STPINZ, ERBNDZ, STPMXZ, SRCOER, SRCSS, SRCCUT,  

$ HCTCUT, ERNOBL, NOBLPT, CRFGER, EPSILON  

$ PARM/U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$ ALP/ALPHA,alpha1  

$ alphcom/ ialpfl,alpco  

C  

REAL*8 ML,K  

C  

EXTERNAL ALPHI  

C  

PSI = PSIF(Z0,ML)  

USTAR = U0*K/(dLOG((Z0+ZR)/ZR) - PSI)  

C  

if(u0 .eq. 0.) then  

    alpha = 0.  

    ustar = 0.  

    return  

endif  

C  

if(ialpfl.eq. 0) then  

    alpha = alpco  

    return  

endif  

C  

C*** ZBRENT USED TO DETERMINE THE ROOT OF THE REQUIRED INTEGRAL EQUATION  

C  

IER = 0  

C  

CALL zbrent(alpha, ALPHI, XLI, XRI, EPS, IER)  

IF( IER .NE. 0 ) CALL trap(19,IER)  

C  

RETURN  

END  

C  

C.....  

C  

C      FUNCTION TO EVALUATE THE WEIGHTED EUCLIDEAN NORM OF THE

```

```

C      ERROR ASSOCIATED WITH THE POWER LAW FIT OF THE WIND PROFILE.
C
C      FUNCTION ALPHI(X)

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

C      include 'sys$degadis:DEGADIS1.dec'
C
C      COMMON
$/ERROR/STPIN,ERBND,STPMX,WTRG,WTtm,WTya,WTyc,WTeb,WTmb,WTuh,XLI,
$ XRI,EPS,ZLOW,STPINZ,ERBNDZ,STPMXZ,SRCOER,srcss,srccut,
$ htcut,ERNOBL,NOBLpt,crfger,epsilon
$/PARM/U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW
$/ALP/ALPHA,alpha1

C      REAL*8 ML,K

C      DIMENSION Y(1),DERY(1),PRMT(5),AUX(8)
C      EXTERNAL ARG,ARGOUT

C      ALPHA = X

C      PRMT(1) = Z0
C      PRMT(2) = dmax1(ZLOW,zr)          ! to take care of large zr
C      PRMT(3) = STPINZ
C      PRMT(4) = ERBNDZ
C      PRMT(5) = stpmxz

C      Y(1) = 0.0D00

C      DERY(1) = 1.0D00

C      NDIM = 1
C      IHLF = 0

C      CALL RKGST(PRMT,Y,DERY,NDIM,IHLF,ARG,ARGOUT,AUX)

C      IF(IHLF .GE. 10) CALL trap(18,IHLF)
C      ALPHA = Y(1)
C      RETURN
C      END

C
C.....FUNCTION TO EVALUATE THE ARGUMENT OF THE INTEGRAL EXPRESSION
C
C      SUBROUTINE ARG(Z,Y,D,PRMT)

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

```

```

C
      include 'sys$degadis:DEGADIS1.dec'
C
      COMMON
      $/PARM/U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW
      $/ALP/ALPHA,alpha1
      $/alphcom/ ialpf1,alpco
C
      REAL*8 ML,K
C
      DIMENSION Y(1),D(1),PRMT(1)
C
C*** WEIGHT FUNCTION USED
C
      W = 1.000/(1.000 + Z)
      if(ialpf1.eq. 2) w= 1.000
C
C*** WIND VELOCITY @ Z -- BEST FIT
C
      UBST = USTAR/K*(dLOG((Z+ZR)/ZR) - PSIF(Z,ML))
C
C*** WIND VELOCITY @ Z -- POWER LAW APPROXIMATION
C
      *      UALP = U0 * (Z/Z0) ** ALPHA
C
      D(1) = W * (UBST - UALP) * dLOG(Z/Z0) * UALP
      RETURN
      END
C
C
      SUBROUTINE ARGOUT
      RETURN
      END
#####

```

```

C.....  

C  

C      SUBROUTINE TO CREATE RADG,QSTR,srcden,srcwc,srcwa,srcenth DATA LISTS  

C  

C      PARAMETERS  --  TABLE - WORKSPACE VECTOR  

C                  NTAB - DIMENSION OF TABLE DIVIDED BY iout_src  

C                  RER - RELATIVE ERROR BOUND OF CREATED  

C                          DATA PAIRS BY LINEAR INTERPOLATION  

C  

C      VALUES OF TIME, RADG, height, QSTR, S20, yc, ya, rho, Ri,  

C      wc,wa,enthalpy,temp  

C      ARE READ INTO  

C      TABLE(1) TO TABLE(13) RESPECTIVELY.  

C  

C.....  

C  

SUBROUTINE CRFG(TABLE,NTAB,rer)  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

DIMENSION TABLE(1)  

C  

include 'sys$degadis:DEGADIS1.dec'  

parameter (zero= 1.e-20)  

C  

COMMON  

$/GEN3/ radg(2,maxl),qstr(2,maxl),srcden(2,maxl),srcwc(2,maxl),  

$ srcwa(2,maxl),srcenth(2,maxl)  

$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

$/NEND/ POUNDN,POUND  

C  

character*4 pound  

C  

LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

C  

DATA NI/1/  

C  

data iti/1/      ! time - element no 1 in record  

data irg/2/      ! Radg - element no 2 in record  

data iqs/4/      ! Qstar - element no 4 in record  

data idn/8/      ! rho - element no 8 in record  

data iwc/10/     ! wc - element no 10 in record  

data iwa/11/     ! wa - element no 11 in record  

data ien/12/     ! enthalpy - element no 12 in record  

C  

C  

C      OUTPUT CREATED VECTORS TO A PRINT FILE

```

```

C
C
      READ(9,*) (TABLE(J),J=1,iout_src)
C
      WRITE(8,1111)
      WRITE(8,1105)
      if(isofl.eq. 1) then
      WRITE(8,1102)
      WRITE(8,1103)
      WRITE(8,1140) (TABLE(J),J=1,6),table(8),table(9)
      else
      WRITE(8,1100)
      WRITE(8,1104)
      WRITE(8,1140) (TABLE(J),J=1,6),table(8),table(13),table(9)
      endif
      ispace = 1
C
1100   FORMAT(/,5X,'Time',5X,2x,'Gas Radius',2x,4X,'Height',4X,
      $4x,'Qstar',5x,2x,'SZ(x=L/2.)',2x,1X,'Mole frac C',2x,
      $3x,'Density',4x,1x,'Temperature',2x,3x,'Rich No.',3x)
1102   FORMAT(/,5X,'Time',5X,2x,'Gas Radius',2x,4X,'Height',4X,
      $4x,'Qstar',5x,2x,'SZ(x=L/2.)',2x,1X,'Mole frac C',2x,
      $3x,'Density',4x,3x,'Rich No.',3x)
1103   FORMAT(1H ,4X,'sec',6X,6X,'m',7X,6X,'m',7X,
      $2X,'kg/m**2/s',3X,6X,'m',7X,14x,3x,'kg/m**3',4x,14x,/)
1104   FORMAT(1H ,4X,'sec',6X,6X,'m',7X,6X,'m',7X,
      $2X,'kg/m**2/s',3X,6X,'m',7X,14x,3x,'kg/m**3',4x,6x,'K',7x,14x,/)
1105   FORMAT(1H ,23X,'*****',21X,'CALCULATED SOURCE PARAMETERS',21X,
      $'*****')
C
      RADG(1,1) = 0.
      RADG(2,1) = TABLE(2)
      QSTR(1,1) = 0.
      QSTR(2,1) = TABLE(4)
      srcden(1,1) = 0.
      srcden(2,1) = table(8)
      srcwc(1,1) = 0.
      srcwc(2,1) = table(10)
      srcwa(1,1) = 0.
      srcwa(2,1) = table(11)
      srcenth(1,1) = 0.
      srcenth(2,1) = table(12)
C
      READ(9,*) (TABLE(J),J=1,iout_src)
      L = 2
C
C*** L IS THE NUMBER OF RECORDS WHICH HAVE BEEN READ
C
100   CONTINUE
      DO 120 I=2,NTAB
C

```

```

C*** MOVE LAST RECORD READ INTO THE LAST ACTIVE POSITION OF TABLE
C
      DO 130 J = 1,iout_src
      KK = iout_src * (I-1) + J
130   TABLE(KK) = TABLE(J)
      KK = iout_src * I
C
C*** READ THE NEXT RECORD.  INCREMENT L.
C
      L = L + 1
      READ(9,* ,END=900) (TABLE(J),J=1,iout_src)
C
C
      DO 140 Kkk = 2,I
C
      KT     = iout_src*(Kkk-1) + iti
      KRG    = iout_src*(Kkk-1) + irg
      KQSTR  = iout_src*(Kkk-1) + iqs
      KCA    = iout_src*(Kkk-1) + idn
      Kwc    = iout_src*(Kkk-1) + iwc
      Kwa    = iout_src*(Kkk-1) + iwa
      Ken    = iout_src*(Kkk-1) + ien
C
      timeslot = radg(1,ni)
      ratio = (table(kt) - timeslot) / (table(iti) - timeslot)
C
      ANSRG = (TABLE(irg) - RADG(2,NI)) * ratio + RADG(2,NI)
      ANSQ = (TABLE(iqs) - QSTR(2,NI)) * ratio + QSTR(2,NI)
      ANSCA = (TABLE(idn) - srccden(2,NI)) * ratio + srccden(2,NI)
      ANSWC = (TABLE(iwc) - srccwc(2,NI)) * ratio + srccwc(2,NI)
      ANSwa = (TABLE(iwa) - srccwa(2,NI)) * ratio + srccwa(2,NI)
      ANSen = (TABLE(ien) - srcenth(2,NI)) * ratio + srcenth(2,NI)
C
      ERRG = ABS(ANSRG - TABLE(KRG))/TABLE(KRG)
      ERQSTR = ABS(ANSQ - TABLE(KQSTR))/(TABLE(KQSTR)+zero)
      ERO = dMAX1(ERRG,ERQSTR)
C
      ERCA = ABS(ANSCA - TABLE(KCA))/TABLE(KCA)
      ERO = dMAX1(ERO,ERCA)
C
      ERWC = ABS(ANSWC - TABLE(KWC))/(TABLE(KWC)+ zero)
      ERO = dMAX1(ERO,ERWC)
      ERWA = ABS(ANSWa - TABLE(Kwa))/(TABLE(Kwa)+ zero)
      ERO = dMAX1(ERO,ERWA)
      ERen = ABS(ANSen - TABLE(Ken))/(TABLE(Ken)+ zero)
      ERO = dMAX1(ERO,ERen)
C
      IF(ERO .GT. RER) GO TO 150
140   CONTINUE
120   CONTINUE
C

```

```

        WRITE(lunlog,1110)
C
C*** RECORD NEXT DATA PAIR. SINCE ERROR EXCEEDED, RECORD THE LAST
C*** DATA PAIR WHICH SATISFIED THE ERROR CRITERIA WHICH IS STORED
C*** IN TABLE(KK-(iout_src-1)) TO TABLE(KK)
C
150   NI = NI + 1
C
KT      = KK - iout_src + iti
KRG     = KK - iout_src + irg
QQSTR   = KK - iout_src + iqs
KCA     = KK - iout_src + idn
KwC     = KK - iout_src + iwc
Kwa    = KK - iout_src + iwa
Ken    = KK - iout_src + ien
C
IF(NI .GT. MAXL) then
    write(lunlog,*) ' CRFG? Time out: ',table(kt)
    CALL trap(5)
    endif
C
RADG(1,NI)  = TABLE(KT)
RADG(2,NI)  = TABLE(KRG)
QSTR(1,NI)  = TABLE(KT)
QSTR(2,NI)  = TABLE(QQSTR)
srcden(1,NI) = TABLE(KT)
srcden(2,NI) = TABLE(KCA)
srcwc(1,NI)  = TABLE(KT)
srcwc(2,NI)  = TABLE(KWC)
srcwa(1,NI)  = TABLE(KT)
srcwa(2,NI)  = TABLE(KWA)
srcenth(1,NI) = TABLE(KT)
srcenth(2,NI) = TABLE(KEN)
C
C*** WRITE THE POINTS JUST RECORDED TO UNIT=8
C
if(isofl.eq. 1) then
    WRITE(8,1140) (TABLE(J),J=kt,kt+5),table(kt+7),table(kt+8)
else
    WRITE(8,1140) (TABLE(J),J=kt,kt+5),table(kt+7),
    1           table(kt+12),table(kt+8)
endif
ispace = ispace + 1
if(ispace.eq. 3) then
    ispace = 0
    write(8,1111)
endif
C
GO TO 100
C
900   CONTINUE          ! EOF encountered

```

```

C
NI = NI + 1
IF(NI+1 .GT. MAXL) then
    write(lunlog,*) ' CRFG? Time out: ',table(iti)
    CALL trap(5)
endif

C
RADG(1,NI) = TABLE(iti)
RADG(2,NI) = TABLE(irg)
QSTR(1,NI) = TABLE(iti)
QSTR(2,NI) = TABLE(iqs)
srcden(1,NI) = TABLE(iti)
srcden(2,NI) = TABLE(idn)
srcwc(1,NI) = TABLE(iti)
srcwc(2,NI) = TABLE(iwc)
srcwa(1,NI) = TABLE(iti)
srcwa(2,NI) = TABLE(iwa)
srcenth(1,NI) = TABLE(iti)
srcenth(2,NI) = TABLE(ien)

C
if(isofl.eq. 1) then
WRITE(8,1140) (TABLE(J),J=1,6),table(8),table(9)
else
WRITE(8,1140) (TABLE(J),J=1,6),table(8),table(13),table(9)
endif
ispace = ispace + 1
if(ispace.eq. 3) then
    ispace = 0
    write(8,1111)
endif

C
NI = NI + 1
DO 910 I =1,2
RADG(I,NI) = POUNDN
QSTR(I,NI) = POUNDN
srcden(I,NI) = POUNDN
srcwc(I,NI) = POUNDN
srcwa(I,NI) = POUNDN
srcenth(I,NI) = POUNDN
910 CONTINUE
C
RETURN
C
1110 FORMAT(' ?CRFG? TABLE exceeded without point selection ',
$'-- execution continuing')
1111 FORMAT(1H )
c1140 FORMAT(1H ,<iout_src>(1PG13.6,1X))
1140 FORMAT(1H ,9(1PG13.6,1X))
END
#####

```

```
C.....  
C  
C      DIMENSIONS/DECLARATIONS for DEGADIS1  
C  
C          include 'sys$degadis:DEGADISIN.dec'  
C  
c  maxl is the length of the /GEN3/ output vectors  
C  
C      parameter (    lunlog= 6,  
$                  sqrtpi= 1.77 245 3851D0,           ! sqrt(pi)  
$                  maxl= 800,  
$                  maxl2= 2*maxl,  
$                  iout_src= 13)  
C  
####
```

PROGRAM DEGADIS1

C*****
C*****
C*****
C
C Program description:
C
C DEGADIS1 estimates the ambient wind profile power alpha and
C characterizes the primary gas source.
C
C
C Program usage:
C
C Consult Volume III of the Final Report to U. S. Coast Guard
C contract DT-CG-23-80-C-20029 entitled "Development of an
C Atmospheric Dispersion Model for Heavier-than-Air Gas Mixtures".
C
C J. A. Havens
C T. O. Spicer
C
C University of Arkansas
C Department of Chemical Engineering
C Fayetteville, AR 72701
C
C April 1985
C
C
C This project was sponsored by the U. S. Coast Guard and the Gas
C Research Institute under contract DT-CG-23-80-C-20029.
C
C
C Disclaimer:
C
C This computer code material was prepared by the University of
C Arkansas as an account of work sponsored by the U. S. Coast Guard
C and the Gas Research Institute. Neither the University of Arkansas,
C nor any person acting on its behalf:
C
C a. Makes any warranty or representation, express or implied,
C with respect to the accuracy, completeness, or usefulness
C of the information contained in this computer code material,
C or that the use of any apparatus, method, numerical model,
C or process disclosed in this computer code material may not
C infringe privately owned rights; or
C
C b. Assumes any liability with respect to the use of, or for
C damages resulting from the use of, any information,
C apparatus, method, or process disclosed in this computer
C code material.

```

C
C*****DIMENSIONS/DECLARATIONS
C
C
Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

      include 'sys$degadis:DEGADIS1.dec'

c
c ntab is the dimension of table divided by iout_src
c
      parameter (      ntab0=910,
$                  ntab=ntab0/iout_src)
c
      include '($ssdef)'

c
c BLOCK COMMON
c
      COMMON
$/GEN3/ radg(2,maxl),qstr(2,maxl),srcden(2,maxl),srcwc(2,maxl),
$ srcwa(2,maxl),srcenth(2,maxl)
$/TITL/ TITLE
$/GEN1/ PTIME(igen), ET(igen), R1T(igen), PWC(igen), PTEMP(igen),
$ PFRACV(igen), PENTH(igen), PRHO(igen)
$/GEN2/ DEN(5,igen)
$/ITI/ T1,TINP,TSRC,TOBS,TSRT
$/ERROR/ STPIN,ERBND,STPMX,WTRG,WTtm,WTya,wtyc,wteb,wtmb,wtuh,XLI,
$ XRI,EPS,ZLOW,STPINZ,ERBNDZ,STPMXZ,SRCOER,srcss,srccut,
$ hcut,ERNOBL,NOBLpt,crfger,epsilon
$/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW
$/SZFC/ szstp0,szerr,szstpmx,szs0
$com_gprop/ gas_nw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc
$/PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND
$/com_ss/ ess,slen,swid,outcc,outsz,outb,outl,swcl,swal,senl,srhl
$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
$/vocom/ vua,vub,vuc,vud,vudelta,vuflag
$/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag
$/com_ENTHAL/ H_masrte,H_airrte,H_watrte
$/NEND/ POUNDN,POUND
$/ALP/ ALPHA,alpha1
$/alphcom/ ialpf1,alpco
$/phicom/ iphifl,dellay
$/sprd_con/ ce, delrhomin
$/COM_SURF/ HTCUTS

```

```

./oomsin/ oodist,avtime
c
c
      character*80 TITLE(4)
c
      character*4 pound
      character*24 TSRC,TINP,TOBS,TSRT
      character*3 gas_name
c
      real*4 tt1
      REAL*8 ML,K
      LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
      logical vuflag
      logical reflag
c
      REAL*8 L,LO
      DIMENSION PRMT(25),Y(7),DERY(7),AUX(8,7)
      EXTERNAL SRC1,SRC10
      character*40 opnrup1
      character OPNRUP(40)
      character*4 INP,ER1,SCD,TR2,scl
c
      dimension table(ntab0)
c
      equivalence(opnrup(1),opnrup1)
c
c.....-----
c
c     DATA
c
      DATA POUNDN/-1.D-20/,POUND'/'//  /
      DATA USTAR/0.D0/,GAMMAF/0.D0/
      DATA G/9.81D0/, K/0.35D0/
      DATA GAMMAF/0.D0/
      DATA PRMT/25*0.D0/
      DATA Y/7*0.D0/, DERY/7*0.D0/
      DATA TIME0/0.D0/,NDIM/0/
      DATA EMAX/0.D0/,TSC1/0.D0/
      DATA PTIME/igen*0.D0/
      DATA ET/igen*0.D0/,R1T/igen*0.D0/
      DATA PWC/igen*0.D0/,PTEMP/igen*0.D0/
      DATA PFRACV/igen*0.D0/,PENTH/igen*0.D0/
      DATA PRHO/igen*0.D0/
      data DEN/igen*0.D0,igen*0.D0,igen*0.D0,igen*0.D0/
c
      data reflag/.true./
c
      DATA INP/'.inp'/,ER1/'.er1'/
      DATA SCD/'.scd'/,TR2/'.tr2'/
      data scl/'.scl'/
c

```

```

C.....  

C  

C      MAIN  

C  

C*** GET THE EXECUTION TIME  

C  

t1 = secnds(0.)  

istat = lib$date_TIME(TSRC)  

if(istat .ne. ss$_normal) stop'lib$date_time failure'  

C  

C*** GET THE FILE NAME FOR FILE CONTROL  

C  

c      WRITE(lunlog,1130)  

c1130 FORMAT(5X,'Enter the file name being used for this run: ',\$)  

      read(5,1000) nchar,opnrup      ! unit 5 gets command file too  

1000  format(q,40a1)  

c  

      opnrup1 = opnrup1(1:nchar) // er1(1:4)  

c  

      CALL ESTRT1(OPNRUP1)  

HTCUTS = HTCUT  

c  

      opnrup1 = opnrup1(1:nchar) // inp(1:4)  

CALL IO(tend,gmass0,OPNRUP1)  

if(check4) srcoer = 2.8*srcoer  

CALL ALPH  

c  

      alpha1 = alpha+1.  

      WRITE(lunlog,1105) ALPHA  

1105 FORMAT(5X,'THE VALUE OF ALPHA IS ',F6.4)  

C  

C  

      GAMMAF = GAMMA(1./ALPHA1)  

TSC1 = TEND  

C  

C  

c*** set the density and enthalpy functions in TPROP  

c  

      call setenthal(h_masrte,h_airrte,h_watrte)  

      call setden(1.000,0.000,h_masrte)  

C  

C.....  

C  

C      SOURCE INTEGRATION (CA = RHOE)  

C  

      opnrup1 = opnrup1(1:nchar) // scd(1:4)  

c  

      OPEN(UNIT=9,NAME=OPNRUP1,recl=202,TYPE='SCRATCH',  

$ carriagecontrol='list',  

$ recordtype='variable')  

c

```

```

gmass = gmass0 .
C
C.....-----
C
C      START THE GAS BLANKET?
C
L = 2.*AFGEN2(PTIME,R1T,TIME0,'R1T-MN')
QSTRE = AFGEN2(PTIME,ET,TIME0,'ET-MN')/(pi*L**2/4.)
PWCP = AFGEN2(PTIME,PWC,TIME0,'PWC')
HPRIM = AFGEN2(PTIME,PENTH,TIME0,'PENTH')
RHOP = AFGEN2(PTIME,PRHO,TIME0,'PRHO')
CCP = PWCP*RHOP

C
C
qstar = 0.
if(u0 .ne. 0.)
$  qstar = CCP*k*ustar*alpha1*dellay/(dellay-1.)/phihat(RHOP,L)
C
C
write(lunlog,3010) time0,l,sz,qstar,qstre
IF(QSTRE.lt.QStAR .and. gmass0.eq.0.) then
    tsc1 = 0.
    GOTO 105
    endif
C
100  CONTINUE
check3 = .false.
C
C*** INITIAL CONDITIONS
C
if(time0.ne. 0.) then
    L0 = 2.*AFGEN2(PTIME,R1T,TIME0,'R1T-MN')
    QSTRE = AFGEN2(PTIME,ET,TIME0,'ET-MN')/(pi*L0**2/4.)
    PWCP = AFGEN2(PTIME,PWC,TIME0,'PWC')
    HPRIM = AFGEN2(PTIME,PENTH,TIME0,'PENTH')
    RHOP = AFGEN2(PTIME,PRHO,TIME0,'PRHO')
    CCP = PWCP*RHOP

C
qstar =0.
if(u0.ne. 0.)
$  qstar = CCP*k*ustar*alpha1*dellay/(dellay-1.)/phihat(RHOP,L0)
C
endif
C
C*** SET UP INTEGRATION PARAMETERS
C
C*** VARIABLE      SUBSCRIPT
C*** -----      -----
C***   RG          Y(1)
C***   mass        Y(2)
C***   massc       Y(3)

```

```

C***      massa          Y(4)
C***      Enthalpy       Y(5)
C***      moe           Y(6)
C***      TIME          X
C
C      PRMT(1) = TIME0
C      PRMT(2) = 6.023E23
C      PRMT(3) = STPIN
C      PRMT(4) = ERBND
C      PRMT(5) = 4.D0*STPMX
C      PRMT(6) = EMAX -- OUTPUT
C
C      do iii = 6,25
C      prmt(iii) = 0.
C      enddo
C
C      Y(1) = AFGEN2(PTIME,R1T,TIME0,'R1T-MN')
C      rmax = y(1)
C      Y(2) = dmax1( gmass, (pi*Y(1)**2*1.1*srccut*RHOP))
C      vuflag = .false.
C      htod = gmass/RHOP/2./pi/Y(1)**3
C      prmt(22) = htod * 2.*Y(1)      ! initial height of the tail
C      prmt(24) = prmt(22)
C      prmt(23) = 0.                  ! initial height of the head
C      prmt(25) = prmt(23)
C      if(htod .gt. 0.1) vuflag = .true.
C      Y(3) = Y(2) * PWCP
C      PWAP = (1. - PWCP)/(1. + HUMID)      ! water from humidity only
C      Y(4) = Y(2) * PWAP
C      Y(5) = Y(2) * HPRIM
C      y(6) = 0.0
C
C      DERY(1) = WTRG
C      DERY(2) = WTtm
C      DERY(3) = WTyC
C      DERY(4) = WTyA
C      DERY(5) = WTeB
C      dery(6) = wtmb
C
C      NDIM = 6
C
C***  PERFORM INTEGRATION
C
C      WRITE(lunlog,1145)
1145  FORMAT(5X,'Beginning Integration Step - Gas blanket')
C
C      CALL RKGST(PRMT,Y,DERY,NDIM,IHLF,SRC1,SRC10,AUX)
C
C      IF(IHLF .GT. 10) CALL trap(1,IHLF)
C      IF(CHECK3) then
C          TEND = TSC1

```

```

        GO TO 110
    end if
c
c.....-----
c
c      RESTART THE GAS BLANKET?
c
    TIME0 = TSC1
    L = 2.*AFGEN2(PTIME,R1T,TIME0,'R1T-MN')
    QSTRE = AFGEN2(PTIME,ET,TIME0,'ET-MN')/(pi*L**2/4.)
    PWCP = AFGEN2(PTIME,PWC,TIME0,'PWC')
    RHOP = AFGEN2(PTIME,PRHO,TIME0,'PRHO')
    CCP = PWCP*RHOP
c
    qstar = 0.
    if(u0.ne. 0.)
    $  qstar = CCP*k*ustar*alpha1*dellay/(dellay-1.)/phihat(RHOP,L)
c
    write(lunlog,3010) time0,l,sz,qstar,qstre
3010  format(//,' time0: ',1pg13.5,t40,'l: ',1pg13.5,
     $/, ' sz0: ',1pg13.5,/, ' qstar: ',1pg13.5,t40,'qstre: ',1pg13.5)

c
c*** before restarting, check to see if this may be a problem child.
c      If the current time is less than SRCSS, this may be a problem
c      simulation. The scheme is to scan the primary source flux for the
c      rest of time. If the flux is never higher, disable the blanket
c      restart feature in NOBL with the logical flag REFLAG, and
c      force the source to use NOBL.
c
    IF(QSTRE .Gt. QSTAR .and. time0.ge.srcss) then
        goto 100

    else if(QSTRE .GT. QSTAR) then

        do iii=1,igen
        if(ptime(iii) .eq. poundn) goto 100
        if(time0 .lt. ptime(iii)) goto 101
        enddo
        goto 100

101      do ii=iii,igen
        if(et(ii) .eq. 0.00) goto 102
        flux = et(ii)/(pi*r1t(ii)**2)
        if(flux .gt. qstre) goto 100
        enddo

102      reflag = .false.
        write(lunlog,1146)
1146      FORMAT(' Forcing NOBL...')

    endif

```

```

C.....  

C  

C      SOURCE INTEGRATION -- NO GAS BLANKET  

C  

105   continue  

      WRITE(lunlog,1147)  

1147   FORMAT(5X,'Source calculation - No Gas blanket')  

C  

      CALL NOBL(timeout, reflag)  

C  

      if(check3) then      ! restart blanket calculation  

          time0 = timeout  

          goto 100  

      end if  

C  

C  

110   RMAX = 1.01*RMAX      ! GUARANTEE A GOOD VALUE  

      aleph = 0.  

      if(u0 .ne. 0.) ALEPH = U0/GAMMAF*(SZM/Z0)**ALPHA  

      $      /(SQRTPI/2.*RM +RMAX)**(ALPHA/ALPHA1)/alpha1  

C  

C  

      rewind (unit=9)  

      opnrup1 = opnrup1(1:nchar) // scl(1:4)  

      open(unit=8,name=opnrup1, type='new',  

      $ carriagecontrol='fortran', recordtype='variable')  

C  

      call head(gmass0)  

      call crfg(table,ntab,crfger)  

      call head(gmass0)  

C  

      CLOSE(UNIT=9)  

      close(unit=8)  

C  

      opnrup1 = opnrup1(1:nchar) // tr2(1:4)  

      CALL TRANS(OPNRUP1)  

C  

C*** CALCULATE EXECUTION TIME  

C  

      tt1 = t1  

      t1 = Secnds(tt1)/60.  

      WRITE(lunlog,2000) TSRC  

      WRITE(lunlog,2010) T1  

2000  FORMAT(1X,'BEGAN AT ',A24)  

2010  FORMAT(5X,' ***** ELAPSED TIME ***** ',1pg13.5,' min ')  

C  

      STOP  

      END  

#####

```

```
C.....  
C  
C  
C      DIMENSIONS/DECLARATIONS for DEGADIS2  
C  
C      include 'sys$degadis:DEGADIS1.dec/list'  
C  
C      MAXNOB IS THE MAXIMUM NUMBER OF OBSERVERS ALLOWED.  
C  
C      parameter(    maxnob = 50,  
$          RT2= 1.41 421 3562D0,   ! sqrt(2.0)  
$          sqpio2= 1.25 331 4137D0)      ! sqrt(pi/2.)  
C  
####
```

PROGRAM DEGADIS2

C*****
C*****
C*****
C*****
C
C Program description:
C
C DEGADIS2 performs the downwind dispersion portion of the calculation
C for each of several observers released successively over the transient
C gas source described by DEGADIS1.
C
C
C Program usage:
C
C Consult Volume III of the Final Report to U. S. Coast Guard
C contract DT-CG-23-80-C-20029 entitled "Development of an
C Atmospheric Dispersion Model for Heavier-than-Air Gas Mixtures".
C
C J. A. Havens
C T. O. Spicer
C
C University of Arkansas
C Department of Chemical Engineering
C Fayetteville, AR 72701
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C April 1985
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C apparatus, method, or process disclosed in this computer
C code material.

```

C
C
C***** ****
C***** ****
C***** ****
C
C
C
c
c
c
C      DIMENSIONS/DECLARATIONS
C
C

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

include 'sys$degadis:DEGADIS2.dec/list'
include '($ssdef)'

C
      COMMON
$/GEN3/ radg(2,maxl),qstr(2,maxl),srcden(2,maxl),srcwc(2,maxl),
$ srcwa(2,maxl),srcenth(2,maxl)
$/SSCON/ NREC(maxnob,2),T0(maxnob),XV(maxnob)
$/TITLE/ TITLE
$/GEN1/ PTIME(igen), ET(igen), R1T(igen), PWC(igen), PTEMP(igen),
$ PFRACV(igen), PENTH(igen), PRHO(igen)
$/GEN2/ DEN(5,igen)
$/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW
$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$/IT1/ T1,TINP,TSRC,TOBS,TSRT
$/ERROR/SY0ER,ERRO,SZ0ER,WTAIO,WTQOO,WTSZO,ERRP,SMXP,
$ WTSZP,WTSYP,WTBEP,WTDH,ERRG,SMXG,ERTDNF,ERTUPF,WTRUH,WTDHG
$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc
$/PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND
$/STP/ STPO,STPP,ODLP,ODLLP,STPG,ODLG,ODLLG
$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
$/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag
$/NEND/ POUNDN,POUND
$/ALP/ ALPHA,alpha1
$/phicom/ iphifl,dellay
$/sprd_con/ ce, delrhomin
$/COM_SURF/ HTCUT
$/STOPIT/ TSTOP
$/CNOBS/ NOBS
./oomsin/ oodist,avtime

C
      character*80 TITLE(4)
C

```

```

character*4 pound
character*24 TINP,TSRC,TOBS,TSRT
character*3 gas_name

c
real*4 tt1
REAL*8 K,ML,L
LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5

c
character*4 TR2,ER2,PSD,TR3, obs
character OPNRUP(40)
character*40 OPNRUP1
equivalence (opnrup(1),opnrup1)

c
c.....
c
c      DATA
c
DATA TSTOP/0./
DATA POUND'//  '//,POUNDN/-1.E-20/
c
DATA TIME0/0./,NDIM/0/
DATA RADG/maxl2*0./,QSTR/maxl2*0./,srcden/maxl2*0./
DATA NREC/maxnob*0,maxnob*0/,T0/maxnob*0./,XV/maxnob*0./

c
DATA TR2/'.TR2',ER2/'.ER2'/
DATA PSD/'.PSD',TR3/'.TR3', obs/'.OBS'/

c
c.....
c
c      MAIN
c
      T1 = SECNDS(0.)
      istat = lib$date_TIME(TOBS)
      if(istat .ne. ss$_normal) stop'lib$date_time failure'

c
C*** GET THE FILE NAME FOR FILE CONTROL
c
c      WRITE(5,1130)
c1130  FORMAT(' Enter the file name being used for this run: ',\$)
      read(5,1130) nchar,opnrup
      1130  format(q,40a1)
c
      opnrup1 = OPNRUP1(1:nchar) // er2(1:4)
      CALL Estrt2(OPNRUP1)

c
C*** GET THE COMMON VARIABLES CARRIED FROM DEGADIS1
c
      opnrup1 = OPNRUP1(1:nchar) // tr2(1:4)
      CALL Strt2(OPNRUP1,H_masrte)
c
c

```

```

C.....  

C  

C      PSEUDO STEADY STATE CALCULATIONS  

C      INTEGRATION IN SUBROUTINE SUPERVISOR  

C  

C          opnrup1 = OPNRUP1(1:nchar) // psd(1:4)  

C          OPEN(UNIT=9,TYPE='NEW',NAME=OPNRUP1,  

C          $ carriagecontrol='list',  

C          $ recordtype='variable')  

C  

C          opnrup1 = OPNRUP1(1:nchar) // obs(1:4)  

C          OPEN( UNIT=12, TYPE='NEW',NAME=OPNRUP1,  

C          $ carriagecontrol='list',  

C          $ recordtype='variable')  

C  

C          CALL SSSUP(H_masrte)  

C  

C.....  

C  

C          CLOSE(UNIT=9)  

C  

C  

C          call setden(1.00,0.00,H_masrte) ! adiabatic mixing w/ pure stuff  

C  

C  

C          opnrup1 = OPNRUP1(1:nchar) // tr3(1:4)  

C          CALL TRANS(OPNRUP1)  

C  

C          tt1 = t1  

C          T1 = SECNDS(tt1)  

C          T1 = T1/60.  

C          WRITE(lunlog,4000) TOBS  

C          WRITE(lunlog,4010) T1  

4000  FORMAT(3X,'BEGAN AT ',A40)  

4010  FORMAT(3X,'*** ELAPSED TIME *** ',1pG13.5,' min')  

C  

STOP  

END  

#####

```

```
c.....  
c  
c      declarations for DEGADIS3  
c  
c      include 'sys$degadis:DEGADIS2.dec/list'  
c  
c      parameter (      maxnt=40,  
c                      maxtnob=maxnt*maxnob)  
c  
#####
```

PROGRAM DEGADIS3

C
C*****
C*****
C*****
C
C Program description:
C
C DEGADIS3 sorts the downwind dispersion calculation made for each of
C the several observers in DEGADIS2. The output concentrations at
C several given times may then be corrected for along-wind dispersion
C as desired.
C
C
C Program usage:
C
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C contract DT-CG-23-80-C-20029 entitled "Development of an
C Atmospheric Dispersion Model for Heavier-than-Air Gas Mixtures".
C
C J. A. Havens
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```

C      code material.
C
C
C***** *****
C***** *****
C***** *****
C
C
C
C

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

include 'sys$degadis:DEGADIS3.dec/list'
include '($ssdef)'

C
C*** MINIMUM DIMENSION ON TABLE IS 6 * MAXNOB + 1
C
C      parameter (ntab0=10*maxnob+1)
C
C      COMMON
$/$ORT/ TCA(maxnob,maxnt),TCASTR(maxnob,maxnt),
$   Tyc(maxnob,maxnt),Trho(maxnob,maxnt),
$   Tgamma(maxnob,maxnt),Ttemp(maxnob,maxnt),
$   TSY(maxnob,maxnt),TSZ(maxnob,maxnt),TB(maxnob,maxnt),
$   TDIST0(maxnob,maxnt),TDIST(maxnob,maxnt),KSUB(maxnt)
$/$SSCON/ NREC(maxnob,2),T0(maxnob),XV(maxnob)
$/$ORTIN/ TIM(maxnt),NTIM,ISTRRT
$/$GEN2/ DEN(5,igen)
$/$PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW
$/$com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$   gas_ufl,gas_lfl,gas_zsp,gas_name
$/$ITI/ T1,TINP,TSRC,TOBS,TSRT
$/$comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
$   humsrc
$/$PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND
$/$PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
$/$com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag
$/$ERROR/ ERT1,ERDT,ERNTIM
$/$NEND/ POUNDN,POUND
$/$ALP/ ALPHA,alpha1
$/$CNOBS/ NOBS
$./oomsin/ oodist,avtime
C
LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
REAL*8 ML,K
DIMENSION TABLE(ntab0)
C
character*24 tsrc,tinp,tobs,tsrt
C
character*3 gas_name

```

```

character*4 TR3,PSD,Er3,SR3,Tr4
character*40 opnrup1
character opnrup(40)

C
      EQUIVALENCE (OPNRUP(1),opnrup1)
C.....
C
C      DATA
C
      DATA POUNDN/-1.E-20/,POUND//   /
      DATA TCA/maxtnob*0./,TCASTR/maxtnob*0./,TSY/maxtnob*0./
      data TSZ/maxtnob*0./,KSUB/maxnt*0/
      DATA TB/maxtnob*0./,TDIST0/maxtnob*0./,TDIST/maxtnob*0./

C
      DATA TR3'!.TR3!',PSD'!.PSD'/
      DATA Er3'!.Er3!',SR3'!.SR3!',Tr4'!.Tr4'/

C
C*** UNITS
C*** 8 -- OUTPUT TO A PRINT FILE
C*** 9 -- I/O WITH DISK
C
      T1 = SECNDS(0.)
      istat = lib$date_time(tsrt)
      if(istat .ne. ss$_normal) stop'lib$date_time failure'

C
C
C*** GET THE FILE NAME FOR FILE CONTROL
C
c      WRITE(5,1130)
c1130  FORMAT(' Enter the file name used for this run: ',$)
      read(5,1130) nchar,opnrup
      1130  format(q,40a1)

C
C*** GET THE VERSION NUMBER
C
c 100  WRITE(5,1140)
c1140  FORMAT(' Enter the version number (between 00 and 99) for',
c     '$' this sort: ',$)
      CALL GTLIN(DUMMY)
      NCAR = LEN(DUMMY)
      IF(NCAR .EQ. 0) GO TO 110
      C
      C      IF(IVERIF(DUMMY,STRING) .NE. 0) GO TO 110
      C      IF(NCAR-2) 130,140,120
      C
      c 110  WRITE(5,1150)
c1150  FORMAT(' ?DEGADIS3? - Invalid character for version number')
      c      GO TO 100
      C
      c 120  WRITE(5,1160)
c1160  FORMAT(' ?DEGADIS3? - Too many characters in the version number')

.
.
```

```

c      GO TO 100
C
c 130  DOT(1) = "060
c      DOT(2) = DUMMY(1)
c      GO TO 150
C
c 140  DOT(1) = DUMMY(1)
c      DOT(2)'= DUMMY(2)
C
c 150  CONTINUE
c      CALL CONCAT(Er3,DOT,Erf3)
c      CALL CONCAT(Sr3,DOT,Sr3)
c      CALL CONCAT(Tr4,DOT,Tr4)
C
C*** NOW, REPLACE THE FILE NAME IN OPNRUP
C
c      CALL SCOPY(BFILE,OPNRUP)
C
C*** THATS IT
C
        opnrup1 = opnrup1(1:nchar) // tr3(1:4)
        CALL STRT3(OPNRUP1)
C
        opnrup1 = opnrup1(1:nchar) // er3(1:4)
        CALL ESTRT3(OPNRUP1)
C
        opnrup1 = opnrup1(1:nchar) // psd(1:4)
        OPEN(UNIT=9,NAME=OPNRUP1,TYPE='OLD')
C
C.....
C
C      TIME SORT SUPERVISOR -- CALCULATE DOWNWIND DISPERSION CORRECTION
C
        CALL SORTS(TABLE)
C
        CLOSE(UNIT=9)
C
C.....
C
C      OUTPUT SORTED PARAMETERS
C
        opnrup1 = opnrup1(1:nchar) // SR3(1:4)
        CALL SRTOUT(OPNRUP1, table)
C
        opnrup1 = opnrup1(1:nchar) // tr4(1:4)
        CALL TRANS(OPNRUP1)
C
        STOP
        END
#####

```

```
c.....  
c  
c      declarations for DEGADIS4  
c  
c      include 'sys$degadis:DEGADIS3.dec/list'  
c  
c      parameter (ndos=10)  
c  
c#####
```

PROGRAM DEGADIS4

C
C*****
C*****
C*****
C
C Program description:
C
C DEGADIS4 sorts the downwind dispersion calculation made for each of
C the several observers in DEGADIS2. The output concentrations at
C several given times may then be corrected for along-wind dispersion
C as desired. DEGADIS4 also outputs the concentration as a function
C of time at a given position.
C
C
C Program usage:
C.
C Consult Volume III of the Final Report to U. S. Coast Guard
C contract DT-CG-23-80-C-20029 entitled "Development of an
C Atmospheric Dispersion Model for Heavier-than-Air Gas Mixtures".
C
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C University of Arkansas
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C Fayetteville, AR 72701
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C April 1985
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```

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C
C
C***** *****
C***** *****
C***** *****
C
C
C
C

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

include 'sys$degadis:DEGADIS4.dec/list'
include '($ssdef)'

C*** MINIMUM DIMENSION ON TABLE IS 6 * MAXNOB + 1
C
parameter (ntab0=10*maxnob+1)
C
COMMON
$/SORT/ TCC(maxnob,maxnt),TCcSTR(maxnob,maxnt),
$  Tyc(maxnob,maxnt),Trho(maxnob,maxnt),
$  Tgamma(maxnob,maxnt),Ttemp(maxnob,maxnt),
$  TSY(maxnob,maxnt),TSZ(maxnob,maxnt),TB(maxnob,maxnt),
$  TD1ST0(maxnob,maxnt),TD1ST(maxnob,maxnt),KSUB(maxnt)
$/SSCON/ NREC(maxnob,2),T0(maxnob),XV(maxnob)
$/cdos/ idos, dosdisx(ndos), dosdis(4,2,ndos)
$/SORTIN/ TIM(maxnt),NTIM,ISTRT
$/GEN2/ DEN(5,igen)
$/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW
$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$  gas_ufl,gas_lfl,gas_zsp,gas_name
$/ITI/ T1,TINP,TSRC,TOBS,TSRT
$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
$  humsrc
$/PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND
$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
$/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag
$/ERROR/ ERT1,ERDT,ERNTIM
$/NEND/ POUNDN,POUND
$/ALP/ ALPHA,alpha1
$/CNOBS/ NOBS

C
LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
REAL*8 ML,K
DIMENSION TABLE(ntab0)

C
character*24 tsrctinp,tobs,tsrt
C

```

```

character*3 gas_name
character*4 TR3,PSD,Er3,SR4,Tr4
character*40 opnrup1
character opnrup(40)

C
EQUIVALENCE (OPNRUP(1),opnrup1)
C.....
C
C      DATA
C
DATA POUNDN/-1.E-20/,POUND'//   '
DATA TCc/maxtnob*0./,TCcSTR/maxtnob*0./,TSY/maxtnob*0./
data TSZ/maxtnob*0./,KSUB/maxnt*0/
DATA TB/maxtnob*0./,TDIST0/maxtnob*0./,TDIST/maxtnob*0./

C
DATA TR3'!.TR3'/,PSD'!.PSD'/
DATA Er3'!.Er3'/,SR4'!.SR4'/,Tr4'!.Tr4'/

C
C*** UNITS
C*** 8 -- OUTPUT TO A PRINT FILE
C*** 9 -- I/O WITH DISK
C
      T1 = SECNDS(0.)
      istat = lib$date_time(tsrt)
      if(istat .ne. ss$_normal) stop'lib$date_time failure'

C
C
C*** GET THE FILE NAME FOR FILE CONTROL
C
      WRITE(lunlog,1130)
1130  FORMAT(' Enter the file name used for this run: ',$)
      read(5,1131) nchar,opnrup
1131  format(q,40a1)

C
C
      opnrup1 = opnrup1(1:nchar) // tr3(1:4)
      CALL STRT3(OPNRUP1)

C
      opnrup1 = opnrup1(1:nchar) // er3(1:4)
      CALL ESTRT3(OPNRUP1)

C
C
C.....
C
C      input the position to calculate the concentration histograms
C
      write(6,*) 'Enter the number of downwind distances desired:'
      write(6,*) ' max of ',ndos,
      1           ' downwind distances; 4 positions at each distance'
      read(5,*) jdos
      if(jdos .gt. ndos) jdos=ndos

```

```

do ii=1,jdos
write(6,*) ''
write(6,*) 'enter the x coordinate:'
read(5,*) dosdisx(ii)
      write(6,*)
1      ' enter the y and z coordinate pairs at this distance:'
      do ij=1,4
      read(5,*) (dosdis(ij,jj,ii),jj=1,2)
      if(dosdis(ij,2,ii).le.0. .and.
1              dosdis(ij,1,ii).le.0.) goto 100
      enddo
100   continue
      enddo
c
c
      opnrup1 = opnrup1(1:nchar) // SR4(1:4)
      open(unit=8,name=opnrup1, type='new',carriagecontrol='fortran')

      do idos=1,jdos
c
c.....-----
c
c      TIME SORT SUPERVISOR -- CALCULATE DOWNWIND DISPERSION CORRECTION
c
      opnrup1 = opnrup1(1:nchar) // psd(1:4)
      OPEN(UNIT=9,NAME=OPNRUP1,TYPE='OLD')
c
      do ij=1,maxnt
      KSUB(ij) = 0
      do ijk=1,maxnob
      TSZ(ijk,ij) = 0.
      enddo
      enddo
c
c
      CALL SORTS(TABLE)
c
      CLOSE(UNIT=9)
c
c.....-----
c
c      OUTPUT SORTED PARAMETERS
c
      CALL dosOUT(table)
c
      enddo
c
      close(unit=8)
c
c

```

```
C.....  
C  
c      declarations for DEGADISIN  
C  
C  
      parameter(  igen= 30, ! dimension of /gen1/ and /gen2/  
$          pi= 3.14 159 265 358 2400)  
c  
####
```

C*****
C*****
C*****
C
C PROGRAM DEGADISIN
C
C Program description:
C
C DEGADISIN acts as an interactive input module to the programs
C which make up the DEGADIS model. The user is guided through a
C series of questions which supply the model with the necessary
C input information.
C
C
C Program usage:
C
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C contract DT-CG-23-80-C-20029 entitled "Development of an
C Atmospheric Dispersion Model for Heavier-than-Air Gas Mixtures".
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```

C      code material.
C
C
C***** *****
C***** *****
C***** *****
C
C
C
C      INITIAL INPUT FOR DEGADIS ROUTINES
C
C      note: this series of programs relies on the system wide
C             logical symbol SYS$DEGADIS which denotes the source
C             and executable code for these images.
C
C
C      PROGRAM DEGADISIN

      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

C
C      include 'SYS$DEGADIS:degadisin.dec'
C
C
C      COMMON
C/TITLE/ TITLE
C/GEN1/ PTIME(igen), ET(igen), R1T(igen), PWC(igen), PTEMP(igen),
C       PFRACV(igen), PENTH(igen), PRHO(igen)
C/GEN2/ DEN(5,igen)
C/ITI/ T1,TINP,TSRC,TOBS,TSRT
C/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW
C/com_gprop/ gas_mm,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
C   gas_ufl,gas_lfl,gas_zsp,gas_name
C/com_ss/ ess,slen,swid,outcc,outsz,outb,outl
C/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
C/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag
C/NEND/ POUNDN,POUND
C   ./oomsin/ oodist,avtime
C
C      character*80 TITLE(4)
C
C      character*3 gas_name
C      character*4 pound
C      character*24 TSRC,TINP,TOBS,TSRT
C
C      REAL*8 ML,K
C      LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
C
C      c check1
C      c check2=t      cloud type release with no liquid source; SRC1  DEGADIS1
C      c again        local communications in SSSUP                   SSSUP
C      c check3        local communications between SRC1 and NOBL    DEGADIS1

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```

c check4=t      steady state simulation          DEGADISIN
c check5=t      operator sets sort parameters    ESTRT3
c
c           data CHECK1/.false./,CHECK2/.false./,AGAIN/.false./
c           data CHECK3/.false./,CHECK4/.false./,CHECK5/.false./
c
c           character*100 OPNRUP
c           character OPNRUP1(100)
c           equivalence (opnrup1(1),opnrup)
c           character*4 INP,er1,er2,er3,com,scl,sr3,lis
c           character*4 dummy
c           character*3 plus
c           character*2 con
c           DATA POUND//  '//,POUNDN/-1.E-20/
c
c           DATA PTIME/igen*0.D0/
c           DATA ET/igen*0.D0/,R1T/igen*0.D0/
c           DATA PWC/igen*0.D0/,PTEMP/igen*0.D0/
c           DATA PFRACV/igen*0.D0/,PENTH/igen*0.D0/
c           DATA PRHO/igen*0.D0/
c           data DEN/igen*0.,igen*0.,igen*0.,igen*0.,igen*0.//
c           DATA INP//.INP'/,er1'/.er1/,er2'/.er2/,er3'/.er3'/
c           data scl'/.scl'/.sr3'/.sr3'/.lis'/.lis'/
c           data com'/.com'/
c           data plus'/' + '/,con'/' - '/
c
c
c*** GET THE FILE NAME TO BE USED BY ALL OF THE ROUTINES
c
c           WRITE(6,800)
c           WRITE(6,810)
c           READ(5,820) NCHAR,opnrup
c           opnrup = opnrup(1:nchar) // inp(1:4)
c
c
c*** NOW GET THE REST OF THE DESIRED INFORMATION
c
c           CALL IOT(OPNRUP)
c           WRITE(6,1000)
c           if(check4) then
c               write(6,1001) ! continuous
c           else
c               if(u0 .eq. 0.) then
c                   write(6,1009)
c               else
c                   WRITE(6,1002) ! transient
c               endif
c           endif
c           write(6,1010)
c
c
c*** FORMATS
c
c           800   FORMAT(//,16X,'DEnse GAs DISpersion Model input module.')

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```

810  FORMAT(/, ' Enter the simulation name',
      $' : [DIR]RUNNAME ', $)
820  FORMAT(Q,A40)
1000 FORMAT(' ',,
      $' In addition to the information just obtained,'.
      $' DEGADIS',/, ' requires a series of numerical parameter',
      $' files which use',/, ' the',
      $' same name as [DIR]RUNNAME given above. ',/,
      $' For convenience, example parameter files are included for',/,
      $' each step. They are: ')
1001 FORMAT(10X,'EXAMPLE.ER1 and',/,10X,'EXAMPLE.ER2')
1002 format(10X,'EXAMPLE.ER1',/,10X,'EXAMPLE.ER2, and',/,
      $10X,'EXAMPLE.ER3')
1009 format(10x,'EXAMPLE.ER1')
1010 format(' Note that each of',
      $' these files can be edited during the course of the',/,
      $' simulation if a parameter proves to be out of specification.',/)
c
c
      write(6,1200)
1200 format(' Do you want a command file to be generated to execute',
      $' the procedure? <Y or n> ', $)
      REad(5,1210) dummy
1210 format(a4)
      if(dummy.eq.'n' .or. dummy.eq.'N') goto 3000
      opnrup = opnrup(1:nchar) // com(1:4)
      write(6,1220) opnrup
1220 format(' The command file will be generated under the file',
      $' name:',/,10x,a40)
c
      open(unit=8,name=opnrup,type='new',
      $ carriagecontrol='list',recordtype='variable')
c
      opnrup = opnrup(1:nchar) // er1(1:4)
c
      write(8,1250) (opnrup1(i),i=1,nchar+4)
1250 format('$ copy/log SYSS$DEGADIS:example.er1 ',40a1)
      IF(u0 .eq. 0.) then
          write(8,1280)
          write(8,1290) (opnrup1(i),i=1,nchar)
          goto 1340
      endif
      opnrup = opnrup(1:nchar) // er2(1:4)
c
      write(8,1260) (opnrup1(i),i=1,nchar+4)
1260 format('$ copy/log SYSS$DEGADIS:example.er2 ',40a1)
      opnrup = opnrup(1:nchar) // er3(1:4)
c
      if(.not.check4) then           ! transient
c
      write(8,1270) (opnrup1(i),i=1,nchar+4)

```

```

1270      format('$ copy/log SYSS$DEGADIS:example.er3 ',40a1)
c
1280      write(8,1280)
1290      format('$ run SYSS$DEGADIS:DEGADIS1')
write(8,1290) (opnrup1(i),i=1,nchar)
1290      format(40a1)
1300      write(8,1300)
1300      format('$ run SYSS$DEGADIS:DEGADIS2')
write(8,1290) (opnrup1(i),i=1,nchar)
1320      write(8,1320)
1320      format('$ run SYSS$DEGADIS:DEGADIS3')
write(8,1290) (opnrup1(i),i=1,nchar)
c
1     opnrup = opnrup(1:nchar) // scl(1:4) //
plus(1:3) // opnrup(1:nchar) // sr3(1:4) // con(1:2)
1370      write(8,1370) (opnrup1(i),i=1,2*nchar+13)
format('$ copy/log ',100a1)
c
1     opnrup = opnrup(1:nchar) // lis(1:4)
write(8,1390) (opnrup1(i),i=1,nchar+4)
1390      format(' ',40a1)
else
    write(8,1280)
    write(8,1290) (opnrup1(i),i=1,nchar)
c
1     write(8,1330)
1330      format('$ run SYSS$DEGADIS:DEGADIS2')
write(8,1290) (opnrup1(i),i=1,nchar)
c
1     opnrup = opnrup(1:nchar) // scl(1:4) //
plus(1:3) // opnrup(1:nchar) // sr3(1:4) // con(1:2)
write(8,1370) (opnrup1(i),i=1,2*nchar+13)
opnrup = opnrup(1:nchar) // lis(1:4)
write(8,1390) (opnrup1(i),i=1,nchar+4)
endif
c
1340  close(unit=8)
write(6,1350)
1350  format(/,' Do you wish to initiate this procedure? ',
'$ <y or N> ',$)
REad(5,1210) dummy
if(dummy.eq.'y' .or. dummy.eq.'Y') goto 2000
goto 3000
2000  opnrup = 'a' // opnrup(1:nchar) // ' '
istat = lib$do_command(opnrup)
write(6,2100)
2100  format(/,' ?DEGADISIN? command file failed to start.')
c
3000  continue
CALL EXIT
END

```

```

SUBROUTINE dosOUT(table)
c
Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

include 'sys$degadis:DEGADIS4.dec/list'
c
COMMON /SORT/TCc(maxnob,maxnt),TCcSTR(maxnob,maxnt),
$ Tyc(maxnob,maxnt),Trho(maxnob,maxnt),
$ Tgamma(maxnob,maxnt),Ttemp(maxnob,maxnt),
$ TSY(maxnob,maxnt),TSZ(maxnob,maxnt),TB(maxnob,maxnt),
$ TDIST0(maxnob,maxnt),TDIST(maxnob,maxnt),KSUB(maxnt)
$/cdos/ idos, dosdisx(ndos), dosdis(4,2,ndos)
$/SORTIN/TIM(maxnt),NTIM,ISTRTRT
$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$/comatm/ istab,tamb,pamb,humid,isoftl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc
$/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag
$/alp/ alpha,alpha1
c
dimension table(1)
dimension chist(4,maxnt)
c
logical cflag,cflag1
c
character*3 gas_name
c
if(sigx_flag.eq. 0.) then
    write(8,1102)
else
    write(8,1104)
    write(8,1105) sigx_coeff,sigx_pow,sigx_min_dist
endif
c
cflag = isoftl.eq. 1.or. ihtfl.eq. 0
cflag1= isoftl.eq.1
if(cflag) then
    call adiabat(2,wc,wa,gas_lfl,ya,cc_lfl,r,w,t,tt)
    call adiabat(2,wc,wa,gas_ufl,ya,cc_ufl,r,w,t,tt)
endif
c
WRITE(8,1110) dosdisx(idos)
c
WRITE(8,1119)
WRITE(8,1119)
if(cflag1) then
    WRITE(8,1116) gas_zsp,(100.*gas_lfl),(100.*gas_ufl)
    WRITE(8,1118)
else
    WRITE(8,1115) gas_zsp,(100.*gas_lfl),(100.*gas_ufl)

```

```

        WRITE(8,1117)
        endif
    WRITE(8,1119)
    ip = 0
c
    DO 110 I=ISTRRT,NTIM
c
    II = KSUB(I)
c
    j = 1
    disto = tdist(j,i)
    cco = tccstr(j,i)
    rho = Trho(j,i)
    yco = Tyc(j,i)
    tempo = Ttemp(j,i)
    gammao = Tgamma(j,i)
    bo = tb(j,i)
    szo = tsz(j,i)
    syo = tsy(j,i)
    if( disto .gt. dosdisx(idos) ) then
        write(lunlog,*) ' Extrapolated point for time: ',Tim(i)
        write(8,*) ' Records for ',tim(i),' s are missing - see source'
        goto 110
        endif
c
    DO 120 J=2,II
c
    dist = tdist(j,i)
    cc = tccstr(j,i)
    rho = Trho(j,i)
    yc = Tyc(j,i)
    temp = Ttemp(j,i)
    gamma = Tgamma(j,i)
    b = tb(j,i)
    sz = tsz(j,i)
    sy = tsy(j,i)
    blfl = 0.
    bufl = 0.

    if( dist .lt. dosdisx(idos) ) goto 119
    dist = (dosdisx(idos) - disto)/(dist-disto)
    cc = dist*(cc-cco) + cco
    rho = dist*(rho-rho) + rho
    yc = dist*(yc-yco) + yco
    temp = dist*(temp-tempo) + tempo
    gamma = dist*(gamma-gammao) + gammao
    b = dist*(b-bo) + bo
    sz = dist*(sz-szo) + szo
    sy = dist*(sy-syo) + syo
c
    if(.not.cflag) then

```

```

call adiabat(-2,wc,wa,gas_lfl,ya,cc_lfl,r,w,gamma,tt)
call adiabat(-2,wc,wa,gas_ufl,ya,cc_ufl,r,w,gamma,tt)
endif
c
c*** calc the concentration time histories
c
do ij=1,4
    chist(ij,i) = 0.
    if(dosdis(ij,2,idos) .ge. 0.) then
        arg = (dosdis(ij,2,idos)/sz)**alpha1
        if(dosdis(ij,1,idos) .gt. b) arg = arg +
            1           ((dosdis(ij,1,idos)-b)/sy)**2
        if(arg .lt. 80.) then
            chist(ij,i) = cc/exp(arg)
        if(cflag) then
            call adiabat(0,xx,aa,ycc,yaa,chist(ij,i),rr,ww,gg,tt)
        else
            call adiabat(-1,xx,aa,ycc,yaa,chist(ij,i),rr,ww,gamma,tt)
        endif
        chist(ij,i) = ycc
        endif
    endif
enddo
c
c
arg = (gas_zsp/sz)**alpha1
if(arg .ge. 80.) then
    if(cflag1) then
        WRITE(8,1120) tim(I),yc,Cc,rho,temp,B,SZ,SY
    else
        WRITE(8,1120) tim(I),yc,Cc,rho,gamma,temp,B,SZ,SY
    endif
    goto 600
endif
c
ccz = cc/exp(arg)
if(ccz .lt. cc_lfl) then
    if(cflag1) then
        WRITE(8,1120) tim(I),yc,Cc,rho,temp,B,SZ,SY
    else
        WRITE(8,1120) tim(I),yc,Cc,rho,gamma,temp,B,SZ,SY
    endif
    goto 600
endif
arg = -(dlog(cc_lfl/cc) + (gas_zsp/sz)**alpha1)
blfl = sqrt(arg)*sy + b
c
if(ccz .lt. cc_ufl) then
    if(cflag1) then
        WRITE(8,1120) tim(I),yc,Cc,rho,temp,B,SZ,SY,blfl
    else

```

```

      WRITE(8,1120) tim(I),yc,Cc,rho,gamma,temp,B,SZ,SY,blfl
                      endif
      goto 600
      endif
      arg = -(dlog(cc_ufl/cc) + (gas_zsp/sz)**alpha1) .
      bufl = sqrt(arg)*sy + b
      if(cflag1) then
      WRITE(8,1120) tim(I),yc,Cc,rho,temp,B,SZ,SY,blfl,bufl
      else
      WRITE(8,1120) tim(I),yc,Cc,rho,gamma,temp,B,SZ,SY,blfl,bufl
      endif
c
      600 continue
      goto 121
      119 continue
      disto = tdist(j,i)
      cco   = tccstr(j,i)
      rho0  = Trho(j,i)
      yco   = TyC(j,i)
      tempo = Ttemp(j,i)
      gamma0 = Tgamma(j,i)
      bo    = tb(j,i)
      szo   = tsz(j,i)
      syo   = tsy(j,i)
      120 CONTINUE
c
      121 ip = ip + 1
      if(ip .eq. 3) then
          ip = 0
          write(8,1119)
      endif
      110 CONTINUE
c
c*** output the concen time histories
c
      write(8,1119)
      write(8,1119)
      iii = 0
      do ij=1,4
          if(dosdis(ij,2,idos).gt.0. .and.
1                   dosdis(ij,1,idos).gt.0.) iii = ij
      enddo

      if(iii .eq. 0) return
      ip = 0

      write(8,2100) ((dosdis(ij,ijj,idos),ij=1,4),ijj=1,2)
2100  format(1H0,11x,'Time',11x,4(4x,'Mole fraction at:',5x),/,
1           1H ,26x,4(4x,'y= ',1pg13.5,' m',4x),/,
1           1H ,11x,'(s)',12x,4(4x,'z= ',1pg13.5,' m',4x))
      DO I=ISTRRT,NTIM

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      write(8,2200) tim(i),(chist(ij,i),ij=1,iii)
      ip = ip + 1
      if(ip .eq. 3) then
          ip = 0
          write(8,1119)
          endif
      enddo
2200  format(1h ,5(6x,1pg14.7,6x))
C
C
1102  format(1H0,5x,'X-Direction correction was NOT applied.')
1104  format(1H0,5x,'X-Direction correction was applied.')
1105  format(1h ,5x,5x,'Coefficient:      ',1pg13.5,/,
           1      1h ,5x,5x,'Power:            ',1pg13.5,/,
           1      1h ,5x,5x,'Minimum Distance: ',1pg13.5' m')
1110  FORMAT(1H0,5X,'Centerline values for the position -->',/,
           1      ' x: ',1pg13.5,' m',/)
1115  FORMAT(1H0,1X,' Time ',2x,3x,'Mole',3x,
           1      'Concentration',1x,'Density',2x,3x,'Gamma',3x,
           1      'Temperature',3x,'Half',4x,4x,'Sz',5x,4x,'Sy',5x,
           1      'Width at z=',0pf6.2,' m to:',/,1x,11x,1x'Fraction',2x,
           1      11x,11x,11x,3x,'Width',3x,11x,9x,
           1      2(1pg9.3,'mole%',1x))
1116  FORMAT(1H0,1X,' Time ',2x,3x,'Mole',3x,
           1      'Concentration',1x,'Density',2x,
           1      'Temperature',3x,'Half',4x,4x,'Sz',5x,4x,'Sy',5x,
           1      'Width at z=',0pf6.2,' m to:',/,1x,11x,1x'Fraction',2x,
           1      11x,11x,11x,3x,'Width',3x,11x,9x,
           1      2(1pg9.3,'mole%',1x))
1117  FORMAT(1H ,4X,'(s)',4x,11x,
           1      2(1X,'(kg/m**3)',1x),11x,4x,'(K)',
           1      5(8X,'(m)'))
1118  FORMAT(1H ,4X,'(s)',4x,11x,
           1      2(1X,'(kg/m**3)',1x),4x,'(K)',
           1      5(8X,'(m)'))
1119  FORMAT(1H )
1120  FORMAT(1H ,3(1X,1PG9.3,1X),2x,0pf7.4,2x,1X,1PG10.3,1X,
           1      6(1X,1PG9.3,1X))

C
      RETURN
      END
#####

```

```

C.....  

C  

C      ROUTINE TO GET RUN PARAMETERS FROM A FILE  

C  

C      SUBROUTINE ESTRT1(OPNRUP)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      include 'sys$degadis:DEGADIS1.dec'  

C  

parameter( iend= 22,  

           1         iend1= iend+1,  

           1         iiiend= 2,  

           1         iiiend1= iiiend+1,  

           1         iiend= 2,  

           1         iiend1= iiend+1,  

           1         jend= 4,  

           1         jend1= jend+1,  

           1         mend= 5,  

           1         mend1= mend+1)  

C  

C      BLOCK COMMON  

C  

COMMON  

$ERROR/STPIN,ERBND,STPMX,WTRG,WTtm,WTya,WTyc,WTeb,WTmb,WTuh,XLI,  

$ XRI,EPS,ZLOW,STPINZ,ERBNDZ,STPMXZ,SRCOER,srss,srcut,  

$ hcut,ERNOBL,NOBLpt,crfger,epsilon  

$vucom/ vua,vub,vuc,vud,vuedelta,vuflag  

$szfc/ szstp0,szerr,szstpmx,szs0  

$/alphcom/ ialpfl,alpc0  

$/phicom/ iphifl,delay  

$/sprd_con/ ce, delrhomin  

C  

EQUIVALENCE  

$(RLBUF(1),STPIN), !MAIN - RKGST - INITIAL STEP SIZE  

$(RLBUF(2),ERBND), !MAIN - RKGST - ERROR BOUND  

$(RLBUF(3),STPMX), !MAIN - RKGST - MAXIMUM STEP SIZE  

$(RLBUF(4),WTRG), !MAIN - RKGST - WEIGHT FOR RG  

$(RLBUF(5),WTtm), !MAIN - RKGST - WEIGHT FOR Total mass  

$(RLBUF(6),WTya), !MAIN - RKGST - WEIGHT FOR Ya  

$(RLBUF(7),WTyc), !MAIN - RKGST - WEIGHT FOR Yc  

$(RLBUF(8),WTeb), !MAIN - RKGST - WEIGHT FOR Energy Balance  

$(RLBUF(9),WTmb), !MAIN - RKGST - WEIGHT FOR Momentum Balance  

$(RLBUF(10),WTuh), !MAIN - RKGST - WEIGHT FOR Ueff*Heff  

$(RLBUF(11),XLI), !ALPH - LOWER LIMIT OF SEARCH FOR ALPHA  

$(RLBUF(12),XRI), !ALPH - UPPER LIMIT OF SEARCH FOR ALPHA  

$(RLBUF(13),EPS)  !ALPH - ERROR BOUND USED BY "RTMI"  

      equivalence  

$(RLBUF(14),ZLOW), !ALPHI - BOTTOM HEIGHT FOR FIT OF ALPHA  

$(RLBUF(15),STPINZ), !ALPHI - INITIAL RKGST STEP <0.

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$(RLBUF(16),ERBNDZ), !ALPHI - ERROR BOUND FOR RKGST
$(RLBUF(17),STPMXZ), !ALPHI - MAXIMUM STEP FOR RKGST
$(RLBUF(18),SRCOER), !SRC10 - OUTPUT error criteria
$(RLBUF(19),SRCSS), !SRC10 - min time for Steady;4*STPMX
$(RLBUF(20),SRCcut), !SRC10 - min height for blanket
$(RLBUF(21),htcut), !SRC1 - min height for blanket heat transfer
$(RLBUFa(iend),ERNOBL), !NOBL - CONVERGENCE CRITERIA
$(RLBUFi(iend),crfger), !CRFG - Error criteria for building tables
$(RLBUFi(iend),epsilon)!SRC1 - Coefficient in Air entrainment
c
      equivalence
$(rlbufa(1),ce), !SRC1 - Coefficient gravity slumping EQ
$(RLBUFa(iend),delrhomin) ! stop spread for delrho<delrhomin
c
      equivalence
$(rlbuf1(1),szstp0), !SZF - Initial step size
$(rlbuf1(2),szerr), !SZF - Error criteria
$(rlbuf1(3),szstpmx), !SZF - Maximum step size
$(rlbuf1(4),szsz0) !SZF - Initial value of rho*dellay*UHeff
c
c
      equivalence
$(rlbuf4(1),vua), ! Constant Av in SRC1
$(rlbuf4(2),vub), ! Constant Bv in SRC1
$(rlbuf4(3),vuc), ! Constant Ev in SRC1
$(rlbuf4(4),vud), ! Constant Dv in SRC1
$(rlbuf4(5),vudelta) ! Constant DELTAv in SRC1
c
      character*40 OPNRUP
      character DUMMY(1:132)
      DIMENSION RLBUF(iend), rlbufi(iend), rlbufa(iend)
      dimension rlbuf1(jend)
      dimension rlbuf4(mend)
c
      logical vuflag
c
      OPEN(UNIT=9,NAME=OPNRUP,TYPE='OLD',err=2000)
c
C*** READ A LINE AND DETERMINE ITS PURPOSE
c
      I = 1
100   CONTINUE
      READ(9,1000,END=350) NCHAR,DUMMY
      IF(DUMMY(1) .EQ. '!') GO TO 100
      DECODE(20,1010,DUMMY,ERR=400) RLBUF(I)
      I = I + 1
      if(i .eq. iend1) goto 110
      GO TO 100

110   CONTINUE
      READ(9,1000,END=350) NCHAR,DUMMY

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IF(DUMMY(1) .EQ. '!!') GO TO 110
DECODE(20,1010,DUMMY,ERR=400) PTNOBL
NOBLPT = INT(PTNOBL)

I = 1
120 CONTINUE
READ(9,1000,END=350) NCHAR,DUMMY
IF(DUMMY(1) .EQ. '!!') GO TO 120
DECODE(20,1010,DUMMY,ERR=400) RLBUFi(I)
I = I + 1
if(i .eq. iiend1) goto 140
GO TO 120

C
C*** READ A LINE AND DETERMINE ITS PURPOSE for /sprd_con/
C
140 I = 1
150 CONTINUE
READ(9,1000,END=350) NCHAR,DUMMY
IF(DUMMY(1) .EQ. '!!') GO TO 150
DECODE(20,1010,DUMMY,ERR=400) RLBUFa(I)
I = I + 1
if(i .eq. iiend1) goto 190
GO TO 150

C
C*** READ A LINE AND DETERMINE ITS PURPOSE to fill szfc
C
190 I = 1
200 CONTINUE
READ(9,1000,END=350) NCHAR,DUMMY
IF(DUMMY(1) .EQ. '!!') GO TO 200
DECODE(20,1010,DUMMY,ERR=400) RLBUF1(I)
I = I + 1
if(i .eq. jend1) goto 230
GO TO 200

C
C*** READ A LINE AND DETERMINE ITS PURPOSE to fill /alphcom/
C
230 I = 1
240 CONTINUE
READ(9,1000,END=350) NCHAR,DUMMY
IF(DUMMY(1) .EQ. '!!') GO TO 240
DECODE(20,1010,DUMMY,ERR=400) Ralpf1

ialpf1 = int(ralpf1)

250 CONTINUE
READ(9,1000,END=350) NCHAR,DUMMY
IF(DUMMY(1) .EQ. '!!') GO TO 250
DECODE(20,1010,DUMMY,ERR=400) alpco

C
C*** READ A LINE AND DETERMINE ITS PURPOSE to fill /phicom/

```

```

C
260  I = 1
270  CONTINUE
      READ(9,1000,END=350) NCHAR,DUMMY
      IF(DUMMY(1) .EQ. '!') GO TO 270
      DECODE(20,1010,DUMMY,ERR=400) Rphifl

      iphifl = int(rphifl)

275  CONTINUE
      READ(9,1000,END=350) NCHAR,DUMMY
      IF(DUMMY(1) .EQ. '!') GO TO 275
      DECODE(20,1010,DUMMY,ERR=400) dellay

C
C*** READ A LINE AND DETERMINE ITS PURPOSE to fill /vocom/
C
280  I = 1
290  CONTINUE
      READ(9,1000,END=350) NCHAR,DUMMY
      IF(DUMMY(1) .EQ. '!') GO TO 290
      DECODE(20,1010,DUMMY,ERR=400) RLBUF4(I)
      I = I + 1
      if(i .eq. mend1) goto 300
      GO TO 290

C
C*** EXIT THE PROCEEDINGS
C
300  CONTINUE
      CLOSE(UNIT=9)
      RETURN

C
350  call trap(20)
C
400  CONTINUE
      CALL trap(21)
C
1000 FORMAT(Q,132A1)
1010 FORMAT(10X,G10.4)
C
2000 call trap(22)
      END

#####

```

```

C.....  

C  

C      ROUTINE TO GET RUN PARAMETERS FROM A FILE  

C  

C      SUBROUTINE ESTRT2(OPNRUP)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      include 'sys$degadis:DEGADIS2.dec/list'  

C  

parameter ( ienda= 18,  

           1          ienda1= ienda+1,  

           2          iendb= 7,  

           3          iendb1= iendb+1)  

C  

COMMON  

$/ERROR/SY0ER,ERRO,SZOER,WTAIO,WTQOO,WTSZO,ERRP,SMXP,  

$ WTSZP,WTSYP,WTBEP,WTDH,ERRG,SMXG,ERTDNF,ERTUPF,WTRUH,WTDHG  

$/STP/STPO,STPP,ODLP,ODLLP,STPG,ODLG,ODLLG  

$/CNOBS/NOBS  

C  

EQUIVALENCE  

$(RLBUF(1),SY0ER), !SSSUP - RKGST - INITIAL SY  

$(RLBUF(2),ERRO), !SSSUP - RKGST(OBS) - ERROR BOUND  

$(RLBUF(3),SZOER), !SSSUP - RKGST(OBS) - INITIAL SZ  

$(RLBUF(4),WTAIO), !SSSUP - RKGST(OBS) - WEIGHT FOR AI  

$(RLBUF(5),WTQOO), !SSSUP - RKGST(OBS) - WEIGHT FOR Q  

$(RLBUF(6),WTSZO), !SSSUP - RKGST(OBS) - WEIGHT FOR SZ  

$(RLBUF(7),ERRP), !SSSUP - RKGST(PSS) - ERROR BOUND  

$(RLBUF(8),SMXP), !SSSUP - RKGST(PSS) - MAXIMUM STEP  

$(RLBUF(9),WTSZP), !SSSUP - RKGST(PSS) - WEIGHT FOR SZ  

$(RLBUF(10),WTSYP), !SSSUP - RKGST(PSS) - WEIGHT FOR SY  

$(RLBUF(11),WTBEP), !SSSUP - RKGST(PSS) - WEIGHT FOR BEFF  

$(RLBUF(12),WTDH), !SSSUP - RKGST(PSS) - WEIGHT FOR DH  

$(RLBUF(13),ERRG), !SSSUP - RKGST(SSG) - ERROR BOUND  

$(RLBUF(14),SMXG), !SSSUP - RKGST(SSG) - MAXIMUM STEP SIZE  

$(RLBUF(15),ERTDNF), !TDNF - CONVERGENCE CRITERIA  

$(RLBUF(16),ERTUPF), !TUPF - CONVERGENCE CRITERIA  

$(RLBUF(17),WTRUH), !SSSUP - RKGST(SSG) - WEIGHT FOR RUH  

$(RLBUF(ienda),WTdhg)!SSSUP - RKGST(SSG) - WEIGHT FOR DH  

C  

EQUIVALENCE  

$(RLBUF1(1),STPO), !SSSUP - RKGST(OBS) - INITIAL STEP  

$(RLBUF1(2),STPP), !SSSUP - RKGST(PSS) - INITIAL STEP  

$(RLBUF1(3),ODLP), !SSSUP - RKGST(PSS) - RELATIVE OUTPUT DELTA  

$(RLBUF1(4),ODLLP), !SSSUP - RKGST(PSS) - MAXIMUM DISTANCE TO OUT  

$(RLBUF1(5),STPG), !SSSUP - RKGST(SSG) - INITIAL STEP  

$(RLBUF1(6),ODLG), !SSSUP - RKGST(SSG) - RELATIVE OUTPUT DELTA  

$(RLBUF1(iendb),ODLLG)!SSSUP - RKGST(SSG) - MAXIMUM DISTANCE TO OUT
C

```

```

character*40 OPNRUP
character dummy(1:132)
DIMENSION RLBUF(ienda),RLBUF1(iendb)

C
OPEN(UNIT=9,NAME=OPNRUP,TYPE='OLD')

C
C*** FIRST, FILL RLBUF
C
C*** READ A LINE AND DETERMINE ITS PURPOSE
C
I = 1
100 CONTINUE
READ(9,1000,END=300) NCHAR,DUMMY
IF(DUMMY(1) .EQ. '!!') GO TO 100
DECODE(20,1010,DUMMY,ERR=400) RLBUF(I)
I = I + 1
IF(I .EQ. ienda1) GO TO 200
GO TO 100

C
C*** NOW, FILL RLBUF1
C
200 I = 1
210 CONTINUE
READ(9,1000,END=300) NCHAR,DUMMY
IF(DUMMY(1) .EQ. '!!') GO TO 210
DECODE(20,1010,DUMMY,ERR=400) RLBUF1(I)
I = I + 1
IF(I .EQ. iendb1) GO TO 260
GO TO 210

C
C*** NOW, PICK UP NOBS
C
260 CONTINUE
READ(9,1000,END=300) NCHAR,DUMMY
IF(DUMMY(1) .EQ. '!!') GO TO 260
DECODE(20,1010,DUMMY,ERR=400) RBUF
NOBS = INT(RBUF)

C
C*** EXIT THE PROCEEDINGS
C
CLOSE(UNIT=9)
RETURN

C
300 call trap(20)
400 CALL trap(21)

C
1000 FORMAT(Q,132A1)
1010 FORMAT(10X,G10.4)
END

#####

```

```

C.....  

C  

C      ROUTINE TO GET RUN PARAMETERS FROM A FILE  

C  

C          SUBROUTINE ESTRT2SS(OPNRUP)  

C  

C          Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C          include 'sys$degadis:DEGadis2.dec/list'  

C  

C          parameter(      ienda= 18,  

C                         1      ienda1= ienda+1,  

C                         2      iendb= 7,  

C                         3      iendb1= iendb+1)  

C  

C          COMMON  

C          $/ERROR/SY0ER,ERRP,SMXP,WTSZP,WTSYP,WTBEP,WTDH,ERRG,SMXG,  

C          $ WTRUH,WTDHG  

C          $/STP/STPP,ODLP,ODLLP,STPG,ODLG,ODLLG  

C  

C          character*40 OPNRUP  

C          character DUMMY(1:132)  

C          DIMENSION RLBUF(ienda),RLBUF1(iendb)  

C  

C          OPEN(UNIT=9,NAME=OPNRUP,TYPE='OLD')  

C  

C*** FIRST, FILL RLBUF  

C  

C*** READ A LINE AND DETERMINE ITS PURPOSE  

C  

C          I = 1  

100    CONTINUE  

        READ(9,1000,END=350) NCHAR,DUMMY  

        IF(DUMMY(1) .EQ. '!!') GO TO 100  

        DECODE(20,1010,DUMMY,ERR=400) RLBUF(I)  

        I = I + 1  

        IF(I.EQ. ienda1) GOTO200  

        GO TO 100  

C  

C*** NOW, FILL RLBUF1  

C  

200    i = 1  

210    READ(9,1000,END=350) NCHAR,DUMMY  

        IF(DUMMY(1) .EQ. '!!') GO TO 210  

        DECODE(20,1010,DUMMY,ERR=400) RLBUF1(I)  

        I = I + 1  

        if(i.eq. iendb1) goto 300  

        GO TO 210
C

```

```

*** EXIT THE PROCEEDINGS
C
300  CONTINUE
    sy0er = rdbuf(1) !SSSUP - RKGST - INITIAL SY
    errp = rdbuf(7) !SSSUP - RKGST(PSS) - ERROR BOUND
    smxp = rdbuf(8) !SSSUP - RKGST(PSS) - MAXIMUM STEP
    wtszp = rdbuf(9) !SSSUP - RKGST(PSS) - WEIGHT FOR SZ
    wtsyp = rdbuf(10) !SSSUP - RKGST(PSS) - WEIGHT FOR SY
    wtbepl = rdbuf(11) !SSSUP - RKGST(PSS) - WEIGHT FOR BEFF
    wtDH = rdbuf(12) !SSSUP - RKGST(PSS) - WEIGHT FOR BEFF
    errg = rdbuf(13) !SSSUP - RKGST(SSG) - ERROR BOUND
    smxg = rdbuf(14) !SSSUP - RKGST(SSG) - MAXIMUM STEP SIZE
    wtruh = rdbuf(17) !SSSUP - RKGST(PSS) - WEIGHT FOR BEFF
    wtdhg = rdbuf(18) !SSSUP - RKGST(PSS) - WEIGHT FOR BEFF
C
    stpp = rdbuf1(2) !SSSUP - RKGST(PSS) - INITIAL STEP
    odlp = rdbuf1(3) !SSSUP - RKGST(PSS) - RELATIVE OUTPUT DELTA
    odllp = rdbuf1(4) !SSSUP - RKGST(PSS) - MAXIMUM DISTANCE TO OUT
    stpg = rdbuf1(5) !SSSUP - RKGST(SSG) - INITIAL STEP
    odlg = rdbuf1(6) !SSSUP - RKGST(SSG) - RELATIVE OUTPUT DELTA
    odllg = rdbuf1(7) !SSSUP - RKGST(SSG) - MAXIMUM DISTANCE TO OUT
C
    CLOSE(UNIT=9)
    RETURN
C
350  call trap(20) ! premature EOF
C
400  CALL trap(21)
C
1000 FORMAT(Q,132A1)
1010 FORMAT(10X,G10.4)
C
END
#####

```

```

C.....  

C  

C      ROUTINE TO GET RUN PARAMETERS FROM A FILE  

C  

C      SUBROUTINE ESTRT3(OPNRUP)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

COMMON  

$PHLAG/CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

$/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag  

$/ERROR/ERT1,ERDT,ERNTIM  

C  

EQUIVALENCE  

$(RLBUF(1),ERT1),    !FIRST SORT TIME - USER OPTION  

$(RLBUF(2),ERDT),    !SORT TIME DELTA - USER OPTION  

$(RLBUF(3),ERNTIM)   !NUMBER OF SORT TIMES - USER OPTION  

C  

LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

character DUMMY(1:132)  

character*40 opnrup  

DIMENSION RLBUF(3),RBUF(6)  

C  

OPEN(UNIT=9,NAME=OPNRUP,TYPE='OLD')  

C  

C*** READ A LINE AND DETERMINE ITS PURPOSE  

I = 1  

100  CONTINUE  

      READ(9,1000,END=300) NCHAR,DUMMY  

      IF(DUMMY(1) .EQ. '!') GO TO 100  

      DECODE(20,1010,DUMMY,ERR=400) RBUF(I)  

      I = I + 1  

      GO TO 100  

C  

C*** EXIT THE PROCEEDINGS AND DETERMINE CHECK5  

300  CONTINUE  

C  

DO 310 I = 1,3  

310  RLBUF(I) = RBUF(I)  

      CHECK5 = .FALSE.           ! IN ORDER FOR FLAG TO WORK  

      IF(RBUF(4) .EQ. 1.) CHECK5 = .TRUE.  

C  

      sigx_flag = rbuf(5)  

      CLOSE(UNIT=9)  

      RETURN  

C  

400  CALL trap(21)  

1000 FORMAT(Q,132A1)  

1010 FORMAT(10X,G10.4)  

      END  

#####

```

```

C
C ..... .
C
C      SUBROUTINE GAMMA
C
C..... .
C
C      This routine was originally supplied by Digital Equipment
C      Corporation as part of the Scientific Subroutine Package
C      available for RT-11 as part of the Fortran Enhancement
C      Package. It was upgraded for use in this package.
C
C..... .
C
C      PURPOSE
C          COMPUTES THE GAMMA FUNCTION FOR A GIVEN ARGUMENT
C
C      USAGE
C          GF = GAMMA(XX)
C
C      DESCRIPTION OF PARAMETERS
C          XX -THE ARGUMENT FOR THE GAMMA FUNCTION
C
C      IER-RESULTANT ERROR CODE WHERE
C          IER=0  NO ERROR
C          IER=1  XX IS WITHIN .000001 OF BEING A NEGATIVE INTEGER
C          IER=2  XX GT 34.5, OVERFLOW
C          IF IER .NE. 0 PROGRAM TAKES A DIP IN THE POOL!
C
C      REMARKS
C          NONE
C
C      SUBROUTINES AND FUNCTION SUBPROGRAMS REQUIRED
C          NONE
C
C      METHOD
C          THE RECURSION RELATION AND POLYNOMIAL APPROXIMATION
C          BY C.HASTINGS,JR., 'APPROXIMATIONS FOR DIGITAL COMPUTERS',
C          PRINCETON UNIVERSITY PRESS, 1955
C
C      MODIFIED TO FUNCTION FORM FROM ORIGINAL SUBROUTINE FORM
C
C .. .
C
C      FUNCTION GAMMA(XX)

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

        IF(XX-34.5)6,6,4
4     IER=2
        GAMMA=1.E38

```

```

      GO TO 1000
6   X=XX
     ERR=1.0E-6
     IER=0
     GAMMA=1.0
     IF(X>2.0)50,50,15
10  IF(X<2.0)110,110,15
15  X=X-1.0
     GAMMA=GAMMA*X
     GO TO 10
50  IF(X<1.0)60,120,110
C
C      SEE IF X IS NEAR NEGATIVE INTEGER OR ZERO
C
60  IF(X-ERR)62,62,80
62  Y=FLOAT(INT(X))-X
     IF(ABS(Y)-ERR)130,130,64
64  IF(1.0-Y-ERR)130,130,70
C
C      X NOT NEAR A NEGATIVE INTEGER OR ZERO
C
70  IF(X>1.0)80,80,110
80  GAMMA=GAMMA/X
     X=X+1.0
     GO TO 70
110 Y=X-1.0
     GY=1.0+Y*(-0.5771017+Y*(+0.9858540+Y*(-0.8764218+Y*
$(+0.8328212+Y*(-0.5684729+Y*(+0.2548205+Y*(-0.05149930)))))))
     GAMMA=GAMMA*GY
120 RETURN
130 IER=1
1000 CONTINUE
     IF(IER.EQ.1) WRITE(5,1100)
     IF(IER.EQ.2) WRITE(5,1110)
1100 FORMAT(5X,'?GAMMA?--ARGUMENT LESS THAN 0.000001')
1110 FORMAT(5X,'?GAMMA?--ARGUMENT GREATER THAN 34.5--OVERFLOW')
     CALL EXIT
     END
####
```

```

C.....  

C  

C      SUBROUTINE TO ESTABLISH THE TIME SORT PARAMETERS  

C  

C          SUBROUTINE GETTIM  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C          include 'sys$degadis:DEGADIS3.dec/list'  

C  

C          COMMON  

$SSCON/ NREC(maxnob,2),T0(maxnob),XV(maxnob)  

$SORTIN/ TIM(maxnt),NTIM,ISTRTR  

$PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

$ERROR/ ERT1,ERDT,ERNNTIM  

$ALP/ ALPHA,alpha1  

$CNOBS/ NOBS  

C  

LOGICAL CHECK1,CHECK2,CHECK3,CHECK4,CHECK5, AGAIN  

C  

DATA T1/0./,DT/0./,TF/0./  

C  

C*** IF CHECK5 IS SET, GET THE TIME SORT PARAMETERS FROM /ERROR/  

C  

IF(.NOT.CHECK5) GO TO 90  

C  

T1 = ERT1  

DT = ERDT  

NTIM = INT(ERNNTIM)  

GO TO 95  

C  

C  

C*** This subroutine sets the default time sort windows.  

C  

C*** The first sort time is set for potential low wind speed cases,  

C*** while the last sort time is set for potential high wind speed  

C*** cases. The first sort time is taken to be when the first  

C*** observer passes through x=RMAX. The last sort time is taken  

C*** to be when the last observer passes through x=6*RMAX.  

C*** The default value for the number of sort times is set to 10.  

C*** Obviously, these values generate some sort times which will be  

C*** useless; hopefully, these values will show the user where to  

C*** look on the next go around.  

C*** The number of times has been doubled to 20 and the time interval  

C*** has been doubled in order to give more information for lower  

C*** concentrations (for the toxic gas problem). tos,5mar86  

C  

90    CONTINUE  

C

```

```
T1 = T0(1) + (2.*RMAX)**(1./ALPHA1)/ALEPH
TF = T0(NOBS) + (6.*RMAX)**(1./ALPHA1)/ALEPH
NTIM = 20          ! for toxic gas problem
C
c DT = (TF-T1)/FLOAT(NTIM-1)
DT = 2.*(TF-T1)/FLOAT(NTIM-1)  ! for toxic gas problem
DT = FLOAT(INT(DT+.5))
C
IF(DT .GE. 5.) GO TO 95
DT = 5.
NTIM = INT((TF - T1)/DT) + 1
C
95  CONTINUE
C
T1 = FLOAT(INT(T1))      !MAKE T1 AN INTEGER VALUE
C
II = max( ntim,2)        ! fill in the lowest two times at least
DO 100 I = 1,II
TIM(I) = DT*FLOAT(I-1) + T1
100  CONTINUE
C
RETURN
END
#####

```

```

C.....  

C  

C      SUBROUTINE TO ESTABLISH THE TIME SORT PARAMETERS  

C  

C      SUBROUTINE GETTIM  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

include 'sys$degadis:DEGADIS4.dec/list'  

C  

COMMON  

$SSCON/ NRECC(maxnob,2),T0(maxnob),XV(maxnob)  

$cdos/ idos, dosdisx(ndos), dosdis(4,2,ndos)  

$SORTIN/ TIM(maxnt),NTIM,ISTRTR  

$PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

$ERROR/ ERT1,ERDT,ERNNTIM  

$ALP/ ALPHA,alpha1  

$CNOBS/ NOBS  

C  

LOGICAL CHECK1,CHECK2,CHECK3,CHECK4,CHECK5, AGAIN  

C  

DATA T1/0./,DT/0./,TF/0./  

C  

C*** This subroutine sets the time sort windows for concentration as  

C      a function of time at a given position.  

C  

dist = dosdisx(idos)  

C  

T1 = ts( T0(1), dist) ! time first observer crosses dist  

TF = ts( T0(nobs), dist) ! time last observer crosses dist  

NTIM = 40  

C  

DT = (TF-T1)/FLOAT(NTIM-1)  

DT = dmax1( DBLE(INT(DT+.5)), 1.D0)  

NTIM = min( INT((TF - T1)/DT) + 1, 40)  

C  

T1 = FLOAT(INT(T1)) !MAKE T1 AN INTEGER VALUE  

write(lunlog,1000) t1, tf, dt, dist  

1000 format(' t1: ',1pg13.5,' tf: ',1pg13.5,' dt: ',1pg13.5,  

$' dist: ',1pg13.5)  

C  

II = max( ntim,2) ! fill in the lowest two times at least  

DO 100 I = 1,II  

TIM(I) = DT*FLOAT(I-1) + T1  

100 CONTINUE  

C  

RETURN  

END  

#####

```

```

SUBROUTINE HEAD(gmass0)

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

C
      include 'sys$degadis:DEGADIS1.dec'
      include '($ssdef)'

C
      COMMON
$/GEN3/ RADG(2,maxl),QSTR(2,maxl),srcden(2,maxl),srcwc(2,maxl),
$ srcwa(2,maxl),srcenth(2,maxl)
$/TITLE/ TITLE
$/GEN1/ PTIME(igen), ET(igen), R1T(igen), PWC(igen), PTEMP(igen),
$      PFRACV(igen), PENTH(igen), PRHO(igen)
$/GEN2/ DEN(5,igen)
$/IT1/ T1,TINP,TSRC,TOBS,TSRT
$/ERROR/ STPIN,ERBND,STPMX,WTRG,WTtm,WTya,WTyc,WTeb,WTmb,WTuh,XLI,
$ XRI,EPS,ZLOW,STPINZ,ERBNDZ,STPMXZ,SRCDER,srcss,srccut,
$ htcut,ERNOBL,NOBLpt,crfger,epsilon
$/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW
$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc
$/com_ss/ ess,sten,swid,outcc,outsz,outb,outl,swcl,swal,senl,srhl
$/phflag/ check1,check2,again,check3,check4,check5
$/NEND/ POUNDN,POUND
$/ALP/ ALPHA,alpha1
$/alphcom/ ialpfl,alpc0
$/phicom/ iphifl,dellay
$/sprd_con/ ce, delrhomin
./oomsin/ oodist,avtime

C
      character*80 TITLE(4)
C
      character*4 pound
      character*24 TINP,TSRC,TOBS,TSRT
      character*3 gas_name
      character*1 stabil(6)
      character*24 id
C
      logical check1,check2,again,check3,check4,check5
C
      REAL*8 K,ML
C
      data stabil/'A','B','C','D','E','F'/
      data iparm/0/
C
      if(iparm .eq. 1) goto 190
      WRITE(8,1100)
1100  FORMAT(1HO,*****9X,'U O A _ D E G A D I S ',
```

```

$2X,'M O D E L ',2X,'O U T P U T ',2X,'- - ',2X,'V E R S I O N ',
$2X,'2.1',8X,'*****')
C
C
      WRITE(8,1111)
C
      WRITE(8,1102) tsrc
1102  FORMAT(1H ,*****',23X,
$*****',1X,a24,1X,
$*****',23X,*****')
C
      WRITE(8,1111)
C
      WRITE(8,1112) TINP
      WRITE(8,1114) TSRC
      IF(tOBS(1:2).NE.' ' .and. .not.check4) WRITE(8,1116) TOBS
      IF(tOBS(1:2).NE.' ' .and. check4) WRITE(8,1117) TOBS
      IF(tSRT(1:2) .NE. ' ') WRITE(8,1118) TSRT
1112  FORMAT(1H ,'Data input on',22X,a24)
1114  FORMAT(1H ,'Source program run on',14X,a24)
1116  FORMAT(1H ,'Pseudo Steady-State program run on ',a24)
1117  FORMAT(1H ,'Steady-State program run on',7X,a24)
1118  FORMAT(1H ,'Time sort program run on',11X,a24)
      WRITE(8,1111)
C
      WRITE(8,1110)
      WRITE(8,1111)
C
1110  FORMAT(1H0,10X,'TITLE BLOCK')
1111  FORMAT(1H )
C
      DO 100 I = 1,4
      WRITE(8,1120) TITLE(I)
100   CONTINUE
C
1120  FORMAT(1H ,A80)
C
      WRITE(8,1111)
      WRITE(8,1130) U0
      WRITE(8,1140) Z0
      WRITE(8,1150) ZR
      write(8,1155) stabil(istab)
      if(ml.ne. 0.) then
          WRITE(8,1160) ML
      else
          write(8,1161)
      endif
      write(8,1168) avtime
      WRITE(8,1170) DELTA
      WRITE(8,1180) BETA
      WRITE(8,1190) ALPHA

```

```

      WRITE(8,1192) USTAR
      WRITE(8,1194) tamb
      if(isofl.eq.0 .and. ihtfl.ne.0) write(8,1195) tsurf
      WRITE(8,1196) pamb
      WRITE(8,1198) humid
      vaporp = 6.0298e-3* exp(5407.* (1./273.15- 1./tamb)) ! atm
      relhumiid = 100.* humid/(0.622*vaporp / (pamb- vaporp))
      write(8,1199) relhumiid

C
1130  FORMAT(1H ,5X,'Wind velocity at reference height ',20X,F6.2,2X,
      '$'m/s')
1140  FORMAT(1H ,5X,'Reference height ',37X,F6.2,2X,'m')
1150  FORMAT(1H0,5X,'Surface roughness length ',25X,1PG10.3,2X,'m')
1155  FORMAT(1H0,5X,'Pasquill Stability class ',25X,4x,a1)
1160  FORMAT(1H0,5X,'Monin-Obukhov length ',29X,1PG10.3,2X,'m')
1161  FORMAT(1H0,5X,'Monin-Obukhov length ',31X,'infinite')
1168  FORMAT(1H ,5X,'Gaussian distribution constants ',/,
      1H ,5x,8x,9x,'Specified averaging time',10X,F9.2,2X,'s')
1170  FORMAT(1H ,5X,32x,4X,'Delta',10X,F9.5)
1180  FORMAT(1H ,5X,32X,4X,' Beta',10X,F9.5)
1190  FORMAT(1H0,5X,'Wind velocity power law constant',4X,'Alpha',
      $10X,F9.5)
1192  FORMAT(1H ,5X,'Friction velocity',15X,4X,5X,10X,F9.5,2X,'m/s')
1194  FORMAT(1H0,5X,'Ambient Temperature ',34X,F6.2,2X,'K')
1195  FORMAT(1H0,5X,'Surface Temperature ',34X,F6.2,2X,'K')
1196  FORMAT(1H ,5X,'Ambient Pressure ',37X,F6.3,2X,'atm')
1198  FORMAT(1H ,5X,'Ambient Absolute Humidity',25X,1PG10.3,2X,
      '$'kg/kg BDA')
1199  FORMAT(1H ,5X,'Ambient Relative Humidity',25X,4X,F6.2,2X,'%')

C
      WRITE(8,1111)

C
      if(isofl .eq. 0) goto 135
      WRITE(8,1200)
      WRITE(8,1205)
      ii = -1
      DO 130 I = 1,igen
      IF(DEN(1,I).gt. 1.) goTO 148
      ii = ii+1
      if(ii.eq. 3) then
          write(8,1211)
          ii = 0
          endif
130   WRITE(8,1210) DEN(1,I),DEN(2,I),den(3,i)
      goto 148
135   write(8,1207)
      write(8,1208)
      ii = -1
      DO 138 I = 1,igen
      IF(DEN(1,I).gt. 1.) goTO 148
      ii = ii+1

```

```

        if(ii.eq. 3) then
            write(8,1211)
            ii = 0
            endif
138  WRITE(8,1212) DEN(1,I),DEN(2,I),den(3,i),den(4,i),den(5,i)
148  continue
C
1200 FORMAT(1H ,5X,'Input:    ',6X,3x,'Mole fraction',4x,
1           'CONCENTRATION OF C',6X,'GAS DENSITY')
1205 FORMAT(1H ,14X,20x,2(13X,'kg/m**3'))
1207 FORMAT(1H ,5X,'Adiabatic Mixing:',3x,'Mole fraction',3x,
1           'CONCENTRATION OF C',6X,'GAS DENSITY',5x,
1           6x,'Enthalpy',6x,4x,'Temperature')
1208 FORMAT(1H ,14X,20x,2(13X,'kg/m**3'),7x,8x,'J/kg',8x,9x,'K')
1210 FORMAT(1H ,14X,3(12X,F8.5))
1211 format(1H )
1212 FORMAT(1H ,14X,3(12X,F8.5),6x,3x,1pg13.5,7x,1pg13.5)
C
        WRITE(8,1111)
        write(8,1233) gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp
        WRITE(8,1111)
        WRITE(8,1220) gmass0
        WRITE(8,1230)
C
        DO 150 I=1,IGEN
        IF(PTIME(I).EQ.POUNDN) GO TO 160
150  WRITE(8,1231) PTIME(I),ET(I),R1T(I), PWC(I),PTEMP(I),
$      PENTH(I)
160  CONTINUE
C
        .
1220 FORMAT(1H , 'Source input data points',//,
$           1h ,15x,'Initial mass in cloud: ',1pg13.5,//,
$           1h ,7X,'Time',10X,'Contaminant',
$           7X,'Source Radius',2x,3x,'Contaminant',4x,
$           3x,'Temperature',4x,5x,'Enthalpy',5x,/,
$           1x,18x,4x,'Mass Rate',5x,18x,2x,'Mass Fraction',3x)
1230 FORMAT(1H ,9X,'s',15X,'kg/s',16X,'m',8x,' kg contam/kg mix ',8x,
$           'K',9x,7x,'J/kg',7x)
1231 FORMAT(1H ,6(3X,1PG12.5,3X))
1233 Format(1H0.5x,'Specified Gas Properties://,
110x,'Molecular weight:',T65,1pg13.5,/,
110x,'Storage temperature:',T65,1pg13.5,'K',/,
110x,'Density at storage temperature and ambient pressure:',
1           T65,1pg13.5,'kg/m**3',/,
110x,'Mean heat capacity constant:',T65,1pg13.5,/,
110x,'Mean heat capacity power:',T65,1pg13.5,/,
110x,'Upper mole fraction contour:',T65,1pg13.5,/,
110x,'Lower mole fraction contour:',T65,1pg13.5,/,
110x,'Height for isopleths:',T65,1pg13.5,'m',/)
1241 format(1h0.5x,'Calculation procedure for ALPHA: ',I2)

```

```

1242 format(1h0,5x,'Entrainment prescription for PHI: ',I2)
1244 format(1h0,5x,'Layer thickness ratio used for average depth: ',
1      1pg13.5)
1250 format(1h0,5x,'Air entrainment coefficient used: ',f5.3)
1251 format(1h0,5x,'NON Isothermal calculation')
1252 format(1h0,5x,'Gravity slumping velocity coefficient used: ',f5.3)
1253 format(1h0,5x,'Heat transfer calculated with fixed coefficient: ',
1      1pg13.5,' J/m**2/s/K')
1254 format(1h0,5x,'Heat transfer not included')
1255 format(1h0,5x,'Heat transfer calculated with correlation: ',I2)
1256 format(1h0,5x,'Isothermal calculation')
1257 format(1h0,5x,'Water transfer calculated with fixed coefficient: ',
1      1pg13.5,' /m**2/s/atm')
1258 format(1h0,5x,'Water transfer not included')
1259 format(1h0,5x,'Water transfer calculated with correlation')

c
      WRITE(8,1111)
      write(8,1241) ialpfl
      write(8,1242) iphilf
      write(8,1244) dellay
      write(8,1250) epsilon
      write(8,1252) ce
      if(isofl.eq. 0) write(8,1251)
      if(isofl.ne. 0) write(8,1256)
      if(ihtfl.lt. 0) write(8,1253) htco
      if(ihtfl.eq. 0) write(8,1254)
      if(ihtfl.gt. 0) write(8,1255) ihtfl
      if(iwtfl.lt. 0) write(8,1257) wtco
      if(iwtfl.eq. 0) write(8,1258)
      if(iwtfl.gt. 0) write(8,1259)
      WRITE(8,1111) '
      iparm =1
      return

c
190  continue
      if(.not.check4) return
      RAD = SQRT(2.*SLEN*SWID/pi)
      WRITE(8,1300) ESS,RAD
      WRITE(8,1320) SLEN,SWID
      qstar = ess/outl/outb/2.
      WRITE(8,1340) OUTCc,OUTS2,qstar
      write(8,1350) swcl,swal,sem1,srhl
      WRITE(8,1360) OUTL,OUTB

c
c
c
1300 FORMAT(1H0,'Source strength [kg/s] : ',18X,1PG13.5,T60,
     $'Equivalent Primary source radius [m] : ',5X,1PG13.5)
1320 FORMAT(1H , 'Equivalent Primary source length [m] : ',4X,1PG13.5,
     $T60,'Equivalent Primary source half-width [m] : ',1X,1PG13.5)
1340 FORMAT(/,' Secondary source concentration [kg/m**3] : ',

```

```
$1PG13.5,T60,'Secondary source SZ [m] : ',18X,1PG13.5,//,
      1 ' Contaminant flux rate: ',1pg13.5,/)
1350  format(//,' Secondary source mass fractions... contaminant: ',
      1       1pg13.6,2x,' air: ',1pg13.5,/,',',10x,' Enthalpy: ',
      1       1pg13.5,5x,' Density: ',1pg13.5,/)
1360  FORMAT(1H , 'Secondary source length [m] : ',13X,1PG13.5,T60,
      $'Secondary source half-width [m] : ',10X,1PG13.5)

C
C
      RETURN
      END
####
```

```

C.....  

c  

c      gamma and incomplete gamma function  

c  

C.....  

c  

c      routines included in this file:  

c          GAMINC      Incomplete gamma function  

c          GAMMLN      Natural log of the gamma function  

c          GSER        computation procedure  

c          GCF         computation procedure  

c  

C.....  

c  

c      These functions return the value for the incomplete Gamma function  

c      with the arguments (ALPHA,BETA).  The functional values are  

c      calculated with either a series representation or a continued  

c      fraction representation.  These routines are based on:  

c  

c      Press, W.H., B.P. Flannery, S.A. Teukolsky, and W.T. Vetterling,  

c      "Numerical Recipes", Cambridge University Press, Cambridge, 1986.  

c  

c  

c      function gaminc(alpha,beta)  

    implicit Real*8 (a-h, o-z), integer*4 (i-n)  

c  

c  

c*** ensure the arguments are within the proper bounds.  

c  

    if( beta.lt.0. .or. alpha.le.0.) then  

        write(6,9000) alpha,beta  

9000    format(' GAMINC? Arguments are out-of-bounds. ALPHA: ',  

         $ 1PG13.5,' BETA: ',1PG13.5)  

        STOP  

        endif  

c  

c*** determine which of the series or continued fraction representation  

c   is more appropriate.  

c  

    if( beta .lt. alpha+1. ) then  

        ! series  

        call gser( aa, alpha, beta, gln)  

    else  

        ! continued fraction  

        call gcf( aa, alpha, beta, gln)  

        aa = 1. - aa  

    endif  

c  

c*** multiply the result by GAMMA(ALPHA) to get the final value.

```

```

c
gaminc = dexp( dlog(aa) + gln)
return
end

c
c
c
subroutine gser( gamser, alpha, beta, gln)
c
c This routine calculates the incomplete gamma function using
c the series representation.
c
implicit real*8 (a-h,o-z), integer*4 (i-n)
c
parameter (itmax=100, eps=1.d-9)

gln = gammaln(alpha)

if(beta .lt. 0.) then
    write(6,*) 'GSER? BETA is out-of-bounds.'
    stop
endif
if(beta .eq. 0.) then
    gamser = 0.
    return
endif

ap = alpha
sum = 1./alpha
del = sum

do 100 n=1,itmax
    ap = ap + 1.
    del = del* beta /ap
    sum = sum + del
    if( dabs(del) .lt. dabs(sum)*eps) then
        gamser = sum*dexp( -beta+alpha*dlog(beta)-gln)
        return
    endif
100   continue

    write(6,*) 'GSER? ALPHA is too large or ITMAX is too small.'
    stop
end

c
c
c
subroutine gcf( gamcf, alpha, beta, gln)
c
c This routine calculates the incomplete gamma function using
c the continued fraction representation.

```

```

c
      implicit real*8 (a-h,o-z), integer*4 (i-n)
c
      parameter (itmax=100, eps=1.d-9)

      gln = gammln(alpha)

c
c*** INITIALIZE some variables.
c
c                               ! previous value tested for convergence
      gold = 0.
      a0 = 1.
      a1 = beta
      b0 = 0.
      b1 = 1.
c                               ! fac is a "renormalization factor"
      fac = 1.

      do 100 n=1,itmax
          an = dble(n)
          ana = an - alpha
          a0 = (a1 + a0*ana)
          b0 = (b1 + b0*ana)
          anf = an
          a1 = beta *a0 + anf*a1
          b1 = beta *b0 + anf*b1
          if( a1 .ne. 0.) then
              fac = 1./b1
              gg = a1*fac
          a0 = a0*fac
          a1 = a1*fac
          b0 = b0*fac
          b1 = b1*fac
          if( dabs((gg-gold)/gg) .lt .eps) then
              gamcf = dexp(
                           -beta+alpha*dlog(beta)-gln)/gg
              return
          endif
          gold = gg
      endif
100   continue

      write(6,*) 'GCF? ALPHA is too large or ITMAX is too small.'
      stop
      end

c
c
c     function gammln( alpha )
c
3 -- sys$degadis:incgamma.for           6-SEP-1989 16:54:46

```

```

c      This routine calculates ln( gamma(alpha)) for alpha>0.
c
c      implicit real*8 (a-h,o-z), integer*4 (i-n)
c
c      dimension cof(6)

      data cof,stp/76.18009173D0, -86.50532033D0, 24.01409822D0,
$ -1.231739516D0, 0.120858003D-2, -0.536382D-5, 2.50662827465D0/
      data half, one, fpf/0.5D0, 1.0D0, 5.5D0/
c
c      if( alpha .lt. 1) then
c          gam = gamma( alpha )
c          gammln = dlog(gam)
c          return
c          endif
c
c      xx = alpha - one
c      tmp = xx + fpf
c      tmp = (xx + half) * dlog(tmp) - tmp
c      ser = one
c      do 100 j=1,6
c          xx = xx + one
c          ser = ser + cof(j)/xx
c 100   continue
c
c      gammln = tmp + dlog(stp*ser)
c      return
c      end
c####

```

```

C.....  

C  

C      INPUT SUBROUTINE FOR DEGADIS MODEL  

C  

C      SUBROUTINE IO(tend,gmass0,OPNRUP)  

C  

C      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      include 'sys$degadis:DEGADIS1.dec'  

C  

C      BLOCK COMMON  

C  

C      COMMON  

$TITLE/TITLE  

$/GEN1/ PTIME(igen), ET(igen), R1T(igen), PWC(igen), PTEMP(igen),  

$      PFRACV(igen), PENTH(igen), PRHO(igen)  

$/GEN2/ DEN5,igen)  

$/ITI/ T1,TINP,TSRC,TOBS,TSRT  

$/PARM/ U0,Z0,ZR,ML,USTAR,vkc,gg,RHOE,RHOA,DELTAy,BETAY,GAMMAF,  

.          CcLOW  

$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,  

$ gas_ufl,gas_lfl,gas_zsp,gas_name  

$/comatm/ istab,tamb,pamb,humid,isoft,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/com_ss/ ess,slen,swid,outcc,outsz,outb,outl  

$/phlag/check1,check2,again,check3,check4,check5  

$/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag  

$/NEND/ POUNDN,POUND  

$/ICNTL/ IPRINT,IOBS(10)  

./oomsin/ oodist,avtime  

C  

C      character*80 TITLE(4)  

C      character*4 pound  

C      character*24 TSRC,TINP,TOBS,TSRT  

C      character*3 gas_name  

C  

C      logical check1,check2,again,check3,check4,check5  

C  

C      character*40 OPNRUP  

C  

C      OPEN(UNIT=9, NAME=OPNRUP, TYPE='OLD')  

C  

DO 90 I=1,4  

READ(9,2000) TITLE(I)  

90  CONTINUE  

2000 FORMAT(A80)  

C  

READ(9,*) U0,Z0,zr  

read(9,*) istab  

read(9,*) oodist, avtime

```

```

READ(9,*) DELTAy,BETAy,rml
read(9,*) sigx_coeff,sigx_pow,sigx_min_dist
read(9,*) tamb,pamb,humid
humsrc = 0.
read(9,*) isofl,tsurf
read(9,*) ihtfl,htco
read(9,*) iwtfl,wtco
read(9,2020) gas_name
2020 FORMAT(A3)
read(9,*) gas_mw,gas_temp,gas_rhoe
read(9,*) gas_cpk,gas_cpp
read(9,*) gas_ufl,gas_lfl,gas_zsp
C
      if(isofl .eq. 0) then
        rhoe = gas_rhoe
        rhoa = pamb*(1.D0+humid)/(.002833D0+ 0.004553D0*humid)/tamb
        goto 105
      endif
      READ(9,*) NP
      DO 100 I=1,NP
100   READ(9,*) DEN(1,I),DEN(2,I),den(3,I),den(4,i),den(5,i)
      RHOE = DEN(3,NP)
      RHOA = DEN(3,1)
      den(1,np+1) = 2.
C
105   READ(9,*) CcLOW
C
      read(9,*) gmass0
      READ(9,*) NP
      DO 110 I=1,NP
      READ(9,*) PTIME(I),ET(I),R1T(I), PWC(I),PTEMP(I),PFRACV(I)
      WA = (1.D0 - PWCI))/((1.D0 + HUMID)
      PENTH(I) = ENTHAL(PWC(I), WA, PTEMP(I))
      CALL TPROP(-1,PWC(I), WA,PENTH(I), YC,YA,WM, PTEMP(I), PRHO(I),CP)
110   CONTINUE
      TEND = PTIME(NP-2)
      PTIME(NP + 1) = POUNDN
C
      read(9,*) check1,check2,again,check3,check4,check5
C
      tobs = ' '
      tsrt = ' '
      READ(9,2010) TINP
2010  format(a24)
C
      if(check4) read(9,*) ess,slen,swid
C
      CLOSE(UNIT=9)
      RETURN
      END
#####

```

```

SUBROUTINE IOT(OPNRUP)

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

C
      include 'sys$degadis:DEGADISIN.dec'
C
      COMMON
$TITLE/ TITLE
$GEN1/ PTIME(igen), ET(igen), R1T(igen), PWC(igen), PTEMP(igen),
$      PFRACV(igen), PENTH(igen), PRHO(igen)
$GEN2/ DEN(5,igen)
$ITI/ T1,TINP,TSRC,TOBS,TSRT
$PARM/ U0,Z0,ZR,rML,USTAR,vkc,gg,RHOE,RHOA,DELTAY,BETAY,GAMMAF,
.          CcLOW
$com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$com_ss/ ess,slen,swid,outcc,outsz,outb,outl
$PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
$com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag
$NEND/ POUNDN,POUND
./oomsin/ oodist,avtime
C
      character*80 TITLE(4)
      character*3 gas_name
      character*4 pound
      character*24 TSRC,TINP,TOBS,TSRT
C
      LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
C
      character*(*) OPNRUP
      character*40 STRING
      character*4 dummy
C
      WRITE(6,1100)
      WRITE(6,1110)
C
C*** OPEN THE INPUT FILE
C
      OPEN(UNIT=8,NAME=OPNRUP,TYPE='NEW',
$ carriagecontrol='list',recordtype='variable')
C
C*** NOW GET THE TITLE BLOCK
C
      WRITE(6,1120)
      WRITE(6,1130)
C
      DO 100 I=1,4
      READ(5,1134) TITLE(I)
      dummy = title(i)
      IF(dummy(1:4) .EQ. POUND(1:4)) GO TO 110

```

```

        WRITE(8,1135) TITLE(I)
100  CONTINUE
     GO TO 130
C
110  CONTINUE           ! FILL OUT THE BLOCK
     II = I
     DO 120 I = II,4
     TITLE(I) = ' '
     WRITE(8,1135) TITLE(I)
120  CONTINUE
130  CONTINUE
C
c*** Atmospheric parameters:
c
     WRITE(6,1140)
     WRITE(6,1142)
     READ(5,*) U0,Z0,ZR
     WRITE(8,1020) U0,Z0,ZR
C
c*** stability, averaging time for DELTAY, and derived parameters
c
     WRITE(6,1150)
     READ(5,1310) NCHAR,STRING
     write(6,1152)
     read(5,*) avtime
     istab = 4           ! default is D stability
     IF(STRING.eq.'A' .or. string.eq.'a') istab=1
     IF(STRING.eq.'B' .or. string.eq.'b') istab=2
     IF(STRING.eq.'C' .or. string.eq.'c') istab=3
     IF(STRING.eq.'D' .or. string.eq.'d') istab=4
     IF(STRING.eq.'E' .or. string.eq.'e') istab=5
     IF(STRING.eq.'F' .or. string.eq.'f') istab=6
     goto(161,162,163,164,165,166) istab
161   timeav = dmax1( avtime, 18.4D0)          ! A
     deltay = 0.423D0*(timeav/600.D0)**0.2D0
     betay = 0.9D0
     rml = -11.43D0 * zr**0.103D0
     sigx_coeff = 0.02
     sigx_pow = 1.22
     sigx_min_dist = 130.
     goto 170
162   timeav = dmax1( avtime, 18.4D0)          ! B
     deltay = 0.313D0*(timeav/600.D0)**0.2D0
     betay = 0.9D0
     rml = -25.98D0 * zr**0.171D0
     sigx_coeff = 0.02
     sigx_pow = 1.22
     sigx_min_dist = 130.
     goto 170
163   timeav = dmax1( avtime, 18.4D0)          ! C
     deltay = 0.210D0*(timeav/600.D0)**0.2D0

```

```

        betay = 0.900
        rml = -123.4D0 * zr**0.304D0
        sigx_coeff = 0.02
        sigx_pow = 1.22
        sigx_min_dist = 130.
        goto 170
164    timeav = dmax1( avtime, 18.3D0)                      ! D
        deltay = 0.136D0*(timeav/600.D0)**0.2D0
        betay = 0.900
        rml = 0.0           ! used for infinity
        sigx_coeff = 0.04
        sigx_pow = 1.14
        sigx_min_dist = 100.
        goto 170
165    timeav = dmax1( avtime, 11.4D0)                      ! E
        deltay = 0.102D0*(timeav/600.D0)**0.2D0
        betay = 0.900
        rml = 123.4D0 * zr**0.304D0
        sigx_coeff = 0.17
        sigx_pow = 0.97
        sigx_min_dist = 50.
        goto 170
166    timeav = dmax1( avtime, 4.6D0)                      ! F
        deltay = 0.0674D0*(timeav/600.D0)**0.2D0
        betay = 0.900
        rml = 25.98D0 * zr**0.171D0
        sigx_coeff = 0.17
        sigx_pow = 0.97
        sigx_min_dist = 50.
C
170    WRITE(8,1040) istab
        oodist = 0.00
        write(8,1020) oodist,avtime
C
172    if( rml.ne. 0.) then
        WRITE(6,1159) deltay,betay,rml,sigx_coeff,sigx_pow,sigx_min_dist
    else
        WRITE(6,1160) deltay,betay,sigx_coeff,sigx_pow,sigx_min_dist
    endif
    read(5,1310) nchar,string
    if(string.eq.'d' .or. string.eq.'D') then
        write(6,1600)
        read(5,*) deltay
        goto 172
    else if(string.eq.'b' .or. string.eq.'B') then
        write(6,1620)
        read(5,*) betay
        goto 172
    else iff(string.eq.'l' .or. string.eq.'L') then
        write(6,1660)
        read(5,*) rml

```

```

        goto 172
else if(string.eq.'c' .or. string.eq.'C') then
    write(6,1670)
    read(5,*) sigx_coeff
    goto 172
else if(string.eq.'p' .or. string.eq.'P') then
    write(6,1680)
    read(5,*) sigx_pow
    goto 172
else if(string.eq.'m' .or. string.eq.'M') then
    write(6,1690)
    read(5,*) sigx_min_dist
    goto 172
else if(nchar.eq.0 .or. string.eq.'n' .or. string.eq.'N') then
    WRITE(8,1020) DELTAY,BETAY,rml
    WRITE(8,1020) sigx_coeff,sigx_pow,sigx_min_dist
else
    goto 172
endif

c
c*** ambient pressure, temperatures, and humidity
c
    write(6,1500)
    read(5,*) tamb,pamb
    vaporp = 6.0298e-3* exp(5407.* (1./273.15- 1./tamb)) ! atm
    sat = 0.622*vaporp / (pamb- vaporp)
    write(6,1580)
    read(5,1310) nchar,string
    if(string.eq.'a' .or. string.eq.'A') then
        write(6,1585)
        read(5,*) humid
        relhumid = 100.*humid/sat
        write(6,1586) relhumid
        goto 200
        endif
    write(6,1587)
    read(5,*) relhumid
    humid = relhumid/100. * sat
200   rhoa = pamb*(1.+humid)/(.002833+.004553*humid)/tamb
    write(6,1588) rhoa
    write(8,1025) tamb,pamb,humid, relhumid

c
    isofl = 0
    ihtfl = 0
    htco = 0.
    iwtfl = 0
    wtco = 0.
    tsurf = tamb

c
    write(6,2000)
    read(5,1310) nchar,string

```

```

if(string.eq.'Y' .or. string.eq.'y') then
    isofl = 1
    tsurf = tamb
    goto 250
endif

c
write(6,2020)
read(5,1310) nchar,string
if(string.eq.'Y' .or. string.eq.'y') then
    write(6,2030)
    read(5,*) tsurf
220   write(6,2040)
    read(5,1310) nchar,string
    if(string.eq.'V' .or. string.eq.'v') then
        ihtfl = -1      ! constant value
        write(6,2050)
        read(5,*) htco
    else if(string.eq.'C' .or. string.eq.'c' .or. nchar.eq.0) then
        ihtfl = 1      ! local correlation
    else if(string.eq.'L' .or. string.eq.'l') then
        ihtfl = 2      ! LLNL correlation
        htco = 0.0125  ! [=]m/s
        write(6,2043) htco
        read(5,1310) nchar,string
        if(string.eq.'Y' .or. string.eq.'y')
            read(5,*) htco
    1
        else
            goto 220
        endif
    else
        goto 250
    endif

c
write(6,2100)
read(5,1310) nchar,string
if(string.eq.'Y' .or. string.eq.'y') then
    iwtfl = 1
    write(6,2045)
    read(5,1310) nchar,string
    if(string.eq.'V' .or. string.eq.'v') then
        iwtfl = -1
        write(6,2120)
        read(5,*) wtco
    endif
endif

c
250   continue
write(8,1060) isofl,tsurf
write(8,1060) ihtfl,htco
write(8,1060) iwtfl,wtco
c

```

```

C
c*** gas characteristics
c
      write(6,1510)
      read(5,1415) gas_name
      write(8,1415) gas_name
      gas_mw = 16.04
      gas_temp = 111.7
      gas_rhoe = 1.792*pamb ! correct to pamb
      gas_cpk = 2730.
      gas_cpp = 1.00
      gas_ufl= 0.15
      gas_lfl= 0.05
      gas_zsp= 0.5

      if(gas_name.eq.'NH3' .or. gas_name.eq.'nh3') then
          gas_mw = 17.
          gas_temp = tamb
          gas_rhoe = pamb*gas_mw/0.08205/tamb ! ideal gas
          gas_cpk = 3845.           ! to get 807.5 J/kg/K
          gas_cpp = 1.000
          gas_ufl= 2.0E-2
          gas_lfl= 2.00E-3
          gas_zsp= 0.5

      else if(gas_name.eq.'LNG' .or. gas_name.eq.'lng') then
          gas_mw = 16.04
          gas_temp = 111.7
          gas_rhoe = 1.792*pamb ! correct to pamb
          gas_cpk = 5.6e-8
          gas_cpp = 5.00
          gas_ufl= 0.15
          gas_lfl= 0.05
          gas_zsp= 0.5

      else if(gas_name.eq.'LPG' .or. gas_name.eq.'lpg') then
          gas_mw = 44.09
          gas_temp = 231.
          gas_rhoe = 2.400*pamb ! correct to pamb
          gas_cpk = 15.4
          gas_cpp = 2.25
          gas_ufl= 0.10
          gas_lfl= 0.02
          gas_zsp= 0.5

      else if(gas_name.eq.'NO2' .or. gas_name.eq.'no2') then
          gas_mw = 46.
          gas_temp = tamb
          gas_rhoe = pamb*gas_mw/0.08205/tamb ! ideal gas
          gas_cpk = 3845.           ! to get 807.5 J/kg/K
          gas_cpp = 1.000

```

```

gas_ufl= 1000.0E-6
gas_lfl= 500.00E-6
gas_zsp= 0.5

else if(gas_name.eq.'N2O' .or. gas_name.eq.'n2o') then
    gas_mw = 92.
    gas_temp = tamb
    gas_rhoe = pamb*gas_mw/0.08205/tamb      ! ideal gas
    gas_cpk = 40990.              ! to get 807.5 J/kg/K
    gas_cpp = 1.000
    gas_ufl= 1000.0E-6
    gas_lfl= 500.00E-6
    gas_zsp= 0.5

else if(gas_name.eq.'CL2' .or.
1      gas_name.eq.'cl2' .or. gas_name.eq.'Cl2') then
    gas_mw = 70.91
    gas_temp = 238.7
    gas_rhoe = 3.672*pamb   ! correct to pamb
    gas_cpk = 484.2
    gas_cpp = 1.000
    gas_ufl= 0.1D-4
    gas_lfl= 0.3D-5
    gas_zsp= 0.5

else if(gas_name.eq.'MEC' .or.gas_name.eq.'mec') then
    gas_mw = 84.94
    gas_temp = tamb
    gas_rhoe = 3.53*(298.15/tamb)*pamb ! correct to pamb,tamb
    gas_cpk = 2730.          ! Open to question?
    gas_cpp = 1.000
    gas_ufl= 1000.D-6
    gas_lfl= 500.D-6
    gas_zsp= 0.5
endif
270 write(6,1520) gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
1      gas_ufl,gas_lfl,gas_zsp
read(5,1310) nchar,string
if(string.eq.'m' .or. string.eq.'M') then
    write(6,1550)
    read(5,*) gas_mw
    goto 270
else if(string.eq.'t' .or. string.eq.'T') then
    write(6,1530)
    read(5,*) gas_temp
    goto 270
else if(string.eq.'d' .or. string.eq.'D') then
    write(6,1535)
    read(5,*) gas_rhoe
    goto 270
else if(string.eq.'h' .or. string.eq.'H') then

```

```

        write(6,1570)
        read(5,*) gas_cpk
        goto 270
    else if(string.eq.'p' .or. string.eq.'P') then
        write(6,1571)
        read(5,*) gas_cpp
        goto 270
    else if(string.eq.'u' .or. string.eq.'U') then
        write(6,1572)
        read(5,*) gas_ufl
        goto 270
    else if(string.eq.'l' .or. string.eq.'L') then
        write(6,1573)
        read(5,*) gas_lfl
        goto 270
    else if(string.eq.'z' .or. string.eq.'Z') then
        write(6,1574)
        read(5,*) gas_zsp
        goto 270
    else if(nchar.eq. 0 .or. string.eq.'n' .or. string.eq.'N') then
        if(gas_cpp .eq. 0.00) then
            gas_cpp = 1.00
            gas_cpk = gas_cpk*gas_mw - 3.33D4
        endif
        WRITE(8,1020) gas_mw, gas_temp,gas_rhoe
        write(8,1020) gas_cpk,gas_cpp
        WRITE(8,1020) gas_ufl,gas_lfl,gas_zsp
    else
        goto 270
    endif
c
c density curve if isothermal
c
        if(isofl .eq. 0) goto 460
        WRITE(6,1161)
        WRITE(6,1162)
        WRITE(6,1163)
        WRITE(6,1164) rhoa
        WRITE(6,1165)
        WRITE(6,1166)
        goto 320
c
280    write(6,1290)
c
320    LUNIN = 5
        WRITE(6,1300)
        READ(5,1310) NCHAR,STRING
        IF(STRING.EQ.'y' .or. string.eq.'Y') GO TO 360
        GO TO 400
360    WRITE(6,1320)
        READ(5,1310) NCHAR,STRING

```

```

OPEN(UNIT=10,NAME=STRING,TYPE='OLD',err=280)
LUNIN = 10
400 CONTINUE
IF(LUNIN .EQ. 5) WRITE(6,1170) igen
READ(LUNIN,*) NP
WRITE(8,1040) NP
IF(LUNIN .EQ. 5) WRITE(6,1180)

C
DO 440 I=1,NP
den(4,i) = 0.                                ! 0.0 by default for isotherm
den(5,i) = tamb                               ! tamb for isotherm
READ(LUNIN,*) DEN(1,I),DEN(2,I),DEN(3,I)
if(i .eq.1 .and.
1      (den(3,1)/rhoa.gt.1.005 .or. rhoa/den(3,1).gt.1.005)) then
den(3,1) = rhoa
write(6,1340) rhoa
endif
if(i.eq.np) THEN
    IF(   den(2,i)/gas_rhoe .gt. 1.005
1     .or.   gas_rhoe/den(2,i) .gt. 1.005
1     .or.   den(3,i)/gas_rhoe .gt. 1.005
1     .or.   gas_rhoe/den(3,i) .gt. 1.005) then
den(2,i) = gas_rhoe
den(3,i) = gas_rhoe
write(6,1341) gas_rhoe
endif
endif
WRITE(8,1025) DEN(1,I),DEN(2,I),DEN(3,I),Den(4,I),den(5,i)
440 CONTINUE
IF(LUNIN .EQ. 10) CLOSE(UNIT=10)

C
C
460 ccmmax = (gas_lfl/2.) * pamb*gas_mw/0.08205/tamb
WRITE(6,1280) ccmmax
READ(5,*) CcLOW
if(cclow .le. 0.) cclow=0.005 ! don't let 0. get through
if(cclow .gt. ccmmax) cclow=ccmax
WRITE(8,1010) CcLOW

C
C
c*** source description
C
        write(6,1440)
C
        write(6,1460)
read(5,1410) dummy
if(dummy.eq.'d' .or. dummy.eq.'D') goto 730
C
check4 = .false.
write(6,1400)
read(5,1410) dummy

```

```

if(dummy.eq.'y' .or. dummy.eq.'Y') goto 480
goto 520
480  continue
gmass0 = 0.           ! no initial cloud for a SS simulation
write(8,1020) gmass0
write(6,1420)
read(5,*) ess
write(6,1430)
read(5,*) r1ss
np = 4
c
tend = 60230. ! [=] sec
c
PTIME(1) = 0.
et(1) = ess
r1t(1)= r1ss
pwc(1) = 1.
ptemp(1) = gas_temp
pfracv(1) = 1.
PTIME(2) = tend
et(2) = ess
r1t(2)= r1ss
pwc(2) = 1.
ptemp(2) = gas_temp
pfracv(2) = 1.
PTIME(3) = tend + 1.
et(3) = 0.
r1t(3)= 0.
pwc(3) = 1.
ptemp(3) = gas_temp
pfracv(3) = 1.
PTIME(4) = tend + 2.
et(4) = 0.
r1t(4)= 0.
pwc(4) = 1.
ptemp(4) = gas_temp
pfracv(4) = 1.
slen = 2.*r1ss
swid = pi*r1ss**2/slen/2.
check4 = .true.      ! steady state run
goto 790
c
520  continue
c
write(6,1450)
read(5,*) gmass0
write(8,1020) gmass0
c
WRITE(6,1190)
WRITE(6,1200)
WRITE(6,1210)

```

```

      WRITE(6,1220)
      WRITE(6,1221)
      WRITE(6,1165)
      WRITE(6,1223)
      WRITE(6,1230)
      WRITE(6,1240)
      WRITE(6,1250)
      goto 600
C
 560  write(6,1290)
C
 600  LUNIN = 5
      WRITE(6,1330)
      READ(5,1310) NCHAR,STRING
      IF(STRING.eq.'Y' .or. string.eq.'y') goto 640
      goto 680
 640  WRITE(6,1320)
      READ(5,1310) NCHAR,STRING
      OPEN(UNIT=10,NAME=STRING,TYPE='OLD',err=560)
      LUNIN = 10
 680  CONTINUE
      IF(LUNIN .EQ. 5) WRITE(6,1260) igen
      READ(LUNIN,*) NP
      IF(LUNIN .EQ. 5) WRITE(6,1270)
C
 720  DO 720 I=1,NP
      READ(LUNIN,*) PTIME(I),ET(I),R1T(I)
      pwc(i) = 1.
      ptemp(i) = gas_temp
      pfracv(i) = 1.
 720  CONTINUE
      IF(LUNIN .EQ. 10) CLOSE(UNIT=10)
      GOTO 790
C
C*** FOR A DILUTED SOURCE ...
C
 730  check4 = .false.
      write(6,1400)
      read(5,1410) dummy
      if(dummy.eq.'y' .or. dummy.eq.'Y') goto 740
      goto 750
 740  continue
      gmass0 = 0.           ! no initial cloud for a SS simulation
      write(8,1020) gmass0
      write(6,1420)
      read(5,*) ess
      write(6,1430)
      read(5,*) riss
 741  WRITE(6,1470)
      read(5,*) PWC(1)
      if(pwc(1).le.0.D0 .or. pwc(1).gt.1.D0) goto 741

```

```

742  WRITE(6,1480)
      read(5,*) PTEMP(1)
      if(pwc(1).le.0.D0) goto 742
      np = 4
c
      tend = 60230. ! [=] sec
c
      PTIME(1) = 0.
      et(1) = ess
      r1t(1)= r1ss
      PWC(1)= pwc(1)
      PTEMP(1)= ptemp(1)
      PFRACV(1)= 1.0
      PTIME(2) = tend
      et(2) = ess
      r1t(2)= r1ss
      PWC(2)= pwc(1)
      PTEMP(2)=ptemp(1)
      PFRACV(2)= 1.0
      PTIME(3) = tend + 1.
      et(3) = 0.
      r1t(3)= 0.
      PWC(3)= pwc(1)
      PTEMP(3)=ptemp(1)
      PFRACV(3)= 1.0
      PTIME(4) = tend + 2.
      et(4) = 0.
      r1t(4)= 0.
      PWC(4)= pwc(1)
      PTEMP(4)=ptemp(1)
      PFRACV(4)= 1.0
      slen = 2.*r1ss
      swid = pi*r1ss**2/slen/2.
      check4 = .true.      ! steady state run
      goto 790
c
750  continue
c
      write(6,1450)
      read(5,*) gmass0
      write(8,1020) gmass0
c
      WRITE(6,1190)
      WRITE(6,2200)
      WRITE(6,2210)
      WRITE(6,2220)
      WRITE(6,2221)
      WRITE(6,1165)
      WRITE(6,2223)
      WRITE(6,2230)
      WRITE(6,2240)

```

```

        WRITE(6,2250)
        goto 760
C
    755  write(6,1290)
C
    760  LUNIN = 5
        WRITE(6,1330)
        READ(5,1310) NCHAR,STRING
        IF(STRING.eq.'Y' .or. string.eq.'y') goto 765
        goto 770
    765  WRITE(6,1320)
        READ(5,1310) NCHAR,STRING
        OPEN(UNIT=10,NAME=STRING,TYPE='OLD',err=755)
        LUNIN = 10
    770  CONTINUE
        IF(LUNIN .EQ. 5) WRITE(6,1260) igen
        READ(LUNIN,*) NP
        IF(LUNIN .EQ. 5) WRITE(6,1270)
C
    DO 780 I=1,NP
        PFRACV(I) = 1.
        READ(LUNIN,*) PTIME(I),ET(I),R1T(I), PWC(I),PTEMP(I)
    780  CONTINUE
        IF(LUNIN .EQ. 10) CLOSE(UNIT=10)
C
    790  continue
        WRITE(8,1040) NP
        DO 800 I=1,NP
    800  WRITE(8,1030) PTIME(I),ET(I),R1T(I), PWC(I),PTEMP(I),PFRACV(I)
C
        if(et(1).eq.0. .and. gmass0.ne.0.) check2=.true. ! HSE type spill
        write(8,*) check1,check2,again,check3,check4,check5
C
        istat = lib$date_time(tinp)
        WRITE(8,1050) TINP
C
        if(check4) write(8,1020) ess,slen,swid      ! steady state
C
C
        CLOSE(UNIT=8)
C
C
    1010  format(1x,1pg14.7)
    1020  format(3(1x,1pg14.7))
    1025  format(5(1x,1pg14.7))
    1030  format(1x,5(1pg14.7,1x),1pg14.7)
    1040  format(1x,I5)
    1050  format(a24)
    1060  format(1x,i4,1x,1pg14.7)
C
    1100  FORMAT(5X,'INPUT MODULE -- DEGADIS MODEL')

```

```

1110  FORMAT(/,5X,'*****')
1120  FORMAT(5X,'Enter Title Block -- up to 4 lines of 80',
      $' characters')
1130  FORMAT(5X,'To stop, type "///"')
1134  FORMAT(A80)
1135  FORMAT(A80)
1140  FORMAT(5X,'ENTER WIND PARAMETERS -- U0 (m/s), Z0 (m), ',
      $'and ZR(m)')
1142  format(5x,'U0 -- Wind velocity at reference height Z0',
      $/,5X,'ZR -- Surface Roughness')
1150  FORMAT(/,5X,'Enter the Pasquill stability class: (A,B,C',
      $'D,E,F) <D> ',$,)
1152  format(' Enter the averaging time (s) for estimating DELTAY: ',$,)
1159  format(/, ' The values for the atmospheric parameters',
      $' are set as follows:',  

      $/, ' DELTAY:           ',F12.4,  

      $/, ' BETAY:           ',F12.4,  

      $/, ' Monin-Obukhov length:   ',F12.4,' m',  

      $/, ' Sigma X Coefficient:   ',F12.4,  

      $/, ' Sigma X Power:        ',F12.4,  

      $/, ' Sigma X Minimum Distance: ',F12.4,' m',  

      $/, ' Do you wish to change any of these?',  

      $/, ' (No,Delтай,Betай,Length,Coefficient,Power,Minimum) <N> ',$,)
1160  format(/, ' The values for the atmospheric parameters',
      $' are set as follows:',  

      $/, ' DELTAY:           ',F12.4,  

      $/, ' BETAY:           ',F12.4,  

      $/, ' Monin-Obukhov length: infinite',  

      $/, ' Sigma X Coefficient:   ',F12.4,  

      $/, ' Sigma X Power:        ',F12.4,  

      $/, ' Sigma X Minimum Distance: ',F12.4,' m',  

      $/, ' Do you wish to change any of these?',  

      $/, ' (No,Delтай,Betай,Length,Coefficient,Power,Minimum) <N> ',$,)
1161  FORMAT(/,5X,'The density is determined as a function of con',
      $'centration')
1162  FORMAT(5X,'by a listing of ordered triples supplied by the ',
      $'user.')
1163  FORMAT(5X,'Use the following form:')
1164  FORMAT(/,5X,5X,'first point',6X,'-- pure air  y=0.0,Cc=0.,',
      1      'RHOG=RHOA=',F7.5' kg/m**3')
1165  FORMAT(3(15X,'.',/))
1166  FORMAT(5X,5X,'last point',7X,'-- pure gas  y=1.0,Cc=RHOE',
      1      'RHOG=RHOE')
1170  FORMAT(/,5X,'ENTER THE NUMBER OF DATA TRIPLES (max=',i2,','),
      $' FOR THE DENSITY FUNCTION: ',$,)
1180  FORMAT(/,5X,'Enter Mole frac, Cc (kg/m**3), then RHOG ',
      1      '(kg/m**3) by triples')
c
c
1190  FORMAT(/,5X,10X,'Source Description')
1200  FORMAT(1X,/,5X,'The description of the primary source mass')

```

```

1210  FORMAT(5X,'evolution rate E and radius R1 for a transient',//,
      $5x,'release is input by ordered triples as follows:')
1220  FORMAT(/,5X,3X,'first point',8X,
      $'--- t=0, E(t=0), R1(t=0) (initial, ',
      $'nonzero values)')
1221  FORMAT(5X,3X,'second point',7X,
      $'--- t=t1, E(t=t1), R1(t=t1)')
1223  FORMAT(5X,3X,'last nonzero point -- ',
      $'t=TEND, E(t=TEND), R1(t=TEND)')
1230  FORMAT(5X,3X,'next to last point -- t=TEND+1., E=0., R1=0.')
1240  FORMAT(5X,3X,'last point           -- t=TEND+2., E=0., R1=0.')
1250  FORMAT(/,5X,'Note: the final time (TEND) is the last time ',
      $'when E and R1 are non-zero.'//)
1260  FORMAT(/,5X,'Enter the number of triples (max= ',i2,')',
      $' starting with t=0. and ending',/,5X,'with t=TEND+2.',
      $' for the source description: ',$)
1270  FORMAT(/,5X,'Enter TIME (sec), EVOLUTION RATE (kg/s), ',
      $'and POOL RADIUS (m)')
1280  FORMAT(/,5X,'The suggested LOWEST CONCENTRATION OF INTEREST ',
      $'(gas_lfl/2.)',/,5x,' is ',1pg13.5,
      $' kg/m**3. Enter the desired value: ',$)
1290  format(/' This file was not found.')
1300  FORMAT(/' Do you have an input file for the Density ',
      $'function? [y or N] ',$)
1310  FORMAT(Q,A20)
1320  FORMAT(' Enter the file name: [DIR]FILE_NAME.EXT ',$)
1330  FORMAT(' Do you have an input file for the Source ',
      $'Description? [y or N] ',$)
1340  format(/' Air density corrected to ',1pg13.5' kg/m**3',/)
1341  format(/' Contaminant density corrected to ',1pg13.5' kg/m**3',/)

c
c
1400  format(//,' Is this a Steady state simulation? <y or N> ',$)
1410  format(a4)
1415  format(a3)
1420  format(/' Enter the desired evolution rate [=] kg/sec : ',$)
1430  format(' Enter the desired source radius [=] m : ',$)
1440  format(/' Specification of source parameters.',/)
1450  format(//,' Enter the initial mass of pure gas',
      $' over the source. (kg)',/, '(Positive or zero): ',$)
1460  format(/' Is this a release of pure (P) or diluted (d) material',
      $' specified above? <P or d> ',$)
1470  FORMAT(' Enter the desired primary source contaminant mass ',
      $'fraction: ',$)
1480  FORMAT(' Enter the desired primary source temperature [=] K : ')
c
c
1500  format(//,' Enter the ambient temperature(K) and pressure',
      1      '(atm): ',$)
1510  format(//,' Enter the code name of the diffusing species: ',$)
1520  format(//,' The characteristics for the gas are set as follows:',/)


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```

$' Molecular weight: ',f7.2,/
$' Storage temperature [K]: ',1pg13.5,/
$' Density at storage temperature, PAMB [kg/m**3]: ',1pg13.5,/
$' Mean Heat capacity constant ',1pg13.5,/
$' Mean Heat capacity power ',1pg13.5,/
$' Upper Flammability Limit [mole frac] ',1pg13.5,/
$' Lower Flammability Limit [mole frac] ',1pg13.5,/
$' Height of Flammability Limit [m] ',1pg13.5,/
$' Do you wish to change any of these? ',
$'(No,Mole,Temp,Den,Heat,Power,Upper,Lower,Z)',

$' <N> ',$)

1530 format(' Enter the desired Storage Temperature: ',$)
1535 format(' Enter the desired Density at Storage ',
1      'Temperature and', ' ambient pressure: ',$)
1550 format(' Enter the desired Molecular Weight: ',$)
1570 format(' Enter the desired Mean Heat Capacity constant: ',$)
1571 format(' Enter the desired Mean Heat Capacity power: ',$)
1572 format(' Enter the desired Upper Flammability Limit: ',$)
1573 format(' Enter the desired Lower Flammability Limit: ',$)
1574 format(' Enter the desired Height for the flammable limit calcula',
1      'tions: ',$)
1580 format(/,' The ambient humidity can be entered as Relative ',
1      'or Absolute.',/, ' Enter either R or A <R or a>: ',$)
1585 format(' Enter the absolute humidity (kg water/kg BDA): ',$)
1586 format(' This is a relative humidity of ',1pg13.5,' %')
1587 format(' Enter the relative humidity (%): ',$)
1588 format(/,' Ambient Air density is ',1pg15.7,' kg/m**3')

c
c
1600 format(' Enter the desired DELTAY: ',$)
1620 format(' Enter the desired BETAY: ',$)
1660 format(' Note: For infinity, RML = 0.0',/,
$' Enter the desired Monin-Obukhov length: (m) ',$)
1670 format(' Enter the desired Sigma X Coefficient: ',$)
1680 format(' Enter the desired Sigma X Power: ',$)
1690 format(' Enter the desired Sigma X Minimum distance: (m) ',$)

c
2000 format(/,' Is this an Isothermal spill? <y or N> ',$)
2020 format(/,' Is heat transfer to be included in the',
1      ' calculations <y or N> ',$)
2030 format(' Enter the surface temperature [=] K : ',$)
2040 format(' Do you want to use the built in correlation?',
1      ' the LLNL correlation, or',/,' enter',
1      ' a particular value?',/,
1      ' (Corr,LLNLcorr,Value) <C> ',$)
2043 format(/,' The form of the correlation is:',/,
1      ' 6x,Q = (Vh * rho * cp) * area * (tsurf-temp)',//,
1      ' with Vh = ',1pg13.5,' m/s.',//,
1      ' Do you wish to change the value of Vh? (y or N): ',$)
2045 format(' Do you want to use the built in correlation or enter',
1      ' a particular value?',/,' <C or v> ',$)

```

```

2050  format(' Enter the HT coefficient value [=] J/m**2/s/K : ',$)
2100  format(/, ' Is water transfer to be included in the',
1      ' source <y or N> ',$)
2120  format(' Enter the WT coefficient value [=] kg/m**2/s : ',$)
C
2200  FORMAT(1X,/,5X,'The description of the primary source contam-')
2210  FORMAT(5X,'inant mass rate E, radius R1, contaminant mass',/,
$5x,'fraction Wc,s, and temperature Ts for a transient',/,
$5x,'release is input as a function of time as follows:')
2220  FORMAT(/,5X,3X,'first point',8X,
$'-- t=0., E(t=0), R1(t=0), Wc,s(t=0), Ts(t=0)',/,
$      5x,3x,18x,10x,'(initial, ',
$'nonzero values)')
2221  FORMAT(5X,3X,'second point',7X,
$'-- t=t1, E(t=t1), R1(t=t1), Wc,s(t=t1), Ts(t=t1)')
2223  FORMAT(5X,3X,'last nonzero point -- ',
$'t=TEND, E(t=TEND), R1(t=TEND), Wc,s(t=TEND), Ts(t=TEND)')
2230  FORMAT(5X,3X,'next to last point -- t=TEND+1., E=0., R1=0.,',
$      ' Wc,s=1., Ts=Tamb')
2240  FORMAT(5X,3X,'last point          -- t=TEND+2., E=0., R1=0.,',
$      ' Wc,s=1., Ts=Tamb')
2250  FORMAT(/,5X,'Note: the final time (TEND) is the last time ',
$'when E and R1 are non-zero.',/)
2260  FORMAT(/,5X,'Enter the number of times (max= ',i2,')',
$' starting with t=0. and ending',/,5x,'with t=TEND+2.',
$' for the source description: ',$)
2270  FORMAT(/,5X,'Enter TIME (sec), CONTAMINANT MASS RATE (kg C/s), ',
$'POOL RADIUS (m)',/,10X,'CONTAMINANT MASS FRACTION (kg C',
$      '/kg mix), and TEMPERATURE (K)')
2280  FORMAT(/,5X,'The suggested LOWEST CONCENTRATION OF INTEREST ',
$'(gas_lfl/2.)',/,5x,' is ',1pg13.5,
$' kg/m**3. Enter the desired value: ',$)
C
      RETURN
      END
#####

```

```

subroutine limit(func, x0, xinc, xhigh, xlow)
c
c*** subroutine to establish the upper and lower limits for ZBRENT
c
      implicit real*8(a-h,o-z), integer*4(i-n)
      rrr = 1.0D0
      aflag = 1.0D0
      bflag = 1.0D0
      fc = func(x0)
100   continue
      if(aflag .eq. 1.0D0) then
          aaa = x0 + xinc*rrr
          if(aaa .ge. xhigh) then
              aaa = xhigh
              aflag = 0.0D0
          endif
          fa = func(aaa)
          if(sign(1.0D0,fa)*sign(1.0D0,fc) .lt. 0.0D0) then
              xhigh = aaa
              xlow = aaa - xinc*1.01D0
              return
          endif
      endif
      if(bflag .eq. 1.0D0) then
          bbb = x0 - xinc*rrr
          if(bbb .le. xlow) then
              bbb = xlow
              bflag = 0.0D0
          endif
          fb = func(bbb)
          if(sign(1.0D0,fb)*sign(1.0D0,fc) .lt. 0.0D0) then
              xhigh = bbb + xinc*1.01D0
              xlow = bbb
              return
          endif
      endif
      if(sign(1.0D0,fa)*fb .gt. 0.0D0) then
          rrr = rrr + 1.0D0
          if(rrr .gt. 400.0D0) then
              write(6,*) 'limit1? rrr>',rrr
              rrr = 0.
              rrr = 1./rrr
              write(6,*) rrr
              stop 'limit1? rrr>400'
          endif
          goto 100
      endif
      xhigh = aaa
      xlow = bbb
      return
  end
####
```

```

C.....  

C  

C      SUBROUTINE FOR SOURCE EVALUATION WHEN NO GAS BLANKET  

C      IS PRESENT.  

C  

C      SUBROUTINE NOBL(timeout, reflag)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      include 'sys$degadis:DEGADIS1.dec'  

C  

COMMON  

$/GEN1/ PTIME(igen), ET(igen), R1T(igen), PWC(igen), PTEMP(igen),  

$      PFRACV(igen), PENTH(igen), PRHO(igen)  

$/ERROR/ STPIN,ERBND,STPMX,WTRG,WTtm,WTya,WTyc,WTeb,WTmb,WTuh,XLI,  

$ XRI,EPS,ZLOW,STPINZ,ERBNDZ,STPMXZ,SRCOER,srcss,srcscut,  

$ htcut,ERNOBL,NOBLpt,crfger,epsilon  

$/PARM/ UD,ZO,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$/comatm/ istab,tamb,pamb,humid,isoft,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$/com_ss/ ess,slen,swid,outcc,outsz,outb,outl,swal,senl,srhl  

$/phflag/ check1,check2,again,check3,check4,check5  

$/ALP/ ALPHA,alpha1  

$/phicom/ iphifl,delay  

C  

REAL*8 ML,K  

C  

LOGICAL REV  

logical check1,check2,again,check3,check4,check5  

logical reflag  

DATA REV/.TRUE./  

REAL*8 L  

C  

data h/0./,Ri/0./  

data delt_min/0.5/  

C  

DELTAT = (TEND - TSC1)/FLOAT(NOBLPT)  

if(deltat .lt. delt_min) then  

    noblpt = int((tend-tsc1)/delt_min) +1  

    deltat = (tend-tsc1)/float(noblpt)  

endif  

C  

T0 = TSC1  

IF(DELTAT .LT. 2.) GO TO 100  

C  

WRITE(lunlog,1100)  

WRITE(lunlog,*) DELTAT  

1100 FORMAT(5X,'TIME INCREMENT USED ON LAST PORTION OF SOURCE CALC')  

C

```

```

100  CONTINUE
C
C      ESTABLISH LOOP TO FINISH SOURCE
C
DO 110 I = 1,NOBLPT
C
TIME = T0 + FLOAT(I)*DELTAT
IF(I .EQ. NOBLPT) TIME = TEND
L = 2.0*AFGEN2(PTIME,R1T,TIME,'R1T-BL')
erate = AFGEN2(PTIME,ET,TIME,'ET-BL')
flux = EraTe/(pi*L*L/4.00)
PWCP = AFGEN2(PTIME,PWC,TIME,'ET-BL')
PWAP = (1.00 - PWCP)/(1.00 + HUMID)
HPRIM = AFGEN2(PTIME,PENTH,TIME,'ET-BL')
CALL SETDEN(PWCP, PWAP, HPRIM)
RHOP = AFGEN2(PTIME,PRHO,TIME,'RH-BL')
CCP = PWCP * RHOP
C
qstar = CCP * k*ustar*alpha1*dellay/(dellay-1.)/phihat(RHOP,L)
if(abs(flux/qstar) .gt. ernobl .and. reflag) then
    check3 = .true.
    timeout = time
    return
end if
C
call szf(flux,L,PWCP,sz,cclay,wclay,rholay)
cc = cclay*dellay
if(cc.gt.ccp) cc=ccp
C
c... Ensure that the SZF calculated values close the material balance
C
twidth = pi*L/4.00
qqqq = erate/u0/z0*alpha1 * z0**alpha1 / twidth
cc = qqqq/sz**alpha1
if(cc.gt.ccp) cc=ccp
sz = (qqqq/cc)**(1.00/alpha1)
C
call adiabat(0,wc,wa,yc,ya,cc,rho,wm,enthalpy,temp)
C
IF(Erate .LT. EMAX) GO TO 220
EMAX = Erate
RM = AFGEN2(PTIME,R1T,TIME,'R1T-BL')
SZM = SZ
220  CONTINUE
RLIST = AFGEN2(PTIME,R1T,TIME,'R1T-BL')
RMAX = dMAX1(RMAX,RLIST)
C
WRITE(9,2000) TIME,RLIST,h,flux,SZ,yc,ya,rho,Ri,wc,wa,enthalpy,temp
C
if(i.eq.3 .and. check4) goto 500      ! steady
C

```

```
110  CONTINUE
     RETURN
c
c
500  continue           ! steady state completion
     outcc = cc
     swcl = wc
     swal = wa
     senl = enthalpy
     srhl = rho
     outsz = sz
     outl = 2. * rlist
     outb = pi * rlist**2 /outl/2.
     return
2000 format(1pg16.9,1x,1pg16.9,<iout_src-2>(1x,1pg13.6))
     END
#####

```

```

C.....  

C  

C      SUBROUTINES OB AND OABOUT ARE USED IN THE OBSERVER INTEGRATIONS  

C          OVER THE SOURCE.  

C  

C      SUBROUTINE OB(time,Y,D,PRMT)  

C  

C      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      include 'sys$degadis:DEGADIS2.dec'  

C  

C      COMMON  

$/GEN3/ radg(2,maxl),qstr(2,maxl),srcden(2,maxl),srcwc(2,maxl),  

$ srcwa(2,maxl),srcenth(2,maxl)  

$/PARM/U0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$/ALP/ALPHA,alpha1  

$/phicom/ iphifl,dellay  

C  

C      REAL*8 K,ML  

C      logical flag  

C  

C      DIMENSION Y(1),D(1),PRMT(1)  

C      INTEGER HWIDTH,Mrate,Crate,BDARate,Hrate  

C      DATA HWIDTH/1/,Mrate/2/,Crate/3/,BDARate/4/,Hrate/5/  

C  

C*** PASS TO IN PRMT(6)  

C  

C      flag = isofl.eq. 1 .or. ihtfl.eq. 0  

C  

C      TOL = PRMT(6)  

C      XUP = prmt(7)  

C      XI = XIT(TIME,TOL)  

C      RG = AFGEN(RADG,TIME,'RADG')  

C      RLEN = PRMT(13)  

C  

C      BIPR = 0.  

C      D(HWIDTH)= 0.  

C      D(Crate) = 0.  

C      D(Mrate) = 0.  

C      D(BDARate)= 0.  

C      D(Hrate) = 0.  

C  

C*** We must deal with the case when the observer sees no gas, but the  

C*** integration is not over yet. Since the derivatives have been set  

C*** to zero above, no work on the derivatives is needed, but the values  

C*** in PRMT(14-18) must be preserved. Since these will be switched  

C*** in OABOUT, switch them now so the old values will be retained.

```

```

c
      IF(ABS(XI) .GE. RG) then      ! use the last values
          PRMT(8) = prmt(14)        ! cclay
          PRMT(9) = prmt(15)        ! wclay
          PRMT(10) = prmt(16)       ! walay
          PRMT(11) = prmt(17)       ! enthlay
          PRMT(12) = prmt(18)       ! rholay
          RETURN
      endif
      BIPR = sqrt(RG*RG - XI*XI)
c
      UI = UIT(TIME,TOL)
c
      Q    = AFGEN(QSTR,TIME,'QSTR')
      WC   = AFGEN(srcwc,time,'srcwc')
      WA   = AFGEN(srcwa,time,'srcwa')
      ENTH = AFGEN(srcenth,time,'srcenth')
c
      WCLAY = Y(Crate)/Y(Mrate)
      WALAY = Y(BDARATE)/Y(Mrate)
      if(.not.flag) ENTHLAY = Y(Hrate)/Y(Mrate)
c
      call tprop(1,wclay,walay,enthalay,yc,ya,wm,temp,rholay,cp)
      CCLAY = WCLAY*RHO
c
      PRMT(8) = CCLAY
      PRMT(9) = WCLAY
      PRMT(10)= WALAY
      PRMT(11)= ENTHLAY
      PRMT(12)= RHO
c
      CC  = CCLAY*DELLAY
      RHO = DELLAY*(RHO-RHOA) + RHOA
c
      SZOB = 0.01
      ARG = Q*(XI-XUP)/CC/(U0*Z0/ALPHA1)
      if(XI.gt. XUP .and. ARG.gt.0.)
      1 SZOB = ARG***(1./ALPHA1) * Z0
c
      HEFF = GAMMAF/ALPHA1* SZOB
      RISTR=RIF(RHO,HEFF)
      PHI = PHIF(RISTR,0.)
      WELAY = DELLAY * KUSTAR* ALPHA1/PHI
c
      D(HWIDTH)= UI * BIPR / RLEN
      D(CRATE) = D(HWIDTH)*RLEN * Q
      D(MRATE) = (Q/WC + RHOA*WELAY) * D(HWIDTH)*RLEN
      D(BDARATE)= (Q*WA/WC + RHOA*WELAY/(1.+HUMID)) * D(HWIDTH)*RLEN
      if(flag) return
      D(HRATE) = Q * ENTH/WC * D(HWIDTH)*RLEN
c

```

```
1000  FORMAT(' ?OB? -- Value of XI ',1pG13.4,'; Value of RG ',
$ 1pG13.4)
      RETURN
      END

c
c
      SUBROUTINE OBOUT( X, Y, DERY, IHLF, NDIM, PRMT)
c
      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

      DIMENSION X(1), Y(1), DERY(1), PRMT(1)

c
      PRMT(14) = prmt(8)      ! cclay
      PRMT(15) = prmt(9)      ! wclay
      PRMT(16) = prmt(10)     ! walay
      PRMT(17) = prmt(11)     ! enthlay
      PRMT(18) = prmt(12)     ! rholay
      RETURN
      END

#####
```

```

C.....  

C  

C      FUNCTION PSI  

C  

C*** AS PER COLENBRANDER --  

C  

C*** THIS FUNCTION HAS BEEN DERIVED FROM BUSINGER,J.A.  

C*** WORKSHOP ON MICROMETEOROLOGY, CHAPTER 2, HAUGEN,D.A. (ED.)  

C*** AMERICAN METEOROLOGICAL SOCIETY.  

C  

FUNCTION PSIF(Z,ML)

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

C
      include 'sys$degadis:DEGADIS1.dec'
C
      REAL*8 ML
C
      IF( ML ) 10,20,30
C
      10   A = (1.-15.*Z/ML)**.25
            PSIF = 2.*dLOG((1.+A)/2.) + dLOG((1.+A*A)/2.) - 2.*daTAN(A) +
$ PI/2.
            RETURN
C
      20   PSIF = 0.
            RETURN
C
      30   PSIF = -4.7*Z/ML
            RETURN
            END
#####

```

```

C.....  

C  

C      SUBROUTINES FOR PSEUDO-STEADY STATE INTEGRATION.  

C  

C          SUBROUTINE PSS(DIST,Y,DERY,PRMT)  

C  

C              Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C              include 'sys$degadis:DEGADIS2.dec/list'  

C  

C              parameter (zero=1.D-10, rcrit=2.D-3)  

C  

C          COMMON  

C/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,  

$ gas_ufl,gas_lfl,gas_zsp,gas_name  

$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

$/ALP/ ALPHA,alpha1  

$/phicom/ iphifl,delay  

$/sprd_con/ ce, delrhamin  

C  

REAL*8 K,ML  

LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

C  

DIMENSION Y(1),DERY(1),PRMT(1)  

DATA rhoh/1/,SY2/2/,BEFF/3/,dh/4/,Mhi/5/,Mlow/6/  

INTEGER rhoh,SY2,BEFF,dh, Mhi, Mlow  

C  

C*** PRMT I/O SETUP
C***   I      VALUE      IN/OUT
C***   ----  -----  -----
C***   6      E          IN
C***   7      Cc         OUT
C***   8      Bb         OUT
C***   9      CON DERY(BEFF)    IN
C***  10      CON DERY(SZ)    IN
C***  11      NREC(I,1)      OUT -- STARTS OUTPUT COUNTER
C***  12      DIST         OUT
C***  13
C***  14      yc          OUT
C***  15      rho          OUT
C***  16      temp         OUT - if recorded
C***  17      gamma        OUT - if recorded
C***  18
C***  19
C***  20
C***  21      sz
C***  22      sz

```

```

C
      Erate = PRMT(6)
      Bb = Y(BEFF) - SQRTPI/2.*sqrt(Y(SY2))
C
c using the last value for Sz
C
      sz0 = PRMT(22)
      sz = sz0
C
C*** MATERIAL BALANCE
C
      iii = 0
100   Cc = Erate*ALPHA1/2./U0*(Z0/SZ)**ALPHA/SZ/Y(BEFF)
      cclay = cc/dellay
      call addheat(cclay,y(dh),rholay,temp,cp)
      prod = dmax1( Y(rhouh)/rholay/prmt(18), zero)
      sz = ( prod )**(.1./alpha1) * z0
      dif = abs(sz - sz0)/(abs(sz)+abs(sz0)+zero)
      if(dif .gt. rcrit) then
          sz0 = sz
          iii=iii+1
          if(iii .gt. 20) call trap(32)
          goto 100
          endif
      prmt(20) = rholay
      prmt(21) = sz
      HEFF = GAMMAF/ALPHA1*SZ

      call adiabat(0,wc,wa,yc,ya,cc,rho,wm,enth,temp)
      call adiabat(0,wc,wa,yclay,ya,cclay,rholam,wml,enth,temp)
C
      rit = 0.
      if(isofl.eq.0 .and. ihtfl.ne.0) then
          call addheat(cc,dellay*y(dh),rho,temp,cp)
          rit = rift(temp,heff)
          endif
      RISTR = RIF(RHO,HEFF)
      PHI = PHIF(RISTR,rit)
C
C*** CALCULATE DERIVATIVES
C
      DERY(BEFF) = 0.
      delrho = rho-rhoa
      IF(delrho .GT. delrhomin) DERY(BEFF) = PRMT(9)*sqrt(delrho/rhoa)
      $   *(SZ/Z0)**(.5 - ALPHA)
C
      DERY(SY2) = 8.*BETA/PI*Y(BEFF)**2 *
      $   (DELTA*SQRTI02/Y(BEFF)) ** (1./BETA)
C
C
      heigh = heff*dellay

```

```

yw = 1.-yclay-ya
yw = min( max( yw, 0.00 ), 1.00 )
call surface(temlay,heigh,rholay,wml,cp,yw,water,q rte)
if(temp.ge. tsurf .or. temlay.ge. tamb) q rte = 0.
rhouhb = rholay* prmt(18) * (sz/z0)**alpha1 * Y(beff)
d_rhouhb = prmt(19)*y(beff)/phi
dERY(dh) = (q rte*Y(beff)/dellay - Y(dh)*d_rhouhb)/rhouhb
dERY(rhouh) = (d_rhouhb-Y(rhouh)*dERY(beff))/Y(beff)

c
c*** Calculate the derivative for the total mass above the UFL and LFL
c
gamma = (rho-rhoa)/cc                      ! gamma
if(check4) then
  DERY(mlow) = 0.
  DERY(mhi) = 0.
  if( isofl.eq.1 .or. ihtfl.eq.0 ) then
    call adiabat(2,aa,dd,gas_ufl,ee,chi ,gg,hh,pp,oo)
    call adiabat(2,aa,dd,gas_lfl,ee,cflow,gg,hh,pp,oo)
  else
    call adiabat(-2,aa,dd,gas_ufl,ee,chi ,gg,hh,gamma,oo)
    call adiabat(-2,aa,dd,gas_lfl,ee,cflow,gg,hh,gamma,oo)
  endif
  gamhi = 2.00* Cc * Bb * Sz / alpha1
  gammmax = 2.00* Cc * Sz * GAMMAF/alpha1 * (Bb+sqrt(pi)/2.00*sqrt(Y(Sy2)))

  if(cc.gt.cflow) then
    wlow = Dlog(cc/cflow)
    gamlow = gaminc(1.00/alpha1, wlow ) * gamhi
    DERY(mlow)= gamlow + 2.00*cflow*sqrt(Y(Sy2))*Sz/alpha1*series(wlow)
    DERY(mlow)= DMIN1( DERY(mlow), gammmax )
  endif

  if(cc.gt.chi) then
    whi = Dlog(cc/chi )
    gamhi = gaminc(1.00/alpha1, whi ) * gamhi
    DERY(mhi) = gamhi + 2.00*chi *sqrt(Y(Sy2))*Sz/alpha1*series(whi )
    DERY(mhi) = DMIN1( DERY(mhi ), gammmax )
  endif
endif

c
c*** RETURNED VALUES
c
PRMT(7) = Cc
PRMT(8) = Bb
prmt(14)= yc
prmt(15)= rho
prmt(16)= temp
prmt(17)= gamma
RETURN
END
#####

```

```

C.....  

C  

C      SUBROUTINE PSSOUT  

C  

C      SUBROUTINE PSSOUT(X,Y,D,IHLF,NDIM,PRMT)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      include 'sys$degadis:DEGADIS2.dec'  

C  

C      parameter (npss=9, zero=1.e-10)  

C  

C      COMMON  

$/PARM/U0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/STP/STPO,STPP,ODLP,ODLLP,STPG,ODLG,ODLLG  

$/PHLAG/CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

$/STOPIT/TSTOP  

C  

REAL*8 K,ML  

LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

DIMENSION Y(1),D(1),PRMT(1),BKSP(npss),OUT(npss),CURNT(npss)  

C  

C*** OUTPUT PARAMETERS  

C  

C*** FROM PSS          OUTPUT TO MODEL  

C*** -----  

C*** X                  DIST  

C*** PRMT(7)            Cc  

C*** Y(1)                SZ  

C*** Y(2)                SY2  

C*** PRMT(8)            B  

C*** PRMT(13)           TO(I)  

C*** prmt(14)            yc  

C*** prmt(15)            rho  

C*** prmt(16)            temp  

C*** prmt(17)            gamma  

C  

ERM = 0.  

TSL = TS(PRMT(13),X)  

prmt(22) = prmt(21)  

IF(PRMT(11) .NE. 0.) GO TO 90  

C  

C*** STARTUP FOR THE OUTPUT ROUTINE  

C  

RII = -100./STPP  

RI = 0.  

CURNT(1) = X  

curnt(2) = prmt(14)    ! yc

```

```

CURNT(3) = PRMT(7)      ! cc
currnt(4) = prmt(15)    ! rho
currnt(5) = prmt(17)    ! gamma
currnt(6) = prmt(16)    ! temp
CURNT(7) = prmt(21)    ! sz
CURNT(8) = sqrt(Y(2))   ! sy2
CURNT(9) = PRMT(8)      ! b
IF(prmt(8) .LE. 0.) CALL trap(16)
90  CONTINUE
C
C*** STOP INTEGRATION WHEN THE HALF WIDTH B < 0.
C
IF( PRMT(8) .LE. 0.) GO TO 1000
C
C*** SET THE CURRENT AND PREVIOUS RECORD
C
DO 100 II=1,npss
100  BKSP(II) = CURNT(II)
C
CURNT(1) = X
currnt(2) = prmt(14)    ! yc
CURNT(3) = PRMT(7)      ! cc
currnt(4) = prmt(15)    ! rho
currnt(5) = prmt(17)    ! gamma
currnt(6) = prmt(16)    ! temp
CURNT(7) = prmt(21)    ! sz
CURNT(8) = sqrt(Y(2))   ! sy2
CURNT(9) = PRMT(8)      ! b
C
C*** STOP INTEGRATION AND GET A NEW OBSERVER WHEN Cc<CcLOW
C
IF(PRMT(7).GT.CcLOW .OR. TSL.LT.TSTOP) GO TO 95
    if(prmt(11) .lt. 5.) then      ! guarantee 5 records
        erm = odlp                ! force output
        goto 95
    endif
    AGAIN = .TRUE.
    TSTOP = TSL
    GO TO 1000
95  CONTINUE
C
C*** Check the error criteria's
C
R1 = R1 + 1.
II = 2                  ! skip DIST,YC; 1 and 2
110  II = II + 1
    ER1 = ABS( (CURNT(II)-BKSP(II))/(CURNT(II)+zero) )
    ER2 = ABS( (CURNT(II)-OUT(II))/(CURNT(II)+zero) )
    ERM = dMAX1(ER1,ER2,ERM)
    IF(II .EQ. 3) II = II + 1      ! skip RHO;4
    IF(II .EQ. 7) II = II + 1      ! skip SY;8

```

```

IF(II .EQ. 5) II = II + 1      ! skip TEMP;6
IF(II .LT. npss) GO TO 110

C
C*** RECORD POINT IF ODLR IS EXCEEDED OR 80 METERS SINCE LAST RECORD
C*** RECORD FIRST POINT
C
DX = CURNT(1) - OUT(1)
IF( RI.NE.1. .AND. ERM.LT.ODLR .AND. DX.LE.ODLRP) RETURN
C
C*** IF THE NEXT INTEGRATION AFTER A POINT IS RECORDED VIOLATES THE
C*** ERROR BOUND, THE CURRENT POINT MUST BE RECORDED. OTHERWISE, THE
C*** LAST POINT TO SATISFY THE ERROR LIMITS IS RECORDED.
C
DO 120 II=1,npss
IF(RI .EQ. RII+1.) BKSP(II) = CURNT(II)
120   OUT(II) = BKSP(II)
C
RI = RII
PRMT(11) = PRMT(11) + 1.
C
WRITE(9,*) (OUT(II),II=1,npss)
RETURN
C
1000  CONTINUE
C
C*** STOP INTEGRATION
C
PRMT(12) = X
C
IF(CURNT(1) .EQ. OUT(1)) GO TO 130
C
PRMT(11) = PRMT(11) + 1.
WRITE(9,*) (CURNT(II),II=1,npss)
C
130  CONTINUE
PRMT(5) = 1.
RETURN
END
#####

```

```

C.....  

C  

C      SUBROUTINE PSSOUT  

C  

C      SUBROUTINE PSSOUT(X,Y,DERY,IHLF,NDIM,PRMT)  

C  

C      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      include 'sys$degadis:DEGADIS2.dec/list'  

C  

C      parameter (npss=9, zero=1.e-10)  

C  

C      COMMON  

C      $/PARM/U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CCLOW  

C      $/STP/STPP,OOLP,ODLLP,STPG,ODLG,ODLLG  

C      $/PHLAG/CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

C      $/com_fl/ cflag,clfl,cufl  

C      $/ALP/ALPHA,alpha1  

C  

C      logical cflag  

C      LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

C  

C      REAL*8 ML,K  

C  

C      DIMENSION Y(1),DERY(1),PRMT(1)  

C      dimension BKSP(npss),OUT(npss),CURNT(npss)  

C  

C*** OUTPUT PARAMETERS  

C  

C*** FROM PSS          OUTPUT TO MODEL  

C*** -----  

C***   X                  DIST  

C***   PRMT(7)            Cc  

C***   Y(1)                SZ  

C***   Y(2)                SY2  

C***   PRMT(8)            B  

C  

C      ERM = 0.  

C      prmt(22) = prmt(21)  

C  

C      IF(PRMT(11) .NE. 0.) GO TO 90  

C  

C*** STARTUP FOR THE OUTPUT ROUTINE  

C  

C      RII = -100./STPP  

C      RI = 0.  

C      CURNT(1) = X  

C      CURNT(2) = PRMT(14)    ! yc  

C      CURNT(3) = prmt(7)     ! cc  

C      CURNT(4) = prmt(15)    ! rho

```

```

        CURNT(5) = PRMT(17)      ! gamma
        curnt(6) = prmt(16)      ! temp
        curnt(7) = prmt(8)       ! b
        curnt(8) = prmt(21)      ! sz
        curnt(9) = sqrt(Y(2))    ! sy2
C
        90  CONTINUE
C
C*** STOP INTEGRATION WHEN THE HALF WIDTH B < 0.
C
        IF( PRMT(8) .LE. 0.) GO TO 1000
C
C*** STOP INTEGRATION when Cc<CcLOW
C
        IF(PRMT(7).GT.CcLOW) GO TO 95
            if(prmt(11) .lt. 5.) then      ! force output
                erm = odlp
                goto 95
            endif
        AGAIN = .TRUE.
        GO TO 1000
        95  CONTINUE
C
C*** SET THE CURRENT AND PREVIOUS RECORD
C
        DO 100 II=1,npss
        100 BKSP(II) = CURNT(II)
C
        CURNT(1) = X
        CURNT(2) = PRMT(14)      ! yc
        CURNT(3) = prmt(7)       ! cc
        CURNT(4) = prmt(15)      ! rho
        CURNT(5) = PRMT(17)      ! gamma
        curnt(6) = prmt(16)      ! temp
        curnt(7) = prmt(8)       ! b
        curnt(8) = prmt(21)      ! sz
        curnt(9) = sqrt(Y(2))    ! sy2
C
        RI = RI + 1.
        II = 1
        110 II = II + 1
        ER1 = ABS( (CURNT(II)-BKSP(II))/(CURNT(II)+zero) )
        ER2 = ABS( (CURNT(II)-OUT(II))/(CURNT(II)+zero) )
        ERM = dMAX1(ER1,ER2,ERM)
        IF(II .EQ. 3) II = 6      ! skip density,gamma,temp
        IF(II .LT. npss-1) GO TO 110     ! skip sy
C
C*** RECORD POINT IF ODLP IS EXCEEDED OR 80 METERS SINCE LAST RECORD
C*** RECORD FIRST POINT
C
        DX = CURNT(1) - OUT(1)

```

```

      IF( RI.NE.1. .AND. ERM.LT.ODLP .AND. DX.LE.ODLLP) RETURN
C
C*** IF THE NEXT INTEGRATION AFTER A POINT IS RECORDED VIOLATES THE
C*** ERROR BOUND, THE CURRENT POINT MUST BE RECORDED. OTHERWISE, THE
C*** LAST POINT TO SATISFY THE ERROR LIMITS IS RECORDED.
C
      DO 120 II=1,npss
      IF(RI .EQ. RII+1.) BKSP(II) = CURNT(II)
120    OUT(II) = BKSP(II)
C
      RI = RII
      PRMT(11) = PRMT(11) + 1.
C
      call ssout(out)
      RETURN
C
      1000  CONTINUE
C
C*** STOP INTEGRATION
C
      PRMT(12) = X
C
      IF(RI .EQ. 0.) CALL trap(16)
C
      IF(CURNT(1) .EQ. OUT(1)) GO TO 130
C
      PRMT(11) = PRMT(11) + 1.
      call ssout(out)
C
      130  CONTINUE
      PRMT(5) = 1.
C
      RETURN
      END
#####

```

```

C.....  

C  

C      RICHARDSON NUMBER (RI*)  

C  

C      FUNCTION RIF(RHOG,HEFF)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      COMMON  

$ /PARM/U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CCLOW  

C  

REAL*8 ML,K  

C  

RIF = G*(RHOG-RHOA)/RHOA*HEFF/USTAR/USTAR  

C  

RETURN  

END  

C  

C.....  

C  

C      RICHARDSON NUMBER (RIt)  

C  

FUNCTION RIFT(temp,HEFF)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      COMMON  

$ /PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$ /comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$ /alp/ alpha,alpha1  

C  

REAL*8 ML,K  

C  

wind = u0*(heff/z0)**alpha  

RIFT = dmax1(G*(tsurf-temp)/temp*HEFF/USTAR/wind,0.00)  

C  

RETURN  

END  

C  

C.....  

C  

C      PHI FUNCTION  

C  

FUNCTION PHIF(RI,rit)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C

```

```

common /phicom/ iphifl,dellay
c
      phif= 0.
      goto(10,1000,2000,3000,4000,9000),iphifl
      goto 9000
c
      10    IF(RI) 100,200,300
c
      100   PHIF = 0.74/(1. + 0.65*ABS(RI)**.6)
             RETURN
c
      200   PHIF = 0.74
             RETURN
c
      300   PHIF = 0.74 + 0.25*(RI)**0.7 + 1.2E-7*RI*RI*RI
             RETURN
c
c
      1000  IF(RI) 1100,1200,1300
c
      1100  PHIF = 0.88/(1. + 0.65*ABS(RI)**.6)
             RETURN
c
      1200  PHIF = 0.88
             RETURN
c
      1300  PHIF = 0.88 + 9.9e-2*(RI)**1.04 + 1.4E-25*RI**5.7
             RETURN
c
c
c
      2000  corr1 = 0.25* rit**.666666 + 1.
             corr = sqrt(corr1)
             riw = ri/corr1
             IF(RI) 2100,2200,2300
c
      2100  PHIF = 0.88/(1. + 0.65*ABS(RIW)**.6)/corr
             RETURN
c
      2200  PHIF = 0.88/corr
             RETURN
c
      2300  PHIF = (0.88 + 9.9e-2*(RIW)**1.04 + 1.4E-25*RIW**5.7)/corr
             RETURN
c
c
      3000  corr1 = 0.25* rit**.666666 + 1.
             corr = sqrt(corr1)
             riw = ri/corr1
             IF(RI) 3100,3200,3300
c

```

```
3100  PHIF = 0.88/corr
      RETURN
c
3200  PHIF = 0.88/corr
      RETURN
c
3300  PHIF = (0.88 + 9.9e-2*(RIW)**1.04 + 1.4E-25*RIW**5.7)/corr
      RETURN
c
c
4000  PHIF = 0.88
      RETURN
c
c
9000  call trap(29)
      return
      END
c
c
c.....-----
c
c
      function phihat(rho,fetch)

      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

c
      common
$/parm/ u0,z0,zr,ml,ustar,k,g,rhoe,rhoa,delta,beta,gammaf,cclow
$/alp/ alpha,alpha1
$/phicom/ iphifl,dellay
c
      real*8 k,ml
c
      data phic/3.1/
c
      if(rho .le. rhoa) then
          phihat = 0.88
          return
          endif
c
      pow = 1./alpha1
      p1 = 1.04/alpha1
      p1i = 1.00/p1
      p2 = 1. + p1i
      p3 = (alpha - 0.04)/1.04
      p4 = (1.08 - alpha)/1.04
      Ci = g*(rho-rhoa)/rhoa*z0/ustar**2*gammaf/alpha1
      Ci = Ci* (k*ustar*alpha1**2 /u0/z0/ phic*dellay/(dellay-1.)) ** pow
c
      Ri = Ci*fetch**pow
```

```

      zzz = - 0.099*Ri**1.04/0.88
c
      if(abs(zzz) .lt. 1.) then
          phihat = 0.88/gseries(1.00, p1i, p2, zzz)
      else
          zzz = 1.00/zzz
          phihat = 0.88/(-zzz/(1.-p1)*gseries(1.00, -p3, p4, zzz)
                     + gamma(p1i)*gamma(p4)/(p1-1.)* (-zzz)**(p1i))
      endif
c
      return
end

c
c-----.
c
c      function to calculate the hypergeometric series
c
      function gseries( aaa, bbb, ccc, zzz)

      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

      data crit/1.0-5/
c
c*** check "zzz" to see if it is greater than 0.999.  If so, calculate
c      "gseries" for "zzz"=0.999 to avoid excessive execution times.
c
      if( abs(zzz) .gt. 0.999) then
          zzzz = sign(0.99900, zzz)
      else
          zzzz = zzz
      endif

      term = aaa*bbb/ccc*zzzz
      sumo = 1.00 + term

      do k = 1,100000
          term = term * (aaa+k)*(bbb+k)/(ccc+k) * zzzz/(k+1)
          sum = sumo + term
          if( abs(term/sum) .le. crit) goto 1000
          sumo = sum
      enddo

      write(6,*) 'GSERIES? did not converge in RIPHIF'
      stop

1000  gseries = (sum + sumo)/2.00
      return
end
#####

```

C
C
C SUBROUTINE RKGST
C
C.....
C
C This routine was originally supplied by Digital Equipment
C Corporation as part of the Scientific Subroutine Package
C available for RT-11 as part of the Fortran Enhancement
C Package. It was upgraded for use as the integration
C routine in this package.
C
C.....
C
C PURPOSE
C TO SOLVE A SYSTEM OF FIRST ORDER ORDINARY DIFFERENTIAL
C EQUATIONS WITH GIVEN INITIAL VALUES.
C
C USAGE
C CALL RKGST (PRMT,Y,DERY,NDIM,IHLF,FCT,OUTP,AUX)
C PARAMETERS FCT AND OUTP REQUIRE AN EXTERNAL STATEMENT.
C
C DESCRIPTION OF PARAMETERS
C
C PRMT AN INPUT AND OUTPUT VECTOR WITH DIMENSION GREATER
C OR EQUAL TO 5, WHICH SPECIFIES THE PARAMETERS OF
C THE INTERVAL AND OF ACCURACY AND WHICH SERVES FOR
C COMMUNICATION BETWEEN SUBROUTINES OUTP AND FCT
C (FURNISHED BY THE USER) AND SUBROUTINE RKGST.
C EXCEPT PRMT(5) THE COMPONENTS ARE NOT DESTROYED
C BY SUBROUTINE RKGST AND THEY ARE:
C PRMT(1) LOWER BOUND OF THE INTERVAL (INPUT),
C PRMT(2) UPPER BOUND OF THE INTERVAL (INPUT),
C PRMT(3) INITIAL INCREMENT OF THE INDEPENDENT VARIABLE
C (INPUT),
C PRMT(4) UPPER ERROR BOUND (INPUT). IF RELATIVE ERROR IS
C GREATER THAN PRMT(4), INCREMENT GETS HALVED.
C IF RELATIVE ERROR LESS THAN PRMT(4)*EXPAND,
C INCREMENT GETS DOUBLED.
C THE USER MAY CHANGE PRMT(4) BY MEANS OF HIS
C OUTPUT SUBROUTINE.
C PRMT(5) MAXIMUM STEP SIZE ORDER OF MAGNITUDE (INPUT).
C SUBROUTINE RKGST INITIALIZES
C PRMT(5)=0. IF THE USER WANTS TO TERMINATE
C SUBROUTINE RKGST AT ANY OUTPUT POINT, HE HAS TO
C CHANGE PRMT(5) TO NON-ZERO BY MEANS OF SUBROUTINE
C OUTP. FURTHER COMPONENTS OF VECTOR PRMT ARE
C FEASIBLE IF ITS DIMENSION IS DEFINED GREATER
C THAN 5. HOWEVER SUBROUTINE RKGST DOES NOT REQUIRE
C AND CHANGE THEM. NEVERTHELESS THEY MAY BE USEFUL
C FOR HANDING RESULT VALUES TO THE MAIN PROGRAM

C (CALLING RKGST) WHICH ARE OBTAINED BY SPECIAL
 C MANIPULATIONS WITH OUTPUT DATA IN SUBROUTINE OUTP.
 C Y INPUT VECTOR OF INITIAL VALUES. (DESTROYED)
 C LATER, Y IS THE RESULTING VECTOR OF DEPENDENT
 C VARIABLES COMPUTED AT INTERMEDIATE POINTS X.
 C DERY INPUT VECTOR OF ERROR WEIGHTS. (DESTROYED)
 C ERROR WEIGHTS ARE CENTERED AT ONE. IF ONE PARA-
 C METER NEEDS A TIGHTER ERROR CRITERIA, THE WEIGHT IS
 C GREATER THAN ONE. IF A PARAMETER NEED NOT BE DETER-
 C MINED SO PRECISELY, THE WEIGHT SHOULD BE LESS
 C THAN ONE. IN OTHER WORDS,
 C ERROR CRITERIA(I) = PRMT(4) / WEIGHT(I)
 C WHERE I IS THE SUBSCRIPT OF A DEPENDENT VARIABLE.
 C LATER, DERY IS THE VECTOR OF DERIVATIVES, WHICH
 C BELONG TO FUNCTION VALUES Y AT A POINT X.
 C NDIM AN INPUT VALUE, WHICH SPECIFIES THE NUMBER OF
 C EQUATIONS IN THE SYSTEM.
 C IHLF AN OUTPUT VALUE, WHICH SPECIFIES THE NUMBER OF
 C BISECTIONS OF THE INITIAL INCREMENT. IF IHLF BE-
 C COMES GREATER THAN 10, SUBROUTINE RKGST RETURNS THE
 C ERROR MESSAGE IHLF=11 INTO MAIN PROGRAM. ERROR
 C MESSAGE IHLF=12 OR IHLF=13 APPEARS IN CASE
 C PRMT(3)=0 OR IN CASE SIGN(PRMT(3)).NE.SIGN(PRMT(2)-
 C PRMT(1)) RESPECTIVELY.
 C FCT THE NAME OF AN EXTERNAL SUBROUTINE USED. THIS
 C SUBROUTINE COMPUTES THE RIGHT HAND SIDES DERY OF
 C THE SYSTEM TO GIVEN VALUES X AND Y. ITS PARAMETER
 C LIST MUST BE X,Y,DERY,PRMT. SUBROUTINE FCT SHOULD
 C NOT DESTROY X AND Y.
 C OUTP THE NAME OF AN EXTERNAL OUTPUT SUBROUTINE USED.
 C ITS PARAMETER LIST MUST BE X,Y,DERY,IHLF,NDIM,PRMT.
 C NONE OF THESE PARAMETERS (EXCEPT, IF NECESSARY,
 C PRMT(4),PRMT(5),...) SHOULD BE CHANGED BY
 C SUBROUTINE OUTP. IF PRMT(5) IS CHANGED TO NON-ZERO,
 C SUBROUTINE RKGST IS TERMINATED.
 C AUX AN AUXILIARY STORAGE ARRAY WITH 8 ROWS AND NDIM
 C COLUMNS.
 C
 C REMARKS
 C
 C THE PROCEDURE TERMINATES AND RETURNS TO CALLING PROGRAM, IF
 C (1) MORE THAN 10 BISECTIONS OF THE INITIAL INCREMENT ARE
 C NECESSARY TO GET SATISFACTORY ACCURACY (ERROR MESSAGE
 C IHLF=11),
 C (2) INITIAL INCREMENT IS EQUAL TO 0 OR HAS WRONG SIGN
 C (ERROR MESSAGES IHLF=12 OR IHLF=13),
 C (3) THE WHOLE INTEGRATION INTERVAL IS WORKED THROUGH,
 C (4) SUBROUTINE OUTP HAS CHANGED PRMT(5) TO NON-ZERO.
 C
 C SUBROUTINES AND FUNCTION SUBPROGRAMS REQUIRED
 C THE EXTERNAL SUBROUTINES FCT(X,Y,DERY,PRMT) AND

```

C OUTP(X,Y,DERY,IHLF,NDIM,PRMT) MUST BE FURNISHED BY THE USER.
C
C METHOD
C EVALUATION IS DONE BY MEANS OF FOURTH ORDER RUNGE-KUTTA
C FORMULAE IN THE MODIFICATION DUE TO GILL. ACCURACY IS
C TESTED COMPARING THE RESULTS OF THE PROCEDURE WITH SINGLE
C AND DOUBLE INCREMENT.
C SUBROUTINE RKGST AUTOMATICALLY ADJUSTS THE INCREMENT DURING
C THE WHOLE COMPUTATION BY HALVING OR DOUBLING. IF MORE THAN
C 10 BISECTIONS OF THE INCREMENT ARE NECESSARY TO GET
C SATISFACTORY ACCURACY, THE SUBROUTINE RETURNS WITH
C ERROR MESSAGE IHLF=11 INTO MAIN PROGRAM.
C TO GET FULL FLEXIBILITY IN OUTPUT, AN OUTPUT SUBROUTINE
C MUST BE FURNISHED BY THE USER.
C FOR REFERENCE, SEE
C RALSTON/WILF, MATHEMATICAL METHODS FOR DIGITAL COMPUTERS,
C WILEY, NEW YORK/LONDON, 1960, PP.110-120.
C
C SOME NOTES ON THE PROGRAM/RALSTON AND WILF
C
C AUX
C ---
C
C AUX(1,I) -- CURRENT VALUE OF Y
C AUX(2,I) -- CURRENT VALUE OF Y'
C AUX(3,I) -- LAST GOOD VALUES OF Q
C AUX(4,I) -- Y AFTER ONE RK STEP H
C AUX(5,I) -- Y AFTER ONE OR TWO RK STEPS OF H/2.
C AUX(6,I) -- CURRENT VALUES OF Q
C AUX(7,I) -- Y' AFTER ONE OR TWO RK STEPS OF H/2.
C AUX(8,I) -- 2/15 * WEIGHTS
C
C A(4),B(4),C(4)
C -----
C
C Y = Y + A *(K - B *Q )
C   i   i-1   i   i   i-1
C
C Q = Q + 3*(A *(K - B *Q ) - C *K
C   i   i-1   i   i   i-1   i
C
C FOR VALUES OF I BETWEEN 1 AND 4, AND FOR VALUES OF K AS FOLLOWS
C
C K = H * F(X ,Y )
C   1       0 0
C
C K = H * F(X +H/2,Y )
C   2       0   1
C
C K = H * F(X +H/2,Y )
C   3       0   2

```

```

C
C      K = H * F(X +H,Y )
C          4      0   3
C
C      RELATIVE ERROR
C      -----
C
C      AS PER RICHARDSON QUOTED IN RALSTON/WILF (P117),
C
C      ABS ERROR = WEIGHT/15*ABS(Y2 - Y1)
C
C      THEN, RELATIVE ERROR
C
C      REL ERROR = WEIGHT*2/15*ABS(Y2 - Y1)/SUM
C
C      where SUM = ABS(Y2 + Y1)
C
C      The solution tries to use SUM=abs(y2+y1) first. If this is zero,
C      then SUM=.25*ABS(Y1) is used since y1 and y2 must be oposite in
C      sign with equal magnitude. If this
C      quantity is zero as well, the values y2 and y1 both must be zero;
C      therefore, the difference is also zero which satisfies the
C      error criteria.
C
C      .....
C
C      SUBROUTINE RKGST(PRMT,Y,DERY,NDIM,IHLF,FCT,OUTP,AUX)

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

C
C      DIMENSION Y(1),DERY(1),AUX(8,NDIM),A(4),B(4),C(4),PRMT(1)
C
C      DATA ERRSET/1./
C
C
C      DO 10 I=1,NDIM
10    AUX(8,I)=.1333333333333333D0 * DERY(I)
X=PRMT(1)
XEND=PRMT(2)
H=PRMT(3)
stpmmin = abs(h/1024.D0)
stpmmax = abs(prmt(5))
PRMT(5)=0.D0
CALL FCT(X,Y,DERY,PRMT)
C
C*** ERROR TEST
C
IF(H*(XEND-X))380,370,20
C
C*** PREPARATIONS FOR RUNGE-KUTTA METHOD

```

```

C
20  A(1)=.5D0
    A(2)= 1.D0 - DSQRT( 0.5D0 )
    A(3)= 1.D0 + DSQRT( 0.5D0 )
    A(4)= 1.D0/6.D0
    B(1)=2.D0
    B(2)=1.D0
    B(3)=1.D0
    B(4)=2.D0
    C(1)=.5D0
    C(2)= A(2)
    C(3)= A(3)
    C(4)=.5D0
C
C*** PREPARATIONS OF FIRST RUNGE-KUTTA STEP
C
      DO 30 I=1,NDIM
      AUX(1,I)=Y(I)
      AUX(2,I)=DERY(I)
      AUX(3,I)=0.D0
30  AUX(6,I)=0.D0
      IREC=0
      H=H+H
      IHLF=-1
      ISTEP=0
      IEND=0
C
C*** START OF A RUNGE-KUTTA STEP
C*** STEP = 2 * SPECIFIED STEP
C
40  IF((X+H-XEND)*H)70,60,50
50  H=XEND-X
60  IEND=1
C
C*** RECORDING OF INITIAL VALUES OF THIS STEP
C
70  CALL FCT(X,Y,DERY,PRMT)
    CALL OUTP(X,Y,DERY,IREC,NDIM,PRMT)
    IF(PRMT(5))400,80,400
80  ITEST=0
90  ISTEP=ISTEP+1
C
C*** START OF INNERMOST RUNGE-KUTTA LOOP
C
      J=1
100  AJ=A(J)
      BJ=B(J)
      CJ=C(J)
      DO 110 I=1,NDIM
      R1=H*DERY(I)
      R2=AJ*(R1-BJ*AUX(6,I))

```

```

Y(I)=Y(I)+R2
R2=R2+R2+R2
110 AUX(6,I)=AUX(6,I)+R2-CJ*R1
IF(J-4)120,150,150
120 J=J+1
IF(J-3)130,140,130
130 X=X+H/2.
140 CALL FCT(X,Y,DERY,PRMT)
GOTO 100

C
C*** END OF INNERMOST RUNGE-KUTTA LOOP
C
C*** TEST OF ACCURACY
C
150 IF(ITEST)160,160,200
C
C*** IN CASE ITEST=0 THERE IS NO POSSIBILITY FOR TESTING OF ACCURACY
C*** IF(ITEST=0) RK STEP JUST PERFORMED WAS FOR TWICE THE SPECIFIED STEP
C
160 DO 170 I=1,NDIM
170 AUX(4,I)=Y(I)
ITEST=1
ISTEP=ISTEP+ISTEP-2
180 IHLF=IHLF+1
X=X-H
H=H/2.
DO 190 I=1,NDIM
Y(I)=AUX(1,I)
DERY(I)=AUX(2,I)
190 AUX(6,I)=AUX(3,I)
GOTO 90

C
C*** IN CASE ITEST=1 TESTING OF ACCURACY IS POSSIBLE ONLY IF EACH
C*** HALF OF THE INTERVAL IS DONE(I.E.,IFF ISTEP IS EVEN)
C
200 IMOD=ISTEP/2
IF(ISTEP-IMOD-IMOD)210,230,210
210 CALL FCT(X,Y,DERY,PRMT)
DO 220 I=1,NDIM
AUX(5,I)=Y(I)
220 AUX(7,I)=DERY(I)
GOTO 90

C
C*** ORIGINAL VERSION; absolute error
C
C      COMPUTATION OF TEST VALUE DELT
C 230 DELT=0.          !Good so far
C      DO 240 I=1,NDIM
C 240 DELT=DELT+AUX(8,I)*ABS(AUX(4,I)-Y(I))
C      IF(DELT-PRMT(4))280,280,250
C

```

```

C*** RELATIVE ERROR
C
230  DELT = 0.
      DO 240 I=1,NDIM
          ARG = ABS(AUX(4,I) + Y(I))
          IF(ARG .EQ. 0.) ARG = .25*ABS(AUX(4,I))
c--- if the next statement is true, aux(4,i)=y(i)=0.0; rer=0.
          IF(ARG .EQ. 0.) ARG = ERRSET
          RER = AUX(8,I)*ABS(AUX(4,I) + Y(I))/ARG
240  DELT = dMAX1(DELT,RER)
      IF(DELT-PRMT(4)) 280,280,250
C
C*** ERROR IS TOO GREAT
C
250  if(abs(h) .lt. stpmin) goto 360
      DO 270 I=1,NDIM
          AUX(4,I)=AUX(5,I)
270  WRITE(5,1200) DELT
C 1200 FORMAT(' ?RKGST? -- ERROR TOO GREAT',G13.5)
      ISTEP=ISTEP+ISTEP/4
      X=X-H
      IEND=0
      GOTO 180
C
C*** RESULT VALUES ARE GOOD
C
280  CALL FCT(X,Y,DERY,PRMT)
      DO 290 I=1,NDIM
          AUX(1,I)=Y(I)
          AUX(2,I)=DERY(I)
          AUX(3,I)=AUX(6,I)
          Y(I)=AUX(5,I)
290  DERY(I)=AUX(7,I)
      CALL FCT(X-H,Y,DERY,PRMT)
      CALL OUTP(X-H,Y,DERY,IHLF,NDIM,PRMT)
      IF(PRMT(5))400,300,400
300  DO 310 I=1,NDIM
          Y(I)=AUX(1,I)
310  DERY(I)=AUX(2,I)
          IREC=IHLF
          IF(IEND) 320,320,390
C
C*** INCREMENT GETS DOUBLED TO KEEP UP WITH HALF STEPPING
C
320  IHLF=IHLF-1
      ISTEP=ISTEP/2
      H=H+H
C
C*** ALLOW THE PROGRAM TO EXPAND BEYOND ORIGINAL STEP SIZE SPECIFICATION
C*** UP TO THE MAXIMUM
C

```

```

IF(abs(h).ge.stpmax) goto 40
C
C*** EXPAND H DUE TO LOW ERROR VALUE after Press et al. Since a 'new'
c      step size is being chosen, reset IHLF and ISTEP to starting values.
c      STPMIN will still stop simulations which bisect the original interval
c      more than 10 times.
C
      if(delt.ne. 0.00) then
          trial = .900 * abs(h) * (abs(prmt(4)/delt)) ** .25D0
      else
          trial = 10.00 * abs(h)
      endif
      if(trial .ge. abs(h)) then
          h = sign( min(trial, stpmax), h)
          IHLF=-1
          ISTEP=0
      endif
      GOTO 40
C
C*** RETURNS TO CALLING PROGRAM
C
      360  IHLF=11
            CALL FCT(X,Y,DERY,PRMT)
            GOTO 390
      370  IHLF=12
            GOTO 390
      380  IHLF=13
      390  CALL FCT(X,Y,DERY,PRMT)
            CALL OUTP(X,Y,DERY,IHLF,NDIM,PRMT)
      400  RETURN
            END
#####

```

PROGRAM SDEGADIS2

C
C*****
C*****
C*****
C
C Program description:
C
C SDEGADIS2 is a simplification of DEGADIS2 which performs the downwind
C dispersion portion of the calculation for a steady state source
C described by DEGADIS1.
C
C
C Program usage:
C
C Consult Volume III of the Final Report to U. S. Coast Guard
C contract DT-CG-23-80-C-20029 entitled "Development of an
C Atmospheric Dispersion Model for Heavier-than-Air Gas Mixtures".
C
C J. A. Havens
C T. O. Spicer
C
C University of Arkansas
C Department of Chemical Engineering
C Fayetteville, AR 72701
C
C April 1985
C
C
C This project was sponsored by the U. S. Coast Guard and the Gas
C Research Institute under contract DT-CG-23-80-C-20029.
C
C
C Disclaimer:
C
C This computer code material was prepared by the University of
C Arkansas as an account of work sponsored by the U. S. Coast Guard
C and the Gas Research Institute. Neither the University of Arkansas,
C nor any person acting on its behalf:
C
C a. Makes any warranty or representation, express or implied,
C with respect to the accuracy, completeness, or usefulness
C of the information contained in this computer code material,
C or that the use of any apparatus, method, numerical model,
C or process disclosed in this computer code material may not
C infringe privately owned rights; or
C
C b. Assumes any liability with respect to the use of, or for
C damages resulting from the use of, any information,
C apparatus, method, or process disclosed in this computer
C code material.

```

C
C
C*****
C*****
C*****
C

      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

      include 'sys$degadis:DEGADIS2.dec'
      include '($ssdef)'

C
      EXTERNAL PSS,PSSOUT,SSG,SSGOUT
C
      COMMON
      $/TITL/ TITLE
      $/GEN2/ DEN(5,igen)
      $/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CCLLOW
      $/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
      $ gas_ufl,gas_lfl,gas_zsp,gas_name
      $/ITI/ T1,TINP,TSRC,TOBS
      $/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
      $ humsrc
      $/ERROR/ SYOER,ERRP,SMXP,WTSZP,WTSYP,WTBEP,WTDH,ERRG,SMXG,
      $ WTRUH,WTDHG
      $/STP/ STPP,ODLP,ODLLP,STPG,ODLG,ODLLG
      $/com_ss/ ESS,SLEN,SWID,OUTCC,OUTSZ,OUTB,OUTL,SWCL,SWAL,SEN1,SRHL
      $/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
      $/com_fl/ cflag,clfl,cufl
      $/NEND/ POUNDNP,POUND
      $/ALP/ ALPHA,alpha1
      $/phicom/ iphifl,dellay
      $/sprd_con/ ce, delrhomin
      $/COM_SURF/ HTCUT
      ./oomsin/ oodist,avtime

C
C
C      DIMENSIONS/DECLARATIONS
C
      logical cflag

      real*4 tt1
      REAL*8 K,ML,L
      LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5

C
      character*24 tinp,tsrc,tobs
      character*80 title(4)
      character*4 pound
      character*3 gas_name

C
      character*4 TR2,ER2,Sr3,SSD,TR3

```

```

C
      character*40 opnrup1
      character opnrup(40)
      EQUIVALENCE (OPNRUP(1),opnrup1)
C
      dimension prmt(22),y(6),dery(6),aux(8,6)
C
C.....  

C
C       DATA
C
      DATA POUND//' //,POUNDN/-1.D-20/
C
      DATA TIME0/0./,NDIM/0/
C
      DATA TR2/'.TR2'/,ER2/'.ER2'/
      DATA Sr3/'.Sr3'/
      DATA SSD/'.SSD'/,TR3/'.TR3'/
C
C
C.....  

C
C       MAIN
C
      T1 = SECNDS(0.)
      istat = lib$date_time(TOBS)
      if(istat .ne. ss$_normal) stop'lib$date_time failure'
C
C*** GET THE FILE NAME FOR FILE CONTROL
C
      read(5,1135) nchar,opnrup
 1135  format(q,40a1)
C
      opnrup1 = opnrup1(1:nchar) // ER2(1:4)
      CALL ESTRT2ss(OPNRUP1)
C
C*** GET THE COMMON VARIABLES CARRIED FROM DEGADIS1
C
      opnrup1 = opnrup1(1:nchar) // tr2(1:4)
      CALL STRT2(OPNRUP1,M_masrte,CCP)
C
      opnrup1 = opnrup1(1:nchar) // sr3(1:4)
      OPEN(UNIT=8,TYPE='NEW',NAME=OPNRUP1,
      $ CARRIAGECONTROL='FORTRAN')
C
      cflag = isofl.eq. 1.or. ihtfl.eq. 0
C
      WRITE(8,1119)
      if(cflag) then
          WRITE(8,1116) gas_zsp,(100.*gas_lfl),(100.*gas_ufl)
          WRITE(8,1118)

```

```

    else
        WRITE(8,1115) gas_zsp,(100.*gas_lfl),(100.*gas_ufl)
        WRITE(8,1117)
        endif
    WRITE(8,1119)

c
c
c
1115  FORMAT(1HO,1X,'Distance',2x,3x,'Mole',3x,
1      'Concentration',1x,'Density',2x,3x,'Gamma',4x,
1      'Temperature',3x,'Half',4x,4x,'Sz',5x,4x,'Sy',5x,
1      'Width at z=',0pf6.2,' m to:',/,1x,11x,1x'Fraction',2x,
1      11x,11x,11x,3x,'Width',3x,11x,9x,
1      2(1pg9.3,'mole%',1x))
1116  FORMAT(1HO,1X,'Distance',2x,3x,'Mole',3x,
1      'Concentration',1x,'Density',2x,
1      'Temperature',3x,'Half',4x,4x,'Sz',5x,4x,'Sy',5x,
1      'Width at z=',0pf6.2,' m to:',/,1x,11x,1x'Fraction',2x,
1      11x,11x,11x,3x,'Width',3x,11x,9x,
1      2(1pg9.3,'mole%',1x))
1117  FORMAT(1H ,4X,'(m)',4x,11x,
1      2(1X,'(kg/m**3)',1x),12x,4x,'(K)',
1      5(8X,'(m)'))
1118  FORMAT(1H ,4X,'(m)',4x,11x,
1      2(1X,'(kg/m**3)',1x),4x,'(K',
1      5(8X,'(m)'))
1119  FORMAT(1H )

c
c
c
c.....-----
c
c      STEADY STATE CALCULATIONS
c
c
        opnrup1 = opnrup1(1:nchar) // ssd(1:4)
c      OPEN(UNIT=9,TYPE='NEW',NAME=OPNRUP1)
c
c
        AGAIN = .FALSE.
c
c
        L = OUTL
        B = OUTB
        S20 = OUTSZ
        Erate = ESS
        QSTRO = Erate/2.00/L/B
        Cc = OUTCc
        wc = swcl
        wa = swal
        enth = senl

```

```

rho = srhl

c
if(cc.gt. ccp) then
    write(lunlog,1126) cc,ccp
1126  format(//,' ',10('****'),/, ' cc: ',1pg13.5,' is greater',
1           ' than ccp: ',1pg13.5.,/, ' ',10('****'),/)
    cc =ccp
endif

c
ratio1= u0*z0/alpha1/ z0**alpha1 * cc /b/qstr0/l
ratio = ratio1* sz0**alpha1 * (B + sqrt(pi)/2.00*sy0er)
if(ratio.lt. 1.00) then
    sy0er = (1.00/(RATIO1*sz0**alpha1) - b)*2.00/sqrt(pi)
else
    sz0 = (1.00/((B+ sqrt(pi)/2.00*sy0er)*ratio1))** (1.00/alpha1)
endif

humsrc = (1.00-wc-wa*(1.00+humid))/wc
call setden(wc,wa,enth)

if(cflag) then
    call adiabat(2,twc,twa,gas_lfl,ya,clfl,r,w,t,tt)
    call adiabat(2,twc,twa,gas_ufl,ya,cufl,r,w,t,tt)
endif

c
cclay = cc/dellay
call adiabat(0,wclay,walay,yclay,yalay,cclay,rholay,w,enthalay,t)
c
c
C*** let everyone know
c
        WRITE(lunlog,1170) L,B
        WRITE(lunlog,1180) QSTR0,SZ0
        write(lunlog,1185) wclay,walay,rholay,cclay,t
        write(lunlog,1186) wc,wa,rho,cc

c
1170  FORMAT(' LENGTH: ',1pG13.5,' BEFF: ',1pG13.5)
1180  FORMAT(' TAKEUP FLUX: ',1pG13.5,' SZ0: ',1pG13.5)
1185  format(' wclay: ',1pg12.5,' walay: ',1pg12.5,
1           ' rholay: ',1pg12.5,' Cclay: ',1pg12.5./,
1           ' temlay: ',1pg13.5)
1186  format(' wc: ',1pg12.5,' wa: ',1pg13.5./,
1           ' rho: ',1pg12.5,' Cc: ',1pg12.5)

c
C*** PREPARE FOR STEADY STATE INTEGRATION.
c
PRMT(1) = L/2.00
PRMT(2) = 6.023D13
PRMT(3) = STPP

```

D=127

```
PRMT(4) = ERRP
PRMT(5) = SMXP
PRMT(6) = Erate
PRMT(7) = Cc      ! OUTPUT
PRMT(8) = B       ! OUTPUT
C
C*** PRMT(9) & PRMT(10) ARE CONSTANTS FOR D(SY) & D(SZ)
C
PRMT(9) = Ce*sqrt(G*Z0/ALPHA1*GAMMAF) *GAMMAF/U0
PRMT(10)= Z0**ALPHA*K*USTAR*ALPHA1 * ALPHA1/U0
C   PRMT(11)= NREC
C   PRMT(12)=
C   PRMT(13)=
prmt(18)= u0*z0/alpha1
prmt(19)= rhoa*k*ustar*alpha1
prmt(20)= rholay
prmt(21)= sz0
prmt(22)= sz0
C
Y(1) = rholay*prmt(18)*(SZ0/z0)**alpha1 ! rholay*u0*heff
Y(2) = SY0ER*sy0er
Y(3) = B + sqrt(pi/2.0)*sy0er
Y(4) = 0.                      ! added heat
Y(5) = 0.                      ! mass above UFL
Y(6) = 0.                      ! mass above LFL
C
DERY(1) = WTSZP
DERY(2) = WTSYP
DERY(3) = WTBEPE
dery(4) = wtch
dery(5) = 1.00
dery(6) = 1.00
C
NDIM = 4
ndim=6      ! to integrate the mass above LFL and UFL
C
WRITE(lunlog,1130)
1130 FORMAT(' Entering Integration Step -- B > 0. ')
C
C*** PERFORM INTEGRATION
C
CALL RKGST(PRMT,Y,DERY,NDIM,IHLF,PSS,PSSOUT,AUX)
C
IF(IHLF .GE. 10) CALL trap(9,IHLF)
C
NREC = INT(PRMT(11))
WRITE(lunlog,1100)NREC
1100 FORMAT(3X,'NUMBER OF RECORDS IN PSS = ',I10)
C
IF(AGAIN) THEN
    Y(3) = Y(5)          ! mass above UFL
```

D=128

```
Y(4) = Y(6)           ! mass above LFL
GO TO 120
ENDIF
C
C*** GAUSIAN COMPLETION OF THE INTEGRATION
C
C*** PSSOUT FORCES THE ABOVE INTEGRATION TO FINISH WHEN B<0 FOR THE
C*** FIRST TIME. THE STEP BEFORE THIS OCCURS IS RECORDED ON UNIT 7.
C*** THE STEP WHEN B GOES NEGATIVE IS CURRENTLY IN Y.
C
C*** THE CALCULATION METHOD CHANGES THE CURRENT VALUE OF SY TO A VALUE
C*** CALCULATED AS IF BEFF=SY RETAINING THE LAST VALUE OF Cc IN THE
C*** MATERIAL BALANCE.
C
heat = Y(4)
rholay = prmt(20)
Cc = PRMT(7)
rhouh = Y(1)
SZ = ( rhouh/rholay/prmt(18) )**(1.00/alpha1) * z0
SYT = Erate*ALPHA1 *(Z0/SZ)**ALPHA/U0/SZ/Cc/SQRTPI
C
XT = PRMT(12)
XV = (SYT/RT2/DELTA)**(1.00/BETA) - XT
C
C*** SET UP INTEGRATION FOR THE GAUSSIAN DISPERSION PHASE.
C
do i=1,22
prmt(i) = 0.00
enddo
C
PRMT(1) = XT
PRMT(2) = 6.023D23
PRMT(3) = STPG
PRMT(4) = ERRG
PRMT(5) = SMXG
PRMT(6) = Erate
PRMT(7) = Cc          !-- OUTPUT
PRMT(8) = XV
C
PRMT(9) = "BLANK"
C
PRMT(10)= "BLANK"
C
PRMT(11)= "BLANK"
C
PRMT(12)= DIST AT COMPLETION -- OUTPUT
C
prmt(18) = u0*z0/alpha1
prmt(19) = rhoa*k*ustar*alpha1
prmt(20) = rholay
prmt(21) = sz
prmt(22) = sz
C
Y(1) = rhouh
Y(2) = heat
```

```

      Y(3) = Y(5)          ! mass above UFL
      Y(4) = Y(6)          ! mass above LFL
C
      DERY(1) = wtruh
      dery(2) = wtdhg
      dery(3) = 1.00
      dery(4) = 1.00
C
c NDIM = 2
      ndim=4           ! to integrate the mass above LFL and UFL
C
      WRITE(lunlog,1140)
1140  FORMAT(' Entering Gaussian Stage of Integration ')
C
C*** PERFORM INTEGRATION
C
      CALL RKGST(PRMT,Y,DERY,NDIM,IHLF,SSG,SSGOUT,AUX)
C
      IF(IHLF .GE. 10) CALL trap(10,IHLF)
C
      NREC = INT(PRMT(11))
C
      120  CONTINUE
C
c*** summarize the information about the mass above the LFL,UFL
C
      write(8,8000) 100.*gas_ufl,100.*gas_lfl,Y(4)-Y(3),Y(4)
8000  format(//,' For the UFL of ',1pg13.5,' mole percent, and',
     $' the LFL of ',1pg13.5,' mole percent:','
     $//,' The mass of contaminant between the UFL and LFL is:'
     $,1pg13.5,' kg.',/, ' The mass of contaminant above the LFL is: ',
     $1pg13.5,' kg.')
C
C
C.....-----
C
      CLOSE(UNIT=9)
      CLOSE(UNIT=8)
C
      opnrup1 = opnrup1(1:nchar) // tr3(1:4)
      CALL TRANS(OPNRUP1)
C
      tt1 = t1
      T1 = SECNDS(tt1)/60.
      WRITE(lunlog,4000) TOBS
      WRITE(lunlog,4010) T1
4000  FORMAT(//,'SDEGADIS2 -->',//,3X,'BEGAN AT ',A40)
4010  FORMAT(3X,'*** ELAPSED TIME *** ',1pG13.5,' min')
C
      STOP
      END
#####

```

```

function series(arg)
c
c*** This function estimates the infinite series used in the evaluation
c*** of the mass above a certain concentration level.
c
c***** ****
c
c*** WARNING: This routine will overflow for arguments greater than 13.8
c
c***** ****
c
      implicit real*8 (a-h,o-z), integer*4 (i-n)
c
      parameter (kmax= 100, error=1.d-4)
c
      common/alp/ alpha,alpha1
c
c*** initialize some variables
c
      pp = 1.D0/alpha1
      pprod = pp      ! for p*(p+1)*...*(p+k)
c
c*** calculate the first term
c
      coef = 2.D0/3.D0
      series = coef / pp * arg**1.5D0
      old = series
c
c
      do 200 kk=1,kmax
      pow = DBLE(kk) + 1.5D0
      plus = DBLE(kk+1)

      pprod = (pp + DBLE(kk)) * pprod
      coef = 2.D0 * plus**2 / DBLE(2*kk + 3)/ plus * coef

      series = series + coef / pprod * arg**pow

      denom = dmax1( (series+old), error)
      if( abs(series-old)/denom .le. error) return

      old = series
      200 continue
c
      write(6,900) arg
900  format(' SERIES? Number of iterations too small for argument: ',
$ 1pg13.5)
      call exit
      end
#####

```

```

C.....  

C  

C      TIME SORT SUPERVISOR  

C  

C      SUBROUTINE SORTS(TABLE)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      include 'sys$degadis:DEGADIS3.dec/list'  

C  

C      COMMON  

$ /SORT/  TCc(maxnob,maxnt),TCcSTR(maxnob,maxnt),  

$   Tyc(maxnob,maxnt),Trho(maxnob,maxnt),  

$   Tgamma(maxnob,maxnt),Ttemp(maxnob,maxnt),  

$   TSY(maxnob,maxnt),TSZ(maxnob,maxnt),TB(maxnob,maxnt),  

$   TDIST0(maxnob,maxnt),TDIST(maxnob,maxnt),KSUB(maxnt)  

$ /SSCON/ NRECC(maxnob,2),TO(maxnob),XV(maxnob)  

$ /SORTIN/ TIM(maxnt),NTIM,ISTRTR  

$ /PARM/ UO,ZO,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$ /CNOBS/ NOBS  

C  

C      DIMENSION TABLE(1)  

C  

C      REAL*8 ML,K  

C  

C      CALL GETTIM  

C  

C  

C*** TABLE(I) VALUES  

C***    I          PARAMETER  

C***    --  

C***    1  11        DIST      1 TO 10 CURRENT READ  

C***    2  12        Yc  

C***    3  13        Cc          11 TO 20 PREVIOUS READ  

C***    4  14        rho  

C***    5  15        gamma  

C***    6  16        temp  

C***    7  17        SZ  

C***    8  18        SY  

C***    9  19        B  

C***   10 20        TS  

C  

C***   21          DIST0  

C***   22          INTERPOLATION FRACTION  

C  

DO 100 I = 1,NOBS  

C  

IT = 0  

C  

DO 105 J=1,20

```

```

105  TABLE(J) = 0.
C
    II = NREC(I,1)

    if( ii .eq. 0) goto 130
C
C*** read first record
C
    read(9,*), (table(k1),k1=1,9)
    table(10) = ts( t0(I), table(1) )
    table(21) = table(1)
C
C*** loop through and read each record even if not pertinent
C
    DO 110 J = 2,II
C
        DO K1 = 1,10
        KK = K1 + 10
        TABLE(KK) = TABLE(K1)
        enddo
C
        READ(9,*), (TABLE(K1),K1=1,9)
        TABLE(10) = TS( T0(I), TABLE(1) )
C
        itl = int( (table(10)-tim(1)) / (tim(2)-tim(1)) + 0.9999999 )
        itl = min( ntim, itl)
        itf = int( (table(20)-tim(1)) / (tim(2)-tim(1)) + 0.9999999 ) + 1
        itf = max( 1, itf)

        do it = itf, itl, 1           ! do all points in range
C
C*** RECORD AN INTERPOLATED TIME SORTED POINT.
C
        KSUB(IT) = KSUB(IT) + 1
C
        TABLE(22) = (TIM(IT) - TABLE(20))/(TABLE(10) - TABLE(20))
C
        TDIST0(I,IT) = TABLE(21)
        TDIST(I,IT) = TABLE(11) + (TABLE(1) - TABLE(11)) * TABLE(22)
        Tyc(I,IT) = TABLE(12) + (TABLE(2) - TABLE(12)) * TABLE(22)
        TCC(I,IT) = TABLE(13) + (TABLE(3) - TABLE(13)) * TABLE(22)
        Trho(I,IT) = TABLE(14) + (TABLE(4) - TABLE(14)) * TABLE(22)
        Tgamma(I,IT) = TABLE(15) + (TABLE(5) - TABLE(15)) * TABLE(22)
        Ttemp(I,IT) = TABLE(16) + (TABLE(6) - TABLE(16)) * TABLE(22)
        TSZ(I,IT) = TABLE(17) + (TABLE(7) - TABLE(17)) * TABLE(22)
        TSY(I,IT) = TABLE(18) + (TABLE(8) - TABLE(18)) * TABLE(22)
        TB(I,IT) = TABLE(19) + (TABLE(9) - TABLE(19)) * TABLE(22)
C
        enddo
110  CONTINUE
C

```

```

C
130  II = NREC(I,2)
      IF(II .EQ. 0) GO TO 100

      DO 200 J=1,II
C
      DO K1 = 1,10
      KK = K1 + 10
      TABLE(KK) = TABLE(K1)
      enddo

C
      READ(9,*) (TABLE(K1),K1=1,7)
C
      TABLE(8) = RT2*DELTA*(TABLE(1) + XV(I))**BETA
      TABLE(9) = 0.
      TABLE(10) = TS(TO(I),TABLE(1))
C
      itl = int( (table(10)-tim(1)) / (tim(2)-tim(1)) + 0.9999999 )
      itl = min( ntim, itl)
      itf = int( (table(20)-tim(1)) / (tim(2)-tim(1)) + 0.9999999 ) + 1
      itf = max( 1, itf)

      do it = itf, itl, 1           ! do all points in range
C
C*** RECORD A TIME SORTED VALUE
C
      KSUB(IT) = KSUB(IT) + 1
      TABLE(22) = (TIM(IT) - TABLE(20))/(TABLE(10) - TABLE(20))
C
      TDIST0(I,IT) = TABLE(21)
      TDIST(I,IT) = TABLE(11) + (TABLE(1) - TABLE(11)) * TABLE(22)
      Tyc(I,IT) = TABLE(12) + (TABLE(2) - TABLE(12)) * TABLE(22)
      TCc(I,IT) = TABLE(13) + (TABLE(3) - TABLE(13)) * TABLE(22)
      Trho(I,IT) = TABLE(14) + (TABLE(4) - TABLE(14)) * TABLE(22)
      Tgamma(I,IT) = TABLE(15) + (TABLE(5) - TABLE(15)) * TABLE(22)
      Ttemp(I,IT) = TABLE(16) + (TABLE(6) - TABLE(16)) * TABLE(22)
      TSZ(I,IT) = TABLE(17) + (TABLE(7) - TABLE(17)) * TABLE(22)
      TSY(I,IT) = TABLE(18) + (TABLE(8) - TABLE(18)) * TABLE(22)
      TB(I,IT) = TABLE(19) + (TABLE(9) - TABLE(19)) * TABLE(22)
C
      enddo
200  CONTINUE
C
100 CONTINUE
C
      CALL SORTS1(TABLE)
C
      RETURN
      END
#####

```

```

C.....  

C  

C  

SUBROUTINE SORTS1(TABLE)  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

include 'sys$degadis:DEGADIS3.dec/list'  

C  

COMMON  

$/$ORT/ TCc(maxnob,maxnt),TCcSTR(maxnob,maxnt),  

$ Tyc(maxnob,maxnt),Trho(maxnob,maxnt),  

$ Tgamma(maxnob,maxnt),Ttemp(maxnob,maxnt),  

$ TSY(maxnob,maxnt),TSZ(maxnob,maxnt),TB(maxnob,maxnt),  

$ TDIST0(maxnob,maxnt),TDIST(maxnob,maxnt),KSUB(maxnt)  

$/$SSCON/ NREC(maxnob,2),T0(maxnob),XV(maxnob)  

$/$ORTIN/ TIM(maxnt),NTIM,ISTRRT  

$/$PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$/$com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,  

$ gas_ufl,gas_lfl,gas_zsp,gas_name  

$/$comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/$PARMSC/RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$/$com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag  

$/$ALP/ ALPHA,alpha1  

$/$CNOBS/NOBS  

C  

DIMENSION TABLE(1)  

C  

REAL*8 ML,K  

logical cflag  

C  

C*** DETERMINE IF ANY TIME VECTORS HAVE NO ENTRIES  

C  

DO 192 I=1,NTIM  

192 IF(KSUB(I).GT. 2) GO TO 194      ! 1or2 points is of little value  

call trap(23)  

194 ISTRRT = I  

DO 196 I=ISTRRT,NTIM  

196 IF(KSUB(I).LE. 2) GO TO 198  

GO TO 199  

198 NTIM = I - 1  

199 CONTINUE  

C  

C*** REVERSE TIME SORTED VECTORS  

C  

C*** At this point, the results from Observer #1 is located in the  

C*** first row of each vector. But the information contributed by  

C*** Observer #1 is the information which is the farthest (TDIST)  

C*** from the source. If the output is printed by time, the

```

```

c*** distances will decrease instead of increase. So, we need to
c*** reverse each of the columns so that the downwind distances
c*** increase as you move down the column. At the same time,
c*** ensure that all of the nonzero entries in a column are at
c*** the top of the column.
c
      DO 200 K1 = ISTRT,NTIM
c
      II = KSUB(K1)
      DO 170 J = 1,NOBS
c
c      IF(TDIST(J,K1).NE.0. .OR. TB(J,K1).NE.0.) GO TO 180
c      IF( TSZ(J,K1).NE.0. ) GO TO 180
170  CONTINUE
180  II = II + J + 1
c
      DO 190 J = 1,II
c
      TABLE(II +      + 1 - J) =    TCc(J,K1)
      TABLE(II + NOBS + 1 - J) =    Tyc(J,K1)
      TABLE(II + 2*NOBS + 1 - J) =   Trho(J,K1)
      TABLE(II + 3*NOBS + 1 - J) =   Tgamma(J,K1)
      TABLE(II + 4*NOBS + 1 - J) =   Ttemp(J,K1)
      TABLE(II + 5*NOBS + 1 - J) =   TSY(J,K1)
      TABLE(II + 6*NOBS + 1 - J) =   TSZ(J,K1)
      TABLE(II + 7*NOBS + 1 - J) =   TB(J,K1)
      TABLE(II + 8*NOBS + 1 - J) =   TDIST0(J,K1)
      TABLE(II + 9*NOBS + 1 - J) =   TDIST(J,K1)
c
190  CONTINUE
c
      DO 210 J = 1,II
c
      TCc(J,K1) = TABLE(J)
      Tyc(J,K1) = TABLE(J + NOBS)
      Trho(J,K1) = TABLE(J + 2*NOBS)
      Tgamma(J,K1) = TABLE(J + 3*NOBS)
      Ttemp(J,K1) = TABLE(J + 4*NOBS)
      TSY(J,K1) = TABLE(J + 5*NOBS)
      TSZ(J,K1) = TABLE(J + 6*NOBS)
      TB(J,K1) = TABLE(J + 7*NOBS)
      TDIST0(J,K1) = TABLE(J + 8*NOBS)
      TDIST(J,K1) = TABLE(J + 9*NOBS)
c
210  CONTINUE
200  CONTINUE
c
c
      if(sigx_flag.eq. 0.) then           ! no correction
c
      , write(lunlog,*) ' No X-direction dispersion correction'

```

```

DO 220 K1 = ISTRT,NTIM
II = KSUB(K1)
DO 220 I = 1,II
TCcSTR(I,K1) = TCc(i,k1)
220    CONTINUE
goto 400
endif

C
C*** GENERATE TCcSTR -- CENTER LINE CONCENTRATION CORRECTED FOR
C*** DOWNDRAFT DISPERSION.

C
DO 230 K1 = ISTRT,NTIM
C
II = KSUB(K1)
DO 240 I = 1,II
C
c calculation for XP = TDIST(I,K1)
c
TCcSTR(I,K1) = 0.
C
DO 260 J = 1,II
C
TABLE(J) = 0.
DIST = TDIST(J,K1) - Tdist0(J,K1)
deltax = ABS(tdist(i,k1) - tdist(j,k1))
C
if(dist.lt. sigx_min_dist) then
  if(i.eq. j) then      ! i.e. deltax = 0.
    table(j) = (tdist(j+1,k1)- tdist(j-1,k1))/2.
    if(j.eq.1)table(j)= (tdist(2,k1)- tdist(1,k1))/2.
    if(j.eq. ii)table(j)=tdist(ii,k1)- tdist(ii-1,k1)
    table(j) = TCc(j,k1) / table(j) * RT2*SQRTPI
  endif
  goto 260
endif
C
SX = sigx_coeff* DIST**sigx_pow
C
ARG = (deltax/SX)**2/2.
C
IF(ARG .EQ. 0.) TABLE(J) = TCc(J,K1)/SX
IF(ARG .NE. 0. .AND. ARG .LE. 80.)
$           TABLE(J) = TCc(J,K1)/SX/EXP(ARG)
C
260    CONTINUE
C
III = KSUB(K1)
TCcSTR(I,K1) = TABLE(1)* (TDIST(2,K1)- TDIST(1,K1))/2.
TCcSTR(I,K1) = TABLE(iii)* (TDIST(iii,K1)- TDIST(iii-1,K1))
1
+   TCcSTR(I,K1)
iii = ksub(k1) - 1

```

```

C
DO 280 J = 2,III
C
TCcSTR(I,K1) = TABLE(J)* (TDIST(J+1,K1)- TDIST(J-1,K1))/2.
1
+ TCcSTR(I,K1)
C
280 CONTINUE
C
TCcSTR(I,K1) = TCcSTR(I,K1)/RT2/SQRTPI
C
c*** correct yc, rho, and temp values
C
      cc = Tccstr(i,k1)
      if(isofl.eq. 1 .or. ihtfl.eq. 0) then
          call adiabat(0,wc,wa,yc,ya,cc,rho,wm,enth,temp)
      else
          enth = Tgamma(i,K1)
          call adiabat(-1,wc,wa,yc,ya,cc,rho,wm,enth,temp)
      endif
      TyC(i,K1) = yc
      Trho(i,K1)= rho
C
240 CONTINUE
C
230 CONTINUE
C
C
C*** Estimate the mass between the UFL and LFL
C
400 continue
C
      cflag = isofl.eq.1 .or. ihtfl.eq.0
      if(cflag) then
          call adiabat(2,aa,dd,gas_ufl,ee,chi ,gg,hh,pp,oo)
          call adiabat(2,aa,dd,gas_lfl,ee,cflow,gg,hh,pp,oo)
      endif
C
      DO 430 K1 = ISTRT,NTIM
      kk  = k1+2*maxnob
      kk1 = k1+3*maxnob
      table(kk) = 0.
      table(kk1)= 0.
C
      II = KSUB(K1)
      DO 460 J = 1,II ! evaluate the function at each point in space
C
c initialize some values
C
      TABLE(J) = 0.
      jj = j+maxnob
      TABLE(JJ) = 0.

```

```

cc = TCcstr(j,k1)
bb = tb(j,k1)
sy = tsy(j,k1)
sz = tsz(j,k1)
gamma = tgamma(j,k1)

if(.not.cfleg) then
    call adiabat(-2,aa,dd,gas_ufl,ee,chi ,gg,hh,gamma,oo)
    call adiabat(-2,aa,dd,gas_lfl,ee,clow,gg,hh,gamma,oo)
endif

c
c*** Calculate the derivative for the total mass above the UFL and LFL
c
        gamhi = 2.00* Cc * Bb * Sz / alpha1
        gammax = 2.00 *Cc *Sz *GAMMAF /alpha1 *(Bb +sqrt(pi)/2.00 *Sy)

        if(cc.gt.clow) then
            wlow = Dlog(cc/clow)
            gamlow = gaminc(1.00/alpha1, wlow ) * gamhi
            table(j)= gamlow + 2.00*clow*Sy*Sz/alpha1*series(wlow)
            table(j)= DMIN1( table(j), gammax )
        endif

        if(cc.gt.chi) then
            whi = Dlog(cc/chi )
            gamhi = gaminc(1.00/alpha1, whi ) * gamhi
            table(jj) = gamhi+ 2.00*chi *Sy*Sz/alpha1*series(whi)
            table(jj)= DMIN1( table(jj), gammax )
        endif

c
460    continue
c
c*** now, finish the integration
c
        DO 450 J = 2,II      ! integrate in space at one value of time
        xx = (tdist(j,k1) - tdist(j-1,k1))
        arglow = (table(j) + table(j-1))/2.
        arghi = (table(j+maxnob) + table(j-1+maxnob))/2.
        table(kk) = arghi *xx + table(kk)
        table(kk1) = arglow*xx + table(kk1)
450    continue
c
430    continue
c
        RETURN
        END

#####

```

C SOURCE EQUATIONS -- Gas Blanket present

SUBROUTINE SRC1(time,Y,D,PRMT)

Implicit Real*8 (A-H, O-Z), Integer*4 (I-N)

include 'sys\$degadis:DEGADIS1.dec'

```
parameter(      delt= 0.1D0,
1             delto2= delt/2.D0,
2             zero= 1.D-10,
3             rcrit= 0.002D0)
```

COMMON

```
$/GEN1/ PTIME(igen), ET(igen), R1T(igen), PWC(igen), PTEMP(igen),
$      PFRACV(igen), PENTH(igen), PRHO(igen)
$/ERROR/STPIN,ERBND,STPMX,WTRG,WTtm,WTya,WTyc,WTeb,WTmb,WTUH,XLI,
$ XRI,EPS,ZLOW,STPINZ,ERBNDZ,STPMXZ,SRCOER,srccss,srccut,
$ htcut,ERNOBL,NOBLpt,crfger,epsilon
$/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW
$/PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND
$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc
$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
$/vocom/ vua,vub,vue,vud,vudelta,vuflag
$/com_enthal/ h_masrte,h_airrte,h_watrte
$/ALP/ ALPHA,alpha1
$/phicom/ iphifl,dellay
$/sprd_con/ ce, delrhomin
```

```
LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
logical vuflag
```

```
REAL*8 ML,K
REAL*8 L,masrte,mole
INTEGER R,mass,massc,massa,ebal,mbal
DIMENSION Y(7),D(7),PRMT(25)
DATA R/1.,mass/2.,massc/3.,massa/4./,ebal/5./,mbal/6./
```

```
if( prmt(20).lt. 0.D0) vuflag = .false.
```

```
if(Y(mass) .le. 0.D0) then
    wc = dmax1(prmt(15),1.d-10)
    if(wc.gt. 1.) wc=1.d-10
    wa = 1.D0 - wc
    enthalpy = wc*h_masrte           ! air contributes nothing
else
    wc = Y(massc)/Y(mass)
    wa = Y(massa)/Y(mass)
```

```

        enthalpy = Y(ebal)/Y(mass)
    endif

    humsrc = (1.00 - wc - wa*(1.00+humid))/wc
    call tprop(1,wc,wa,enthalpy,yc,ya,mole,temp,rho,cp)

    RADP = AFGEN2(PTIME,R1T,TIME,'R1TSRC')
    hei = dmax1( Y(mass)/pi/Y(r)/Y(r)/rho , 0.000 )
    delrho = rho-rhoa
    if(delrho .lt. 0.00) delrho = 0.00
    gprime = g*delrho/rhoa *hei

C*** CALCULATE D(R),airrte,vel

    D(R) = 0.00
    vel = 0.00
    airrte = 0.00
    Ri = 0.00
    D(mbal) = 0.00

    IF(Gprime.GT. 0.00) then
        slump = Ce*sqrt(Gprime)

        if(vuflag) then           ! momentum balance
            iii = 0              ! initialize loop counter
            vel = prmt(14)         ! old velocity value
            velmin = 0.00
            velmax= dmax1( slump, 0.100, vel)

100       hh = vel*vel/Ce/Ce/g/ (delrho/rhoa)
            rh = Y(r)-vua*vub*hh
            value = Y(r)**2/rh**2

            if(prmt(25).ge. prmt(24)) then      ! hh .ge. ht

                ht = 2.00*(value*hei - vua*hh*(value-1.00)) + hh
                velc = Y(mbal)/(2.00/3.00*pi*rho*(2.00/3.00*ht + hh)*rh**3/Y(r))
1               + 2.00/3.00*pi*vua*rho*hh* (Y(r)**2 - rh*rh*rh/Y(r))
2               + vue*pi*Y(r)*hei**2*rhoa
                D(mbal) = pi*g*delrho*Y(r)*ht**2
1               - vua*vud*pi*rhoa*Y(r)*hh*vel**2
                else

                    ht = value*hei - vua*hh*(value-1.00)
                    velc = Y(mbal)/(2.00/3.00*pi*rho*ht*rh**3/Y(r))
1                   + 2.00/3.00*pi*vua*rho*hh* (Y(r)**2 - rh*rh*rh/Y(r))
2                   + vue*pi*Y(r)*hei**2*rhoa
                    D(mbal) = pi*g*delrho*(rh*ht**2 + vua*vub*hh*hh*hh)
1                   - vua*vud*pi*rhoa*Y(r)*hh*vel**2
                endif

```

```

dif = abs(vel-velc)           ! convergence check
sum = abs(vel) + abs(velc) + zero

if(dif/sum .le. rcrit) then
    vel = (vel+velc)/2.00
    prmt(13) = vel

    if(vel .gt. 0.00) then
        Ri = gprime / vel**2
        airrte= 2.*pi* epsilon/Ri *rhoa*Y(r)*hei* vel
        D(r) = vel
        prmt(20) = slump
    endif
else

    dif = vel-velc

    if(velc.lt.velmin) velmin= dmax1(velc, 0.00)
    if(velc.gt.velmax) velmax=velc

    if(dif .gt. 0.00) then
        velmax = vel
        vel = 0.5D0*(velmax-velmin) + velmin
        vel = 0.382D0*(velmax-velmin) + velmin
    else
        velmin = velc
        vel = (1.00-0.5D0)*(velmax-velmin) + velmin
    vel = (1.00-0.382D0)*(velmax-velmin) + velmin
    endif

    iii = iii+1
    if(iii .gt. 40) then
        vel = min(velmin,velmax)
        if(vel .gt. slump) then
            vel = slump
            prmt(13) = vel
            if(vel .gt. 0.00) then
                Ri = gprime / vel**2
                airrte= 2.*pi* epsilon/Ri
                *rhoa*Y(r)*hei* vel
                D(r) = vel
                prmt(20) = slump
            endif
            goto 200
        endif
        write(6,*) 'Time, vel, slump: ',time,vel,slump
        write(6,*) 'prmt(14), velc: ',prmt(14),velc
        stop'SRC1 velocity loop'
    endif
    goto 100
endif

```

```

else

    vel= slump      ! gravity slumping
    hh = hei .
    ht = hei
    Ri = gprime / vel**2
    airrte= 2.00*pi* epsilon/Ri *rhoa*Y(r)*hei* vel
    D(r) = vel
endif
endif

c
200  IF(delrho.lt.delrhamin .and. u0.ne.0.) then
    D(R) = 0. ! not for windless cases
    airrte = 0.
endif

c
area = pi * (Y(r)**2 - radp**2)
IF(Y(R).le. RADP) THEN
    AREA = 0.00
    Y(R) = RADP + zero
    IF(time.gt. delto2) then ! delt; num prob
        D(R) = dmax1(0.00,
1           ((AFGEN2(PTIME,R1T,TIME+delto2,'R1TSRD')-
2           AFGEN2(PTIME,R1T, (TIME-delto2),'R1TSRe'))/ delt))
    else
        D(R) = dmax1(0.00,
1           ((AFGEN2(PTIME,R1T,delto2,'R1TSRD')-
2           AFGEN2(PTIME,R1T, 0.00,'R1TSRe'))/ delto2))
    endif
ENDIF
c
c   calculate totrteout
c
masrte = AFGEN2(PTIME,ET,TIME,'src1')
PWCP = AFGEN2(PTIME,PWC,TIME,'src1')
TOTPR = MASRTE/PWCP
HPRIM = AFGEN2(PTIME,PENTH,TIME,'src1')
L = 2.000 * Y(R)
c
cc = wc*rho
c
c
qstrmx = 0.00
if(u0 .ne. 0.00)
1 qstrmx = cc*K*USTAR*ALPHA1*dellay/(dellay-1.00)/phihat(rho,L)
c
c
qstrll = qstrmx * pi*L*4.00

```

```

totrteout = qstrll/wc
c
c      surface effects
c
      watrte = 0.00
      surface_q = 0.00
      yw = 1.00-ya-yc
      yw = min( max(yw, 0.000), 1.000)
      call surface(temp,hei,rho,mole,cp,yw,watrte,surface_q)
      surface_q = area * surface_q
      if(surface_q.lt. 0.00) surface_q = 0.00 ! don't let the cloud cool
      watrte = area * watrte

c
500   totrtein = airrte + TOTPRT + watrte
c
      IF(totrtein.lt.totrteout .and. .not.check2
      1           ) then ! check2 is True for HSE type spills
      D(R) = 0.
      if(hei.gt.srccut .and. Y(r).gt.srccut) then
      dHdt = (totrtein - totrteout)/3.00/pi/ Y(r)/Y(r)/rho
      D(R) = Y(R)/Hei * dHdt
      ! Let cloud radius shrink when...
      endif      ! cloud height is decreasing unless...
      if( Y(R).le.0.0100*rmax .and. masrte.eq.0.00)
      $      D(R) = 0. ! the primary source has stopped. tos;6mar86
      if( Y(R).le.hei .and. masrte.eq.0.00)
      $      D(R) = 0. ! the cloud height > the cloud radius. tos;16nov87
      endif

c
c      CALCULATE D(mass),D(massc),D(massa),D(anything left)
c
      D(mass) = totrtein - totrteout
      D(massc) = masrte - qstrll
      D(massa) = (airrte + MASRTE*(1.00/PWCP - 1.00))/(1.00+humid)
      $          - wa/wc*qstrll
      D(ebal) = 0.00
      if(ihtfl.ne. 0) ! equivalent to adiabatic mixing from TPROP for ihtfl=0
      $      D(ebal) = HPRIM*TOTPRT + h_airrte*airrte
      $          + h_watrte*watrte - enthalpy*totrteout + surface_q

c
c
      uheff = qstrmx*L/cc
      sz = 0.00
      if(u0 .ne. 0.00) sz = ( uheff*alpha1/u0/z0 )**((1.00/alpha1) * z0
c
c
      PRMT(6) = QSTRMX
      prmt(7) = sz
      prmt(8) = hei
      prmt(9) = rho

```

```

prmt(10)= Ri
prmt(11)= yc
prmt(12)= ya
prmt(13)= D(r)
prmt(16) = wc
prmt(17) = wa
prmt(18) = enthalpy
prmt(19) = temp
prmt(21) = masrte
prmt(22) = ht
prmt(23) = hh
RETURN
END

c
c
C.....
C
C      SUBROUTINE FOR OUTPUT FROM SOURCE in the presence of a Blanket
C
C      SUBROUTINE SRC10(TIME,Y,DERY,IHLF,NDIM,PRMT)

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

C
include 'sys$degadis:DEGADIS1.dec'
C
COMMON
$ERROR/STPIN,ERBND,STPMX,WTRG,WTtm,WTya,WTyc,WTeb,WTmb,WTuh,XLI,
$ XRI,EPS,ZLOW,STPINZ,ERBNDZ,STPMXZ,SRCOER,srcss,srccut,
$ htcut,ERNOBL,NOBLpt,crfger,epsilon
$PARM/ UQ,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW
$PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND
$/com_ss/ ess,slen,swid,outcc,outsz,outb,outl,swcl,swal,sent,srhl
$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
$/ALP/ ALPHA,alpha1
C
LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
C
DIMENSION Y(6),DERY(6),PRMT(25)
DIMENSION CURNT(iout_src),BKSP(iout_src),OUTP(iout_src)
C
DATA I/O/,III/O/
DATA EMAX/0./,tlast/0./
C
REAL*8 ML,K
INTEGER R,mass,massc,masa,ebal
DATA R/1./,mass/2./,massc/3./,masa/4./,ebal/5/
C
data nrec1/0/
C
I = I + 1

```

```

III = III + 1
C
qstr = prmt(6)
sz = prmt(7)
hei = prmt(8)
rho = prmt(9)
Ri = prmt(10)
yc = prmt(11)
ya = prmt(12)
vel = prmt(13)
prmt(14) = vel
if(vel .gt. prmt(20)) prmt(20) = -prmt(20)
prmt(15) = prmt(16)      ! wc
wc = prmt(16)
cc = wc * rho
wa = prmt(17)
enthalpy = prmt(18)
temp = prmt(19)
prmt(24) = prmt(22)      ! ht
prmt(25) = prmt(23)      ! hh
C
IF(hei .Le. 0.0D0) GO TO 1000
C
QSAV = PI*Y(R)*Y(R)*qstr
IF(QSAV .LT. EMAX) GO TO 110
EMAX = QSAV
RM = Y(R)
SZM = SZ
110 CONTINUE
RMAX = dMAX1(RMAX,Y(R))
C
IF(hei .Le. srccut) GO TO 1000
if(cc .le. cclow .and. u0 .eq. 0.) goto 1000      ! no wind
if(time.gt.tend+1. .and. u0.eq.0. .and. vel.eq.0.)goto 1000!no wind LNG
C
IF(I .NE. 1) GO TO 115
CURNT(1) = TIME
CURNT(2) = Y(R)
CURNT(3) = hei
CURNT(4) = qstr
CURNT(5) = sz
CURNT(6) = yc
CURNT(7) = ya
CURNT(8) = rho
CURNT(9) = ri
CURNT(10)= wc
CURNT(11)= wa
CURNT(12)= enthalpy
CURNT(13)= temp
III = 1
GO TO 125

```

```

115 IF(I .EQ. 0) RETURN
C
116 DO 116 II=1,iout_src
      BKSP(II) = CURNT(II)
C
      CURNT(1) = TIME
      CURNT(2) = Y(R)
      CURNT(3) = hei
      CURNT(4) = qstr
      CURNT(5) = sz
      CURNT(6) = yc
      CURNT(7) = ya
      CURNT(8) = rho
      CURNT(9) = ri
      CURNT(10)= wc
      CURNT(11)= wa
      CURNT(12)= enthalpy
      CURNT(13)= temp
C
      ERM = 0.
      ermss = 0.
      DO 120 II=2,iout_src
      div = curnt(ii)
      if(div .eq. 0.) div = srcoer
      ER1 = ABS( (CURNT(II)-BKSP(II))/div )
      ER2 = ABS( (CURNT(II)-OUTP(II))/div )
      if(ii.ne.3 .and. ii.ne.9 .and. ii.ne.12 .and. ii.ne.7 .and. ii.ne.11)
      1      ermss = dMAX1(ER1,ER2,ERMss) ! ex hei,QSTR,Ri,enth,wa,ya for SS
      120   ERM = dMAX1(ER1,ER2,ERM)
C
      if(check4) then           ! steady state
          if( .not. (vel.eq. 0..and.time.gt.srcess)) goto 124
      122       check3 = .true.
          outcc = wc * rho
          swcl = wc
          swal = wa
          srhl = rho
          senl = enthalpy
          outl = 2.000 * Y(r)
          Qstar = prmt(21)/pi/Y(r)**2
          if(u0.ne. 0.) sz= (alpha1/u0/z0*Qstar*outl/outcc)**(1.00/alpha1)* z0
          outsz = sz
          outb = pi*Y(r)**2 /outl/2.00
          goto 1000
      124       if(ermss .gt. srcoer) goto 125
          if( time-tlast .gt. srcess) goto 122
      return
      endif
C

```

```

      IF(ERM .LT. SRCOER) RETURN
C
125  CONTINUE
      tlast = time
      DO 130 II=1,iout_src
      IF(III.EQ.1) BKSP(II) = CURNT(II)
130  OUTP(II) = BKSP(II)
C
      III = 0
      NREC1 = NREC1 + 1
      WRITE(9,2000) (OUTP(II),II=1,iout_src)
      RETURN
C
1000 CONTINUE
      I = -1
      IF(TIME .GE. TEND) CHECK3 = .TRUE.
      NREC1 = NREC1 + 1
      WRITE(lunlog,1100)
      WRITE(lunlog,*) Hei,TIME
      TSC1 = TIME
      if(hei .le. 0.) then
          hei = 0.
          y(r) = dmin1(rmax,y(r))
          endif
      WRITE(9,2000)
      1      TIME,Y(R),hei,qstr,sz,yc,ya,rho,ri,wc,wa,enthalpy,temp
      WRITE(lunlog,1110) NREC1
C
      PRMT(5) = 1.
C
      RETURN
1100 FORMAT(5X,'VALUE OF Hei AT SOURCE TERMINATION -- @ TIME')
1110 FORMAT(5X,'NUMBER OF LINES --> ',I8)
2000 format(1pg16.9,1x,1pg16.9,<iout_src-2>(1x,1pg13.6))
      END
#####

```

```

SUBROUTINE SRTOUT(OPNRUP, table)
c
Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

include 'sys$degadis:DEGADIS3.dec/list'
c
COMMON /SORT/TCc(maxnob,maxnt),TCcSTR(maxnob,maxnt),
$ Tyc(maxnob,maxnt),Trho(maxnob,maxnt),
$ Tgamma(maxnob,maxnt),Ttemp(maxnob,maxnt),
$ TSY(maxnob,maxnt),TSZ(maxnob,maxnt),TB(maxnob,maxnt),
$ TDIST0(maxnob,maxnt),TDIST(maxnob,maxnt),KSUB(maxnt)
$/SORTIN/TIM(maxnt),NTIM,Istrt
$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$/com_atm/ istab,tamb,pamb,humid,isoftl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc
$/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag
$/alp/ alpha,alpha1
./oomsin/ oodist,avtime
c
dimension table(1)
c
logical cflag,cflag1
c
character*3 gas_name
character*40 OPNRUP
c
OPEN(UNIT=8,TYPE='NEW',NAME=OPNRUP,CARRIAGECONTROL='FORTRAN')
c
WRITE(8,1100)
if(sigx_flag.eq. 0.) then
    write(8,1102)
else
    write(8,1104)
    write(8,1105) sigx_coeff,sigx_pow,sigx_min_dist
endif
c
cflag = isoftl.eq. 1.or. ihtfl.eq. 0
cflag1= isoftl.eq.1
if(cflag) then
    call adiabat(2,wc,wa,gas_lfl,ya,cc_lfl,r,w,t,tt)
    call adiabat(2,wc,wa,gas_ufl,ya,cc_ufl,r,w,t,tt)
endif
c
DO 110 I=Istrt,NTIM
c
WRITE(8,1119)
WRITE(8,1119)
WRITE(8,1110) TIM(I)
if(cflag1) then

```

```

        WRITE(8,1116) gas_zsp,(100.*gas_lfl),(100.*gas_ufl)
        WRITE(8,1118)
    else
        WRITE(8,1115) gas_zsp,(100.*gas_lfl),(100.*gas_ufl)
        WRITE(8,1117)
        endif
    WRITE(8,1119)
    ip = 0
    II = KSUB(I)

c
    DO 120 J=1,II
c
    dist  = tdist(j,i) + oodist
    cc    = tccstr(j,i)
    rho   = Trho(j,i)
    yc    = TyC(j,i)
    temp  = Ttemp(j,i)
    gamma = Tgamma(j,i)
    b     = tb(j,i)
    sz    = tsz(j,i)
    sy    = tsy(j,i)
    blfl  = 0.
    bufl  = 0.

c
    if(.not.cflag) then
        call adiabat(-2,wc,wa,gas_lfl,ya,cc_lfl,r,w,gamma,tt)
        call adiabat(-2,wc,wa,gas_ufl,ya,cc_ufl,r,w,gamma,tt)
        endif
c
    arg = (gas_zsp/sz)**alpha1
    if(arg .ge. 80.) goto 600
c
    ccz = cc/exp(arg)
    if(ccz .lt. cc_lfl) then
        if(cflag1) then
        WRITE(8,1120) DIST,yc,Cc,rho,temp,B,SZ,SY
            else
        WRITE(8,1120) DIST,yc,Cc,rho,gamma,temp,B,SZ,SY
            endif
        goto 600
        endif
    arg = -(dlog(cc_lfl/cc) + (gas_zsp/sz)**alpha1)
    blfl = sqrt(arg)*sy + b

c
    if(ccz .lt. cc_ufl) then
        if(cflag1) then
        WRITE(8,1120) DIST,yc,Cc,rho,temp,B,SZ,SY,blfl
            else
        WRITE(8,1120) DIST,yc,Cc,rho,gamma,temp,B,SZ,SY,blfl
            endif
        goto 600

```

```

        endif
        arg = -(dlog(cc_ufl/cc) + (gas_zsp/sz)**alpha1)
        bufl = sqrt(arg)*sy + b
        if(cflag1) then
        WRITE(8,1120) DIST,yc,Cc,rho,temp,B,SZ,SY,bufl,bufl
        else
        WRITE(8,1120) DIST,yc,Cc,rho,gamma,temp,B,SZ,SY,bufl,bufl
        endif
c
600  continue
ip = ip + 1
if(ip .eq. 3) then
    ip = 0
    write(8,1119)
endif
120  CONTINUE
c
c*** summarize the mass above the UFL and LFL
c
aufl = table(i+2*maxnob)
alfl = table(i+3*maxnob)
write(8,8000) 100.*gas_ufl,100.*gas_lfl,alfl-aufl,alfl
8000  format(//,' For the UFL of ',1pg13.5,' mole percent, and',
     $' the LFL of ',1pg13.5,' mole percent:','
     $//,' The mass of contaminant between the UFL and LFL is:'
     $,1pg13.5,' kg.',/,,' The mass of contaminant above the LFL is: ',
     $1pg13.5,' kg.')
110  CONTINUE
c
CLOSE(UNIT=8)
c
c
1100 FORMAT(1H0,5X,'Sorted values for each specified time.')
1102 format(1H0,5x,'X-Direction correction was NOT applied.')
1104 format(1H0,5x,'X-Direction correction was applied.')
1105 format(1h ,5x,5x,'Coefficient:      ',1pg13.5/,,
     1      1h ,5x,5x,'Power:          ',1pg13.5/,,
     1      1h ,5x,5x,'Minimum Distance: ',1pg13.5' m')
1110 FORMAT(1H0,5X,'Time after beginning of spill ',G14.7,' sec')
1115 FORMAT(1H0,1X,'Distance',2x,3x,'Mole',3x,
     1      'Concentration',1x,'Density',2x,3x,'Gamma',3x,
     1      'Temperature',3x,'Half',4x,4x,'Sz',5x,4x,'Sy',5x,
     1      'Width at z=',0pf6.2,' m to:',/,1x,11x,1x'Fraction',2x,
     1      11x,11x,11x,3x,'Width',3x,11x,9x,
     1      2(1pg9.3,'mole%',1x))
1116 FORMAT(1H0,1X,'Distance',2x,3x,'Mole',3x,
     1      'Concentration',1x,'Density',2x,
     1      'Temperature',3x,'Half',4x,4x,'Sz',5x,4x,'Sy',5x,
     1      'Width at z=',0pf6.2,' m to:',/,1x,11x,1x'Fraction',2x,
     1      11x,11x,11x,3x,'Width',3x,11x,9x,
     1      2(1pg9.3,'mole%',1x))

```

```
1117 FORMAT(1H ,4X,'(m)',4X,11X,
1           2(1X,'(kg/m**3)',1X),11X,4X,'(K)',
1           5(8X,'(m)'))
1118 FORMAT(1H ,4X,'(m)',4X,11X,
1           2(1X,'(kg/m**3)',1X),4X,'(K)',
1           5(8X,'(m)'))
1119 FORMAT(1H )
1120 FORMAT(1H ,3(1X,1PG9.3,1X),2X,0pf7.4,2X,1X,1PG10.3,1X,
1           6(1X,1PG9.3,1X))
C
RETURN
END
#####
4 -- sys$degadis:srtout.for          6-SEP-1989 17:55:16
```

```

C.....  

C  

C  

SUBROUTINE SSG(DIST,Y,Dery,PRMT)  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

DIMENSION Y(1),Dery(1),PRMT(1)  

C  

include 'sys$degadis:DEGADIS2.dec'  

C  

parameter (zero=1.D-10, rcrit=2.5D-3)  

C  

COMMON  

$/PARM/U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,  

$ gas_ufl,gas_lfl,gas_zsp,gas_name  

$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

$/ALP/ ALPHA,alpha1  

$/phicom/ iphifl,dellay  

C  

REAL*8 K,ML  

LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

C  

INTEGER rhoun,du, mhi, mlow  

DATA rhoun/1/,du/2/, mhi/3/, mlow/4/  

C  

C*** PRMT(I) I/O  

C***   I          VALUE        IN/OUT  

C***   --         -----       -----  

C***   6           E          IN  

C***   7           Cc         OUT  

C***   8           XV(I)      IN  

C***   9           TO(I)      IN  

C***   10          -          -  

C***   11          NREC(I,2)   OUT -- STARTS OUTPUT UNIT=9  

C***   12          DIST       OUT  

C***   13          sz          -  

C***   14          yc          out  

C***   15          rho         out  

C***   16          temp        out  

C***   17          gamma       out  

C***   18          -          -  

C***   19          -          -  

C***   20          rholay     -  

C***   21          sz          -  

C

```

```

XVL = PRMT(8)
SY = RT2*DELTA*(DIST + XVL)**BETA
Erate = PRMT(6)

c
c
sz0 = prmt(22)
sz = sz0

c
C*** MATERIAL BALANCE
c
      iii = 0
100   Cc = Erate*ALPHA1*(Z0/SZ)**ALPHA/U0/SZ/SQRTPI/SY
c
      cclay = cc/dellay
      call addheat(cclay,Y(dh),rholay,temp,cp)
      prod = dmax1( Y(rhouh)/rholay/prmt(18), zero)
      sz = ( prod ) **(1./alpha1) * z0
      dif = abs(sz - sz0)/(abs(sz)+abs(sz0)+zero)
      if(dif.gt. rcrit) then
          iii = iii+1
          if(iii.gt. 20) call trap(33)
          sz0 = sz
          goto 100
          endif
      prmt(20) = rholay
      prmt(21) = sz
      HEFF = GAMMAF*SZ/ALPHA1

      call adiabat(0,wc,wa,yc,ya,cc,rho,wm,enth,temp)
      call adiabat(0,wc,wa,yclay,ya,cclay,rholam,wml,enth,temp)
      rit = 0.

c
      if(isofl.eq.0 .and. ihtfl.ne.0) then
          call addheat(cc,dellay*Y(dh),rho,temp,cp)
          rit = rift(temp,heff)
          endif
c
      RISTR = RIF(RHO,HEFF)
      PHI = PHIF(RISTR,rit)
c
      dery(rhouh) = prmt(19)/phi
      heigh = heff*dellay
      yw = 1.-yclay-ya
      yw = min( max( yw, 0.000 ), 1.000 )
      call surface(temp,heigh,rholay,wml,cp,yw,water,qrte)
      if(temp.ge. tsurf .or. temp.ge. tamb) qrte = 0.
      dery(dh) = ( qrte/dellay-Y(dh)*Dery(rhouh) )/Y(rhouh)
c
c
c*** Calculate the derivative for the total mass above the UFL and LFL
c

```

```

gamma = (rho-rhoa)/cc           ! gamma

if(check4) then
  DERY(mlow) = 0.
  DERY(mhi)  = 0.
  if( isofl.eq.1 .or. ihtfl.eq.0 ) then
    call adiabat(2,aa,dd,gas_ufl,ee,chi ,gg,hh,pp,oo)
    call adiabat(2,aa,dd,gas_lfl,ee,clow,gg,hh,pp,oo)
  else
    call adiabat(-2,aa,dd,gas_ufl,ee,chi ,gg,hh,gamma,oo)
    call adiabat(-2,aa,dd,gas_lfl,ee,clow,gg,hh,gamma,oo)
  endif

gammax = sqrt(pi * Cc * Sz * Sy * GAMMAF / alpha1

if(cc.gt.clow) then
  wlow = Dlog(cc/clow)
  DERY(mlow)= 2.00*clow*Sz/alpha1*series(wlow)
  DERY(mlow)= DMIN1( DERY(mlow), gammax )
endif

if(cc.gt.chi ) then
  whi = Dlog(cc/chi )
  DERY(mhi) = 2.00*chi *Sz/alpha1*series(whi )
  DERY(mhi) = DMIN1( DERY(mhi ), gammax )
endif
endif

c
PRMT(7) = Cc
PRMT(12) = DIST
prmt(14) = yc
prmt(15) = rho
prmt(16) = temp
prmt(17) = gamma
RETURN
END

#####

```

```

C.....  

C  

C      SUBROUTINE SSGOUT(X,Y,D,IHLF,NDIM,PRMT)  

C  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

include 'sys$degadis:DEGADIS2.dec'  

C  

parameter (nssg=7, zero=1.e-10)  

C  

DIMENSION Y(1),D(1),PRMT(1),BKSP(nssg),OUT(nssg),CURNT(nssg)  

C  

COMMON  

$/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/STP/ STPO,STPP,ODLP,ODLLP,STPG,ODLG,ODLLG  

$/STOPIT/ TSTOP  

C  

REAL*8 K,ML  

C  

C*** PARAMETER OUTPUT  

C  

C*** FROM SSG           OUTPUT TO MODEL .  

C*** -----  

C*** X                  DIST  

C*** PRMT(7)            Cc  

C*** Y(1)                SZ  

C*** prmt(14)           yc  

C*** prmt(15)           rho  

C*** prmt(16)           temp  

C*** prmt(17)           gamma  

C  

ERM = 0.  

TOL = PRMT(9)  

TSL = TS(TOL,X)  

prmt(22) = prmt(21)    ! sz  

C  

IF(PRMT(11) .NE. 0.) GO TO 90  

C  

C*** STARTUP FOR OUTPUT ROUTINE  

C  

RII = -100./STPG  

RI = 0.  

CURNT(1) = X  

curnt(2) = prmt(14)    ! yc  

CURNT(3) = PRMT(7)    ! cc  

curnt(4) = prmt(15)    ! rho  

curnt(5) = prmt(17)    ! gamma  

curnt(6) = prmt(16)    ! temp

```

```

        CURNT(7) = prmt(21)      ! sz
C
90    CONTINUE
C
C*** RECORD THE CURRENT AND PREVIOUS RECORDS
C
        RI = RI + 1.
C
        DO 100 II=1,nssg
100    bksp(ii) = curnt(ii)
C
        CURNT(1) = X
        curnt(2) = prmt(14)      ! yc
        CURNT(3) = PRMT(7)      ! cc
        curnt(4) = prmt(15)      ! rho
        curnt(5) = prmt(17)      ! gamma
        curnt(6) = prmt(16)      ! temp
        CURNT(7) = prmt(21)      ! sz
C
C*** stop integration when cc<cclow and time is satisfied
C
        IF(PRMT(7).LT.CCLOW .AND. TSL.GE.TSTOP) GO TO 1000
C
C*** CHECK FOR OUTPUT
C
        DO 110 II=3,nssg
        ER1 = ABS( (CURNT(ii)-BKSP(ii))/(CURNT(ii)+zero) )
        ER2 = ABS( (CURNT(ii)-OUT(ii))/(CURNT(ii)+zero) )
110    ERM = dMAX1(ER1,ER2,ERM)
C
C*** OUTPUT RECORD IF ODLG IS EXCEEDED OR 100 METERS SINCE LAST OUTPUT
C
        DX = CURNT(1) - OUT(1)
        IF( RI.NE.1. .AND. ERM.LT.ODLG .AND. DX.LE.ODLLG) RETURN
C
C*** RECORD THE LAST POINT TO BE UNDER THE ERROR CRITERIA. IN CASE
C*** THE FIRST POINT AFTER A RECORD EXCEEDS THE ERROR BOUND, RECORD
C*** THAT POINT AS WELL.
C
        DO 120 II=1,nssg
        IF(RI .EQ. RII+1.) BKSP(ii) = CURNT(ii)
120    OUT(ii) = BKSP(ii)
C
        RI = RII
        PRMT(11) = PRMT(11) + 1.
C
        WRITE(9,*) (OUT(ii),II=1,nssg)
        RETURN
C
1000   CONTINUE
C

```

```
C*** STOP INTEGRATION
C
PRMT(12) = X
TSTOP = TSL
PRMT(11) = PRMT(11) + 1.
WRITE(9,*) (CURNT(II),II=1,nssg)
C
PRMT(5) = 1.
C
RETURN
END
#####
3 .. sys$degadis:ssgout.for          6-SEP-1989 18:05:18
```

```

C.....  

C  

C      SUBROUTINE SSGOUT(X,Y,D,IHLF,NDIM,PRMT)  

C  

C      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      parameter (nssg=9, zero=1.e-10)  

C  

C      DIMENSION Y(1),D(1),PRMT(1),BKSP(nssg),OUT(nssg),CURNT(nssg)  

C  

C      include 'sys$degadis:DEGADIS2.dec/list'  

C  

C      COMMON  

$/PARM/U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$/STP/STPP,ODLP,ODLLP,STPG,ODLG,ODLLG  

$/ALP/ALPHA,alpha1  

C  

C  

C      REAL*8 ML,K  

C  

C*** PARAMETER OUTPUT  

C  

C*** FROM SSG          OUTPUT TO MODEL  

C*** -----  

C*** X                  DIST  

C*** PRMT(7)            Cc  

C*** Y(1)                SZ  

C*** PRMT(8)            XV  

C  

C      ERM = 0.  

C      prmt(22) = prmt(21)  

C  

C      IF(PRMT(11) .NE. 0.) GO TO 90  

C  

C*** STARTUP FOR OUTPUT ROUTINE  

C  

RII = -100./STPG  

RI = 0.  

CURNT(1) = X  

CURNT(2) = PRMT(14)    ! yc  

CURNT(3) = prmt(7)     ! cc  

CURNT(4) = PRMT(15)    ! rho  

curnt(5) = prmt(17)    ! gamma  

curnt(6) = prmt(16)    ! temp  

curnt(7) = 0.0          ! b  

curnt(8) = prmt(21)    ! sz  

curnt(9) = rt2*delta*(x+prmt(8)**beta ! sy  

C  

90  CONTINUE  

C

```

```

C*** RECORD THE CURRENT AND PREVIOUS RECORDS
C
      RI = RI + 1.
C
      DO 100 II=1,nssg
100   BKSP(II) = CURNT(II)
      CURNT(1) = X
      CURNT(2) = PRMT(14)      ! yc
      CURNT(3) = prmt(7)       ! cc
      CURNT(4) = PRMT(15)      ! rho
      curnt(5) = prmt(17)      ! gamma
      curnt(6) = prmt(16)      ! temp
      curnt(7) = 0.0            ! b
      curnt(8) = prmt(21)      ! sz
      curnt(9) = rt2*delta*(x+prmt(8))**beta ! sy
C
C*** STOP INTEGRATION WHEN Cc < CcLOW
C
      IF(PRMT(7).LT.CcLOW) GO TO 1000
C
C*** CHECK FOR OUTPUT
C
      DO 110 II=2,nssg
      if(ii.eq.7) goto 110
      ER1 = ABS( (CURNT(II)-BKSP(II))/(CURNT(II)+zero) )
      ER2 = ABS( (CURNT(II)-OUT(II))/(CURNT(II)+zero) )
110   ERM = dMAX1(ER1,ER2,ERM)
C
C*** OUTPUT RECORD IF ODLG IS EXCEEDED OR 100 METERS SINCE LAST OUTPUT
C
      DX = CURNT(1) - OUT(1)
      IF( RI.NE.1. .AND. ERM.LT.ODLG .AND. DX.LE.ODLLG) RETURN
C
C*** RECORD THE LAST POINT TO BE UNDER THE ERROR CRITERIA. IN CASE
C*** THE FIRST POINT AFTER A RECORD EXCEEDS THE ERROR BOUND, RECORD
C*** THAT POINT AS WELL.
C
      DO 120 II=1,nssg
      IF(RI .EQ. RII+1.) BKSP(II) = CURNT(II)
120   OUT(II) = BKSP(II)
C
      RI = RII
      PRMT(11) = PRMT(11) + 1.
C
      call ssout(out)
      RETURN
C
      1000  CONTINUE
C
C*** STOP INTEGRATION
C

```

D-160

```
PRMT(12) = X
PRMT(11) = PRMT(11) + 1.

C
call ssout(currnt)
C
PRMT(5) = 1.

C
RETURN
END

####
```

```

subroutine ssout(out)
c
Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

dimension out(9)
c
common
$ /com_gprop/ gas_mm,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$/com_fl/ cflag,clfl,cufl
$/alp/ alpha,alpha1
./oomsin/ oodist,avtime
c
character*3 gas_name
c
data ip/0/
c
logical cflag
c
c
dist = out(1) + oodist
yc = out(2)
cc = out(3)
rho = out(4)
gamma = out(5)
temp = out(6)
b = out(7)
sz = out(8)
sy = out(9)
c
if(.not.cflag) then
    call adiabat(-2,wc,wa,gas_lfl,ya,clfl,r,w,gamma,tt)
    call adiabat(-2,wc,wa,gas_ufl,ya,cufl,r,w,gamma,tt)
endif
c
arg = (gas_zsp/sz)**alpha1
if(arg .ge. 80.) then
    if(cflag) then
        WRITE(8,1120) DIST,yc,Cc,rho,temp,B,SZ,SY
        else
        WRITE(8,1125) DIST,yc,Cc,rho,gamma,temp,B,SZ,SY
        endif
        goto 600
    endif
c
ccz = cc/exp(arg)
if(ccz .lt. clfl) then
    if(cflag) then
        WRITE(8,1120) DIST,yc,Cc,rho,temp,B,SZ,SY
        else

```

```

      WRITE(8,1125) DIST,yc,Cc,rho,gamma,temp,B,SZ,SY
      endif
      goto 600
      endif
      arg = -(dlog(clfl/cc) + (gas_zsp/sz)**alpha1)
      blfl = sqrt(arg)*sy + b
      c
      if(ccz .lt. cufl) then
          if(cflag) then
              WRITE(8,1120) DIST,yc,Cc,rho,temp,B,SZ,SY,blfl
          else
              WRITE(8,1125) DIST,yc,Cc,rho,gamma,temp,B,SZ,SY,blfl
          endif
          goto 600
      endif
      arg = -(dlog(cufl/cc) + (gas_zsp/sz)**alpha1)
      bufl = sqrt(arg)*sy + b
      if(cflag) then
          WRITE(8,1120) DIST,yc,Cc,rho,temp,B,SZ,SY,blfl,bufl
      else
          WRITE(8,1125) DIST,yc,Cc,rho,gamma,temp,B,SZ,SY,blfl,bufl
      endif
      c
      600  continue
      ip = ip + 1
      if(ip .eq. 3) then
          ip = 0
          write(8,1119)
      endif
      c
      c
      1119  FORMAT(1H )
      1120  FORMAT(1H ,11(1X,1PG9.3,1X))
      1125  FORMAT(1H ,3(1X,1PG9.3,1X),2x,0PF7.4,2x,1X,1PG10.3,1X,
      1       6(1X,1PG9.3,1X))
      c
      return
      end
####
```

```

C.....  

C  

C      PSEUDO-STEADY STATE SUPERVISOR  

C  

C      SUBROUTINE SSSUP(H_masrte)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

include 'sys$degadis:DEGADIS2.dec/nolist'  

C  

COMMON  

$/GEN3/ radg(2,maxl),qstr(2,maxl),srcden(2,maxl),srcwc(2,maxl),  

$ srcwa(2,maxl),srcenth(2,maxl)  

$/SSCON/ NREC(maxnob,2),TO(maxnob),XV(maxnob)  

$/GEN1/ PTIME(igen), ET(igen), R1T(igen), PWC(igen), PTEMP(igen),  

$      PFRACV(igen), PENTH(igen), PRHO(igen)  

$/gen2/ den(5,igen)  

$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,  

$ gas_ufl,gas_lfl,gas_zsp,gas_name  

$/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CCLOW  

$/ERROR/SY0ER,ERRO,SZ0ER,WTA10,WTQOO,WTSZ0,ERRP,SMXP,  

$ WTSZP,WTSYR,WTBEP,WTDH,ERRG,SMXG,ERTDNF,ERTUPF,WTRUH,WTDHG  

$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$/STP/ STPO,STPP,ODLP,ODLLP,STPG,ODLG,ODLLG  

$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

$/nend/ poundn, pound  

$/ALP/ ALPHA,alpha1  

$/phicom/ iphifl,delay  

$/sprd_con/ ce, delrhomin  

$/STOPIT/ TSTOP  

$/CNOBS/ NOBS  

C  

REAL*8 K,ML,L  

LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

logical pup,pdn  

C  

character*4 pound  

character*3 gas_name  

C  

EXTERNAL PSS,PSSOUT,SSG,SSGOUT,OB,OBOUT  

C  

DIMENSION PRMT(22),Y(5),DERY(5),AUX(8,5)  

C  

DATA RTOT/0./  

data wma/28.96/, wmw/18.0/  

C  

C  

c*** Estimate the earliest and latest time an observer can be released

```

```

c*** over the source.
c
      R = AFGEN(RADG, 0.0D0, 'RADG')
      T01 = TOOB(R, 0.0D0)
c
c** For low wind speed cases which form a blanket, earlier times
c** than T01 may be possible. Check each of the points in RADG.
c
      do i=2,max1
        if ( radg(1,i).eq.poundn .and. radg(2,i).eq.poundn) goto 20
        TOF = TOOB( radg(2,i), radg(1,i) )
        if ( TOF .lt. T01) T01 = TOF
      enddo
c
c** Now, calculate the last possible time an observer can be released.
c
      20   continue
      XEND = AFGEN(RADG,TEND,'RADG')
      TOF = TOOB(-XEND,TEND)
c
c*** Now, divide the total possible time among the observers.
c
      DTOB = (TOF-T01)/FLOAT(NOBS)
      T01 = T01 + DTOB/2.0D0
c
c*** perform the calculation for each observer
c
      write(12,1162)
c
      DO 120 I = 1,NOBS
c
c*** RESET AGAIN
c
      AGAIN = .FALSE.
c
      T0(I) = DTOB*dble(I-1) + T01
      pup = .true.
      pdn = .true.
c
c*** IF(XEND .GT. XIT(TEND,T0(I))) -- IS
c*** TRUE WHEN THE SOURCE WILL TERMINATE BEFORE THE OBSERVER
c*** CAN REACH THE DOWNWIND EDGE.
c
      IF(XEND .GT. XIT(TEND,T0(I))) then
        pdn = .false.
        TDOWN = TEND
      endif
c
c*** IF(XIT(0.0D0,T0(I)) .gt. -R) -- IS
c*** TRUE WHEN THE SOURCE WILL begin after THE OBSERVER
c*** has passed THE DOWNWIND EDGE @ t=0.0

```

```

C
      R = AFGEN(RADG,0.0D0,'RADG')
      IF(t0(i).le.0. .and. XIT(0.0D0,T0(I)).gt.-R) then
          pup = .false.
          TUP = 0.0D0
          endif
C
      if(pup) TUP = TUPF(T0(I))
      if(pdin) TDOWN = TDNF(T0(I))
C
      XDOWN = XIT(TDOWN,T0(I))
      XUP = XIT(TUP,T0(I))
      WRITE(lunlog,1160) TUP,XUP,TDOWN,XDOWN
C
C*** SET UP INTEGRATION PARAMETERS FOR EACH OBSERVER.
C
      do ijk=1,22
          prmt(ijk) = 0.0D0
      enddo
      do ijk=1,5
          y(ijk) = 0.0D0
          dery(ijk)= 0.0D0
      do ijkl=1,8
          aux(ijkl,ijk) = 0.0D0
      enddo
      enddo
C
      PRMT(1) = TUP
      PRMT(2) = TDOWN
      PRMT(3) = STPO
      PRMT(4) = ERRO
      PRMT(5) = dMAX1(1.0D0,(TDOWN-TUP)/50.0D0)
      PRMT(6) = T0(I)
      prmt(7) = xup
      PRMT(13)= XDOWN - XUP
C
      Y(1) = sz0er    ! tos;3mar86
      Y(2) = sz0er    ! Mrate
      y(3) = sz0er * AFGEN(srcwc,tup,'IC')    ! Crate
      y(4) = sz0er * AFGEN(srcwa,tup,'IC') + 1.0-6    ! BDARate
      y(5) = sz0er * AFGEN(srcenth,tup,'IC') ! Hrate
C
      DERY(1) = WTAIO
      DERY(2) = WTQOO
      DERY(3) = WTSZ0
      DERY(4) = 1.0D0
      DERY(5) = 1.0D0
C
      NDIM = 4
      if(isofl.eq. 0 .and. ihtfl.ne. 0) ndim=5
C

```

```

C*** PERFORM INTEGRATION.
C
      WRITE(lunlog,1120) I
1120  FORMAT(//,' Entering Observer Integration Step for Observer # ',
      $   I3)
C
      CALL RKGST(PRMT,Y,DERY,NDIM,IHLF,OB,OBOUT,AUX)
C
      IF(IHLF .GE. 10) CALL trap(8,IHLF)
C
      write(lunlog,1125)
1125  format(' ',10x,'Observer Integration complete...')

c
c
c*** Establish initial conditions
c
      cclay = prmt(14)
      cc = cclay*dellay
      wclay = prmt(15)
      walay = prmt(16)
      enthlay=prmt(17)
      rholay = prmt(18)

C
      L = XDOWN - XUP
      B = Y(1)
      AREA = B*L
      QSTR0 = Y(3)/area
      Erate = 2.00*qstr0*L*b
      cc = min(cc, rhoe)
      WCP = AFGEN2(PTIME,PWC,TDOWN,'SSS-WC')
      RHOP = AFGEN2(PTIME,PRHO,TDOWN,'SSS-RH')
      CCP = WCP*RHOP
      if(cc .gt. ccp) cc = ccp

      sz0 = (qstr0*L/cc * alpha1/u0/z0)**(1.00/alpha1) * z0

C
      ratio1= u0*z0/ALPHA1/ Z0**ALPHA1 *Cc /B/qstr0/L
      ratio = ratio1* sz0**alpha1 * (B + sqrt(pi)/2.00*sy0er)
      if(ratio.le. 1.00) then
          sy0er = (1.00/(RATIO1*sz0**alpha1) - b)*2.00/sqrt(pi)
      else
          sz0 = (1.00/((B+ sqrt(pi)/2.00*sy0er)*ratio1))**(1.00/alpha1)
      endif

C
c*** Establish the thermodynamic properties of mixtures of air and
c*** the gas mixture (WCLAY,WALAY,ENTHLAY) assuming adiabatic
c*** mixing. This is accomplished with the call to SETDEN.
c*** Then, extrapolate the properties to the centerline,
c*** ground level concentration.
c

```

```

humsrc = (1.00-wclay-walay*(1.00+humid))/wclay
call setden(wclay,walay,enthalay)
if(isofl.eq. 1) goto 200

c
c*** Scan through the array DEN for the last value, and establish a new
c*** final value based on the centerline, ground level concentration.
c
do iii= 1,igen
  if(den(1,iii) .gt. 1.00) then
    ii = iii+1
    rholay = den(3,iii-1)
    temlay = den(5,iii-1)
    if(ii .gt. igen .or. iii.le.3) call trap(2)
    cc = cclay*dellay      ! exact due to profile assumptions
    if(cc.gt. rhoe) then
      write(lunlog,1126) cc,rhoe
1126      format(1,1,10('** **'),1,1,cc: ',1pg13.5,', 'is greater',
      $           ' than rhoe: ',1pg13.5,1,1,10('** **'),1)
      cc = rhoe
    endif
    rho = cc*(rhohay-rhoa)/cclay + rhoa      ! assumes gamma=con
    wc = cc/rho
    w2 = den(2,iii-2)/den(3,iii-2)
    w1 = den(2,iii-1)/den(3,iii-1)
    wa = (1.00-(1.00+humsrc)*wc)/(1.00+humid)
    wm = 1.00/(wc/gas_mw + wa/wma + (1.00-wa-wc)/wmw)
    yc = wm/gas_mw * wc
    Yc= dmax1(0.000, dmin1(1.000, yc) )
    den(1,iii) = Yc
    den(2,iii) = cc
    den(3,iii) = rho

c
c*** now, determine the enthalpy and temperature of such a mixture.
c*** Base both enthalpy and temperature on the fact that
c*** (yc/rho) is proportional to temperature (and therefore enthalpy).
c
denom = den(1,iii-2)/den(3,iii-2)-den(1,iii-1)/den(3,iii-1)
if(denom.ne. 0.00) then
  slope = (yc/rho-den(1,iii-1)/den(3,iii-1))/denom
  den(4,iii) = slope*(den(4,iii-2)-den(4,iii-1))+den(4,iii-1)
  den(4,iii) = dmin1( dmax1(h_masrte,den(4,iii)), 0.00)
  den(5,iii) = slope*(den(5,iii-2)-den(5,iii-1))+den(5,iii-1)
  den(5,iii) = dmin1( dmax1(gas_temp,den(5,iii)), tamb)
else
  den(4,iii) = den(4,iii-1)
  den(5,iii) = den(5,iii-1)
endif
temp = den(5,iii)
den(1,ii) = 2.00      ! end-of-record
c.   if(cc.gt. rhoe) call trap(31)
      goto 200

```

```

        endif
    enddo
C
200   CONTINUE

    IF(Cc .GT. RHOE) then
        WRITE(lunlog,1127) QSTRO,SZ0,Cc,rhoe
1127      format(1 1,10('****'),/, ' qstr0: ',1pg13.5,
      $      ' sz0: ',1pg13.5,/,
      $      ' cc: ',1pg13.5,' is greater',
      $      ' than rhoe: ',1pg13.5,/,1 1,10('****'),/)
C      call trap(30)
        endif
C
C*** SHOW THE OPERATOR WHAT IS GOING ON
C
        WRITE(lunlog,1160) TUP,XUP,TDOWN,XDOWN
        WRITE(lunlog,1170) AREA,L,B
        WRITE(lunlog,1180) QSTRO,SZ0,sy0er
        write(lunlog,1185) wclay,walay,rholay,cclay,temlay
        write(lunlog,1186) wc,rho,cc,temp

        write(12,1161) i,t0(i), tup,tdown, xdown,L, B,Y(3), wc,temp,Sz0

1160  FORMAT(1 TUP: ',1pG13.5,' XUP: ',1pG13.5,' TDOWN: ',
      $ 1pG13.5,' XDOWN: ',1pG13.5)
1161  format(1 ,I3,1x,f8.1,1x,6(1x,f6.1,1x),1x,f5.3,1x,1x,
      $ f5.1,1x,1x,f5.2)
1162  format(1 ,1x,f8.1,1x,6(1x,f6.1,1x),1x,f5.3,1x,1x,
      $ 1x,'Xdown',2x,1x,'Length',1x,1x,'Hwidth',1x,1x,'E rate',1x,
      $ 'Mass fr',1x,'Temp',2x,2x,'Sz0',//)
1170  FORMAT(' Half AREA: ',1pG13.5,' LENGTH: ',1pG13.5,' B: ',1pG13.5)
1180  FORMAT(' TAKEUP FLUX: ',1pG12.5,' SZ0: ',1pG12.5,
      $ ' sy0: ',1pg12.5)
1185  format(' wclay: ',1pg12.5,' walay: ',1pg12.5,
      $      ' rholay: ',1pg12.5,' cclay: ',1pg12.5,/,
      $      ' temlay: ',1pg13.5)
1186  format(' wc: ',1pg12.5,
      $      ' rho: ',1pg12.5,' cc: ',1pg12.5,' temp: ',1pg12.5)
C
C*** PREPARE FOR PSEUDO-STEADY STATE INTEGRATION.
C
        do ijk=1,22
            prmt(ijk) =0.00
        enddo
        do ijk=1,5
            y(ijk) = 0.00
            dery(ijk)= 0.00
        do ijkl=1,8
            aux(ijkl,ijk) = 0.00
        enddo

```

```

enddo

c
PRMT(1) = XDOWN
PRMT(2) = 6.023023
PRMT(3) = STPP
PRMT(4) = ERRP
PRMT(5) = SMXP
PRMT(6) = Erate
PRMT(7) = Cc      ! -- OUTPUT
PRMT(8) = B       ! -- OUTPUT

c
C*** PRMT(9) & PRMT(10) ARE CONSTANTS FOR D(SY) & D(SZ)
c
PRMT(9) = Ce*sqrt(G*Z0/ALPHA1*GAMMAF)*GAMMAF/U0
PRMT(10)= Z0**ALPHA*K*USTAR*ALPHA1*ALPHA1/U0
PRMT(11)= NREC(I,1)
c
PRMT(12)= DIST AT COMPLETION -- OUTPUT
PRMT(13)= TO(I)
c
prmt(14)= yc    ! output
c
prmt(15)= rho   ! output
c
prmt(16)= temp ! output; not recorded if isofl=1
c
prmt(17)= gamma ! output; not recorded if isofl=1 .or. ihtfl=0
prmt(18)= u0*z0/alpha1
prmt(19)= rhoa*k*ustar*alpha1
prmt(20)= rholay
prmt(21)= sz0
prmt(22)= sz0

c
Y(1) = rholay*prmt(18)*(S20/z0)**alpha1           ! rho*ueff*heff
Y(2) = SYOER*SYOER
Y(3) = B + sqrt(pi)/2.D0*sy0er
y(4) = 0.          ! added heat

c
DERY(1) = WTSZP
DERY(2) = WTSYP
DERY(3) = WTBEPE
dery(4) = WTDH

c
NDIM = 4

c
WRITE(lunlog,1130)
1130 FORMAT(' Entering Integration Step -- B > 0. ')
c
C*** PERFORM INTEGRATION
c
CALL RKGST(PRMT,Y,DERY,NDIM,IHLF,PSS,PSSOUT,AUX)
c
IF(IHLF .GE. 10) CALL trap(9,IHLF)
c
NREC(I,1) = INT(PRMT(11))
WRITE(lunlog,1100) NREC(I,1),TO(I)

```

```

1100  FORMAT(3X,'NUMBER OF RECORDS IN PSS = 'I10,' FOR T0='1pg13.5)
C
      IF(AGAIN) GO TO 119
C
C*** GAUSIAN COMPLETION OF THE INTEGRATION
C
C*** PSSOUT FORCES THE ABOVE INTEGRATION TO FINISH WHEN B<0 FOR THE
C*** FIRST TIME. THE STEP BEFORE THIS OCCURS IS RECORDED ON UNIT 7.
C*** THE STEP WHEN B GOES NEGATIVE IS CURRENTLY IN Y.
C
C*** THE CALCULATION METHOD CHANGES THE CURRENT VALUE OF SY TO A VALUE
C*** CALCULATED AS IF BEFF=sqrtpi*SY/2. RETAINING THE LAST VALUE OF Cc IN THE
C*** MATERIAL BALANCE.
C
      heat = y(4)
      rholay = prmt(20)
      Cc = PRMT(7)
      rhouh = Y(1)
      sz = ( rhouh/rholay/prmt(18) )**(1.00/alpha1) * z0
      SYT = Erate*ALPHA1*(Z0/SZ)**ALPHA/U0/SZ/Cc/SQRTPI
C
      XT = PRMT(12)
      XV(I) = (SYT/RT2/DELTA)**(1.00/BETA) - XT
C
C*** SET UP INTEGRATION FOR THE GAUSSIAN DISPERSION PHASE.
C
      do ijk=1,22
      prmt(ijk) =0.00
      enddo
      do ijk=1,5
      y(ijk) = 0.00
      dery(ijk)= 0.00
      do ijkl=1,8
      aux(ijkl,ijk) = 0.00
      enddo
      enddo
C
      PRMT(1) = XT
      PRMT(2) = 6.023D23
      PRMT(3) = STPG
      PRMT(4) = ERRG
      PRMT(5) = SMXG
      PRMT(6) = Erate
      PRMT(7) = Cc      ! -- OUTPUT
      PRMT(8) = XV(I)
      PRMT(9) = TO(I)
C      PRMT(10)= "BLANK"
      PRMT(11)= NREC(I,2)
C      PRMT(12)= DIST AT COMPLETION -- OUTPUT
C      prmt(13)= "blank"
C      prmt(14)= yc          ! output

```

```

c      prmt(15)= rho          ! output
c      prmt(16)= temp         ! output
c      prmt(17)= gamma        ! output
      prmt(18)= u0*z0/alpha1
      prmt(19)= rhoa*k*ustar*alpha1
      prmt(20)= rholay
      prmt(21)= sz
      prmt(22)= sz

c
      Y(1) = rhouh
      Y(2) = heat

c
      DERY(1) = WTRUH
      dery(2) = WTDHG

c
      NDIM = 2

c
      WRITE(lunlog,1140)
1140  FORMAT(' Entering Gaussian Stage of Integration ')
c
c*** PERFORM INTEGRATION
c
      CALL RKGST(PRMT,Y,DERY,NDIM,IHLF,SSG,SSGOUT,AUX)

c
      IF(IHLF .GE. 10) CALL trap(10,IHLF)

c
      NREC(I,2) = INT(PRMT(11))
      RTOT = RTOT + FLOAT(NREC(I,1) + NREC(I,2))
      WRITE(lunlog,1110) RTOT,I
1110  FORMAT(5X,'TOTAL NUMBER OF RECORDS = ',1pg13.4,' THROUGH',
      $' OBS # ',I3)
c
      IF(RTOT .GT. 120000.) CALL trap(11)

c
119   CONTINUE
      write(lunlog, 1150) tstop, Prmt(12)
1150  format(/, ' Last time Observer was active: ',1pg13.5,' s at ',
      $ 1pg13.5,' m')
120   CONTINUE

c
      RETURN
      END
#####

```

```

C.....  

C  

SUBROUTINE STRT2(OPNRUP,H_masrte)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

include 'sys$degadis:DEGADIS2.dec'  

C  

COMMON  

$/GEN3/ radg(2,maxl),qstr(2,maxl),srcden(2,maxl),srcwc(2,maxl),  

$ srcwa(2,maxl),srcenth(2,maxl)  

$/TITLE/ TITLE  

$/GEN1/ PTIME(igen), ET(igen), R1T(igen), PWC(igen), PTEMP(igen),  

$ PFRACV(igen), PENTH(igen), PRHO(igen)  

$/GEN2/ DEN(5,igen)  

$/ITI/ T1,TINP,TSRC,TOBS,TSRT  

$/ERROR/SY0ER,ER0,SZ0ER,WTAIO,WTQ00,WTSZ0,ERRP,SMXP,  

$ WTSZP,WTSYP,WTBEP,WTDH,ERRG,SMXG,ERTDNF,ERTUPF,WTRUH,WTDHG  

$/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,  

$ gas_ufl,gas_lfl,gas_zsp,gas_name  

$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

$/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag  

$/NEND/ POUNDN,POUND  

$/ALP/ ALPHA,alpha1  

$/phicom/ iphifl,dellay  

$/sprd_con/ ce, delrhomin  

$/COM_SURF/ HTCUT  

./oomsin/ oodist,avtime  

C  

character*80 TITLE(4)  

C  

character*4 pound  

character*24 TINP,TSRC,TOBS,TSRT  

character*3 gas_name  

C  

REAL*8 K,ML  

LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

C  

character*40 OPNRUP  

C  

OPEN(UNIT=9,NAME=OPNRUP,TYPE='OLD')  

C  

DO 90 I = 1,4  

90 READ(9,1000) TITLE(I)  

1000 FORMAT(A80)  

C

```

```

      READ(9,*) NP
      DO 100 I=1,NP
100   READ(9,*) PTIME(I),ET(I),R1T(I), PWC(I), PTEMP(I),
     $      PFRACV(I), PENTH(I), PRHO(I)
      PTIME(NP + 1) = POUNDN
c
      READ(9,*) NP
      DO 220 I=1,NP
220   READ(9,*) DEN(1,I),DEN(2,I),den(3,I),den(4,i),den(5,i)
      den(1,np+1) = 2.
c
      READ(9,*) NP
      DO 300 I=1,NP
      READ(9,*) radg(1,I),radg(2,I),qstr(2,I),srcden(2,I),srcwc(2,i),
     1      srcwa(2,i),srcenth(2,i)
      qstr(1,I) = radg(1,I)
      srcden(1,I) = radg(1,I)
      srcwc(1,i) = radg(1,i)
      srcwa(1,i) = radg(1,i)
      srcenth(1,i) = radg(1,i)
300   continue
      I = NP + 1
      radg(1,I) = POUNDN
      radg(2,I) = POUNDN
      qstr(1,I) = POUNDN
      qstr(2,I) = POUNDN
      srcden(1,I) = POUNDN
      srcden(2,I) = POUNDN
      srcwc(1,I) = POUNDN
      srcwc(2,I) = POUNDN
      srcwa(1,I) = POUNDN
      srcwa(2,I) = POUNDN
      srcenth(1,I) = POUNDN
      srcenth(2,I) = POUNDN
c
      READ(9,1010) TINP,TSRC
      READ(9,1010) tobs,TSRT
c
      read(9,*) oodist,avtime
c
      READ(9,*) U0,Z0,ZR,ML,USTAR
      read(9,*) K,G,RHOE,RHOA,DELTA
      read(9,*) BETA,GAMMAF,CcLOW
c
      READ(9,*) RM,SZM,EMAX,RMAX,TSC1
      read(9,*) ALEPH,TEND
c
      READ(9,*) CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
c
      READ(9,*) ALPHA
      alpha1 = alpha + 1.

```

```

c
read(9,1020) gas_name
read(9,*) gas_mw,gas_temp,gas_rhoe
read(9,*) gas_cpk,gas_cpp
read(9,*) gas_ufl,gas_lfl,gas_zsp
c
read(9,*) istab
read(9,*) tamb,pamb,humid
humsrc = 0.00
read(9,*) isofl,tsurf
read(9,*) ihtfl,htco
read(9,*) iwtfl,wtco
c
read(9,*) sigx_coeff,sigx_pow,sigx_min_dist
c
read(9,*) iphifl,dellay
c
H_masrte = 0.
if(isofl.eq. 0) read(9,*) H_masrte
c
READ(9,*) HTCUT, ce, delrhomin
c
1010  format(2(a24,1x))
1020  format(a3)
c
CLOSE(UNIT=9)
c
RETURN
END
#####

```

```

C.....  

C  

C      SUBROUTINE STRT2(OPNRUP,H_msrte,CCP)  

C  

C      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

C      INCLUDE 'sys$degadis:DEGADIS2.DEC/LIST'  

C  

C.....  

C  

C      BLOCK COMMON  

C  

C          COMMON  

C/TITL/ TITLE  

C/GEN2/ DEN(5,IGEN)  

C/ITI/ T1,TINP,TSRC,TOBS  

C/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

C/com_ss/ ESS,SLEN,SWID,OUTCc,OUTSZ,OUTB,OUTL,swcl,swal,snl,srhl  

C/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

C/com_gprop/ gas_mm,gas_temp,gas_rhoe,gas_cpk,gas_cpp,  

C/gas_ufl,gas_lfl,gas_zsp,gas_name  

C/comatm/ istab,tamb,pamb,humid,isoftl,tsurf,ihtfl,htco,iwtfl,wco,  

C/humsrc  

C/NEND/ POUNDN,POUND  

C/ALP/ ALPHA,alpha1  

C/phicom/ iphifl,dellay  

C/sprd_con/ ce, delrhomin  

C/COM_SURF/ HTCUT  

./oomsin/ oodist,avtime  

C  

C          character*80 TITLE(4)  

C          character*24 TSRC,TINP,TOBS  

C          character*40 OPNRUP  

C          character*3 gas_name  

C          character*4 pound  

C  

C          REAL*8 K,ML  

C          LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

C  

C          OPEN(UNIT=9,NAME=OPNRUP,TYPE='OLD')  

C  

C          DO 90 I = 1,4  

90      READ(9,1000) TITLE(I)  

1000     FORMAT(A80)  

C  

C          read(9,*) np  

C          do 100 i=1,np  

C          read(9,*) dummy1,dummy2,dummy3,PWC,PTEMP,PFRACV,PENTH,PRHO  

100      IF(I .EQ. 1) CCP = PWC*PRHO  

C  

C          READ(9,*) NP

```

```

      DO 120 I=1,NP
120  READ(9,*) DEN(1,I),DEN(2,I),den(3,i),den(4,i),den(5,i)
      I = NP + 1
      DEN(1,I) = 2.

c
      .
      read(9,*) np
      do 140 i=1,np
140  read(9,*) dummy1,dummy2,dummy3,dummy4,dum5,dum6,dum7
c
      read(9,1100) tinp,tsc
      read(9,1100) tobs,tsrt
1100 format(a24,1x,a24)
c
      read(9,*) oodist,avtime
c
      READ(9,*) U0,Z0,ZR,ML,USTAR
      Read(9,*) K,G,RHOE,RHOA,DELTA
      read(9,*) BETA,GAMMAF,CcLOW
c
      read(9,*) dummy1
      read(9,*) dummy1
c
      READ(9,*) CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
c
      READ(9,*) ALPHA
      alpha1 = alpha + 1.

c
      read(9,1200) gas_name
      read(9,*) gas_mw,gas_temp,gas_rhoe
      read(9,*) gas_cpK,gas_cpp
      read(9,*) gas_ufl,gas_lfl,gas_zsp
c
      read(9,*) istab
      read(9,*) tamb,pamb,humid
      humsrc = 0.00
      read(9,*) isofl,tsurf
      read(9,*) ihtfl,htco
      read(9,*) iwtfl,wtco
      read(9,*) dummy1

      READ(9,*) ESS,SLEN,SWID
      read(9,*) OUTCc,OUTSZ,OUTB,OUTL
      read(9,*) swcl,swal,senl,srhl
c
      read(9,*) iphifl,dellay
      h_masrte = 0.
      if(isofl.eq. 0) read(9,*) H_masrte
c
      READ(9,*) HTCUT, ce, delrhomin
      CLOSE(UNIT=9)
      RETURN

```

```

C.....  

C  

SUBROUTINE STRT3(OPNRUP)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

include 'sys$degadis:DEGADIS3.dec/list'  

C  

BLOCK COMMON  

C  

COMMON  

$/SSCON/ NREC(maxnob,2),T0(maxnob),XV(maxnob)  

$/GEN2/ DEN(5,igen)  

$/PARM/ UO,ZO,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,  

$ gas_ufl,gas_lfl,gas_zsp,gas_name  

$/ITI/ T1,TINP,TSRC,TOBS,TSRT  

$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

$/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag  

$/NEND/ POUND,POUND  

$/ALP/ ALPHA,alpha1  

$/CNOBS/ NOBS  

./oomsin/ oodist,avtime  

C  

character*3 gas_name  

character*40 OPNRUP  

character*24 TINP,TSRC,TOBS,TSRT  

C  

REAL*8 K,ML  

LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

C  

OPEN(UNIT=9,NAME=OPNRUP,TYPE='OLD')  

C  

READ(9,*) NOBS  

DO 125 I=1,NOBS  

125 READ(9,*) NREC(I,1),NREC(I,2),T0(I),XV(I)  

C  

READ(9,*) Npts  

DO 140 I=1,Npts  

140 READ(9,*) den(1,i),den(2,i),den(3,i),den(4,i),den(5,i)  

den(1,npts+1) = 2.  

C  

read(9,*) oodist,avtime  

C  

READ(9,*) UO,ZO,ZR,ML,USTAR  

read(9,*) K,G,RHOE,RHOA,DELTA  

read(9,*) BETA,GAMMAF,CcLOW

```

```

c
      READ(9,1010) TINP,TSRC
      read(9,1010) TOBS,TSRT
1010  format(2(a24,1x))

c
      READ(9,*) RM,SZM,EMAX,RMAX,TSC1
      read(9,*) ALEPH,TEND

c
      read(9,1020) gas_name
      read(9,*) gas_mw,gas_temp,gas_rhoe
      read(9,*) gas_cpk,gas_cpp
      read(9,*) gas_ufl,gas_lfl,gas_zsp
      read(9,*) istab
      read(9,*) tamb,pamb,humid
      humsrc = 0.00
      read(9,*) isofl,tsurf
      read(9,*) ihtfl,htco
      read(9,*) iwtfl,wtco
      read(9,*) sigx_coeff,sigx_pow,sigx_min_dist
1020  format(a3)

c
      READ(9,*) CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
      READ(9,*) ALPHA
      alpha1 = alpha + 1.

c
      CLOSE(UNIT=9)

c
      RETURN
      END
#####
```

```

C      Surface effects
C
C      SUBROUTINE Surface(temp,height,rho,mole,cp,yw,watrte,q rte)
C
C      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )
C
C      include 'sys$degadis:DEGADIS1.dec'
C
C      COMMON
C      $/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW
C      $/comatm/ istab,tamb,pamb,humid,isoftl,tsurf,ihtfl,htco,iwtfl,wtco,
C      $ humsrc
C      $/ALP/ ALPHA,alpha1
C      $/phicom/ iphifl,dellay
C      $/COM_SURF/ HTcut
C
C      REAL*8 ML,K
C      REAL*8 L,masrte,mole
C
C      vapor_p(txxx) = 6.0298e-3* exp(5407. *(1./273.15- 1./txxx))
C
C
C      wrte = 0.
C      q rte = 0.
C      if(isoftl.eq.1 .or. ihtfl.eq.0) return
C      if(height.le. htcut) return
C      delta_t = tsurf - temp
C
C      if(delta_t .lt. 0.) return
C      hhh = dmax1( height/2.00, zr)
C      top_vel = u0 * (hhh/z0)**alpha
C      prod_nat = ((rho/mole)**2 * abs(delta_t)) ** 0.333333
C      if(ihtfl .eq. 1) then           ! local correlation
C          hn = 18. * prod_nat
C          hf = 0.
C          hf = 1.22 * rho*cp * ustard**2/top_vel
C          ho = dmax1(hn,hf)
C      else if(ihtfl .eq. 2) then       ! LLNL correlation
C          ho = htco* rho* cp
C      else if(ihtfl .eq. 3) then       ! Colenbrander's method
C          av_temp= (tsurf + temp)/2.
C          hn = 89.* (delta_t/av_temp**2)**.333333
C          u10 = u0*(10./z0)**alpha
C          hf = 1.22 * rho*cp * ustard**2/u10
C          ho = dmax1(hn,hf)
C      else
C          ho = htco           ! ihtfl=-1

```

```
        endif
        qrte = ho * delta_t
        if(qrte .lt. 0.) qrte = 0.
c           ! since correlations are not valid for qrte<0.
        wrate = 0.
        if(iwtfl .eq. 0) return
        fo = wtco
        if(iwtfl .gt. 0) then
          fn = 9.9e-3 * prod_nat
          ff = 20.7 * ho /cp /mole
          fo = dmax1(fn,ff)
        endif
        wrate = min( vapor_p(temp), yw*pamb )
        wrate = fo * (vapor_p(tsurf) - wrate)/pamb
        if(wrate .le. 0.) wrate = 0.
c
        return
      end
#####
.
```

```

c
c      FUNCTION TO RETURN SZ0 CALCULATED over the source without
c      a blanket present underneath
c
c      NOTE: Uses the integration package RKGST and cannot be used
c            with any other routine without a local copy of RKGST.
c
c      subroutine SZF(Q,L,WCP,sz,cclay,wclay,rholay)
c
c      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )
c
c      external szlocal,szloco
c
c      REAL*8 L
c
c      include 'sys$degadis:DEGADIS1.dec'
c
c      COMMON
c      $/szfc/ szstp0,szerr,szstpmx,szszo
c
c      dimension Y(1),D(1),PRMT(17),aux(8,1)
c
c      prmt(1) = 0.
c      prmt(2) = L
c      prmt(3) = szstp0
c      prmt(4) = szerr
c      prmt(5) = szstpmx
c      prmt(6) = Q
c      PRMT(7) = WCP
c
c      Y(1) = 0.00      ! rho*dellay*u0*z0/(1.+alpha)*(sz/z0)**(1.+alpha)
c      D(1) = 1.00
c
c      ndim = 1
c
c      call rkgst(prmt,y,d,ndim,ihlf,szlocal,szloco,aux)
c
c      if(ihlf.ge. 10) call trap(3,ihlf)
c
c      cclay = prmt(13)
c      wclay = prmt(14)
c      rholay = prmt(15)
c      cc = prmt(16)
c      sz = prmt(17)
c
c      RETURN
c      END
c
c      subroutine szlocal(x,y,d,prmt)
c

```

```

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

dimension y(1),d(1),prmt(1)
c
common
$/parm/ u0,z0,zr,ml,ustar,k,g,rho,rhoa,delta,beta,gammaf,cclow
$/alp/ alpha,alpha1
$/phicom/ iphifl,dellay
c
real*8 ml,k
integer rho,lay/1/
c
Q = prmt(6)
WCP = PRMT(7)
c
c... for start up...
c
if(Y(rhouhlay) .le. 0.00) then
    wclay = 0.00
    rholay = rhoa
    cclay = 0.00
    cc     = 0.00
    rho    = rhoa
else
c
c... from the contaminant material balance over the source, determine WCLAY
c
wclay = Q*x/Y(rhouhlay)
c
call adiabat(1,wclay,wa,yc,ya,cclay,rholay,wm,enth,temp)
cc = cclay*dellay
call adiabat(0,wc,wa,yc,ya,cc,rho,wm,enth,temp) ! centerline
c
endif
c
uheff = Y(rhouhlay)/rholay/dellay
sz = ( uheff/u0/z0*(alpha1) )**(.1./alpha1) * z0
heff = gammaf*sz/alpha1
ristar = rif(rho,heff)
phi = phif(ristar,0.00)
wel = dellay * k*ustar*alpha1/phi
D(rhouhlay) = wel*rhoa + Q/WCP
c
prmt(8) = cclay
prmt(9) = wclay
prmt(10)= rholay
prmt(11)= cc
prmt(12) = sz
return
end
c

```

```
c
c      subroutine szloco(x,y, dery,ihlf,ndim, prmt)
c
c      Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )
c
c      dimension y(1), dery(1), prmt(1)
c
c      prmt(13) = prmt(8)
c      prmt(14) = prmt(9)
c      prmt(15) = prmt(10)
c      prmt(16) = prmt(11)
c      prmt(17) = prmt(12)
c      return
c      end
c
#####
```

```

subroutine tprop(ifl,wc,wa,enthalpy,yc,ya,wm,temp,rho,cp)
c
c      subroutine to return:
c          mole fractions (y's)
c          molecular weight (wm)
c          temperature (temp[=]K)
c          density (rho[=]kg/m**3)
c          heat capacity (cp[=]J/kg/K)
c
c      for a mixture from:
c          mass fractions (w's)
c          temperature (K)      for ifl.lt.0
c
c      for a mixture from:
c          mass fractions (w's)
c          enthalpy (J/kg)      for ifl.ge.0
c
c          adiabatic mixing of: emitted gas @ gas_temp
c                                entrained ambient humid air @ tamb
c                                entrained water from surface @ tsurf
c                                for ifl.eq.0 calculate and return
c
c          adiabatic lookup CALL ADIABAT
c              for isofl.eq.1 .or. ihtfl.eq.0.and.ifl.eq.1
c
c
Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

include 'sys$degadis:DEGADIS1.dec'
c
parameter (acrit=0.02D0)
c
common
$/GEN2/ DEN(5,igen)
$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$/comatm/ istab,tamb,pamb,humid,isoftl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc
./ctprop/ cwc,cwa,centh
c
character*3 gas_name

external enthal0
c
c*** data for air/water sys
c
data wma/28.96D0/      ! molecular weight of air
data wmw/18.02D0/      ! molecular weight of water
data rho_water/1000.D0/ ! liquid water density [=] kg/m**3
data cpa/1.0063D3/     ! heat capacity of air [=] J/kg/K
data cpw/1865.D0/       ! heat capacity of water vapor [=] J/kg/K

```

```

data dhvap/2.5023D6/      !latent heat of vap [=]J/kg water
data dhfus/0.33D6/        !latent heat of fus [=]J/kg water

data rgas/0.08205D0/     ! gas constant for P[=]atm, T[=]K
c
c
vapor_p(txxx) = 6.0298D-3* exp(5407.00 *(1.00/273.15D0- 1.00/txxx))
c
c
ww = 1.00-wc-wa
wm = 1.00/(wc/gas_mw + wa/wma + ww/wmw)
yc = wm/gas_mw *wc
ya = wm/wma *wa
yw = 1.00 - yc - ya
c
c
if(isofl.eq. 1) then
    call adiabat(1,wc,wa,yc,ya,cc,rho,wm,enthalpy,temp)
    return      ! interp density from wc
endif
c
c
if(ifl.eq. 0)
$      enthalpy = wc*cpc(gas_temp)*(gas_temp - tamb)
$      + (ww - wa*humid)*cpw*(tsurf - tamb)
c $      + wa*humid*cpw*(tamb - tamb)
c $      + wa*cpa*(tamb - tamb)           ! TR=tamb
c
c
if(ifl.eq. 1 .and. ihtfl.eq.0) then
    call adiabat(1,wc,wa,yc,ya,cc,rho,wm,enthalpy,temp)
    return      ! interp density from wc
endif
c
c
if(ifl .eq. -1) goto 400
c
c
tmin = dmin1(gas_temp, tsurf, tamb)
tmax = dmax1(gas_temp, tsurf, tamb)

elow = enthal(wc,wa,tmin)
if(enthalpy.lt. elow) then
    temp = tmin
    enthalpy = elow
    goto 400
endif

elow = enthal(wc,wa,tmax)
if(enthalpy.gt. elow) then

```

```

        temp = tmax
        enthalpy = elow
        goto 400
    endif
c
    cwc = wc
    cwa = wa
    centh = enthalpy

    call zbrent(temp, enthal0, tmin, tmax, acrit, ierr)
    if(ierr .ne. 0) call trap(24)
c
c
c
400  continue           ! density calculation
    vp = vapor_p(temp)
    ywsat = dmax1( vp/pamb, 0.00)
    wwsat = wmw/wm * ywsat * (1.00 - (yw-ywsat))
    conden = dmax1( 0.00, ww-wwsat)

    rho = 1.00/(      wa / (pamb*wma/rgas/temp)
                  + (ww-conden) / (pamb*wmw/rgas/temp)
                  + conden / rho_water
                  + wc * temp/gas_temp/gas_rho )
c
    tmin = temp + 10.
    if(tmin .gt. tmax0) tmin = temp - 10.
    if(tmin .lt. tmin0) tmin = temp + .1
c
    tmax = enthal(wc,wa,tmin)
    cp = (enthalpy - tmax)/(temp - tmin)
    if(cp .lt. cpa) cp = cpa           ! nominal value of air
c
    return
end

c
c
function cpc(temp)
c

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

common
$com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
c
data con/3.33D4/
c
character*3 gas_name
c
cpc = con + gas_cpk*gas_cpp * gas_temp**(gas_cpp-1.00)

```

```

if(temp .ne. gas_temp) then
    cpc = con + gas_cpk*
    (temp**gas_cpp + gas_temp**gas_cpp)/(temp- gas_temp)
    endif
cpc = cpc/gas_mm
return
end

c
c
function enthal(wc,wa,temp)      ! used by TPROP
c

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

parameter (delta=10.00)

c
common
$com_gprop/ gas_mm,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$comatm/ istab,tamb,pamb,humid,isoftl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc

c
character*3 gas_name
c

data cpa/1006.300/      ! heat capacity of air [=]J/kg/K
data cpw/1865.00/       ! heat capacity of water vapor [=]J/kg/K
data dhvap/2.502306/    !latent heat of vap [=]J/kg water
data dhfus/0.3306/      !latent heat of fus [=]J/kg water
data wma/28.9600/        ! molecular weight of air
data wmw/18.0200/        ! molecular weight of water

c
c
ww = 1.00-wa-wc
wm = 1.00/(wc/gas_mm + wa/wma + ww/wmw)
yw = ww * wm/wmw

vp = 6.02980-3* exp(5407.00 *(1.00/273.1500- 1.00/temp))

c
dh = dhvap
frac = 0.
if(temp .lt. 273.1500) frac = dmin1( (273.1500-temp)/delta,1.00)
dh = dhvap + dhfus*frac
ywsat = vp/pamb
wwsat = wmw/wm * ywsat * (1.00 - (yw-ywsat))
conden = dmax1( 0.00, ww-wwsat)

c
1000 enthal = wc*cpc(temp)*(temp - tamb)
$      - conden*dh
$      + ww*cpw*(temp - tamb)
$      + wa*cpa*(temp - tamb)      ! TR=tamb

```

```

c
      return
end

c
c
      function enthal0( temp )
c
      implicit real*8 (a-h, o-z), integer*4(i-n)
c
      include 'sys$degadis:degadisin.dec'

      common /ctprop/ cwc,cwa,centh

c
      enthal0 = centh - enthal(cwc,cwa,temp)
      return
end

c
c
      subroutine adiabat(ifl,wc,wa,yc,ya,cc,rho,wm,enthalpy,temp)

c
c
      subroutine to return:
      mass fractions (w's)
      mole fractions (y's)
      concentration (cc [=] kg/m**3)
      density (rho [=] kg/m**3)
      molecular weight (wm)
      enthalpy ([=] J/kg)
      temperature (temp [=] K)

c
c
      for a mixture from DEN lookup of adiabatic mixing calculation
      den(1,i)      mole fraction (yc)
      den(2,i)      concentration (cc [=] kg c/m**3)
      den(3,i)      mixture density (rho [=] kg mix/m**3)
      den(4,i)      mixture enthalpy (enthalpy [=] J/kg)
      den(5,i)      mixture temperature (temp [=] K)

c
c
      ifl indicates given information:
      -2)mole fraction (Yc) and assumption of constant gamma in enthalpy
      -1)concentration (cc) and assumption of constant gamma in enthalpy
      0) concentration (cc)
      1) mass fraction c (wc)
      2) mole fraction (Yc)
c

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

      include 'sys$degadis:DEGADISIN.dec'
c
      common
$/GEN2/ DEN(5,igen)

```

```

$/parm/ u0,z0,zr,ml,ustar,k,g,rhoe,rhoa,delta,beta,gammaf,cclow
$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc

c
      character*3 gas_name
      real*8 ml,k

c
c*** data for air/water sys
c
      data wma/28.96D0/           ! molecular weight of air
      data wmw/18.02D0/           ! molecular weight of water
c
c
      if(ifl.ne. 0) goto 1000
      ccl = cc
      if(cc .lt. 0.) ccl=0.
      i = 2
30      if(den(1,i) .gt. 1.) then
             i=i-1
             if(cc.gt. den(2,i)) ccl=den(2,i)
             goto 50
             endif
             if(cc.le. den(2,i)) goto 50      ! lookup in concentration
             i=i+1
             goto 30
50      slope = (den(3,i)-den(3,i-1)) / (den(2,i)-den(2,i-1)) ! interp in conc
      rho = (ccl - den(2,i-1))*slope + den(3,i-1)
      wcl = ccl / rho
      wc = wcl
      wa = (1.00-(1.00+humsrc)*wc)/(1.00+humid)
      ww = 1.00 - wa - wc
      wm = 1.00/(wc/gas_mw + wa/wma + ww/wmw)
      yc = wm/gas_mw * wc
      ya = wm/wma * wa
      goto 8000

c
c
1000   if(ifl.ne. -1) goto 1500
      ccl = cc
      if(ccl.lt. 0.) ccl=0.
      gamma = enthalpy
      wc = ccl/(rhoa+ccl*gamma)
      wa = (1.00-(1.00+humsrc)*wc)/(1.00+humid)
      ww = 1.00-wa-wc
      wm = 1.00/(wc/gas_mw + wa/wma + ww/wmw)
      yc = wm/gas_mw * wc
      ya = wm/wma * wa
      rho = ccl/wc
      return

```

```

c
c
1500  if(ifl.ne. -2) goto 1700
      ycl = yc
      if(ycl.lt. 0.) ycl=0.00
      gamma = enthalpy
      ya = (1.00-(1.00+gas_mw*humsrc/wmw)*ycl)/(1.00+humid*wma/wmw)
      yw = 1.00-ya-ycl
      wm = ycl*gas_mw + ya*wma + yw*wmw
      wc = gas_mw/wm * ycl
      wa = wma/wm * ya
      cc = wc*rhoa/(1.00 - gamma*wc)
      rho = cc/wc
      return

c
c
1700  if(ifl.ne. 2) goto 2000
      ycl = yc
      if(yc .lt. 0.) then
          ycl = 0.
          wa = 1.00/(1.00+humid)
          ww = 1.00-wa
          wm = 1.00/(wma/wa + wmw/ww)
          ya = wm/wma * wa
          endif
      if(yc .gt. 1.) then
          ycl = 1.00
          ya = 0.00
          endif
      i = 2
1730  if(den(1,i) .gt. 1.) then
          i = i-1
          goto 1750           ! extrapolate
          endif
      if(ycl.le. den(1,i)) goto 1750 ! lookup in mole frac
      i=i+1
      goto 1730
1750  continue
      wm = ycl*gas_mw + (1.00-ycl)*wma*wmw*(1.00+humid) / (wmw + wma*humid)
      wc = ycl*gas_mw / wm
      wa = (1.00-wc)/(1.00 + humid)
      ww = 1.00 - wc - wa

      slope = (den(3,i)-den(3,i-1)) / (den(2,i)-den(2,i-1)) ! interp in cc
      cc = wc*(den(3,i-1) - slope*den(2,i-1))/(1.00 - wc*slope)
      rho = cc/wc

      w1 = den(2,i-1)/den(3,i-1)
      w2 = den(2,i)/den(3,i)
      slope = (den(4,i)-den(4,i-1)) / (w2 - w1)           ! interp in w

```

```

        enthalpy = (wc - w1) *slope + den(4,i-1)
        slope = (den(5,i)-den(5,i-1)) / (w2 - w1)      ! interp in w
        temp = (wc - w1) *slope + den(5,i-1) .

c
        return
c
c
2000  if(ifl.ne. 1) goto 9000
        wcl = wc
        if(wc .lt. 0.) then
          wcl = 0.00
          wa = 1.00/(1.00+humid)
        endif
        if(wc .gt. 1.) then
          wcl = 1.00
          wa = 0.00
        endif
        ww = 1.00-wa-wcl
        wm = 1.00/(wcl/gas_mw + wa/wma + ww/wmw)
        yc = wm/gas_mw *wcl
        ya = wm/wma *wa
        i = 2
2030  if(den(1,i) .gt. 1.) then
          i = i-1
          goto 2050           ! extrapolate
        endif
        if(yc.le. den(1,i)) goto 2050   ! lookup in mole frac
        i=i+1
        goto 2030
2050  slope = (den(3,i)-den(3,i-1)) / (den(1,i)-den(1,i-1))
        rho = (yc-den(1,i-1))*slope + den(3,i-1)
        slope = (den(2,i)-den(2,i-1)) / (den(1,i)-den(1,i-1))
        cc = (yc-den(1,i-1))*slope + den(2,i-1)
        i = 2
2060  if(den(1,i) .gt. 1.) then
          i = i-1
          goto 8000           ! extrapolate
        endif
        cwc = den(2,i)/den(3,i)
        if(wcl.le. cwc) goto 8000       ! lookup in mass frac
        i=i+1
        goto 2060
c
c
8000      w1 = den(2,i-1)/den(3,i-1)
        w2 = den(2,i)/den(3,i)
        slope = (den(4,i)-den(4,i-1)) / (w2 - w1)      ! interp in w
        enthalpy = (wcl - w1) *slope + den(4,i-1)
        slope = (den(5,i)-den(5,i-1)) / (w2 - w1)      ! interp in w
        temp = (wcl - w1) *slope + den(5,i-1)
c

```

```

        return
c
9000  call trap(26)
      end
c
c      subroutine setenthal(h_masrte,h_airrte,h_watrte)
c
c      subroutine to load /com_ENTHAL/ through passed arguments if needed
c
c

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

include 'sys$degadis:DEGADISIN.dec'
c
      common
$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc
c
      character*3 gas_name
c
c*** data for air/water sys
c
      data cpa/1.0063D3/      ! heat capacity of air [=]J/kg/K
      data cpw/1865.D0/       ! heat capacity of water vapor [=]J/kg/K
c
      h_masrte = 0.00
      h_airrte = 0.00
      h_watrte = 0.00
c
      if(isofl.eq. 1) return
c
      h_masrte = cpc(gas_temp)*(gas_temp - tamb)      ! TR=tamb
c
c h_airrte = cpa*(tamb - tamb) = 0.
c
      if(iwatfl .eq. 0) return
      h_watrte = cpw*(tsurf - tamb)
c
      return
      end
c
c
c      subroutine setden(wc,wa,enthalpy)
c
c      subroutine to load /GEN2/ as needed
c

```

```

c      adiabatic mixing of:    WC
c                           WA
c                           WW @ specified enthalpy
c
c      with ambient humid air @ tamb
c
c      den(1,i)      mole fraction (yc)
c      den(2,i)      concentration (cc [=] kg c/m**3)
c      den(3,i)      mixture density (rho [=] kg mix/m**3)
c      den(4,i)      mixture enthalpy (enthalpy [=] J/kg)
c      den(5,i)      mixture temperature (temp [=] K)
c
c

      implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

      include 'sys$degadis:DEGADISIN.dec'
c
      parameter (tcrit=0.002D0, zero=1.D-20)
      parameter (iils=200, iis=iils-1, iback=25)
c
      common
$/GEN2/ DEN(5,igen)
$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc
c
      character*3 gas_name
c
      dimension curnt(5),backsp(5,iback)
c
c*** data for air/water sys
c
      data wma/28.96D0/           ! molecular weight of air
      data wmw/18.02D0/           ! molecular weight of water
      data cpa/1.0063D3/          ! heat capacity of air [=]J/kg/K
      data cpw/1865.D0/           ! heat capacity of water vapor [=]J/kg/K
c
c
c
      if(isofl.eq. 1) return
c
c
c
      k = 1
      den(1,k) = 0.000           ! yc
      den(2,k) = 0.000           ! cc
      den(3,k) = pamb*(1.D0+humid)/(.002833D0+.004553D0*humid)/tamb ! rhoa
      den(4,k) = 0.000           ! enthalpy of ambient air; TR=tamb
      den(5,k) = tamb
c

```

```

c
do 300 i= ils,1,-1
zbda = (float(i)/float(iils)) / (1.+humid)
zw = zbda*humid
zg = 1.00-zbda-zw

c
ermmix = zg*enthalpy + zbda*cpa*(tamb-tamb) + zw*cpw*(tamb-tamb) ! TR=tamb
ermmix = zg*enthalpy

c
zbda = zbda + zg*wa
zg = zg*wc
call tprop(2,zg,zbda,ermmix,yc,ya,wm,temp,rho,cp)
cc = zg*rho

c
c
curnt(1) = yc
curnt(2) = cc
curnt(3) = rho
curnt(4) = ermmix
curnt(5) = temp

c
if(i .eq. ils) then
    ind = 1
    do 150 jj= 1,5
150      backsp(jj,ind) = curnt(jj)
    goto 300
    endif

c
c      ADIABAT interpolation scheme
c
err = 0.
do 180 iind = 1,ind
yc = backsp(1,iind)
cc = backsp(2,iind)
rho = backsp(3,iind)
ermmix = backsp(4,iind)
temp = backsp(5,iind)
slope = (den(2,k)- curnt(2)) / (den(1,k)- curnt(1))
ccint = (yc - curnt(1))*slope + curnt(2)
err = dmax1(err,2.00* abs(cc - ccint)/(abs(cc + ccint) + zero))
slope = (den(3,k)- curnt(3)) / (den(1,k)- curnt(1))
rhoint = (yc - curnt(1))*slope + curnt(3)
err = dmax1(err,2.00* abs(rho - rhoint)/(abs(rho + rhoint) + zero))
wccal = cc / rhoint
w1 = curnt(2)/curnt(3)
w2 = den(2,k)/den(3,k)
slope = (den(4,k)- curnt(4)) / (w2 - w1)
entint = (wccal - w1)*slope + curnt(4)
err = dmax1(err,2.00* abs(ermmix - entint)/(abs(ermmix + entint) + zero))
slope = (den(5,k) - curnt(5)) / (w2 - w1)
temint = (wccal - w1)*slope + curnt(5)

```

```

      err = dmax1(err,2.D0* abs(temp - temint)/(abs(temp + temint) + zero))
180  continue
c
      if(err .le. tcrit) then
          if(ind .ge. iback) goto 200
          ind = ind + 1
          do 190 jj=1,5
190      backsp(jj,ind) = curnt(jj)
          goto 300
          endif
c
c      record a point in DEN
c
c
200  k = k+1
      if(k.ge. igen) call trap(28)
      do 250 jj=1,5
      den(jj,k) = backsp(jj,ind)
250      backsp(jj,1) = curnt(jj)
      ind = 1
c
300  continue
c
      k = k+1
      if(k.ge. igen) call trap(28)
      if(wc.eq. 1.D00) then
          den(1,k) = 1.D00      ! yc
          den(2,k) = gas_rhoe   ! cc
          den(3,k) = gas_rhoe   ! rhoe
          den(4,k) = enthalpy    ! enthalpy
          den(5,k) = gas_temp    ! temp
      else
          call tprop(2,wc,wa,enthalpy,den(1,k),ya,wm,den(5,k),den(3,k),cp)
          den(2,k) = wc*den(3,k)  ! cc
          den(4,k) = enthalpy
      endif
      den(1,k+1) = 2.           ! .gt. 1. end-of-record indicator
c
      return
end

c
c
c
c
      subroutine addheat(cc,dh,rho,temp,cp)
c

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

include 'sys$degadis:DEGADIS1.dec'
c

```

```

parameter (ascrit=0.02D0)

c
      common
$/GEN2/ DEN(5,igen)
$com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc
./ctprop/ cwc,cwa,centh

c
      character*3 gas_name

      external enthal0

c
c*** data for air/water sys
c
      data wma/28.96D0/      ! molecular weight of air
      data wmw/18.02D0/      ! molecular weight of water
      data rho_water/1000.D0/ ! liquid water density [=] kg/m**3
      data cpa/1.0063D3/     ! heat capacity of air [=] J/kg/K
      data cpw/1865.D0/      ! heat capacity of water vapor [=] J/kg/K
      data dhvap/2.5023D6/   !latent heat of vap [=] J/kg water
      data dhfus/0.33D6/     !latent heat of fus [=] J/kg water

      data rgas/0.08205D0/   ! gas constant for P[=]atm, T[=]K

c
c
c
      vapor_p(txxx) = 6.0298D-3* exp(5407.D0 *(1.D0/273.15D0- 1.D0/txxx))

c
c
      cp = cpa
      rhoa = den(3,1)

c
      call adiabat(0,wc,wa,yc,ya,cc,rho,wm,enthalpy,amt)
      mw = 1.D0 - wc - wa
      yw = 1.D0 - yc - ya
      temp = amt
      IF(isofl.eq.1 .or. ihtfl.eq.0) return ! adiabatic mixing is valid
      if(dh.le. 0.) return           ! catch colder surface temperatures
      enthalpy = enthalpy + dh

c
c
      if(enthalpy.gt. 0.) then
          temp = tamb
          goto 400
      endif

c
c
      tmin = amt      ! adiabatic mixing temp

```

```

tmax = dmax1(gas_temp, tsurf, tamb)
ehi = enthal(wc, wa, tmax)
if(enthalpy.gt. ehi) then
    temp = tmax
    goto 400
endif

cwc = wc
cwa = wa
centh = enthalpy

c
call zbrent(temp, enthal0, tmin, tmax, acrit, ierr)
c
c... If there are troubles, get LIMIT to help
c
if(ierr .ne. 0) then
    tinc = (tmin+tmax)/100.00
    temp = tmin
    tmin = tmin/2.00
    tmax = 2.00*tmax
    call limit(enthal0, temp, tinc, tmax, tmin)
    call zbrent(temp, enthal0, tmin,tmax,acrit,ierr)
    if(ierr .eq. 0) goto 400
    call trap(17)
endif
c
c
400  continue           ! density calculation
vp = vapor_p(temp)
ywsat = dmax1( vp/pamb, 0.00)
wwsat = wmw/wm * ywsat * (1.00 - (yw-ywsat))
conden = dmax1( 0.00, ww-wwsat)

rho = 1.00/(      wa / (pamb*wma/rgas/temp)
               + (ww-conden) / (pamb*wmw/rgas/temp)
               + conden / rho_water
               + wc * temp/gas_temp/gas_rho )
if(temp.ne.amt) cp = dmax1(dh/(temp-amt),cpa)
c
c
return
end
#####

```

```

C.....  

C  

C      FILE NAME TRANS1 -- FOR USE IN DEGADIS1  

C  

C.....  

C  

C      SUBROUTINE TRANS(FILE)  

C  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

C  

include 'sys$degadis:DEGADIS1.dec'  

C  

C      BLOCK COMMON  

C  

COMMON  

$/GEN3/ radg(2,maxl),qstr(2,maxl),srcden(2,maxl),srcwc(2,maxl),  

$ srcwa(2,maxl),srcenth(2,maxl)  

$/TITL/TITLE  

$/GEN1/ PTIME(igen), ET(igen), R1T(igen), PWC(igen), PTEMP(igen),  

$ PFRACV(igen), PENTH(igen), PRHO(igen)  

$/GEN2/ DEN(5,igen)  

$/ITI/T1,TINP,TSRC,TOBS,TSRT  

$/ERROR/STPIN,ERBND,STPMX,WTRG,WTtm,WTya,WTyc,WTeb,WTmb,WTuh,XLI,  

$ XRI,EPS,ZLOW,STPINZ,ERBNDZ,STPMXZ,SRCOER,srcss,srccut,  

$ htcut,ERNOBL,NOBLpt,crfger,epsilon  

$/PARM/U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,  

$ gas_ufi,gas_lfl,gas_zsp,gas_name  

$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$/com_ss/ ess,slen,swid,outcc,outsz,outb,outl,swcl,swal,sent,srhl  

$/PHLAG/CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

$/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag  

$/com_enthal/ h_masrte,H_airrte,H_watrte  

$/NEND/ POUNDN,POUND  

$/ALP/ ALPHA,alpha1  

$/phicom/ iphifl,dellay  

$/sprd_con/ ce, delrhomin  

$/COM_SURF/ HTCUTS  

./oomsin/ oodist,avtime  

C  

character*80 TITLE(4)  

C  

character*4 pound  

character*24 TSRC,TINP,TOBS,TSRT  

character*3 gas_name  

C  

REAL*8 ML,K  

LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5

```

```

c
      character*(*) file
c
      OPEN(UNIT=8,NAME=FILE,TYPE='NEW',
$  carriagecontrol='list',
$  recordtype='variable')
c
      WRITE(8,1000) (TITLE(I),I=1,4)
1000 FORMAT(A80)
c
      DO 100 I=1,igen
100   IF(PTIME(I).EQ.POUNDN) GO TO 105
      write(6,*) ' POUND WAS NOT DETECTED '
105   NP = I - 1
      WRITE(8,1040) NP
      DO 110 I=1,NP
110   WRITE(8,1030) PTIME(I),ET(I),R1T(I), PWC(I),PTEMP(I),
$                 PFRAVC(I), PENTH(I), PRHO(I)
c
      DO 120 I=1,igen
120   IF(DEN(1,I) .gt. 1.) GOTO 125
      DO 122 I=1,igen
122   WRITE(8,1060) DEN(1,I),DEN(2,I),den(3,i),den(4,i),den(5,i)
      write(6,*) ' density function blew the loop'
125   NP = I - 1
      WRITE(8,1040) NP
      DO 130 I=1,NP
130   WRITE(8,1060) DEN(1,I),DEN(2,I),den(3,i),den(4,i),den(5,i)
c
      DO 140 I=1,maxl
c      cc = srcwc(2,i)*srcden(2,i)
c      if(cc.lt. ccclow) then
c          fcc = 0.
c          do ii=i+1,maxl
c              fcc = amax1(srcwc(2,ii)*srcden(2,ii),fcc)
c          enddo
c          if(fcc.ge. cc) goto 140
c          np = i
c          tend = srcwc(1,i)
c          goto 146
c      endif
140   IF(radg(1,I).EQ.POUNDN .AND. radg(2,I).EQ.POUNDN) GO TO 145
      write(6,*) ' POUND WAS NOT DETECTED '
145   NP = I - 1
146   WRITE(8,1040) NP
      DO 150 I=1,NP
150   WRITE(8,1060) radg(1,i),radg(2,i),qstr(2,i),srcden(2,i),srcwc(2,i)
1           ,srcwa(2,i),srcenth(2,i)
c
      1020  format(1x,i4,1x,1pg14.7)
      1030  format(8(1x,1pg14.7))

```

```

1040  format(1x,i4)
1050  format(2(a24,1x))
1060  format(1x,1pg23.16,7(1x,1pg14.7))
1070  format(1x,1pg14.7)
1080  format(a3)

c
      WRITE(8,1050) TINP,TSRC
      write(8,1050) TOBS,TSRT
      write(8,1030) oodist,avtime
      WRITE(8,1060) U0,Z0,ZR,ML,USTAR
      write(8,1060) K,G,RHOE,RHOA,DELTA
      write(8,1030) BETA,GAMMAF,CcLOW
      WRITE(8,1060) RM,SZM,EMAX,RMAX,TSC1
      write(8,1030) ALEPH,TEND
      WRITE(8,*) CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
      WRITE(8,1070) ALPHA
      write(8,1080) gas_name
      write(8,1030) gas_mw,gas_temp,gas_rhoe
      write(8,1030) gas_cpk,gas_cpp
      write(8,1030) gas_ufl,gas_lfl,gas_zsp
      write(8,1040) istab
      write(8,1030) tamb,pamb,humid
      write(8,1020) isofl,tsurf
      write(8,1020) ihtfl,htco
      write(8,1020) iwtfl,wtco
      write(8,1030) sigx_coeff,sigx_pow,sigx_min_dist

c
      if(check4) then
          write(8,1030) ess,slen,swid
          write(8,1060) outcc,outsz,outb,outl
          write(8,1060) swcl,swal,senl,srhl
          end if

c
      write(8,1020) iphifl,dellay
c
      if(isofl.eq. 0) write(8,1030) H_masrte
c
      WRITE(8,1030) HTCUTS, ce, delrhomin
c
      CLOSE(UNIT=8)
c
      RETURN
      END
#####

```

```

C.....  

C  

C      FILE NAME TRANS2 -- USE WITH DEGADIS2  

C  

C      SUBROUTINE TRANS(FILE)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

include 'sys$degadis:DEGADIS2.dec'  

C  

COMMON  

$/SSCON/ NREC(maxnob,2),T0(maxnob),XV(maxnob)  

$/GEN2/ DEN(5,igen)  

$/ITI/ T1,TINP,TSRC,TOBS,TSRT  

$/PARM/ UO,ZO,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW  

$/com_gprop/ gas_mm,gas_temp,gas_rhoe,gas_cpk,gas_cpp,  

$ gas_ufl,gas_lfl,gas_zsp,gas_name  

$/comatm/ istab,tamb,pamb,humid,isofl,tsurf,ihtfl,htco,iwtfl,wtco,  

$ humsrc  

$/PARMSC/ RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

$/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag  

$/nend/ poundn,pound  

$/ALP/ ALPHA,alpha1  

$/CNOBS/ NOBS  

./oomsin/ oodist,avtime  

C  

character*3 gas_name  

character*80 TITLE(4)  

character*24 TINP,TSRC,TOBS,TSRT  

character*(*) file  

C  

REAL*8 K,ML  

LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5  

C  

OPEN(UNIT=9,NAME=FILE,TYPE='NEW',  

$ carriagecontrol='list',  

$ recordtype='variable')  

C  

WRITE(9,1040) NOBS  

DO 125 I=1,NOBS  

125 WRITE(9,1010) NREC(I,1),NREC(I,2),T0(I),XV(I)  

C  

DO 140 I=1,igen  

140 IF(DEN(1,I).gt. 1.) GOTO 145  

      write(6,*) ' density function error in TRANS'  

145  NP = I - 1  

      WRITE(9,1040) NP  

      DO 150 I=1,NP  

150  WRITE(9,1060) DEN(1,I),DEN(2,I),den(3,i),den(4,i),den(5,i)

```

```

C           write(9,1060) oodist,avtime
C
C           WRITE(9,1060) U0,Z0,ZR,ML,USTAR
C           write(9,1060) K,G,RHOE,RHOA,DELTA
C           write(9,1030) BETA,GAMMAF,CcLOW
C
C           WRITE(9,1050) TINP,TSRC
C           write(9,1050) TOBS,TSRT
C
C           WRITE(9,1060) RM,SZM,EMAX,RMAX,TSC1
C           write(9,1020) ALEPH,TEND
C
C           write(9,1080) gas_name
C           write(9,1030) gas_mw,gas_temp,gas_rhoe
C           write(9,1020) gas_cpK,gas_cpp
C           write(9,1030) gas_ufl,gas_lfl,gas_zsp
C           write(9,1040) istab
C           write(9,1030) tamb,pamb,humid
C           write(9,1025) isofl,tsurf
C           write(9,1025) ihtfl,htco
C           write(9,1025) iwtfl,wtco
C           write(9,1030) sigx_coeff,sigx_pow,sigx_min_dist
C
C           WRITE(9,*) CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
C
C           WRITE(9,1070) ALPHA
C
C
1010      format(1x,i8,1x,i8,2(1x,1pg14.7))
1020      format(2(1x,1pg14.7))
1025      format(1x,i4,1x,1pg14.7)
1030      format(3(1x,1pg14.7))
1040      format(1x,i4)
1050      format(2(a24,1x))
1060      format(5(1x,1pg14.7))
1070      format(1x,1pg14.7)
1080      format(a3,1x)
C
C           CLOSE(UNIT=9)
C           RETURN
C           END
#
#####

```

```

C.....  

C  

C      FILE NAME TRANS2 -- USE WITH SDEGADIS2  

C  

C.....  

C  

C      SUBROUTINE TRANS(FILE)  

C  

C  

C  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

COMMON
$/PARM/ U0,Z0,ZR,ML,USTAR,K,G,RHOE,RHOA,DELTA,BETA,GAMMAF,CcLOW
$/com_gprop/ gas_mw,gas_temp,gas_rhoe,gas_cpk,gas_cpp,
$ gas_ufl,gas_lfl,gas_zsp,gas_name
$/comatm/ istab,tamb,pamb,humid,isoftl,tsurf,ihtfl,htco,iwtfl,wtco,
$ humsrc
$/ITI/ t1,TINP,TSRC,TOBS
$/PHLAG/CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
$/ALP/ALPHA,alpha1
C
      character*24 TSRC,TINP,TOBS
      character*3 gas_name
      character(*) file
C
      REAL*8 K,ML
      LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
C
      OPEN(UNIT=9,NAME=FILE,TYPE='NEW')
C
      WRITE(9,1060) U0,Z0,ZR,ML,USTAR
      write(9,1060) K,G,RHOE,RHOA,DELTA
      write(9,1030) BETA,GAMMAF,CcLOW
C
      WRITE(9,1050) TINP,TSRC
      write(9,1050) TOBS
C
      write(9,1080) gas_name
      write(9,1030) gas_mw,gas_temp,gas_rhoe
      write(9,1020) gas_cpk,gas_cpp
      write(9,1030) gas_ufl,gas_lfl,gas_zsp
C
      write(9,1040) istab
      write(9,1030) tamb,pamb,humid
      write(9,1025)isoftl,tsurf
      write(9,1025) ihtfl,htco
      write(9,1025) iwtfl,wtco
C

```

```
      WRITE(9,*) CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
      WRITE(9,1070) ALPHA
C
      CLOSE(UNIT=9)
C
1020  format(2(1x,1pg14.7))
1025  format(1x,i4,1x,1pg14.7)
1030  format(3(1x,1pg14.7))
1040  format(1x,i4)
1050  format(2(a24,1x))
1060  format(5(1x,1pg14.7))
1070  format(1x,1pg14.7)
1080  format(a3,1x)
C
      RETURN
      END
#####
2 .. sys$degadis:trans2ss.for          6-SEP-1989 18:36:57
```

```

C      FILE NAME TRANS3 FOR USE WITH DEGADIS3
C
C.....-----
C
C      SUBROUTINE TRANS(OPNRUP)
C

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

include 'sys$degadis:DEGADIS3.dec/list'

C
COMMON /SORT/TCc(maxnob,maxnt),TCcSTR(maxnob,maxnt),
$ Tyc(maxnob,maxnt),Trho(maxnob,maxnt),
$ Tgamma(maxnob,maxnt),Ttemp(maxnob,maxnt),
$ TSY(maxnob,maxnt),TSZ(maxnob,maxnt),TB(maxnob,maxnt),
$ TDIST0(maxnob,maxnt),TDIST(maxnob,maxnt),KSUB(maxnt)
$/SORTIN/TIM(maxnt),NTIM,ISTRT
$/ITI/T1,TINP,TSRC,TOBS,TSRT

C
character*24 tinp,tsrc,tobs,tsrt

C
character*(*) OPNRUP

C
T0 = TIM(ISTRT)
DT = TIM(ISTRT+1) - TIM(ISTRT)

C
OPEN(UNIT=9,NAME=OPNRUP,TYPE='NEW',
$ carriagecontrol='list',recordtype='variable')

C
C
DO 110 I = ISTRT,NTIM
II = I - ISTRT + 1
110 KSUB(II) = KSUB(I)
NTIM = NTIM - ISTRT + 1
IF(NTIM .EQ. maxnt) GO TO 120
II = NTIM + 1
DO 115 I=II,maxnt
115 KSUB(I) = 0
C
120 CONTINUE
C
WRITE(9,*) T0,DT,NTIM
WRITE(9,*) KSUB
C
CLOSE(UNIT=9)
C
RETURN
END
#####

```

```

C.....  

C  

C      SUBROUTINE TRAP -- DIAGNOSTICS  

C  

C      SUBROUTINE trap(N,N1)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

include 'sys$degadis:DEGADIS2.dec'  

C  

COMMON /ITI/T1,TINP,TSRC,TOBS,TSRT  

real*4 tt1  

C  

character*24 TINP,TSRC,TOBS,TSRT  

C  

character*24 tt  

character*80 dd  

C  

WRITE(lunlog,1100)  

WRITE(lunlog,1110)  

write(lunlog,1115) n  

C  

c*** check to see if the operator is ready to read the text of the error  

c*** message  

C  

        write(lunlog,1000)  

irtn = lib$get_command( dd )      ! get a line from the terminal  

10      write(lunlog,1002)  

C  

IF(N .EQ. 1 ) then  

        WRITE(lunlog,2010) N1  

        WRITE(lunlog,2011)  

else IF(N .EQ. 2 ) then  

        WRITE(lunlog,2020)  

else IF(N .EQ. 3 ) then  

        WRITE(lunlog,2030) N1  

else IF(N .EQ. 4 ) then  

        WRITE(lunlog,2040)  

else IF(N .EQ. 5 ) then  

        WRITE(lunlog,2050)  

else IF(N .EQ. 6 ) then  

        WRITE(lunlog,2060)  

else IF(N .EQ. 7 ) then  

        WRITE(lunlog,2070)  

else IF(N .EQ. 8 ) then  

        WRITE(lunlog,2080) N1  

        WRITE(lunlog,2081)  

else IF(N .EQ. 9 ) then  

        WRITE(lunlog,2090) N1

```

```

        WRITE(lunlog,2091)
else IF(N .EQ. 10) then
        WRITE(lunlog,2100) N1
        WRITE(lunlog,2101)
endif
IF(N .EQ. 11) WRITE(lunlog,2110)
IF(N .EQ. 12) WRITE(lunlog,2120)
IF(N .EQ. 13) WRITE(lunlog,2130)
IF(N .EQ. 14) WRITE(lunlog,2140)
IF(N .EQ. 15) WRITE(lunlog,2150)
IF(N .EQ. 16) WRITE(lunlog,2160)
IF(N .EQ. 17) WRITE(lunlog,2170)
IF(N .EQ. 18) then
        WRITE(lunlog,2180) N1
        WRITE(lunlog,2181)
endif
IF(N .EQ. 19) WRITE(lunlog,2190) N1
IF(N .EQ. 20) WRITE(lunlog,2200)
IF(N .EQ. 21) WRITE(lunlog,2210)
IF(N .EQ. 22) WRITE(lunlog,2220)
IF(N .EQ. 23) then
        WRITE(lunlog,2230) maxnob
        WRITE(lunlog,2231)
endif
IF(N .EQ. 24) WRITE(lunlog,2240)
IF(N .EQ. 25) WRITE(lunlog,2250)
IF(N .EQ. 26) WRITE(lunlog,2260)
IF(N .EQ. 27) WRITE(lunlog,2270)
IF(N .EQ. 28) WRITE(lunlog,2280)
IF(N .EQ. 29) WRITE(lunlog,2290)
IF(N .EQ. 30) WRITE(lunlog,2300)
IF(N .EQ. 31) WRITE(lunlog,2310)
IF(N .EQ. 32) WRITE(lunlog,2320)
IF(N .EQ. 33) WRITE(lunlog,2330)

C
1000 format(//,5x,'Ready for the text of the error message? <CR> ',$,)
1002 format(/)
1100 FORMAT(5X,'The best laid plans of mice and men...')
1110 FORMAT(5X,'You have entered a TRAP -- the land of no RETURN.')
1115 format(' Code: ',i4)

2010 FORMAT(5X,'DEGADIS1? Source integration has returned IHLF=',I3,/,
  ./,' This error occurs during integration of the equations',/,
  .' which describe the gas source. IHLF is an error code',/,
  .' returned by the integration package RKGST.',/,
  .' When IHLF=11, more than 10 bisections of the initial',/,
  .' increment of the independent variable were necessary to make',/,
  .' an integration step within the specified error. Reduce the',/,
  .' initial step size of the independent variable',/,
  .' (STPIN in the ER1 file). If this does not work,',/,
  .' it will be necessary to either increase the error criteria',/

```

```

.' for all of the dependent variables being integrated',//,
.' (ERBND in the ER1 file) or increase the error criteria',//,
.' for the variable violating the criteria by decreasing the',//,
.' error weight for that variable (one of the following: WTRG',//,
.' WTTM, WTYA, WTYC, WTEB, WTMB, or WTUH in the ER1 file).',//)
2011 format(
    .' When IHLF=12, the initial increment of the independent',//,
    .' variable (STPIN) is 0. Correct the ER1 file and execute the',//,
    .' program again',//,
    .' When IHLF=13, the initial increment of the independent',//,
    .' variable (STPIN) is not the same sign as the difference',//,
    .' between the upper bound of the interval and the lower bound',//,
    .' of the interval. STPIN must be positive. Correct the ER1',//,
    .' file and execute the program again.//)

2020 FORMAT(5X,'Reserved')

2030 format(5x,'SZF? Local integration failed; IHLF=',//,
    .' This error occurs during estimation of SZ over the',//,
    .' source when no gas is present. IHLF is an error code',//,
    .' returned by the integration package RKGST.',//,
    .' When IHLF=11, more than 10 bisections of the initial',//,
    .' increment of the independent variable were necessary to make',//,
    .' an integration step within the specified error. Reduce the',//,
    .' initial step size of the independent variable',//,
    .' (SZSTPO in the ER1 file). If this does not work',//,
    .' increase the error criteria for all of the dependent',//,
    .' variables being integrated (SZERR in the ER1 file).',//,
    .' When IHLF=12, the initial increment of the independent',//,
    .' variable (SZSTPO) is 0. Correct the ER1 file and execute the',//,
    .' program again.//,
    .' When IHLF=13, the initial increment of the independent',//,
    .' variable (SZSTPO) is not the same sign as the difference',//,
    .' between the upper bound of the interval and the lower bound',//,
    .' of the interval. SZSTPO must be positive. Correct the ER1',//,
    .' file and execute the program again.//)

2040 format(5x,'SURFACE? Negative QRTE for positive DELTA_T',//,
    .' This is a diagnostic message indicating an error in',//,
    .' estimation of the heat capacity. Check the input',//,
    .' to the model and execute the program again.//)

2050 FORMAT(5X,'CRFG? MORE POINTS FOR GEN3 WERE NEEDED',//,
    .' The COMMON area /GEN3/ stores representative values',//,
    .' of the calculated source parameters. If this message',//,
    .' occurs, relax the CRFG error criteria (CRFGER) in the',//,
    .' ER1 file. If this is a common problem, the length of the',//,
    .' /GEN3/ vectors can be increased by changing the value of',//,
    .' MAXL in DEGADIS1.DEC and reinstalling DEGADIS.//)

2060 FORMAT(5X,'TUPF? OBSERVER CALCULATIONS -- TUPF FAILED',//,

```

```

.'      The trial and error search associated with finding the',/,
.' upwind edge of the gas source for an observer failed.',/,
.' Often, this problem can be avoided by adding one or two',/,
.' additional observers to the present number of observers',/,
.' (which changes the conditions for the trial and error).',/,
.' Another possibility is to increase the error criteria for',/,
.' this function (ERTUPF) in the ER2 file.',//)

2070  FORMAT(5X,'TUPF? OBSERVER CALCULATIONS -- TDNF FAILED',/,
.'      The trial and error search associated with finding the',/,
.' downwind edge of the gas source for an observer failed.',/,
.' Often, this problem can be avoided by adding one or two',/,
.' additional observers to the present number of observers',/,
.' (which changes the conditions for the trial and error).',/,
.' Another possibility is to increase the error criteria for',/,
.' this function (ERTDNF) in the ER2 file.',//)

2080  FORMAT(5X,'SSSUP? OBSERVER INTEGRATION FAILED, IHLF=',I3,/,
.'      This error occurs during integration of the five',/,
.' differential equations which average the source for each',/,
.' observer. IHLF is an error code returned by the',/,
.' integration package RKGST.',/,,
.'      When IHLF=11, more than 10 bisections of the initial',/,
.' increment of the independent variable were necessary to make',/,
.' an integration step within the specified error. Reduce the',/,
.' initial step size of the independent variable',/,
.' (STPO in the ER2 file). If this does not work',/,,
.' it will be necessary to either increase the error criteria',/,
.' for all of the dependent variables being integrated',/,
.' (ERRO in the ER2 file) or increase the error criteria',/,
.' for the variable violating the criteria by decreasing the',/,
.' error weight for that variable (one of the following: WTAIO',/,
.' WTQOO, or WTSZ0 in the ER2 file).',/)

2081  format(
.'      When IHLF=12, the initial increment of the independent',/,
.' variable (STPO) is 0. Correct the ER2 file and execute the',/,
.' program again.',/,,
.'      When IHLF=13, the initial increment of the independent',/,
.' variable (STPO) is not the same sign as the difference',/,
.' between the upper bound of the interval and the lower bound',/,
.' of the interval. STPO must be positive. Correct the ER2',/,
.' file and execute the program again.',//)

2090  FORMAT(5X,'SSSUP/SDEGADIS2? PSEUDO-STEADY INTEG FAILED, IHLF=',I3,
.'//,'      This error occurs during integration of the four',/,
.' differential equations describing the portion of the',/,
.' downwind calculation when b>0. The routine calling TRAP is',/,
.' SSSUP if a transient simulation is being executed; if a',/,
.' steady state simulation is being executed, the calling',/,
.' routine is SDEGADIS2. IHLF is an error code returned by the',/,
.' integration package RKGST.',//)

```

```

.' When IHLF=11, more than 10 bisections of the initial',/,
.' increment of the independent variable were necessary to make',/,
.' an integration step within the specified error. Reduce the',/,
.' initial step size of the independent variable',/,
.' (STPP in the ER2 file). If this does not work,',/,
.' it will be necessary to either increase the error criteria',/,
.' for all of the dependent variables being integrated',/,
.' (ERRP in the ER2 file) or increase the error criteria',/,
.' for the variable violating the criteria by decreasing the',/,
.' error weight for that variable (one of the following: WTSZP',/,
.' WTSYP, WTBEPE, or WTDH in the ER2 file).',/)

2091 format(
.' When IHLF=12, the initial increment of the independent',/,
.' variable (STPP) is 0. Correct the ER2 file and execute the',/,
.' program again.',//,
.' When IHLF=13, the initial increment of the independent',/,
.' variable (STPP) is not the same sign as the difference',/,
.' between the upper bound of the interval and the lower bound',/,
.' of the interval. STPP must be positive. Correct the ER2',/,
.' file and execute the program again.',//)

2100 FORMAT(5X,'SSSUP/SDEGADIS2? GAUSSIAN INTEGRATION FAIL, IHLF=',I3,
.//,' This error occurs during integration of the',/,
.' differential equations describing the portion of the',/,
.' downwind calculation when b=0. The routine calling TRAP is',/,
.' SSSUP if a transient simulation is being executed; if a',/,
.' steady state simulation is being executed, the calling',/,
.' routine is SDEGADIS2. IHLF is an error code returned by the',/,
.' integration package RKGST.',//,
.' When IHLF=11, more than 10 bisections of the initial',/,
.' increment of the independent variable were necessary to make',/,
.' an integration step within the specified error. Reduce the',/,
.' initial step size of the independent variable',/,
.' (STPG in the ER2 file). If this does not work,',/,
.' it will be necessary to either increase the error criteria',/,
.' for all of the dependent variables being integrated',/,
.' (ERRG in the ER2 file) or increase the error criteria',/,
.' for the variable violating the criteria by decreasing the',/,
.' error weight for that variable (either WTRUH or WTDHG',/,
.' in the ER2 file).',/)

2101 format(
.' When IHLF=12, the initial increment of the independent',/,
.' variable (STPG) is 0. Correct the ER2 file and execute the',/,
.' program again.',//,
.' When IHLF=13, the initial increment of the independent',/,
.' variable (STPG) is not the same sign as the difference',/,
.' between the upper bound of the interval and the lower bound',/,
.' of the interval. STPG must be positive. Correct the ER2',/,
.' file and execute the program again.',//)

2110 FORMAT(5X,'SSSUP/SDEGADIS2? TOTAL No. OF RECORDS EXCEED 120000',

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.//,' This is an arbitrary stopping point for the process',//,
.' in order to keep a runaway simulation from filling up disk',//,
.' space. Relax the output specifications (ODLP, ODLLP, ODLG,'//,
.' or ODLLG) in the ER2 file in order to generate less output',//,
.' if the input parameters are valid.'//)

2120 FORMAT(5X,'Reserved')
2130 FORMAT(5X,'Reserved')
2140 FORMAT(5X,'Reserved')
2150 FORMAT(5X,'Reserved')

2160 FORMAT(5X,'PSSOUT/PSSOUTSS? PSS STARTED WITH B<0.'//,
.' This condition is checked at the beginning of the',//,
.' downwind calculation in order to confirm proper handling of',//,
.' the movement to the Gaussian phase of the downwind',//,
.' calculation. Check the initial conditions and execute the',//,
.' program again.'//)

2170 format(5x,'TPROP/ADDHEAT? Enthalpy out of bounds',//,
.' Diagnostic message indicates an enthalpy lower',//,
.' than the adiabatic mixing enthalpy was passed to ADDHEAT.'//,
.' Check the input conditions and execute the program again.'//)

2180 FORMAT(5X,'ALPH? ALPHA INTEGRATION FAILED, IHLF=',I3,//,
.' The integration which determines the integral least',//,
.' squares fit for ALPHA has failed. Note that small values',//,
.' of the Monin-Obukhov length ( ML < 0(1m) ) in combination',//,
.' with stable atmospheric conditions may cause this failure.'//,
.' IHLF is an error code returned by the integration package',//,
.' RKGST.'//,
.' When IHLF=11, more than 10 bisections of the initial',//,
.' increment of the independent variable were necessary to make',//,
.' an integration step within the specified error. Reduce the',//,
.' absolute value of the initial step size of the independent',//,
.' variable (STPINZ in the ER1 file). If this does not work',//,
.' it will be necessary to increase the error criteria',//,
.' (ERBNRZ in the ER1 file).'//)

2181 format(
.' When IHLF=12, the initial increment of the independent',//,
.' variable (STPINZ) is 0. Correct the ER1 file and execute',//,
.' the program again.'//,
.' When IHLF=13, the initial increment of the independent',//,
.' variable (STPINZ) is not the same sign as the difference',//,
.' between the upper bound of the interval and the lower bound',//,
.' of the interval. STPINZ must be negative. Correct the ER1',//,
.' file and execute the program again. This error will also',//,
.' occur if the surface roughness ZR is greater than the',//,
.' reference height Z0.'//)

2190 FORMAT(5X,'ALPH? ZBRENT has failed to locate ALPHA IERR: ',I4,//,
.' The search procedure which determines ALPHA has failed.'//,

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```

.' This error may be the result of an unusual velocity',//,
.' specification such as small values of the Monin-Obukhov',//,
.' length ( ML < 0(1.m) ) or small reference heights',//,
.' ( Z0 < 0(10. * ML) ). IERR is an error code returned by',//,
.' the routine ZBRENT',//,
.' When IERR=1, the search for ALPHA failed after a',//,
.' specified number of iterations. Increase the error bound',//,
.' used by ZBRENT (EPS in ER1 file).',//,
.' When IERR=2, the basic assumption that the function which',//,
.' governs the search for ALPHA changes sign over the specified',//,
.' interval is false. Increase the search interval by',//,
.' decreasing the lower bound of ALPHA (XLI in the ER1 file),//,
.' and increasing the upper bound (XRI in the ER1 file).//)

2200 format(5x,'ESTRT? Premature EOF in RUN_NAME.ER1 or RUN_NAME.ER2.')
     .' The portion of the program which reads ER1 and',//,
     .' ER2 files encountered an end-of-file mark before all of',//,
     .' the information had been read. Confirm these files and',//,
     .' execute the program again. If necessary, copy and edit',//,
     .' the appropriate EXAMPLE file and execute the program again.'//)

2210 FORMAT(5X,'ESTRT1/ESTRT2/ESTRT2SS/ESTRT3? DECODE failed',//,
     .' The portion of the program which reads the ER1, ER2,//,
     .' or the ER3 file failed to understand a numerical entry',//,
     .' The numbers must appear in columns 11-20 of the line with',//,
     .' no alphabetic characters in the field. (Note that',//,
     .' exponential notation is not allowed.) This restriction',//,
     .' does not apply to comment lines which have an exclamation',//,
     .' point (!) in the first column.'//)

2220 format(5x,'ESTRT1? The parameter file RUN_NAME.ER1 ',
     .'was not found.'//,
     .' The ER1 file was not found for the current simulation',//,
     .' (RUN_NAME). Copy the file EXAMPLE.ER1 file to RUN_NAME.ER1',//,
     .' and edit it as necessary. Execute the program again.'//)

2230 format(5x,'SORTS1? Fewer than 3 points sorted for any time.'//,
     .' Only one or two simulation points were applicable for',//,
     .' the sort times specified. There are three possible causes',//,
     .' for this condition://,
     .' (1) If this message appears when the sort times are',//,
     .' defaulted (CHECK5 is set to 0. in the ER3 file), the',//,
     .' number of observers will probably have to be increased',//,
     .' to give a good resolution of the downwind concentration',//,
     .' field. (The number of observers is NOBS in the ER2 file',//,
     .' with a maximum of ',I3,' (MAXNOB) in DEGADIS2.DEC.)',//,
     .' As a rule of thumb, one gets good resolution of the downwind',//,
     .' concentration field if the ratio://,
     .' (secondary source duration / number of observers) is less',//,
     .' than about 10 seconds (or 20 at most).',//,
     .' (2) The sort times specified in the ER3 file were',//,

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```

.' before the simulation had developed significantly.',/
.' This is only applicable when the user is specifying the sort',/
.' times (i.e. when CHECK5 is set to 1. in the ER3 file).',/
.' Increase the time of the first sort (ERT1), and rerun the',/
.' program.',/)
2231 format(
.'      (3) The sort times specified in the ER3 file were',/
.' after the gas was below the lowest concentration of interest',/
.' This is only applicable when the user is specifying the sort',/
.' times (i.e. when CHECK5 is set to 1. in the ER3 file).',/
.' Increase the time of the first sort (ERT1), and rerun the',/
.' program. If additional results are desired for later',/
.' times, restart the simulation and specify a lower',/
.' concentration of interest in the input step ( lower',/
.' CCLOW in DEGADISIN).',/
.'      (4) The downwind distance specified in DEGADIS4 is too',/
.' large for the lowest concentration of interest specified. To',/
.' get a time history at this downwind distance, restart the',/
.' simulation with a lower concentration of interest in the input.',/
.//)

2240 format(5x,'TPROP? Trial and error loop compromised',/
.'      TPROP estimates the temperature of a mixture based',/
.' upon the composition and enthalpy of the mixture. Ensure',/
.' the properties for the diffusing species are entered',/
.' correctly and execute the simulation again.',/)

2250 format(5x,'TPROP? Isothermal density loop compromised',/
.'      This error should never occur, but if it does, rebuild',/
.' the model from the original files and run the simulation',/
.' over.',/)

2260 format(5x,'TPROP? Invalid entry flag in ADIABAT',/
.'      This is a programming diagnostic and should never occur.',/
.' If it does, rebuild the model from the original files.',/)

2270 format(5x,'Reserved')

2280 format(5x,'TPROP? IGEN request too large in SETDEN',/
.'      The subroutine SETDEN (in TPROP) performs a series of',/
.' adiabatic mixing calculations with a specified gas mixture',/
.' and ambient air and places the result in the array',/
.' DEN(5,IGEN). This error indicates more points are needed in',/
.' DEN than were originally requested. Increase the allocation',/
.' for DEN by changing the value of IGEN in DEGADISIN.DEC and',/
.' reinstalling DEGADIS.',/)

2290 format(5x,'PHIF? flag IPHIFL is out of bounds',/
.'      Proper values of IPHIFL are integers between 1 and 5 ',/
.' inclusive. Although values of IPHIFL are entered in the ER1',/
.' file as real numbers, they should be in this range. Check',/)
```

```

.' the ER1 file and execute the program again.',//)

2300  format(5x,'SSSUP/SDEGADIS2? concentration greater than RHOE',//,
.'      If the concentration of the contaminant becomes',//,
.' greater than the pure component density for an isothermal',//,
.' simulation, this error will occur. However, this situation',//,
.' should never occur. Check the input conditions and execute',//,
.' the program again.',//)

2310  format(5x,'SSSUP? concentration greater than RHOE',//,
.'      If the concentration of the contaminant becomes',//,
.' greater than the pure component density for an isothermal',//,
.' simulation, this error will occur. However, this situation',//,
.' should never occur. Check the input conditions and execute',//,
.' the program again.',//)

2320  format(5x,'PSS? Sz convergence failure.',//,
.'      This is a programming diagnostic and should never occur.',//,
.' If it does, check the input conditions and execute the',//,
.' program again.',//)

2330  format(5x,'SSG? Sz convergence failure.',//,
.'      This is a programming diagnostic and should never occur.',//,
.' If it does, check the input conditions and execute the',//,
.' program again.',//)

C
      CLOSE(UNIT=9)

C
c CALL TRANS('trap.DBG') - this isn't really needed
C
      istat = lib$date_time(TT)
      tt1 = t1
      ttime = secnds(tt1)/60.

C
      140  WRITE(lunlog,3000) TT
            WRITE(lunlog,3010) Ttime
      3000 FORMAT(1X,' -- ENDING AT ',A24)
      3010 FORMAT(5X,' ***** ELAPSED TIME ***** ',1pg13.5,' MIN ')
C
      irtn = LIB$DO_COMMAND( 'Exit' ) ! this issues the command EXIT to VMS
c                               ! which should cancel any pending
c                               ! programs in a command file.

      CALL EXIT
      END

#####

```

```
C.....  
C  
C      FUNCTION TO CALCULATE A SPECIFIED TIME  
C  
C      FUNCTION TS(TOL,DIST)  
C  
  
Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  
  
COMMON  
$/PARMSC/RM,S2M,EMAX,RMAX,TSC1,ALEPH,TEND  
$/ALP/ALPHA,alpha1  
C  
TS = TOL + (DIST+RMAX)**(1./ALPHA1) /ALEPH  
C  
RETURN  
END  
####
```

```

C.....  

C  

C      OBSERVER TRIAL AND ERROR FUNCTIONS  

C      --- TUPF ---- TDNF ---  

C  

c      Modified 30 Jan 86 for more general trial and error scheme. ts  

C  

FUNCTION TUPF(TOL)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

include 'sys$degadis:DEGADIS2.dec'  

C  

COMMON  

$/GEN3/ radg(2,maxl),qstr(2,maxl),srcden(2,maxl),srcwc(2,maxl),  

$ srcwa(2,maxl),srcenth(2,maxl)  

$/ERROR/SY0ER,ERRO,S20ER,WTAIO,WTQOO,WTSZO,ERRP,SMXP,  

$ WTSZP,WTSYP,WTBEP,WTDH,ERRG,SMXG,ERTDNF,ERTUPF,WTRUH,WTDHG  

$/PARMSC/RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$/ALP/ALPHA,alpha1  

C  

LOGICAL pflag          ! for diagnostic output  

pflag = .false.  

C  

TMAX = RMAX**(1./ALPHA1)/ALEPH + TOL  

TMIN = TOL  

IF(TOL .LT. 0.) TMIN = 0.  

C  

c*** TMIN and TMAX represent the first and last time this observer could  

c*** encounter the upwind edge of the source. TMAX is the time when the  

c*** observer passes over x=0, and TMIN is the time the observer is  

c*** released (unless set to zero because the spill has not yet begun).  

c*** Now, refine the guess of TMIN and TMAX by dividing the interval  

c*** into 20 segments and checking if the observer crosses the upwind  

c*** edge over the smaller interval starting with TMIN.  

C  

DT = (TMAX - TMIN) / 20.  

TL = TMIN  

DO 10 I=1,19      ! I don't have to check the last interval.  

TL = TL + DT  

XO = XIT(TL, TOL)  

XG = -AFGEN(RADG, TL, 'tupf')  

DIF = XO - XG  

IF(DIF .LT. 0.) GOTO 10 ! observer has not yet reached upwind edge  

        TMAX = TL          ! now observer has reached upwind edge  

        TMIN = TL - DT  

        DIFATMAX = DIF  ! DIF at MAX  

        GOTO 20  

10    CONTINUE  

        TMIN = TMAX - DT

```

```

      DIFATMAX = DIF
20    CONTINUE
c
c*** Now perform bisection search to get desired convergence between new TMAX
c*** and TMIN
c
      TL = (TMAX + TMIN)/2.
c
      DO 100 I = 1,20
      II = 0
110    XG = -AFGEN(RADG,TL,'tupf')
      XO = XIT(TL,TOL)
      IF(XO .LT. 0.) GO TO 120
      TL = (TL+TMIN)/2.
      II = II + 1
      if(pflag) write(6,5020) tl,t0l,xg,xo
5020    format(' tl:',1pg13.5,' t0l:',1pg13.5,' xg:',1pg13.5,
      1 ' xo:',1pg13.5)
      IF(II.EQ. 20) GOTO 101 ! Kill the program.
      GO TO 110
c
120    CONTINUE
      DIF = XO - XG
      sum = abs(xo + xg)/2. + ERTUPF
      IF(ABS(DIF)/sum .LT. ERTUPF) GO TO 1000
      if(pflag) write(6,5040) tmin,tmax,tl,xo,xg
5040    format(' tmin:',1pg13.5,' tmax:',1pg13.5,' tl:',1pg13.5,
      1 ' xo:',1pg13.5,' xg:',1pg13.5)
c
      IF( DIF*DIFATMAX .GT. 0.) THEN
          TMAX = TL           ! a new maximum for the range
          DIFATMAX = DIF       ! a new DIF for the maximum
      ELSE
          TMIN = TL           ! a new minimum for the range
      ENDIF
c
100    TL = (TMAX + TMIN)/2.
c
c*** The above search scheme failed. Before killing the program, check to
c*** see if the desired point falls on a transition from a blanket to a
c*** non-blanket situation.
c
      t1 = TL+.01
      T2 = TL-.01
      XG1 = AFGEN(RADG,T1,'tupf')
      XG2 = AFGEN(RADG,T2,'tupf')
      dif = abs(xg1-xg2)
      if(dif.gt. 100. .AND. (XO.GE.XG2 .AND. XO.LE.XG1)) then
          tupf = t2           ! jump from blanket to non-blanket occurred
          RETURN
      ENDIF

```

```

c
c*** Kill the program.
c
101   if(pflag) write(6,4000) RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND,alpha
4000  format(' rm:',1pg13.5,' szm:',1pg13.5,' emax: ',1pg13.5,,,
           1 ' rmax:',1pg13.5,' tsc1:',1pg13.5,' aleph:',1pg13.5,,,
           2 ' tend:',1pg13.5,' alpha:',1pg13.5)
           if(pflag) write(6,4010) tmax,tmin
4010  format(' tmax: ',1pg13.5,' tmin: ',1pg13.5)
           CALL trap(6)

c
c*** successful completion
c
1000  TUPF = TL
      RETURN
      END

c
c
c
c
FUNCTION TDNF(TOL)
c

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

include 'sys$degadis:DEGADIS2.dec'
c
      COMMON
      $/GEN3/ radg(2,maxl),qstr(2,maxl),srcden(2,maxl),srcwc(2,maxl),
      $ srcwa(2,maxl),srcenth(2,maxl)
      $/ERROR/SY0ER,ERRO,SZOER,WTAIO,WTSQ0,WTSZ0,ERRP,SMXP,
      $ WTSZP,WTSYP,WTBEP,WTDH,ERRG,SMXG,ERTDNF,ERTUPF,WTRUH,WTDHG
      $/PARMSC/RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND
      $/ALP/ALPHA,alpha1
c
      LOGICAL pflag
      pflag = .FALSE.

c
      TMIN = RMAX**(1./ALPHA1)/ALEPH + TOL
      IF(TMIN .LT. 0.) TMIN = 0.
      TMAX = (2.*RMAX)**(1./ALPHA1)/ALEPH + TOL
c
c*** TMIN and TMAX represent the first and last time this observer could
c*** encounter the downwind edge of the source. TMAX is the time when the
c*** observer passes over x=+RMAX, and TMIN is the time the observer
c*** passes over x=0 (unless set to zero because the spill has not begun
c*** yet). Now, refine the guess of TMIN and TMAX by dividing the interval
c*** into 20 segments and checking if the observer crosses the downwind
c*** edge over the smaller interval starting from TMAX.
c
      DT = (TMAX - TMIN) / 20.

```

```

TL = TMAX
if(pflag) write(6,*) 'tmax, tmin, dt: ',tmax, tmin, dt

DO 10 I=1,19 ! I don't have to check the last interval.
TL = TL - DT
XO = XIT(TL, TOL)
XG = AFGEN(RADG, TL, 'tupf')
DIF = XO - XG
if(pflag) write(6,*) 'tl, xo, xg, dif:',tl, xo, xg, dif
IF(DIF .GT. 0.) GOTO 10 ! observer has passed the downwind edge
    TMIN = TL           ! now observer is about to reach downwind edge
    TMAX = TL + DT
    GOTO 20

10   CONTINUE
c
TMAX = TMIN + DT
20   CONTINUE
XO = XIT(TMAX, TOL)
XG = AFGEN(RADG, TMAX, 'tupf')
DIFATMAX = XO - XG
if(pflag) then
    write(6,*) '20: tmax, tmin: ',tmax, tmin
    write(6,*) 'difatmax: ',difatmax
endif
c
*** Now perform bisection search to get desired convergence between new TMAX
*** and TMIN.
c
TL = (TMAX + TMIN)/2.
c
.DO 100 I = 1,20
II = 0
110  XG = AFGEN(RADG, TL, 'tdnf')
XO = XIT(TL,TOL)
c
IF(XO .GT. 0.) GO TO 120
TL = (TMAX + TL)/2.
II = II + 1
if(pflag) write(6,5020) tl,t0l,xg,xo
5020  format(' tl:',1pg13.5,' t0l:',1pg13.5,' xg:',1pg13.5,
1 ' xo:',1pg13.5)
IF(II.EQ. 20) GOTO 101 ! Kill the program.
GO TO 110
c
120  CONTINUE
c
DIF = XO - XG
sum = abs(xo+xg)/2. + ERTDNF
IF(ABS(DIF)/sum .LT. ERTDNF) GO TO 1000
if(pflag) write(6,5040) tmin,tmax,tl,xo,xg
5040  format(' tmin:',1pg13.5,' tmax:',1pg13.5,' tl:',1pg13.5,

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```
1 ' xo:',1pg13.5,' xg:',1pg13.5)
C
IF( DIF*DIFATMAX .GT. 0.) THEN
    TMAX = TL                      ! a new maximum for the range
    DIFATMAX = DIF                  ! a new DIF for the maximum
ELSE
    TMIN = TL                      ! a new minimum for the range
ENDIF
C
100   TL = (TMAX + TMIN)/2.
C
c*** The above search scheme failed. Before killing the program, check to
c*** see if the desired point falls on a transition from a blanket to a
c*** non-blanket situation.
C
t1 = TL+.01
T2 = TL-.01
XG1 = AFGEN(RADG,T1,'tupf')
xg2 = AFGEN(RADG,T2,'tupf')
dif = abs(xg1-xg2)
if(dif.gt. 100. .AND. (XO.LE.XG2 .AND. XO.GE.XG1)) then
    tdnf = t2          ! jump from blanket to non-blanket occurred
    RETURN
ENDIF
C
c*** Kill the program.
C
101   if(pflag) write(6,4000) RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND,alpha
4000   format(' rm:',1pg13.5,' szm:',1pg13.5,' emax: ',1pg13.5,/,
           1 ' rmax:',1pg13.5,' tsc1:',1pg13.5,' aleph:',1pg13.5,/,
           2 ' tend:',1pg13.5,' alpha:',1pg13.5)
    if(pflag) write(6,4010) tmax,tmin
4010   format(' tmax: ',1pg13.5,' tmin: ',1pg13.5)
        CALL trap(7)
C
c*** successful completion
C
1000  TDNF = TL
      RETURN
      END
#####

```

```

C.....  

C  

C      FUNCTIONS ASSOCIATED WITH THE OBSERVER CALCULATIONS  

C  

C.....  

C  

C      FUNCTION TO RETURN OBSERVER VELOCITY AS A FUNCTION OF TIME  

C  

FUNCTION UIT(T,TOL)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

COMMON  

$PARMSC/RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$/ALP/ALPHA,alpha1  

C  

UIT = ALPHA1 * ALEPH**ALPHA1 *(T-TOL)**ALPHA  

C  

RETURN  

END  

C  

C  

C.....  

C  

C      FUNCTION TO RETURN POSITION AS A FUNCTION OF TIME AND TO  

C  

C  

FUNCTION XIT(TL,TOL)  

C  

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )  

COMMON  

$PARMSC/RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND  

$/ALP/ALPHA,alpha1  

C  

xit = -rmax  

arg = tl-tol  

if(arg .le. 0.) return  

XIT = (ALEPH*(TL - TOL))**ALPHA1 - RMAX  

C  

RETURN  

END  

C  

C  

C.....  

C  

C      FUNC TO RETURN A VALUE OF TO BASED ON A POSITION AND TIME  

C

```

```
FUNCTION TOOB(X,T)
C
Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

COMMON
$/PARMSC/RM,SZM,EMAX,RMAX,TSC1,ALEPH,TEND
$/ALP/ALPHA,alpha1
C
ARG = 0.
CHECK = ABS((ABS(X)-ABS(RMAX)))/(ABS(X)+ABS(RMAX))
IF(CHECK .GT. 0.001) ARG = (X + RMAX)**(1./ALPHA1)/ALEPH
TOOB = T - ARG
RETURN
END
#####

```

```

        subroutine zbrent(answer, func, x1, x2, tol, ierr)
c
c... Function to determine the root of FUNC which is between X1 and X2
c      from Press et al., pg 253. Note that the SIGN function has
c      been used here to avoid underflows corrupting the programs
c      logic.
c
        implicit real*8(a-h,o-z), integer*4(i-n)

        parameter (itmax=100, eps=3.D-8)

c
c... check the passed arguments for validity and set up the procedure
c
        aaa = x1
        bbb = x2
        ierr = 0
        fa = func(aaa)
        fb = func(bbb)
        if(sign(1.D0,fa)*sign(1.D0,fb) .gt. 0.D0) then
          ierr = 2
          return
        endif

        fc = fb

        do 11 iter = 1,itmax
c
c... shuffle A,B,C and adjust the bounding interval D
c
        if(sign(1.D0,fb)*sign(1.D0,fc) .gt. 0.D0) then
          ccc = aaa
          fc = fa
          ddd = bbb-aaa
          eee = ddd
        endif

        if(abs(fc) .lt. abs(fb)) then
          aaa = bbb
          bbb = ccc
          ccc = aaa
          fa = fb
          fb = fc
          fc = fa
        endif
c
c... convergence check
c
        tol1 = 2.D0*eps*abs(bbb) + 0.5D0*tol
        xm = 0.5D0*(ccc-bbb)
        if(abs(xm).le. tol1 .or. fb.eq.0.D0) then

```

```

        answer = bbb
        return
    endif
c
c... attempt inverse quadratic interpolation
c
    if(abs(eee).ge.tol1 .and. abs(fa).gt.abs(fb)) then
        sss = fb/fa

        if(aaa.eq.ccc) then
            ppp = 2.D0*xm*sss
            qqq = 1.D0-sss
        else
            qqq = fa/fc
            rrr = fb/fc
            ppp = sss*(2.D0*xm*qqq*(qqq-rrr) - (bbb-aaa)*(rrr-1.D0))
            qqq = (qqq-1.D0)*(rrr-1.D0)*(sss-1.D0)
        endif
c
c..... check to insure in bounds
c
        if(ppp .gt. 0.D0) qqq = -qqq
        ppp = abs(ppp)
        if
            . (2.D0*ppp .lt. min(3.D0*xm*qqq-abs(tol1*qqq),abs(eee*qqq))) then

c..... interpolation valid
            eee = ddd
            ddd = ppp/qqq
        else

c..... interpolation failed, use bisection
            ddd = xm
            eee = ddd
        endif
        else

c..... bounds decreasing too slowly, use bisection.
            ddd = xm
            eee = ddd
        endif
c
c..... move last best guess to A
c
        aaa = bbb
        fa = fb
c
c... evaluate new trial root
c
        if(abs(ddd) .gt. tol1) then
            bbb = bbb+ddd

```

```
    else
        bbb = bbb+sign(tol1,xm)
    endif
    fb = func(bbb)
11    continue
c
c... loop failed to converge the x range
c
    ier = 1
    return
end
#####

```

PROGRAM DEGBRIDGE

C-----
C
C Program description:
C
C DEGBRIDGE is designed to read the input file for the OOMS/DEGADIS
C model input. From this information and the output from the OOMS model,
C DEGBRIDGE prepares the necessary input file to run DEGADIS
C (RUN_NAME.INP).
C-----
C
C-----
C The structure of the RUN_NAME.IN file is outlined in OOMS_IN.
C
C The output file from the OOMS model contains three (3) variables
C as follows (in RUN_NAME.IND):
C
C <DIST> <CC> <HWIDTH>
C
C where:
C
C <DIST> is the distance to the centerline touchdown (m)
C <CC> is the centerline concentration at <DIST>
C <HWIDTH> is jet/plume half width at <DIST>
C
C To establish the initial conditions for DEGADIS, the source
C concentration is assumed to be <CC>, and the source radius is assumed
C to be <HWIDTH>.
C-----
C
C Program usage:
C
C Consult Volume III of the Final Report to U. S. Coast Guard
C contract DT-CG-23-80-C-20029 entitled "Development of an
C Atmospheric Dispersion Model for Heavier-than-Air Gas Mixtures".
C
C J. A. Havens
C T. O. Spicer
C
C University of Arkansas
C Department of Chemical Engineering
C Fayetteville, AR 72701
C
C April 1985
C
C This project was sponsored by the U. S. Coast Guard and the Gas
C Research Institute under contract DT-CG-23-80-C-20029.
C

```

C
C      Disclaimer:
C
C      This computer code material was prepared by the University of
C      Arkansas as an account of work sponsored by the U. S. Coast Guard
C      and the Gas Research Institute. Neither the University of Arkansas,
C      nor any person acting on its behalf:
C
C      a. Makes any warranty or representation, express or implied,
C          with respect to the accuracy, completeness, or usefulness
C          of the information contained in this computer code material,
C          or that the use of any apparatus, method, numerical model,
C          or process disclosed in this computer code material may not
C          infringe privately owned rights; or
C
C      b. Assumes any liability with respect to the use of, or for
C          damages resulting from the use of, any information,
C          apparatus, method, or process disclosed in this computer
C          code material.
C
C
C*****
```

```

Implicit Real*8 ( A-H, O-Z ), Integer*4 ( I-N )

C
C      include 'SYSS$DEGADIS:degadisin.dec'
C
C      COMMON
C/TITL/ TITLE
$/GEN1/ PTIME(igen), ET(igen), R1T(igen), PWC(igen), PTEMP(igen),
$      PFRACV(igen), PENTH(igen), PRHO(igen)
$/GEN2/ DEN(5,igen)
$/ITI/ T1,TINP,TSRC,TOBS,TSRT
$/PARM/ U0,Z0,ZR,rml,USTAR,vkc,gg,gRHOE,RHOA,DELTA,BETA,GAMMAF,
.          CcLOW
$/com_gprop/ gasmw,temp,temjet,rhoec,pk,ccp,gasul,gasll,zll,gasnam
$/com_ss/ ess,slen,swid,outcc,outsz,outb,outl
$/PHLAG/ CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
$/com_sigx/ sigx_coeff,sigx_pow,sigx_min_dist,sigx_flag
$/NEND/ POUNDN,POUND
./oomsin/ oodist,avtime
C
C      character*80 TITLE(4)
C
C      character*3 gasnam
```

```

character*4 pound
character*24 TSRC,TINP,TOBS,TSRT
C
LOGICAL CHECK1,CHECK2,AGAIN,CHECK3,CHECK4,CHECK5
c
c check1
c check2=t    cloud type release with no liquid source; SRC1  DEGADIS1
c again      local communications in SSSUP                  SSSUP
c check3      local communications between SRC1 and NOBL   DEGADIS1
c check4=t    steady state simulation                      DEGADISIN
c check5=t    operator sets sort parameters                ESTRT3
c
data CHECK1/.false./,CHECK2/.false./,AGAIN/.false./
data CHECK3/.false./,CHECK4/.false./,CHECK5/.false./
C
character*100 OPNRUP
character OPNRUP1(100)
equivalence (opnrup1(1),opnrup)
character*4 in,ind,INP
character*100 dummy
DATA POUND/'//  '/',POUNDN/-1.E-20/
C
DATA PTIME/igen*0.D0/
DATA ET/igen*0.D0/,R1T/igen*0.D0/
DATA PWC/igen*0.D0/,PTEMP/igen*0.D0/
DATA PFRACV/igen*0.D0/,PENTH/igen*0.D0/
DATA PRHO/igen*0.D0/
data DEN/igen*0.,igen*0.,igen*0.,igen*0.,igen*0./

DATA .INP/'.INP'/,IN/'.IN '//,IND/'.IND'/
C
c
C... GET THE FILE NAME TO BE USED BY ALL OF THE ROUTINES..... .
C
write(6,*) 'Beginning DEGBRIDGE...'

READ(5,820) NCHAR,opnrup
opnrup = opnrup(1:nchar) // ind(1:4)
C
C... First, get the desired information from RUN_NAME.IND
C
open(unit=8,name=opnrup,type='old')
read(8,*) oodist, cc, hwidth
C
c... If OODIST is zero, discontinue the run
c
if(oodist .eq. 0.D0) then
    write(6,*) ' ...terminating the run since CCLOW is met.'
    dummy = opnrup(1:nchar)
    opnrup = opnrup(1:nchar) // '.com_xxx;1'
    open(unit=1,name=opnrup,type='unknown')

```

```

opnrup = '$ copy '//opnrup(1:nchar)//'.out .lis'
write(1,8020) (opnrup1(iii),iii=1,nchar+16)
opnrup = '$ delete '//dummy(1:nchar)//'.com_xxx;1'
write(1,8020) (opnrup1(iii),iii=1,nchar+19)
opnrup = '$ stop'
write(1,8020) (opnrup1(iii),iii=1,6)
close(unit=1)
opnrup = '2'// dummy(1:nchar) // '.com_xxx;1'//'
istat = lib$do_command(opnrup)
write(6,*) ' ?DEGBRIDGE? EXIT failed to work.'
stop
endif
close(unit=8)
c
c... Now, get the desired information from RUN_NAME.IN
c
opnrup = opnrup(1:nchar) // in(1:4)
open(unit=8,name=opnrup,type='old')

read(8,8000) title(1)
read(8,8000) title(2)
read(8,8000) title(3)
read(8,8000) title(4)
read(8,*) u0, z0
read(8,*) zr

read(8,*) indvel, istab, rml
c
c... Based on INDVEL, set the Monin-Obukhov length as desired.....
c
if( indvel.ne.1 .or. indvel.ne.2) indvel = 1
if( istab.le.0 .or. istab.gt.6) istab = 4
if(indvel .eq. 1) then
    if(istab.eq.1) then
        rml = -11.43D0 * zr**0.103D0
    else if(istab.eq.2) then
        rml = -25.98D0 * zr**0.171D0
    else if(istab.eq.3) then
        rml = -123.4D0 * zr**0.304D0
    else if(istab.eq.4) then
        rml = 0.0D0
    else if(istab.eq.5) then
        rml = 123.4D0 * zr**0.304D0
    else if(istab.eq.6) then
        rml = 25.98D0 * zr**0.171D0
    endif
endif
endif

read(8,*) tamb, pamb, rethum

if( tamb.le.0. ) tamb = tamb+273.15D0

```

```

      if( pamb.gt.1.1 ) pamb = pamb/101325.00
      if( relhum.lt.0. .or. relhum.gt.100. ) relhum = 50.

c
c... Calculate the absolute humidity HUMID
c
      vaporp = 6.0298D-3* exp(5407.00*(1.00/273.15D0 - 1.00/tamb))
      sat = 0.622D0*vaporp/(pamb - vaporp)
      humid = relhum/100.00 * sat

      read(8,*) tsurf
      if( tsurf.lt.250. ) tsurf = tamb

      read(8,8010) gasnam
      read(8,*) gasmw
      read(8,*) avtime
      read(8,*) TEMJET
      read(8,*) gasul, gasll, zll
      if( gasll.le.0. ) gasll = 0.01
      if( gasul.le.gasll ) gasul = dmax1( 1.1D0*gasll, 1.000)

c
c... Now that AVTIME has been set, the value of DELTAY can be fixed. Also
c      set the values of BETAY
c
      goto(161,162,163,164,165,166) istab
161   timeav = dmax1( avtime, 18.4D0)           ! A
      deltay = 0.423D0*(timeav/600.00)**0.2D0
      betay = 0.900
      sigx_coeff = 0.02
      sigx_pow = 1.22
      sigx_min_dist = 130.
      goto 170
162   timeav = dmax1( avtime, 18.4D0)           ! B
      deltay = 0.313D0*(timeav/600.00)**0.2D0
      betay = 0.900
      sigx_coeff = 0.02
      sigx_pow = 1.22
      sigx_min_dist = 130.
      goto 170
163   timeav = dmax1( avtime, 18.4D0)           ! C
      deltay = 0.210D0*(timeav/600.00)**0.2D0
      betay = 0.900
      sigx_coeff = 0.02
      sigx_pow = 1.22
      sigx_min_dist = 130.
      goto 170
164   timeav = dmax1( avtime, 18.3D0)           ! D
      deltay = 0.136D0*(timeav/600.00)**0.2D0
      betay = 0.900
      sigx_coeff = 0.04
      sigx_pow = 1.14

```

```

    sigx_min_dist = 100.
    goto 170
165   timeav = dmax1( avtime, 11.4D0)                      ! E
    deltay = 0.102D0*(timeav/600.D0)**0.200
    betay = 0.9D0
    sigx_coeff = 0.17
    sigx_pow = 0.97
    sigx_min_dist = 50.
    goto 170
166   timeav = dmax1( avtime, 4.6D0)                      ! F
    deltay = 0.0674D0*(timeav/600.D0)**0.200
    betay = 0.9D0
    sigx_coeff = 0.17
    sigx_pow = 0.97
    sigx_min_dist = 50.

c
170   continue

c
c... Recover INDHT, CPK, and CPP. If CPP is set to 0, then CPK contains.....
c      the (constant) heat capacity.
c
read(8,*) indht, CPK, cpp
if(cpp .eq. 0.D0) then
    cpp = 1.D0
    CPK = CPK*gasmw - 3.33D4
endif

c
c... recover NDEN and set ISOFL and DEN accordingly..... .
c
read(8,*) nden
if(nden .lt. -1) nden=-1

if(nden .eq. -1) then
    isofl = 1
    rhoe = pamb*101325.D0*gasmw/8314.D0/TEMJET
    grhoe = rhoe
    rhoa = pamb*
        (1.00+humid)/(0.002833D0 + 0.004553D0*humid)/tamb
    den(1,1) = 0.D0
    den(2,1) = 0.D0
    den(3,1) = rhoa
    den(4,1) = 0.D0
    den(5,1) = tamb
    den(1,2) = 1.D0
    den(2,2) = rhoe
    den(3,2) = rhoe
    den(4,2) = 0.D0
    den(5,2) = tamb
    den(1,3) = 2.D0

```

```

else if(nden .eq. 0) then
    isofl = 0

    rhoe = pamb*101325.D0*gasmw/8314.D0/TEMJET
    grhoe = rhoe
    rhoa = pamb*
        (1.D0+humid)/(0.002833D0 + 0.004553D0*humid)/tamb

    wc = 1.D0
    wa = 0.D0
    enth = cpc(temjet)*(temjet - tamb)
    call setden(wc, wa, enth)

else
    isofl = 1

do iii = 1,nden
    read(8,*) den(1,iii), den(2,iii), den(3,iii)
    den(4,iii) = 0.D0
    den(5,iii) = tamb
    enddo
    den(1,nden+1) = 2.D0
    rhoe = den(3,nden)
    grhoe = rhoe
    rhoa = den(3,1)
endif

nden0 = nden
if(nden .eq. -1) nden0 = 2

read(8,*) erate
read(8,*) elejet, DIAJET
read(8,*) tend
check4 = .true.
if(tend .gt. 0.) check4 = .false.

read(8,*) distmx

close(unit=8)

c
c... It is time to prepare the input file for DEGADIS.....
c
opnrup = opnrup(1:nchar) // inp(1:4)
open(unit=8,name=opnrup,type='new')
c
c... TITLE
c
      DO 200 I=1,4
      WRITE(8,8000) TITLE(I)

```

```

200  CONTINUE
C
c*** Atmospheric parameters:
C
      WRITE(8,1020) U0,Z0,ZR
C
c*** stability, averaging time for DELTAY, and derived parameters
C
      WRITE(8,1040) istab
      write(8,1020) oodist,avtime
      WRITE(8,1020) DELTAY,BETAY,rml
      WRITE(8,1020) sigx_coeff,sigx_pow,sigx_min_dist
C
c*** ambient pressure, temperatures, and humidity
C
      write(8,1025) tamb,pamb,humid
C
      ihtfl = indht
      htco  = 0.
      iwtfl = 0
      wtco  = 0.
C
      write(8,1060) isofl,tsurf
      write(8,1060) ihtfl,htco
      write(8,1060) iwtfl,wtco
C
c*** gas characteristics
C
      write(8,8010) gasnam
      WRITE(8,1020) gasmw, temjet,rhoe
      write(8,1020) cpk,cpp
      WRITE(8,1020) gasll,gaslt,zll
C
c density curve if isothermal
C
      if(isofl .eq. 0) goto 460

      WRITE(8,1040) nden0
      DO 440 I=1,nden0
      WRITE(8,1025) DEN(1,I),DEN(2,I),DEN(3,I),Den(4,I),den(5,i)
440  CONTINUE
C
C
460  ccflow = (gasll/2.00) * pamb*101325.00*gasmw/8314.00/Tamb
      WRITE(8,1010) CcLOW
C
c*** source description FOR A DILUTED SOURCE ...
C
      call adiabat(0,wc,wa,yc,ya,cc,rho,wm,enth,temp)
C
      gmass0 = 0.          ! no initial cloud mass

```

```

        write(8,1020) gmass0
        np = 4
c
        if(check4) tend = 60230. ! [=] sec
c
        ess = erate
        r1ss = hwidth
        slen = 2.D0*hwidth
        swid = pi*r1ss**2/slen/2.D0
        pwc(1) = wc
        ptemp(1) = temp
        pfracv(1) = 1.0
c
        PTIME(1) = 0.D0
        et(1) = ess
        r1t(1) = r1ss
        PWC(1) = pwc(1)
        PTEMP(1) = ptemp(1)
        PFRACV(1)= 1.000

        PTIME(2) = tend
        et(2) = ess
        r1t(2) = r1ss
        PWC(2) = pwc(1)
        PTEMP(2) = ptemp(1)
        PFRACV(2)= 1.000

        PTIME(3) = tend + 1.
        et(3) = 0.
        r1t(3) = 0.
        PWC(3) = pwc(1)
        PTEMP(3) = ptemp(1)
        PFRACV(3)= 1.0D0

        PTIME(4) = tend + 2.
        et(4) = 0.
        r1t(4) = 0.
        PWC(4) = pwc(1)
        PTEMP(4) = ptemp(1)
        PFRACV(4)= 1.0D0

        WRITE(8,1040) NP
        DO 800 I=1,NP
800      WRITE(8,1030) PTIME(I),ET(I),R1T(I), PWC(I),PTEMP(I),PFRACV(I)
C
        write(8,*) check1,check2,again,check3,check4,check5
C
        istat = lib$date_time(tinp)
        write(8,1050) tinp
c
        if(check4) write(8,1020) ess,slen,swid           ! steady state

```

```
c
c
      CLOSE(UNIT=8)
c
c
  820  FORMAT(Q,A40)

  1010  format(1x,1pg14.7)
  1020  format(3(1x,1pg14.7))
  1025  format(5(1x,1pg14.7))
  1030  format(1x,5(1pg14.7,1x),1pg14.7)
  1040  format(1x,I5)
  1050  format(a24)
  1060  format(1x,i4,1x,1pg14.7)

  8000  format(a80)
  8010  format(a3)
  8020  format(80a1)
c

      CALL EXIT
      END
####
```

APPENDIX F
PARTIAL LISTING OF PROGRAM VARIABLES

<u>Variable</u>	<u>Data Type</u>	<u>Symbol</u>	<u>Units</u>	<u>Comments</u>
AGAIN	LOGICAL			Local communications in SSSUP
ALEPH	REAL			Collection of constants to calculate observer position and velocity
ALPHA	REAL	α	n/a	Power law velocity profile power
ALPHA1	REAL	(1.0 + α)	n/a	
BETAY	REAL	β_y	n/a	Lateral similarity power
CCLOW	REAL		kg/m ³	Lowest mixture concentration of interest
CHECK1	LOGICAL			Unused logical flag
CHECK2	LOGICAL			When true, release type without a liquid source
CHECK3	LOGICAL			Local communications flag used in DEGADIS1
CHECK4	LOGICAL			When true, steady-state simulation
CHECK5	LOGICAL			When true, user sets time-sort parameters
DELTAY	REAL	δ_y	m ^{1-β}	Lateral similarity coefficient
DEN(1,I)	REAL	y _c	mole fraction	Contaminant mole fraction
DEN(2,I)	REAL	c _c	kg/m ³	Contaminant concentration for the given mole fraction

<u>Variable</u>	<u>Data Type</u>	<u>Symbol</u>	<u>Units</u>	<u>Comments</u>
DEN(3,I)	REAL	ρ	kg/m ³	Mixture density for the given mole fraction
DEN(4,I)	REAL	h	J/kg	Mixture enthalpy for the given mole fraction
DEN(5,I)	REAL	T	K	Mixture temperature for the given mole fraction
EMAX	REAL		kg/s	Maximum of secondary source mass evolution rate
ESS	REAL	E	kg/s	Steady-state release rate
ET(I)	REAL	E(t)	kg/s	Source mass evolution rate as a function of time PTIME(I)
G	REAL	g	m/s ²	Acceleration due to gravity
GAMMAF	REAL	$\Gamma(1/(1+\alpha))$	n/a	
GAS_CPK	REAL	q_1	$J/kmol K^{p_1 - 1}$	Constant for contaminant heat capacity
GAS_CPP	REAL	p1	n/a	Power for contaminant heat capacity
GAS_LFL	REAL		mole fraction	Lower contaminant concentration level for estimating contours
GAS_MW	REAL	MWC	kg/kmol	Contaminant molecular weight
GAS_NAME	CHARACTER*3			Name of contaminant
GAS_RHOE	REAL	ρ_0	kg/m ³	Saturated vapor density of contaminant at T0
GAS_TEMP	REAL	T0	K	Contaminant storage temperature
GAS_UFL	REAL		mole fraction	Upper contaminant concentration level for estimating contours

<u>Variable</u>	<u>Data Type</u>	<u>Symbol</u>	<u>Units</u>	<u>Comments</u>
GAS_ZSP	REAL		m	Height for estimating contours
GMASS0	REAL		kg	Initial mass of gas over the primary source
HTCO	REAL	h_0	J/m ² sK	Constant coefficient when IHTFL=1
		v_H	m/s	LLNL heat transfer velocity when IHTFL=2
HUMID	REAL		kg water/kg dry air	Ambient absolute humidity
IHTFL	INTEGER			Heat transfer flag: IHTFL=1 constant coefficient IHTFL=0 no heat transfer IHTFL=1 DEGADIS correlation IHTFL=2 LLNL correlation
ISOFL	INTEGER			Isothermal release when ISOFL=1
ISTAB	INTEGER			Pasquill atmospheric stability indicator (ISTAB=1 for A, (ISTAB=2 for B, etc.)
IWTFL	INTEGER			Water transfer flag IWTFL=1 constant coefficient IWTFL=0 no water transfer IWTFL=1 DEGADIS correlation
K	REAL	k	n/a	von Karman's constant, 0.35
LUNLOG	INTEGER			Fortran logical unit number which acts as a simulation log
MAXNOB	INTEGER			Maximum number of observers
ML	REAL	λ	m	Monin-Obukhov length
NOBS	INTEGER			Number of observers for the pseudosteady-state simulation

<u>Variable</u>	<u>Data Type</u>	<u>Symbol</u>	<u>Units</u>	<u>Comments</u>
NREC(I,1)	INTEGER			Number of records generated in PSSOUT for observer I
NREC(I,2)	INTEGER			Number of records generated in SSGOUT for observer I
PAMB	REAL	p	atm	Ambient pressure
POUND	CHARACTER*4			Character string to signal end of data ('//')
POUNDN	REAL			Numerical value to signal end of data (-1.E-20)
QSTR(1,I)	REAL	t	s	Independent variable time for ordered pairs QSTR
QSTR(2,I)	REAL	Q*	kg/m ² s	Atmospheric takeup rate as a function of time
RADG(1,I)	REAL	t	s	Independent variable time for ordered pairs RADG
RADG(2,I)	REAL	R	m	Secondary source radius as a function of time
RELHUMID	REAL		%	Ambient relative humidity
RHOA	REAL	ρ_a	kg/m ³	Ambient air density
RM	REAL	R _m	m	Radius at EMAX (when secondary source mass evolution rate is a maximum)
RMAX	REAL	R _{max}	m	Maximum secondary source radius
RT2	REAL	$\sqrt{2}$.	n/a	Constant
R1SS	REAL	R _p	m	Steady-state primary source radius

<u>Variable</u>	<u>Data Type</u>	<u>Symbol</u>	<u>Units</u>	<u>Comments</u>
R1T(I)	REAL	R_p	m	Primary source radius as a function of time PTIME(I)
SIGX_COEFF	REAL			Along-wind similarity coefficient
SIG_MIN_DIST	REAL		m	Minimum distance to apply x-direction dispersion correction
SIGX_POW	REAL		n/a	Along-wind similarity power
SLEN	REAL	L	m	Steady-state source length
SQPIO2	REAL	$\sqrt{\pi/2}$	n/a	Constant
SQRTPI	REAL	$\sqrt{\pi}$	n/a	Constant
SRCDEN(1,I)	REAL	t	s	Independent variable time for ordered pairs SRCDEN
SRCDEN(2,I)	REAL	ρ	kg/m ³	Secondary source density as a function of time
SRCENTH(1,I)	REAL	t	s	Independent variable time for ordered pairs SRCENTH
SRCENTH(2,I)	REAL	h	J/kg	Secondary source enthalpy as a function of time
SRCWA(1,I)	REAL	t	s	Independent variable time for ordered pairs SRCWA
SRCWA(2,I)	REAL	w_a	mass fraction	Secondary source air mass fraction as a function of time
SRCWC(1,I)	REAL	t	s	Independent variable time for ordered pairs SRCWC

<u>Variable</u>	<u>Data Type</u>	<u>Symbol</u>	<u>Units</u>	<u>Comments</u>
SRCWC(2,I)	REAL	w _c	mass fraction	Secondary source contaminant mass fraction as a function of time
SWID	REAL		m	Steady-state source half-width
SZM	REAL	S _{z0m}	m	Value of S _{z0} at EMAX (when secondary source mass evolution rate is a maximum)
TAMB	REAL	T	K	Ambient temperature
TEND	REAL		s	Termination time of secondary source
TINP	CHARACTER*24			Time DEGADISIN was executed
TITLE(1:4)	CHARACTER*80			Text title block 4 lines of 80 spaces
T0(I)	REAL		s	Time of release for observer I
TSURF	REAL	T _s	K	Surface temperature
USTAR	REAL	u*	m/s	Friction velocity
U0	REAL	u ₀	m/s	Ambient velocity at height z ₀
WTCO	REAL	F	kg/m ² s	Mass transfer coefficient when IWTFL=1
XV(I)	REAL	x _v	m	Virtual source position for estimation of S _y in SSG
Z0	REAL	z ₀	m	Height for velocity u ₀
ZR	REAL	z _R	m	Roughness length

**APPENDIX G
DEGADIS DIAGNOSTIC MESSAGES**

To assist the user in determining the source of any problems, a diagnostic procedure has been included in DEGADIS. The subroutine TRAP is meant to cause an orderly termination of the program for many detected errors. It performs two basic functions: TRAP displays an error code and a single line diagnostic message giving the reason for premature termination, and TRAP forces an output of the COMMON area data sets to the file TRAP.DBG.

The first three lines sent to the execution log (default-TERMINAL) include the TRAP introductory lines and the error code number:

```
The best laid plans of mice and men . . .
You have entered a TRAP--THE LAND OF NO RETURN
CODE: NN
```

where NN represents the code of the error message which follows in the log. The error message begins with the name of the calling routine.

The following is a list of the error codes, error messages, and suggested actions for each problem.

Code: 1

DEGADIS1? Source integration has returned IHLF-NN

Action: This error occurs during integration of the equations which describe the gas source. IHLF is an error code returned by the integration package RKGST.

When IHLF=11, more than 10 bisections of the initial increment of the independent variable were necessary to make an integration step within the specified error. Reduce the initial step size of the independent variable (STPIN in the ER1 file). If this does not work, it will be necessary to either increase the error criteria for all of the dependent variables being integrated (ERBND in the ER1 file) or increase the error criteria for the variable violating the criteria by decreasing the error weight for that variable (one of the following: WTRG, WTTM, WTYA, WTYC, WTEB, WTMB, or WTUH in the ER1 file).

When IHLF=12, the initial increment of the independent variable (STPIN) is 0. Correct the ER1 file and execute the program again.

When IHLF=13, the initial increment of the independent variable (STPIN) is not the same sign as the difference between the upper bound of the interval and the lower bound of the interval. STPIN must be positive. Correct the ERL file and execute the program again.

Code: 2
Reserved

Code: 3
SZF? Local integration failed; IHLF>NN

Action: This error occurs during estimation of SZ over the source when no gas is present. IHLF is an error code returned by the integration package RKGST.

When IHLF=11, more than 10 bisections of the initial increment of the independent variable were necessary to make an integration step within the specified error. Reduce the initial step size of the independent variable (SZSTPO in the ERL file). If this does not work, increase the error criteria for all of the dependent variables being integrated (SZERR in the ERL file).

When IHLF=12, the initial increment of the independent variable (SZSTPO) is 0. Correct the ERL file and execute the program again.

When IHLF=13, the initial increment of the independent variable (SZSTPO) is not the same sign as the difference between the upper bound of the interval and the lower bound of the interval. SZSTPO must be positive. Correct the ERL file and execute the program again.

Code: 4
SURFACE? Negative QRTE for positive DELTA_T.

Action: This is a diagnostic message indicating an error in estimation of the heat capacity. Check the input to the model and execute the program again.

Code: 5
CRFG? More points for GEN3 were needed.

Action: The COMMON area /GEN3/ stores representative values of the calculated source parameters. If this message occurs, relax the CRFG error criteria (CRFGER) in the ERL file. If this is a common problem, the length of the /GEN3/ vectors can be increased by changing the value of MAXL in DEGADIS1.DEC and reinstalling DEGADIS.

Code: 6

TUPF? Observer calculations--TUPF failed

Action: The trial-and-error search associated with finding the upwind edge of the gas source for an observer failed. Often this problem can be avoided by adding one or two additional observers to the present number of observers (which changes the conditions for the trial and error). Another possibility is to increase the error criteria for this function (ERTUPF) in the ER2 file.

Code: 7

TDNF? Observer calculations--TDNF failed

Action: The trial-and-error search associated with finding the downwind edge of the gas source for an observer failed. Often this problem can be solved by adding one or two additional observers to the present number of observers (which changes the conditions for the trial and error). Another possibility is to increase the error criteria for this function (ERTDNF) in the ER2 file.

Code: 8

SSSUP? Observer Integration failed, IHLF>NN

Action: This error occurs during integration of the five differential equations which average the source for each observer. IHLF is an error code returned by the integration package RKGST.

When IHLF=11, more than 10 bisections of the initial increment of the independent variable were necessary to make an integration step within the specified error. Reduce the initial step size of the independent variable (STPO in the ER2 file). If this does not work, it will be necessary to either increase the error criteria for all of the dependent variables being integrated (ERRO in the ER2 file) or increase the error criteria for the variables violating the criteria by decreasing the error weight for that variable (one of the following: WTAIO, WTQOO, or WTSZ0 in the ER2 file).

When IHLF=12, the initial increment of the independent variable (STPO) is 0. Correct the ER2 file and execute the program again.

When IHLF=13, the initial increment of the independent variable (STPO) is not the same sign as the difference between the upper bound of the interval and the lower bound of the interval. STPO must be positive. Correct the ER2 file and execute the program again.

Code: 9

SSSUP/SDEGADIS2? Pseudosteady Integration failed, IHLF-NN

Action: This error occurs during integration of the four differential equations describing the portion of the downwind calculation when $b > 0$. The routine calling TRAP is SSSUP if a transient simulation is being executed; if a steady-state simulation is being executed, the calling routine is SDEGADIS2. IHLF is an error code returned by the integration package RKGST.

When IHLF=11, more than 10 bisections of the initial increment of the independent variable were necessary to make an integration step within the specified error. Reduce the initial step size of the independent variable (STPP in the ER2 file). If this does not work, it will be necessary to either increase the error criteria for all of the dependent variables being integrated (ERRP in the ER2 file) or increase the error criteria for the variable violating the criteria by decreasing the error weight for that variable (one of the following: WTSZP, WTSYP, WTBEPE, or WTDH in the ER2 file).

When IHLF=12, the initial increment of the independent variable (STPP) is 0. Correct the ER2 file and execute the program again.

When IHLF=13, the initial increment of the independent variable (STPP) is not the same sign as the difference between the upper bound of the interval and the lower bound of the interval. STPP must be positive. Correct the ER2 file and execute the program again.

Code: 10

SSSUP/SDEGADIS2? Gaussian Integration fail, IHLF-nn

Action: This error occurs during integration of the differential equations describing the portion of the downwind calculation when $b=0$. The routine calling TRAP is SSSUP if a transient simulation is being executed; if a steady-state simulation is being executed, the calling routine is SDEGADIS2. IHLF is an error code returned by the integration package RKGST.

When IHLF=11, more than 10 bisections of the initial increment of the independent variable were necessary to make an integration step within the specified error. Reduce the initial step size of the independent variable. (STPG in the ER2 file). If this does not work, it will be necessary to either increase the error criteria for all of the dependent variables being integrated (ERRG in the ER2 file) or increase the error criteria for the variable violating the criteria by decreasing the error weight for that variable (either WTRUH or WTDHG in the ER2 file).

When IHLF=12, the initial increment of the independent variable (STPG) is 0. Correct the ER2 file and execute the program again.

When IHLF=13, the initial increment of the dependent variable (STPG) is not the same sign as the difference between the upper bound of the interval and the lower bound of the interval. STPG must be positive. Correct the ER2 file, and execute the program again.

Code: 11

SSSUP/SDEGADIS2 Total No. of Records exceeds 120,000

Action: This is an arbitrary stopping point for the process in order to keep a runaway simulation from filling up disk space. Relax the output specification (ODLP, ODLLP, ODLG, or, DLLG) in the ER2 file in order to generate less output if the input parameters are valid.

Code: 12

Reserved

Code: 13

Reserved

Code: 14

Reserved

Code: 15

Reserved

Code: 16

PSSOUT/PSSOUTSS? PSS started with B < 0.

Action: This condition is checked at the beginning of the downwind calculation in order to confirm proper handling of the movement to the Gaussian phase of the downwind calculation. Check the initial conditions and execute the program again.

Code: 17

TPROP/ADDHEAT? Enthalpy out of bounds

Action: Diagnostic message indicating that an enthalpy lower than the adiabatic mixing enthalpy was passed to ADDHEAT. Check the input conditions and execute the program again.

Code: 18

ALPH? ALPHA integration failed, IHLF>NN

Action: The integration which determines the integral least squares fit for ALPHA has failed. Note that small values of the Monin-Obukhov length ($ML < 0(1m)$) in combination with stable atmospheric conditions may cause this failure. IHLF is an error code returned by the integration package RKGST.

When IHLF=11, more than 10 bisections of the initial increment of the independent variable were necessary to make an integration step within the specified error. Reduce the absolute value of the initial step size of the independent variable (STPINZ in the ER1 file). If this does not work, it will be necessary to increase the error criteria (ERBNDZ in the ER1 file).

When IHLF=12, the initial increment of the independent variable (STPINZ) is 0. Correct the ER1 file and execute the program again.

When IHLF=13, the initial increment of the independent variable (STPINZ) is not the same sign as the difference between the upper bound of the interval and the lower bound of the interval. STPINZ must be negative. Correct the ER1 file and execute the program again. This error will also occur if the surface roughness ZR is greater than the reference height Z0.

Code: 19

ALPH? RTMI has failed to locate ALPHA IERR>NN

Action: The search procedure which determines ALPHA has failed. This error may be the result of an unusual velocity specification such as small values of the Monin-Obukhov length ($ML < 0(1.m)$) or small reference heights ($Z0 < 0(10. * ML)$). IERR is an error code returned by the routine RTMI.

When IERR=1, the search for ALPHA failed after a specified number of iterations. Increase the error bound used by RTMI (EPS in the ER1 file).

When IERR=2, the basic assumption that the function which governs the search for ALPHA changes sign over the specified interval is false. Increase the search interval by decreasing the lower bound of ALPHA (XLI in the ER1 file) and increasing the upper bound (XRI in the ER1 file).

Code: 20

Estrt? Premature EOF in RUN_NAME.ER1 or RUN_NAME.ER2

Action: The portion of the program which reads ER1 and ER2 files encountered an end-of-file mark before all of the information had been read. Confirm these files and execute the program again. If necessary, copy and edit the appropriate EXAMPLE file and execute the program again.

Code: 21

ESTRT1/ESTRT2/ESTRT2SS/ESTRT3? DECODE failed.

Action: The portion of the program which reads the ER1, ER2, or the ER3 file failed to understand a numerical entry. The numbers must appear in columns 11-20 of the line with no alphabetic characters in the field. (Note that exponential notation is not allowed.) This restriction does not apply to comment lines which have an exclamation point (!) in the first column.

Code: 22

ESTRT1? The parameter file RUN_NAME.ER1 was not found.

Action: The ER1 file was not found for the current simulation (RUN_NAME). Copy the file EXAMPLE.ER1 to RUN_NAME.ER1 and edit it as necessary. Execute the program again.

Code: 23

SORTS1? Fewer than 3 points sorted for any time.

Action: Only one or two simulation points were applicable for the sort times specified. There are three possible causes for this condition:

(1) if this message appears when the sort times are defaulted (CHECK5 is set to 0. in the ER3 file), the number of observers will probably have to be increased to give a good resolution of the downwind concentration field. (The number of observers is NOBS in the ER2 file with a maximum (MAXNOB) given in DEGADIS2.DEC.). As a rule of thumb, one gets good resolution of the downwind concentration field if the ratio: (secondary source duration / number of observers) is less than about 10 seconds (or 20 at most).

(2) The sort times specified in the ER3 file were before the simulation had developed significantly. This is only applicable when the user is specifying the sort times (i.e. when CHECK5 is set to 1. in the ER3 file). Increase the time of the first sort (ERT1), and rerun the program.

(3) The sort times specified in the ER3 file were after the gas was below the lowest concentration of interest. This is only applicable when the user is specifying the sort times (i.e. when CHECK5 is set to 1. in the ER3 file). Increase the time of the first sort (ERT1), and rerun the program. If additional results are desired for later times, restart the simulation and specify a lower concentration of interest in the input step (lower CCLOW in DEGADISIN).

Code: 24

TPROP? Trial and error loop compromised.

Action: TPROP estimates the temperature of a mixture based upon the composition and enthalpy of the mixture. Ensure the properties for the diffusing species are entered correctly and execute the simulation again.

Code: 25

TPROP? Isothermal density loop compromised.

Action: This error should never occur, but if it does, rebuild the model from the original files and run the simulation over.

Code: 26

TPROP? Invalid entry flag in ADIABAT.

Action: This is a programming diagnostic and should never occur. If it does, rebuild the model from the original files.

Code: 27

Reserved

Code: 28

TPROP? IGEN request too large in SETDEN.

Action: The subroutine SETDEN (in TPROP) performs a series of adiabatic mixing calculations with a specified gas mixture and ambient air and places the result in the array DEN(5,IGEN). This error indicates more points are needed in DEN than were originally requested. Increase the allocation for DEN by changing the value of IGEN in DEGADISIN.DEC and reinstalling DEGADIS.

Code: 29

PHIF? Flag IPHIFL is out of bounds.

Action: Proper values of IPHIFL are integers between 1 and 5 inclusive. Although values of IPHIFL are entered in the ER1 file as real numbers, they should be in this range. Check the ER1 file and execute the program again.

Code: 30

SSSUP/SDEADIS2? Concentration greater than RHOE.

Action: If the concentration of the contaminant becomes greater than the pure component density for an isothermal simulation, this error will occur. However, this situation should never occur. Check the input conditions and execute the program again.

Code: 31

SSSUP? Concentration greater than RHOE.

Action: If the concentration of the contaminant becomes greater than the pure component density for an isothermal simulation, this error will occur. However, this situation should never occur. Check the input conditions and execute the program again.

Code: 32

PSS? Sz convergence failure.

Action: This is a programming diagnostic and should never occur. If it does, check the input conditions and execute the program again.

Code: 33

SSG? Sz convergence failure.

Action: This is a programming diagnostic and should never occur. If it does, check the input conditions and execute the program again.

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(Please read instructions on the reverse before completing)

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