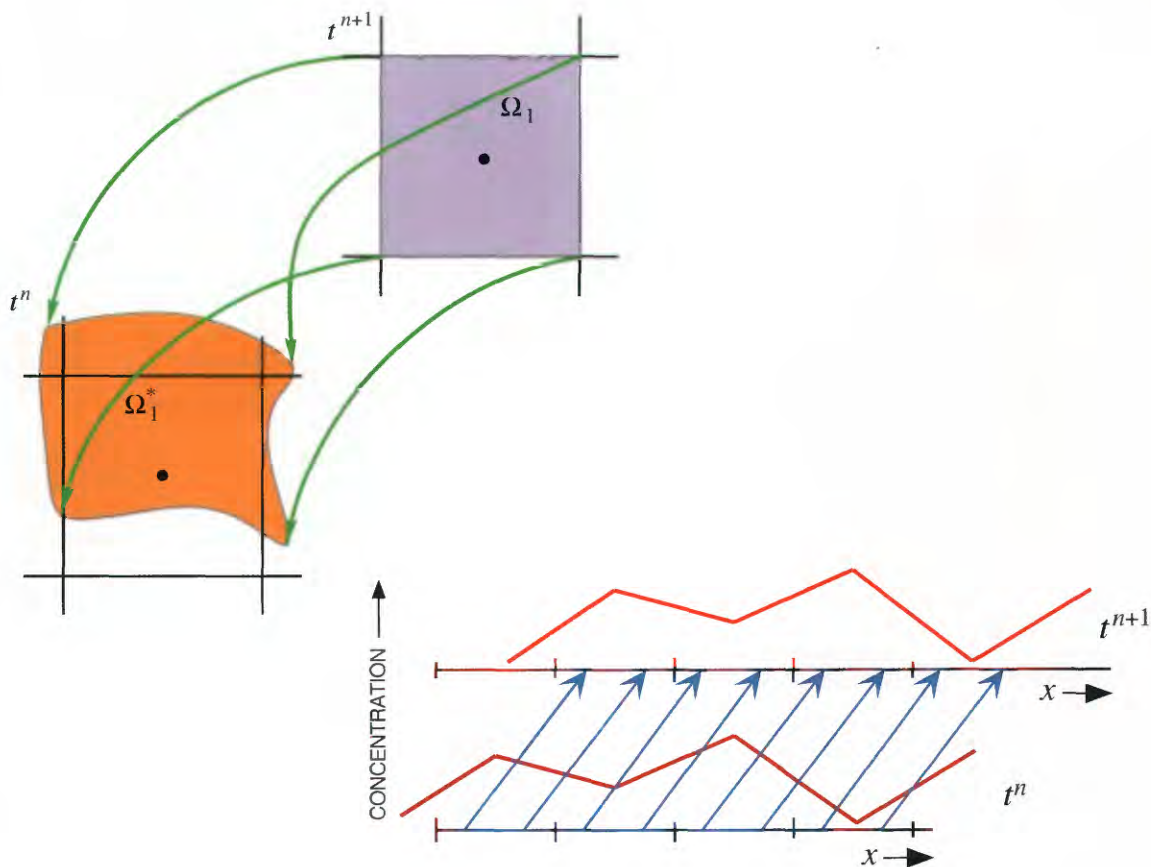


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A Three-Dimensional Finite-Volume Eulerian-Lagrangian Localized Adjoint Method (ELLAM) for Solute-Transport Modeling



U.S. GEOLOGICAL SURVEY
Water-Resources Investigations Report 00-4087

Cover: Illustrations of backtracking and forward tracking approaches for calculating changes in concentration of a solute caused by advective transport (see figures 5 and 6 of this report).

A Three-Dimensional Finite-Volume Eulerian-Lagrangian Localized Adjoint Method (ELLAM) for Solute-Transport Modeling

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U.S. GEOLOGICAL SURVEY

Water-Resources Investigations Report 00-4087

Reston, Virginia
2000



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PREFACE

The *MOC3D* computer code simulates the transport of a single solute in ground water that flows through porous media. The model is a package for the U.S. Geological Survey (USGS) *MODFLOW* ground-water model.

The new algorithm (*ELLAM*) documented in this report solves an integral form of the solute-transport equation, which is an expression of local conservation of mass. The code incorporates an implicit difference approximation in time for the dispersive term. Thus *ELLAM* offers an alternative to previously documented particle-tracking methods of *MOC3D*. It yields a precise global mass balance and can provide qualitatively good results for advection-dominated systems having very high Courant numbers at low computational cost.

The new *MOC3D* code is available for downloading over the Internet from a USGS software repository. The repository is accessible on the World Wide Web (WWW) from the USGS Water-Resources Information Web page at <http://water.usgs.gov/>. The site for the public repository is: <http://water.usgs.gov/software/>. The public anonymous FTP site is on the Water-Resources Information server (water.usgs.gov or 130.11.50.175) in the /pub/software directory. The code and documentation will also be available through an alternative web page for USGS ground-water models at <http://water.usgs.gov/nrp/gwsoftware/>. When this code is revised or updated in the future, new releases will be made available at these same sites. The code that incorporates the *ELLAM* algorithm, as documented in this report, is designated as Version 3.5.

Although extensive testing of the *ELLAM* algorithm indicates that this model will yield reliable calculations for a wide variety of field problems, the user is cautioned that the accuracy and efficiency of the model can be affected significantly for certain combinations of parameter values and boundary conditions.

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rearrange terms. The system of equations to be solved is then:

$$\begin{aligned} & \int_{\Omega_l} (\varepsilon C)^{n+1} d\mathbf{x} - \Delta t \int_{\partial\Omega_l} \frac{1}{R_f} (\varepsilon \mathbf{D} \nabla C)^{n+1} \cdot \mathbf{n} ds = \\ & e^{-\lambda \Delta t} \int_{\Omega_l^*} (\varepsilon C)^n d\mathbf{x} \\ & - \iint_{\text{supp } u_l \cap \Gamma^{n+1}} e^{-\lambda(t^{n+1}-t)} \varepsilon C_{\text{inflow}} \frac{\mathbf{V}}{R_f} \cdot \mathbf{n} dt ds \\ & + \iint_{\text{supp } u_l \cap \text{supp } W} e^{-\lambda(t^{n+1}-t)} \sum C' \frac{W}{R_f} dt d\mathbf{x} \quad (10) \end{aligned}$$

where Ω_l^* means the pre-image in the spatial domain at t^n of Ω_l at t^{n+1} ; $\Gamma^{n+1} \equiv \partial\Omega \times (t^n, t^{n+1})$ is the space-time boundary at time step $n+1$; and $\text{supp } f \equiv \{x \mid f(x) \neq 0\}$. These integral equations are solved for C^{n+1} , the concentration at the new $(n+1)$ time level at each cell center. Note that the right-hand side of eq. 10 consists of advective mass contributions from storage (that is, advection of mass in the domain at the start of the time increment), inflow boundaries, and sources. This is illustrated schematically for a simple case having constant velocity in figure 3, which shows how mass is advected into cell Ω_l at time level t^{n+1} from an inflow boundary, from a fluid source in a nearby cell, and from the mass present at time level t in nearby cells.

Outflow Boundary Equations

The integral in eq. 7 expressing mass crossing the transport subdomain boundary during a time step is:

$$\begin{aligned} & \int_{\Omega} \int_{t^n}^{t^{n+1}} \frac{1}{R_f} \nabla \cdot (u \varepsilon C \mathbf{V} - u \varepsilon \mathbf{D} \nabla C) dt d\mathbf{x} = \\ & \int_{t^n}^{t^{n+1}} \int_{\partial\Omega} \frac{1}{R_f} (u \varepsilon C \mathbf{V} - u \varepsilon \mathbf{D} \nabla C) \cdot \mathbf{n} ds dt \quad (11) \end{aligned}$$

Considering just the outflow portion of the boundary, this becomes

$$\begin{aligned} & \int_{t^n}^{t^{n+1}} \int_{(\partial\Omega)_{\text{outflow}}} \frac{1}{R_f} (u \varepsilon C \mathbf{V} - u \varepsilon \mathbf{D} \nabla C) \cdot \mathbf{n} ds dt \\ & \approx \int_{(\partial\Omega)_{\text{outflow}}} \int_{t^n}^{t^{n+1}} \left(u \varepsilon C_{\text{outflow}} \frac{\mathbf{V}}{R_f} \right) \cdot \mathbf{n} dt ds \quad (12) \end{aligned}$$

where total flux across the boundary is now approximated by advective flux.

We index the outflow boundary faces with ll and define the following test functions:

$$u_{ll} = \begin{cases} e^{-\lambda(t^{n+1}-t)} & \text{on characteristics from } \Omega \\ & \text{at time level } n \text{ into boundary} \\ & \text{area } (\partial\Omega)_{ll} \text{ at any time} \\ & \text{during time step} \\ 0 & \text{otherwise.} \end{cases} \quad (13)$$

The mass across outflow boundary face ll is the mass stored at the previous time level that flows across the face, together with any inflow and source mass that both enters the transport subdomain and leaves through face ll during the time step.

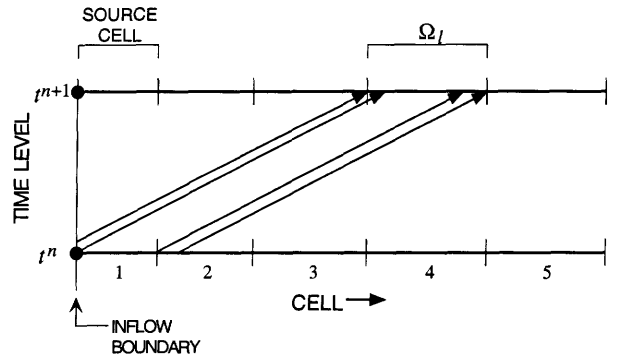


Figure 3. Schematic representation (for a simple case having constant velocity) showing how mass is advected into cell Ω_l at time level t^{n+1} from the inflow boundary, from a fluid source in cell 1, and from storage at time level t^n in cells 1 and 2.

Taking $u = u_{II}$ on the right-hand side of eq. 12, and including those terms from eq. 7 that are appropriate in the context of the

outflow boundary, we can write three terms representing mass contributions from storage, inflow, and sources, as follows:

$$\begin{aligned}
& \int_{(\partial\Omega)_{II}} \int_{t^n}^{t^{n+1}} \left(e^{-\lambda(t^{n+1}-t)} \varepsilon C_{\text{outflow}} \frac{\mathbf{V}}{R_f} \right) \cdot \mathbf{n} dt ds = e^{-\lambda\Delta t} \int_{(\partial\Omega)_{II}^*} (\varepsilon C)^n dx \\
& - \iint_{\text{supp } u_{II} \cap \partial\Omega_{\text{inflow}} \times (t^n, t^{n+1})} e^{-\lambda(t^{n+1}-t)} \varepsilon C_{\text{inflow}} \frac{\mathbf{V}}{R_f} \cdot \mathbf{n} dt ds \\
& + \iint_{\text{supp } u_{II} \cap \text{supp } W \times (t^n, t^{n+1})} e^{-\lambda(t^{n+1}-t)} \sum C' \frac{W}{R_f} dt dx
\end{aligned} \tag{14}$$

where $(\partial\Omega)_{II}$ is a discretized portion of the boundary face, and $(\partial\Omega)_{II}^*$ is its pre-image, in the manner described following eq. 9. The advection of mass to an outflow boundary is illustrated schematically for a simple case having constant velocity in figure 4, where mass is advected to an outflow boundary at time level t^{n+1} from an inflow boundary, from a fluid source in a nearby cell, and from the mass present at time level t in nearby cells.

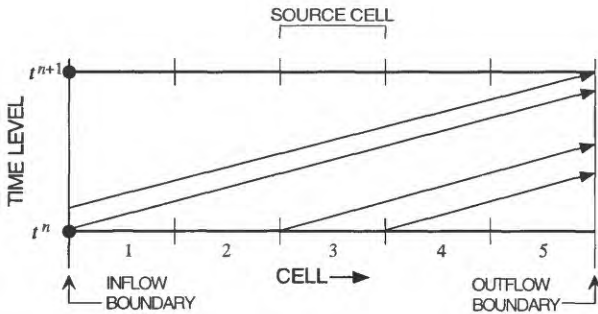


Figure 4. Schematic representation showing how mass is advected to an outflow boundary at time level t^{n+1} from the inflow boundary, from a fluid source in cell 3, and from storage at time level t^n in all five cells.

ELLAM equations are a formulation of mass conservation on each cell. Therefore, approximations to concentrations result that conserve mass locally (on each finite-

difference cell) and globally (on the entire transport subdomain).

Mass Tracking

For each cell in the fixed finite-difference grid, the integrals on the right-hand side of eq. 10 represent solute mass advected into the cell during the time step from storage (that is, advection of mass in the domain at the start of the time increment), the transport subdomain boundary, or a fluid source, respectively.

Advection in flowing ground water is simulated by mass tracking along the characteristic curves determined by the seepage velocity. Calculation of advective movement during a flow time step is based on the specific discharges computed at the end of the step.

As in *MOC3D*, tracking is performed using linear interpolation of velocity in the direction of the component of interest and piecewise-constant interpolation in the other two directions. The approach is to solve a system of three ordinary differential equations to find the characteristic curves $[x = x(t), y = y(t), \text{ and } z = z(t)]$ along which fluid is advected:

$$\frac{dx}{dt} = \frac{V_x}{R_f} \tag{15}$$

$$\frac{dy}{dt} = \frac{V_y}{R_f} \quad (16)$$

$$\frac{dz}{dt} = \frac{V_z}{R_f} \quad (17)$$

This is accomplished by introducing a set of moving points that can be traced within the stationary coordinates of a finite-difference grid. Each point corresponds to one characteristic curve, and values of x , y , and z are obtained as functions of t for each characteristic (Garder and others, 1964). Each point moves through the flow field by the flow velocity acting along its trajectory.

The *ELLAM* equations, eqs. 10 and 14, suggest that mass is tracked backwards along characteristics to the pre-image of each cell or boundary face. It is not possible, however, to exactly locate all of the mass at the previous time level by backtracking a finite number of points (see figure 5). In order to achieve mass

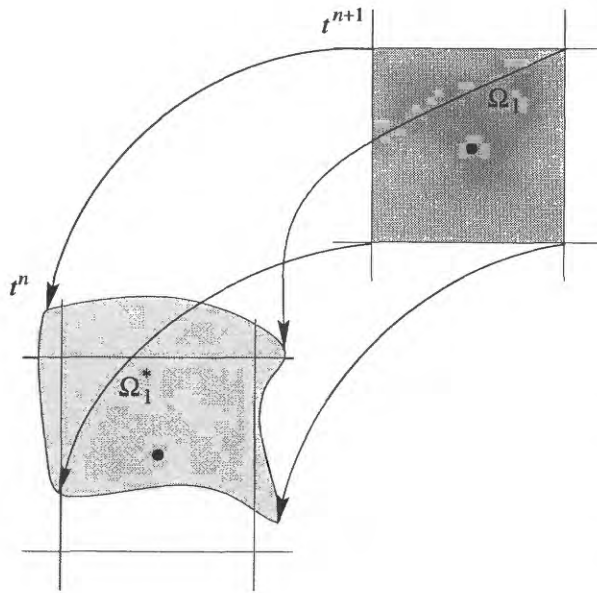


Figure 5. Two-dimensional example illustrating that pre-image of a cell may be irregularly shaped and not easily defined by backtracking from t^{n+1} to t^n .

balance, this implementation of the *ELLAM* algorithm tracks the known mass distribution forward from the old time level to the new time level (see figure 6). The accuracy of point tracking can be related to the Courant number, which is the ratio of (1) the distance a point will move in one time increment (velocity times Δt) to (2) the grid spacing (Δx).

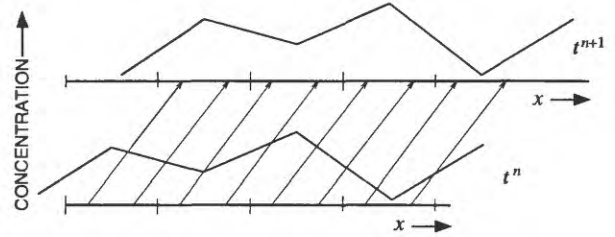


Figure 6. Plots of solute concentration versus distance that illustrate one-dimensional advection of known mass distribution from old time level to new time level (Courant number = 1).

ELLAM tracks points that are the centers of volumes of fluid. Thus, mass in a fluid volume is tracked under advection during a time step, distributed among destination cells, and accumulated to the right hand side storage, inflow, or source integral for each cell.

Decay

When simulating linear decay, all mass in the system at the beginning of each transport time step is decayed over the entire time step by a factor $e^{-\lambda \Delta t}$, where λ is the decay rate. Inflow and source mass are decayed in the same way, where the time interval is now not the entire time step, but the part of it during which new mass is in the transport subdomain.

This decay algorithm has no numerical stability restrictions associated with it. If the half-life is on the order of, or smaller than, the transport time step, however, some accuracy will be lost.

When a solute subject to decay enters the aquifer through a fluid source, it is assumed that the fluid source contains the solute at the concentration specified by C' . *MOC3D* allows decay to occur only within the ground-water system, and not within the source reservoir. In other words, for a given stress period, C' remains constant in time. If the problem being simulated requires that the solute in the source fluid itself undergo decay, then the source code will have to be modified.

Numerical Integration

The numerical treatment of each term in eqs. 10 and 14 will next be

$$\Delta t \int_{\partial \Omega_i} \frac{1}{\epsilon R_f} (\epsilon \mathbf{D} \nabla C)^{n+1} \cdot \mathbf{n} ds \approx \frac{\Delta t}{(R_f)_k (\epsilon b)^{n+1}_{j,i,k}} \left\{ \left[\left(\epsilon b D_{1m} \frac{\partial C}{\partial x_m} \right)^{n+1}_{j+1/2,i,k} - \left(\epsilon b D_{1m} \frac{\partial C}{\partial x_m} \right)^{n+1}_{j-1/2,i,k} \right] \Delta y_i b^{n+1}_{j,i,k} \right. \\ + \left[\left(\epsilon b D_{2m} \frac{\partial C}{\partial x_m} \right)^{n+1}_{j,i+1/2,k} - \left(\epsilon b D_{2m} \frac{\partial C}{\partial x_m} \right)^{n+1}_{j,i-1/2,k} \right] \Delta x_j b^{n+1}_{j,i,k} \\ \left. + \left[\left(\epsilon b D_{3m} \frac{\partial C}{\partial x_m} \right)^{n+1}_{j,i,k+1/2} - \left(\epsilon b D_{3m} \frac{\partial C}{\partial x_m} \right)^{n+1}_{j,i,k-1/2} \right] \Delta x_j \Delta y_i \right\} \quad (18)$$

where $m = 1,2,3$ is a summation index for the dispersion term. Finite-difference approximations to the space derivatives in the dispersion integral are calculated using centered differences as in *MOC3D*, but with *ELLAM* they are modified for varying grid dimensions. (See Konikow and others (1996), p. 64 for $\epsilon b D_m \frac{\partial C}{\partial x_m}$ expansion.)

Mass Storage at New Time Level

The quantity mass/porosity in a cell at the new time level t^{n+1} is expressed using the trapezoidal rule for integration, formulated over each cell octant. Concentrations at octant corners are weighted averages of

discussed. The j,i,k subscripts for a cell Ω_i will denote the spatial finite-difference grid indexing, as discussed previously in the section "Numerical Methods."

The equations are first divided through by porosity, which is represented by piecewise constants in space and time. This is valid because there are no spatial derivatives of porosity in the local *ELLAM* equations.

Dispersion

Time integration is accomplished using a one point in time backward Euler rule. Spatially, a one point integration rule with a seven point stencil is used:

neighboring node concentrations, determined by trilinear interpolation.

For each octant,

$$\frac{\text{mass}}{\text{porosity}} = \frac{1}{64} \Delta x_j \Delta y_i b^{n+1}_{j,i,k} \sum_{corner=1}^8 C_{corner} \\ = \frac{1}{64} \Delta x_j \Delta y_i b^{n+1}_{j,i,k} \sum_{corner=1}^8 \sum_{nbr=1}^8 (weight)_{nbr} C_{nbr} \quad (19)$$

where for an interior octant, $nbr = neighbor$ and nbr is one of the eight grid nodes between which concentration varies trilinearly. In the case of a boundary octant, a boundary face value is needed for calculation, and is taken to be the following:

- Inflow: user input;
- No flow: same as associated interior node; and
- Outflow: calculated using cell parameters, boundary flow rate, and mass tracked across boundary during transport time step.

One important implication of the above procedure, which is designed to be mass conservative, is that the concentration calculated and reported (in the output) for the location of the block-centered node, $C_{j,i,k}$, represents the estimated concentration at that point and not the average concentration in the cell. Thus, unlike many other numerical methods, $C_{j,i,k}$ multiplied by the volume of water in the cell would not necessarily equal the solute mass in the cell.

Coefficients calculated by $((1/8) \times \text{octant volume} \times \text{nodal weight})$ for all nodes neighboring a cell comprise the storage matrix entries for the equation for each cell. Boundary terms are put on the right-hand side of the equation because all boundary face concentrations are determined before the solution of the interior equations.

It should be noted that linear interpolation in the vertical dimension is approximate in the case where adjacent cells in the same layer of the transport subdomain have varying thicknesses, as is allowed by *MODFLOW*. Extreme variations could affect accuracy of the solution.

For an interior cell with all neighbors active and using b at time $n+1$:

$$\begin{aligned}
 \int_{\Omega_i} C^{n+1} d\mathbf{x} = & \left(\left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) + \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) - 4 \right) \left(\left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) + \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) - 4 \right) \\
 & \left(\left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k-1}} \right) + \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k+1}} \right) - 4 \right) C_{j,i,k} \\
 & - \left(\left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) \left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k-1}} \right) C_{j-1,i-1,k-1} + \left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) \left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k+1}} \right) C_{j-1,i-1,k+1} \right. \\
 & + \left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k+1}} \right) C_{j-1,i+1,k+1} + \left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k-1}} \right) C_{j-1,i+1,k-1} \\
 & + \left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k-1}} \right) C_{j+1,i-1,k-1} + \left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k+1}} \right) C_{j+1,i-1,k+1} \\
 & + \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k-1}} \right) C_{j+1,i+1,k-1} + \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k+1}} \right) C_{j+1,i+1,k+1} \Big) \\
 & + \left(\left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) + \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) - 4 \right) \left(\left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k-1}} \right) C_{j,i-1,k-1} + \left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k+1}} \right) C_{j,i-1,k+1} \right. \\
 & + \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k-1}} \right) C_{j,i+1,k-1} + \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k+1}} \right) C_{j,i+1,k+1} \Big) \\
 & + \left(\left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) + \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) - 4 \right) \left(\left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k+1}} \right) C_{j-1,i,k+1} + \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k-1}} \right) C_{j+1,i,k-1} \right. \\
 & + \left. \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k+1}} \right) C_{j+1,i,k+1} + \left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k-1}} \right) C_{j-1,i,k-1} \right)
 \end{aligned}$$

$$\begin{aligned}
& + \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k-1}} + \frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k+1}} - 4 \right) \left(\left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) C_{j+1,i-1,k} + \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) C_{j+1,i+1,k} \right. \\
& + \left. \left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) \left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) C_{j-1,i-1,k} + \left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) C_{j-1,i+1,k} \right) \\
& - \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k-1}} \right) \left(\left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) \left(\left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) + \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) - 4 \right) + \left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) \left(\left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) - 4 \right) \right. \\
& + \left. \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) \left(\left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) - 4 \right) - 4 \left(\left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) - 4 \right) \right) C_{j,i,k-1} \\
& - \left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k+1}} \right) \left(\left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) \left(\left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) + \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) - 4 \right) + \left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) \left(\left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) - 4 \right) \right. \\
& + \left. \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) \left(\left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) - 4 \right) - 4 \left(\left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) - 4 \right) \right) C_{j,i,k+1} \\
& - \left(\left(\frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k-1}} + \frac{b_{j,i,k}}{b_{j,i,k} + b_{j,i,k+1}} - 4 \right) \left(\left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) + \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) - 4 \right) \left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) C_{j,i-1,k} \right. \\
& + \left. \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) C_{j,i+1,k} \right) \left(\left(\frac{\Delta y_i}{\Delta y_{i-1} + \Delta y_i} \right) + \left(\frac{\Delta y_i}{\Delta y_{i+1} + \Delta y_i} \right) - 4 \right) \left(\left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) C_{j-1,i-1,k} + \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) C_{j+1,i,k} \right) \quad (20)
\end{aligned}$$

Mass Storage at Old Time Level

The total mass advected into each cell during a transport time step that was already stored within the system at the old time level is needed for the right-hand side of the *ELLAM* equation. Numerically, this is accomplished by tracking mass forward from the old time level, n , along characteristics. Each cell is divided into subcells determined by parameters NSC, NSR, NSL, specifying the number of subcells in the column, row, and layer direction, respectively. The center of each subcell is tracked through the time step under advection. Depending on the exact location of this point in the destination cell at the new time, all of the mass in the subcell may or may not also be found in that destination cell. In order to mitigate the effects of unwarranted mass lumping, subcell mass is distributed among cells neighboring the destination cell using the “approximate test functions,” w_l , described below. The value of w_l at the subcell center destination point is the fraction of subcell mass to be

distributed to cell Ω_l .

This yields the formulation,

$$\begin{aligned}
& e^{-\lambda \Delta t} \int_{\Omega_i^*} C^n d\mathbf{x} \approx \\
& e^{-\lambda \Delta t} \sum_{j,i,k} \sum_{\substack{p=\text{subcell} \\ \text{center}}} \left[\frac{\Delta x_j \Delta y_i b_{j,i,k}}{(NSC)(NSR)(NSL)} \right. \\
& \quad \left. \left(w_l(p^f) C(p) \right) \right] \quad (21)
\end{aligned}$$

where summation runs through all subcells of each cell in the transport subdomain, and p^f is the image of p under forward tracking to the new time level.

Approximate Test Functions

An approximate test function is defined for each active cell for the purpose of distributing advected mass among neighboring cells. The designation “approximate test

function” is given because the graph of this function looks like a characteristic (indicator) function with slanted sides extending into adjacent cells, whereas the test functions described in the derivation of the governing equation are exactly characteristic functions in space at time t^{n+1} . Examples of approximate test functions are illustrated in figure 7 for one direction. An approximate test function is determined by NSC, NSR, and NSL, the proximity of the transport subdomain boundary, and the active status of neighboring cells. Mass is not split across the transport boundary or into inactive cells.

We define local reference coordinates $\hat{x}, \hat{y}, \hat{z}$ centered around cell Ω_l with node indices j, i, k by

$$\hat{x} = \frac{x - x_j}{\Delta x_j} \quad (22)$$

and similarly for \hat{y} and \hat{z} . For $\hat{x}, \hat{y}, \hat{z} \in \left(-\frac{1}{2}, \frac{1}{2}\right)$, the corresponding point (x, y, z) is in cell Ω_l . For an interior cell on a uniform grid with all surrounding cells active, one approximate test function is

$$w_{jik}(\hat{x}, \hat{y}, \hat{z}) = f(\hat{x})g(\hat{y})h(\hat{z}) \quad (23)$$

where

$$f(\hat{x}) = \begin{cases} 0 & \hat{x} \leq -\frac{1}{2} - \frac{1}{2NSC} \\ NSC\hat{x} + \frac{1}{2}(NSC+1) & -\frac{1}{2} - \frac{1}{2NSC} < \hat{x} < -\frac{1}{2} + \frac{1}{2NSC} \\ 1 & -\frac{1}{2} + \frac{1}{2NSC} \leq \hat{x} \leq \frac{1}{2} - \frac{1}{2NSC} \\ -NSC\hat{x} + \frac{1}{2}(NSC+1) & \frac{1}{2} - \frac{1}{2NSC} < \hat{x} < \frac{1}{2} + \frac{1}{2NSC} \\ 0 & \frac{1}{2} + \frac{1}{2NSC} \leq \hat{x} \end{cases}$$

and similarly for g and h . This function, in one direction on a uniform grid, is shown graphically in figures 7a and 7b. Approximate test functions in each direction are multiplied together (see eq. 23) to get the test

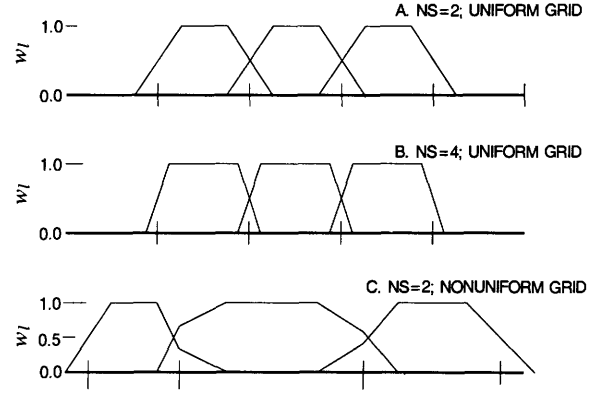


Figure 7. Examples of spatial distribution of approximate test functions (w_l) for selected one-dimensional cases. Vertical ticks represent cell boundaries.

$$\begin{aligned} \hat{x} &\leq -\frac{1}{2} - \frac{1}{2NSC} \\ -\frac{1}{2} - \frac{1}{2NSC} &< \hat{x} < -\frac{1}{2} + \frac{1}{2NSC} \\ -\frac{1}{2} + \frac{1}{2NSC} &\leq \hat{x} \leq \frac{1}{2} - \frac{1}{2NSC} \\ \frac{1}{2} - \frac{1}{2NSC} &< \hat{x} < \frac{1}{2} + \frac{1}{2NSC} \\ \frac{1}{2} + \frac{1}{2NSC} &\leq \hat{x} \end{aligned} \quad (24)$$

functions used to distribute the advected mass.

In the general case of a possibly nonuniform grid, the single variable functions f , g , and h are given by:

$$f(\hat{x}) = \begin{cases} 0 & \hat{x} \leq -\frac{1}{2} - \frac{1}{2NSC} \\ 1 + 2NSC \left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) (1 + \hat{x}) - \left(1 - \frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) - NSC \left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) & -\frac{1}{2} - \frac{1}{2NSC} < \hat{x} < -\frac{1}{2} \\ 2NSC \left(1 - \left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) \right) \hat{x} + \left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) + NSC \left(1 - \left(\frac{\Delta x_j}{\Delta x_{j-1} + \Delta x_j} \right) \right) & -\frac{1}{2} \leq \hat{x} < -\frac{1}{2} + \frac{1}{2NSC} \\ 1 & -\frac{1}{2} + \frac{1}{2NSC} \leq \hat{x} \leq \frac{1}{2} - \frac{1}{2NSC} \\ 2NSC \left(\left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) - 1 \right) \hat{x} + \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) + NSC \left(1 - \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) \right) & \frac{1}{2} - \frac{1}{2NSC} < \hat{x} < \frac{1}{2} \\ 1 + 2NSC \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) (1 - \hat{x}) - \left(1 - \frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) - NSC \left(\frac{\Delta x_j}{\Delta x_{j+1} + \Delta x_j} \right) & \frac{1}{2} < \hat{x} < \frac{1}{2} + \frac{1}{2NSC} \\ 0 & \frac{1}{2} + \frac{1}{2NSC} \leq \hat{x} \end{cases} \quad (25)$$

with g and h defined analogously, where for points outside of Ω_l , $\hat{x}, \hat{y}, \hat{z}$ must be defined according to the scale of the appropriate cell; for example, in the neighboring cell having coordinates $j-1$, we have

$$\hat{x} \in \left(-\frac{3}{2}, -\frac{1}{2}\right), \quad \hat{x} = -1 + \frac{x - x_{j-1}}{\Delta x_{j-1}}. \quad \text{This}$$

function is shown graphically in figure 7c.

In practice, to evaluate all test functions at a given point in any Ω_l , use cell coordinates j, i, k and reference coordinates $\hat{x}, \hat{y}, \hat{z} \in (-\frac{1}{2}, \frac{1}{2})$ for that point within that cell. Then eqs. 25 are used to evaluate f, g, h and eq. 23 to find $w_{j,i,k}$. If $\hat{x}, \hat{y}, \hat{z} > 0$, there are potentially seven more nonzero test functions, with values given by:

$$w_{j+1,i,k} = (1 - f(\hat{x}))g(\hat{y})h(\hat{z}) \quad (26a)$$

$$w_{j,i+1,k} = f(\hat{x})(1 - g(\hat{y}))h(\hat{z}) \quad (26b)$$

$$w_{j,i,k+1} = f(\hat{x})g(\hat{y})(1 - h(\hat{z})) \quad (26c)$$

$$w_{j+1,i+1,k} = (1 - f(\hat{x}))(1 - g(\hat{y}))h(\hat{z}) \quad (26d)$$

$$w_{j,i+1,k+1} = f(\hat{x})(1 - g(\hat{y}))(1 - h(\hat{z})) \quad (26e)$$

$$w_{j+1,i,k+1} = (1 - f(\hat{x}))g(\hat{y})(1 - h(\hat{z})) \quad (26f)$$

$$w_{j+1,i+1,k+1} = (1 - f(\hat{x}))(1 - g(\hat{y}))(1 - h(\hat{z})) \quad (26g)$$

Analogous expressions hold for reference coordinates with different signs. Test functions for neighboring cells are evaluated as in eq. 26, and $w_{l'} \equiv 0$ for all l' indexing cells not adjacent to Ω_l .

The sum of the values of the test functions at any point in the transport domain is one, thus conserving mass for the integral equation. This may be expressed as

$$\sum_l w_l(x) = 1.$$

The test function component in the direction of the boundary extends from the center of a boundary cell to the boundary face with the value of one. Thus, there is no splitting of mass across the boundary.

There is no test function associated with an inactive cell. In a cell adjacent to an inactive cell, the value of the test function that would normally be assigned to the inactive cell is distributed proportionally to other test functions that are nonzero at that point. All test functions are zero in inactive cells.

Extreme variation in cell thickness among neighboring cells in a layer may adversely affect model results. In this case, linear interpolation of concentration in the

vertical direction, or approximate distribution of advected mass, may be inaccurate.

Source Integral

Source and sink integrals correspond to the last term in eq. 10, divided through by porosity as noted above. In general, *MOC3D* assumes that a source or sink is distributed uniformly over the finite-difference cell containing it.

For a source, a time step (t^n, t^{n+1}) is discretized into NT uniform sub-time steps, $t_0 = t^n$, $t_{NT} = t^{n+1}$, $t_m = t^n + \Delta t m/NT$ for $m = 1, 2, \dots, NT-1$. Inflow of mass is integrated using the compound trapezoid rule. In effect, this means that the amount of mass associated with a sub-time step $\Delta t/NT$ is introduced at

$$\iint_{\text{supp } u_l \cap \text{supp } W} e^{-\lambda(t^{n+1}-t)} \sum C' \frac{W}{R_f} dt d\mathbf{x} \approx \frac{1}{\varepsilon_{jik}} \sum_{\substack{ALL \\ SOURCE \\ CELLS}} \sum_{\substack{source \\ p=subcell \\ center}} \sum_{m=0}^{NT} e^{-\lambda(t^{n+1}-t_m)} \frac{\Delta T_m}{(NSC)(NSR)(NSL)} w_l(p^f) \sum_s C'_s \frac{Q_s}{R_f} \quad (27)$$

where summation runs through all subcells of each source cell in the transport subgrid; p^f is the image of p under forward tracking to the new time level; t_m represents the time during the time step at which discretized source mass enters the system ($t_0 = t^n$, $t_{NT} = t^{n+1}$, $t_m = t^n + \Delta t m/NT$); and $\Delta T_m = \Delta t/NT$ or $\Delta T_m = \Delta t/2NT$ if m equals 0 or NT , s indexes all source terms within the source cell of interest, and Q_s is the volumetric flow rate associated with the fluid source having a concentration C'_s . Note that Q_s equals W_s multiplied by the volume of the source cell.

Sink Integral

Analytically, the domain of integration is the support of the space-time test function for a cell Ω_l intersected with any sink cells. To approximate, this term is only formulated if cell Ω_l contains a sink, and the sink concentration is assumed to be the average nodal concentration for the transport time

each of the times $t_1, t_2, \dots, t_{NT-1}$, and half this amount is introduced at each of t_0 and t_{NT} . Starting from its designated time, each packet of mass is tracked forward to time t^{n+1} and accumulated in the same manner as non-source mass that began the time step already in the transport subdomain.

This accumulation for the last term in eq. 10 is done with the following integration for cell Ω_l . To account for all source mass that flows into Ω_l during the time step, all sources that intersect the support of the space-time test function associated with Ω_l are included. Note that the integration determines the source mass flowing into Ω_l , not the source mass originating in Ω_l . Multiple sources within the same cell are summed. This yields:

step, with the exception of a sink related to evapotranspiration, where sink concentration is taken to be zero. Integration rules are a one point in space and a one point backward Euler in time. Multiple sinks within a sink cell are summed. The averaging of concentration results in this integral approximation contributing to both the left and right hand sides of the equation for sink cell Ω_l with coordinates j, i, k :

$$\begin{aligned} & \iint_{\text{supp } u_l \cap \text{supp } W} e^{-\lambda(t^{n+1}-t)} \sum C' \frac{W}{\varepsilon R_f} dt d\mathbf{x} \\ & \approx \Delta t \int_{\Omega_{\text{sink}}} \frac{1}{\varepsilon} \left(C_{\text{average}} \sum \frac{W}{R_f} + C_{\text{ET}} \frac{W_{\text{ET}}}{R_f} \right) d\mathbf{x} \\ & = \frac{\Delta t}{\varepsilon_{jik} R_f} \left[\frac{(C^{n+1} + C^n)}{2} \sum Q_{\text{sink}} + C_{\text{ET}} Q_{\text{ET}} \right] \quad (28) \end{aligned}$$

where the subscript ET refers to evapotranspirative flux.

Inflow Boundary Integral

Inflow boundary integrals correspond to the next-to-last term in eq. 10, divided through by porosity. For an inflow boundary, as for a source, a single time step is discretized into a number of sub-time steps determined by parameter NT . The composite trapezoidal rule is applied in time. At each sub-time step, inflow mass is spatially discretized, tracked, and accumulated, just

like mass already in the system at the start of the transport time step, but for the shorter interval. The only difference in the treatment of the inflow boundary from the treatment of the source is that only the two-dimensional boundary face is discretized, whereas for a source, the entire cell is discretized. For a cell Ω_l , the integration is performed over the intersection of the space-time test function for that cell and the transport subdomain boundary; that is, all mass entering through the boundary and advected to Ω_l during the time step is accumulated to the right-hand side of local equation l , yielding:

$$\begin{aligned} & \iint_{\text{supp } u_l \cap \Gamma^{n+1}} e^{-\lambda(t^{n+1}-t)} C_{\text{inflow}} \frac{\mathbf{V}}{R_f} \cdot \mathbf{n} dt ds \\ & \approx \sum_{\substack{\text{all} \\ \text{inflow} \\ \text{faces}}} \sum_{\substack{\text{face} \\ p=\text{subarea} \\ \text{center}}} \sum_{m=0}^{NT} e^{-\lambda(t^{n+1}-t_m)} \frac{\Delta T_m}{\text{ref area}} w_l(p^f) C_{\text{inflow}} \frac{Q_{\text{inflow}}}{R_f} \end{aligned} \quad (29)$$

where $\text{ref area} = \text{NSC} \times \text{NSR}$, $\text{NSC} \times \text{NSL}$, or $\text{NSR} \times \text{NSL}$, depending on plane of face; p^f is the image of p under forward tracking to the new time level; Q_{inflow} is the volumetric rate of inflow across the face; and t_m and ΔT_m are defined in eq. 27. Summation runs over each p on the transport subgrid boundary, with the approximate test function w_l used to select mass advected to Ω_l .

Outflow Integrals

Concentration is calculated at each outflow boundary face using cell parameters, velocity information from *MODFLOW*, and the amount of mass tracked across the cell boundary determined by *MOC3D*.

On the left-hand side of the system of boundary equations (eq. 14) is an integral approximated using a one point in space, one point backward Euler in time formulation. This time approximation eliminates the exponential factor, as follows:

$$\begin{aligned} & \int_{(\partial\Omega)_{ll}} \int_{t^n}^{t^{n+1}} e^{-\lambda(t^{n+1}-t)} C_{\text{outflow}} \frac{\mathbf{V}}{R_f} \cdot \mathbf{n} dt ds \\ & \approx \Delta t \int_{(\partial\Omega)_{ll}} C_{\text{outflow}} \frac{\mathbf{V}}{R_f} \cdot \mathbf{n} ds \\ & = \Delta t \frac{Q_{\text{outflow}}}{R_f} C_{\text{outflow}} \end{aligned} \quad (30)$$

where ll is the index for boundary faces; and Q_{outflow} is determined using the outflow velocity calculated from *MODFLOW* output and cell parameters. The concentration on face ll is the unknown in the boundary equation.

The right-hand side boundary integrals are constructed from the mass contributions tracked across the boundary from interior cells, sources, and inflow boundaries during the transport time step. All mass associated with a tracked point that reaches the outflow boundary at any time during the time step is

considered to leave the transport subdomain. Test functions are evaluated to distribute mass among neighboring boundary outflow faces.

The three terms on the right-hand side of eq. 14 are approximated in a manner analogous to eqs. 21, 29, and 27, respectively, where now outflow boundary integrals, instead of cell integrals, are being considered. We define test functions W_{ll} associated with outflow boundary faces analogous to the test functions W_l for cells. Thus, $W_{ll}(p^f)$ will be

nonzero if, tracking p^f forward from sub-time step t_m , p^f reaches the given outflow boundary face during time interval $[t_m, t^{n+1}]$. Spatially along the boundary, W_{ll} will have a profile analogous to those in figure 7, thus distributing mass to neighboring boundary faces. Substituting the right-hand side of eq. 30 for the left-hand side of eq. 14, approximating the right-hand side of eq. 14 as described above, and using a trapezoidal rule in time and a cell-midpoint rule in space, we obtain:

$$\begin{aligned} \Delta t \frac{Q_{\text{outflow}}}{R_f} C_{\text{outflow}} \approx & e^{-\lambda \Delta t} \sum_{j,i,k} \sum_{\substack{p=\text{subcell} \\ \text{center}}} \frac{\Delta x_j \Delta y_i b_{j,i,k}}{(NSC)(NSR)(NSL)} w_l(p^f) C(p) \\ & + \sum_{\substack{\text{all} \\ \text{inflow} \\ \text{faces}}} \sum_{\substack{\text{face} \\ p=\text{subarea} \\ \text{center}}} \sum_{m=0}^{NT} e^{-\lambda(t^{n+1}-t_m)} \frac{\Delta T_m}{\text{ref area}} w_l(p^f) C_{\text{inflow}} \frac{Q_{\text{inflow}}}{R_f} \\ & + \frac{1}{\epsilon_{jik}} \sum_{\substack{\text{ALL} \\ \text{SOURCE} \\ \text{CELLS}}} \sum_{\substack{\text{source} \\ p=\text{subcell} \\ \text{center}}} \sum_{m=0}^{NT} e^{-\lambda(t^{n+1}-t_m)} \frac{\Delta T_m}{(NSC)(NSR)(NSL)} w_l(p^f) \sum_s C'_s \frac{Q_s}{R_f} \end{aligned} \quad (31)$$

We thus have a system of equations represented by a diagonal matrix, to be solved for C_{outflow} .

Accuracy Criteria

An accuracy criterion incorporated in *MOC3D* constrains the distance that solute mass is advected during each transport time step. A restriction can be placed on the size of the time step to ensure that the number of grid cells a point moves in the x -, y -, or z -directions does not exceed some maximum. The simulator allows the user to specify this maximum (named *CELDIS* in the code and input instructions). This translates into a limitation on the transport time-step length. If the time step used to solve the flow equation exceeds the time limit, the flow time step will be subdivided into an appropriate number of

equal-sized smaller time increments for solving transport.

For advective transport, a mesh density sufficient so that at least four grid nodes are represented across a solute front (or zone of relatively steep concentration gradient) is needed for good accuracy. Similarly, for advecting a peak concentration, the area of the peak should be represented across at least eight nodes of the grid for good accuracy. In such cases, testing suggests that a peak concentration value can be advected with a very small dissipation of the maximum concentration per time step for a variety of

Courant numbers. With insufficient mesh density, a peak will dissipate rapidly for an initial period of time during which it spreads out and oscillates; thereafter, the numerical decay slows and the oscillations do not worsen. A fine discretization of tracked mass (large NSC, NSR, NSL) reduces the rate of peak decay when modeling with many transport time steps (see sections “Special Problems” and “Input Parameter Values”). Regardless of the solution accuracy, global mass is conserved.

The accuracy of the dispersion calculation is governed in part by the accuracy of the central-difference approximations to the space derivatives, meaning a finer mesh will result in better accuracy. The implicit formulation for the solution of the dispersion equation is unconditionally stable. This allows for large time steps during the simulation. Because *ELLAM* solves for advection along characteristics, thus avoiding large values of the second time derivative of the solution at passage of a steep front, error in calculation of the time derivative may be expected to be small compared to a standard finite-difference solution to an advection-dispersion equation. Some dependence of the accuracy of the

dispersion calculation on the size of the time step is retained, however. Note that stability does not imply accuracy; accuracy of the solution to the dispersion equation decreases as the time step size increases. On the other hand, modeling with many time steps in order to resolve dispersion to the desired accuracy could result in a loss of peak to numerical dispersion inherent in the treatment of advection, an effect that can be reduced by increasing NSC, NSR, and NSL.

One additional difficulty encountered with implicit temporal differencing results from the use of a symmetric spatial differencing for the cross-derivative terms of the dispersion tensor. This creates a potential for overshoot and undershoot in the calculated concentration solution, particularly when the velocity field is oblique to the axes of the grid. A remedy for excessive overshoot and undershoot is to refine the finite-difference mesh. This may, however, increase simulation times.

ELLAM can produce qualitatively good results in a small number of time steps, provided the NT value is sufficient to yield smooth distribution of mass along the inflow path. (See sections “Special Problems” and “Input Parameter Values”).

Mass Balance

As described by Konikow and others (1996), global mass-balance calculations are ordinarily performed to help check the numerical accuracy and precision of the solution. Modifications to the previously described mass-balance calculations have been implemented to assure consistency with the implicit algorithm. In calculating the cumulative mass flux out of the system, the explicit procedure assumes that the concentration associated with a fluid sink is $C_{j,i,k}^n$, the node concentration at the beginning of the time increment (see Konikow and others, 1996, eq. 66). The *ELLAM* code

assumes that the concentration associated with a flux in or out of the system is the average nodal concentration during the time increment, $(C_{j,i,k}^n + C_{j,i,k}^{n+1})/2$.

ELLAM conserves mass globally, regardless of the accuracy of the solution. Mass balance errors of less than 10^{-4} percent can generally be expected (this depends on the value of approximate zero, which is FORTRAN variable AZERO in the code, and solver tolerance, both of which are currently predefined in the code and cannot be specified in the input data).

Special Problems

Fronts too sharp for the given mesh density (grid spacing) may produce negative concentration values and/or numerical dispersion. For the mass in one cell to be positive, the mass in the adjacent cell to be zero, and concentration to vary linearly, the cell with zero mass may show a negative concentration. For non-integer Courant numbers, numerical dispersion results from the solution algorithm being insensitive to the exact location of advected mass in a destination cell. On a well-discretized front, these effects are minimal due to error cancellation. Thus, solving on a fine grid with few time steps may mitigate these difficulties.

Numerical dispersion may also result from tracking subdivisions of mass that are too coarse. The level of discretization of mass tracked and accumulated to the right-hand side vector is determined by parameters, NSC, NSR, and NSL. These parameters define the number of subcells in the column, row, and layer direction, respectively. To increase the resolution of mass tracking under

advection, it may be desirable to increase the values of these parameters.

Parameter NT defines the number of sub-time steps per transport time increment. NT should be large enough so that all cells in the path of flow from the inflow boundary or source to the location of the front at the end of the time step receive incoming mass. This is to avoid artificial mass lumping. See section "Input Parameter Values."

To avoid non-physical accumulation of mass at an outflow boundary, the spatial NS parameter in the direction normal to the boundary must be such that $1/(2NS) < \text{Courant}$. This is to ensure that at least some mass is calculated by the algorithm as reaching the boundary during a time step.

Extreme variation in cell thickness among neighboring cells in a layer may adversely affect model results. This is caused by the inherent geometric inconsistency in the vertical direction between adjacent cells that have different thicknesses. (Also see McDonald and Harbaugh, 1988, figure 9 and related discussion.)

Review of *ELLAM* Assumptions

The assumptions that have been incorporated into the *ELLAM* simulator are very similar to those for *MOC3D* Version 1 and Version 2. They are relevant to both grid design and model application. Efficient and accurate application of *ELLAM* requires the user to be aware of these assumptions. Therefore, the user should review the description of these items as presented by Konikow and others (1996).

Transport subgrid boundaries are assumed to be far enough from the plume that any errors in the treatment of the boundaries will not have a significant effect on the solution. The boundary condition is that the normal component of the

concentration gradient on the boundary is zero, meaning there is no dispersive flux across the transport subdomain boundary.

Unlike the previous *MOC3D* explicit and implicit difference approximations, *ELLAM* does not require a uniform grid spacing within the transport subdomain. Likewise, there is no longer a formal restriction on variations in the product of porosity and thickness within the subdomain.

ELLAM does assume:

- Concentration at an outflow boundary face at the new time level is well approximated by the mass crossing the face during the

time step divided by the fluid volume across the face.

- Mass in or out of the transport subdomain during a time step via a source or sink cell is well approximated by the average nodal concentration during the

time step times the fluid volume through the source or sink. Mass loss in flow through upstream sinks is negligible.

- Cell thicknesses are smoothly varying within a horizontal layer.

COMPUTER PROGRAM

ELLAM is implemented as a package for *MODFLOW*. *ELLAM* uses the flow components calculated by *MODFLOW* to compute velocities across each cell face in the transport domain. The computed velocities are used in an interpolation scheme to move each mass-bearing volume a distance and direction with time to represent advection. An integral formulation of conservation of mass is applied, yielding a dispersion equation including boundary fluxes, fluid sources, and decay.

Because the model is based on the assumption that the fluid properties (such as density and viscosity) are constant and uniform and independent of changes in concentration, the head distribution and flow field are independent of the solution to the solute-transport equation. Therefore, the flow and transport equations can be solved sequentially, rather than simultaneously. Because transport depends on fluid velocity, which is calculated from the solution to the flow equation, the flow equation must be solved first.

A separate executable version of *MODFLOW*, which is adapted to link with and use the *ELLAM* package, must first be created to run the simulations. *MOC3D* is written in standard FORTRAN-77, and it has been successfully compiled and executed on multiple platforms, including Pentium-based personal computers, Macintosh personal computers, and Data General, Sun, and Silicon Graphics Unix workstations. FORTRAN compilers for each of these platforms vary in their characteristics and may require the use of certain options to

compile *MOC3D* successfully. For instance, the compiler should initialize all variables to zero. Depending on the size of the X-array (defined by LENX in the *MODFLOW* source code), options to enable the compiler to handle large-array addressing may be needed. Most real variables in *MOC3D* are defined as single precision variables in the FORTRAN code. In our experience, use of double-precision definitions for these variables has not been necessary.

Implementing *ELLAM* requires the use of a separate "name" file that contains file names, similar to the one used in *MODFLOW*. The principal *ELLAM* input data (such as subgrid dimensions, hydraulic properties, and particle information) are read from the main *ELLAM* data file. Other files are used for observation wells, concentrations in recharge, and several input and output options. Detailed input-data requirements and instructions are presented in Appendix A. *ELLAM* input requirements differ from those of *MOC3D* in that NSCEXP, NSREXP, NSLEXP, and NTEXP values must be provided, whereas parameter values related to fluid particles are omitted.

The input data set used for the first test problem (involving one-dimensional steady flow) is included in Appendix B to provide the reader with an illustrative example.

MOC3D output is routed to a main file, separate from the *MODFLOW* main output file, and optionally to additional output files. Appendix C contains output from the example input data set contained in Appendix B.

Program Segments

MOC3D input and output utilizes the standard *MODFLOW* array reading and writing utilities as much as possible. Konikow and others (1996) describe briefly each of the subroutines in *MOC3D* that are used for ten different categories of functions. Discussion related to table 4 in that report (p. 37) is to some extent irrelevant, inasmuch as it pertains to particle manipulations. Several existing *MOC3D* subroutines were modified. *ELLAM* routines have been added to the main *MOC3D* transport time loop. Table 1 provides a list of the primary subroutines and their descriptions. Table 2 contains secondary subroutines and the calling tree. The *ELMOVE* routine in *ELLAM* uses a linear velocity interpolation. Output routines from earlier versions of the code are retained. A flow chart of the program segments controlling the transport calculations for *ELLAM* is shown in figure 8 (also see figures 12 and 13 of Konikow and others, 1996).

Dispersion coefficients are calculated at cell faces. To improve efficiency, the dispersion coefficients are lumped with the porosity, thickness, and an appropriate grid dimension factor of the cell into combined parameters called "dispersion equation coefficients." Here, the *ELLAM* version of subroutine *DSP6FM* has been modified to calculate the distance between cell nodes in the column or row direction using the newly implemented variable grid dimensions, and to use an appropriate grid dimension factor for the *ELLAM* integral formulation of the problem. For example, the dispersion equation coefficient for the $j+1/2, i, k$ face in the column direction is

$$\frac{2(\epsilon b D_{xx})_{j+1/2, i, k}}{\Delta x_j + \Delta x_{j+1}} \quad (32)$$

These combined coefficients are the ones that are written to the output files.

Table 1. *ELLAM* transport loop

Subroutine	Description
ELLBDY	Create boundary arrays; Track inflow mass, and accumulate to RHS storage or outflow integral
BDYINT	Track boundary layer mass, and accumulate to RHS storage or outflow integral; Calculate LHS storage coefficient, and save value if inflow or outflow face (1 st transport time step)
ELLLHI	Track interior mass, and accumulate to RHS storage or outflow integral; Calculate LHS storage coefficient (1 st transport time step)
ELLDIS	Build LHS dispersion matrix (1 st transport time step)
ELLSRC	Sink: accumulate to dispersion matrix (1 st transport time step) and RHS; Source: track source mass; accumulate to RHS storage or outflow integral
ELIUPD	Move inflow boundary concentrations to solution vector
ELLOUT	Outflow and inflow processing: solve outflow boundary equations; accumulate inflow and outflow boundary mass to RHS; accumulate boundary mass for mass balance
CONVERT	Convert storage plus dispersion matrix into SLAP column format (1 st transport time step)
SMOC5I	Convert storage matrix into SLAP column format (1 st transport time step); Calculate initial mass in system
ELLSLV	Solve interior equations
ELNUPD	Move no-flow boundary concentrations to solution vector
SMOC6BE	Mass-balance calculations
SMOC6C and SOBS5O	Print results

Table 2. Calling tree for *ELLAM* code, showing hierarchy of secondary subroutines for transport calculations. Subroutine to the left calls subroutine(s) to the right. Subroutines in column 1 are called from the main transport loop (subroutine MOC6MVOT).

ELLBDY	BDYENT	BDYRHS	ELMOVE EVALTF BNDYTF	EMOVTIM EVALTF
BDYINT	BDYCON	BUILDNEI BDYCR BDYCL	BINTERP ELMOVE EVALTF BNDYTF DISTRIB EDGWGHT	EDGWGHT EMOVTIM EVALTF
ELLHI	CLAYER BLAYER ELLRHI	CROW HFACED VFACED EDGCOR ELLCR	CINTERP ELMOVE EVALTF BNDYTF	EMOVTIM EVALTF
ELLDIS				
ELLSRC	BDYRHS	ELMOVE EVALTF BNDYTF	EMOVTIM EVALTF	
ELIUPD				
ELLOUT				
CONVERT	GET_IDISP			
SMOC5I	GET_IDISP SMOC5A			
ELLSLV	SSLUGM	solver routines		
ELNUPD				
SMOC6BE	SMOC5A			
SMOC6C SOBS5O	print routines			

MODFLOW source and sink packages contain an option called CBCALLOCATE. When used, the package will save the cell-by-cell flow terms across all faces of every source or sink cell. *MOC3D* uses these fluid fluxes to calculate solute flux to or from the source/sink nodes. Because these individual

solute fluxes are required to compute the solute mass balance, the CBCALLOCATE option must always be selected when using *MOC3D*. Implicit calculations of concentration changes at nodes caused by mixing with fluid sources are controlled by the ELLSRC subroutine.

The *ELLAM* code includes a preconditioned generalized minimum residual method (GMRES) sparse iterative solver with an incomplete lower-upper (ILU) decomposition preconditioner to solve the non-symmetric system of equations. The solver routines are code from the SLATEC Common Mathematical Library, available through the NETLIB public domain repository; see <http://www.netlib.org/slatec/index.html>. Users interested in seeing more details about the numerical solvers should also examine the FORTRAN source code listing for *MOC3D* and its *ELLAM* subroutines, as they are very clearly documented with explanatory “comment” lines throughout the code.

Some mass-balance calculations have been reordered compared to the original *MOC3D* sequence. The initial mass calculation is called during the first transport time step from within the transport loop, after the matrix of storage coefficients has been created.

Mass associated with inflow and outflow boundary faces is accumulated for mass-balance reporting each time step. This mass is calculated using integration weights associated with boundary nodes and boundary cell porosities, along with current boundary concentration values. Total amounts are added to interior and no-flow boundary mass calculated from the storage coefficient matrix and current concentrations.

Similarly, decayed mass from inflow faces and sources is accumulated for mass-balance reporting. This total, and decay of mass stored at the beginning of the time step, are both reported as contributions to mass flux out of the transport domain.

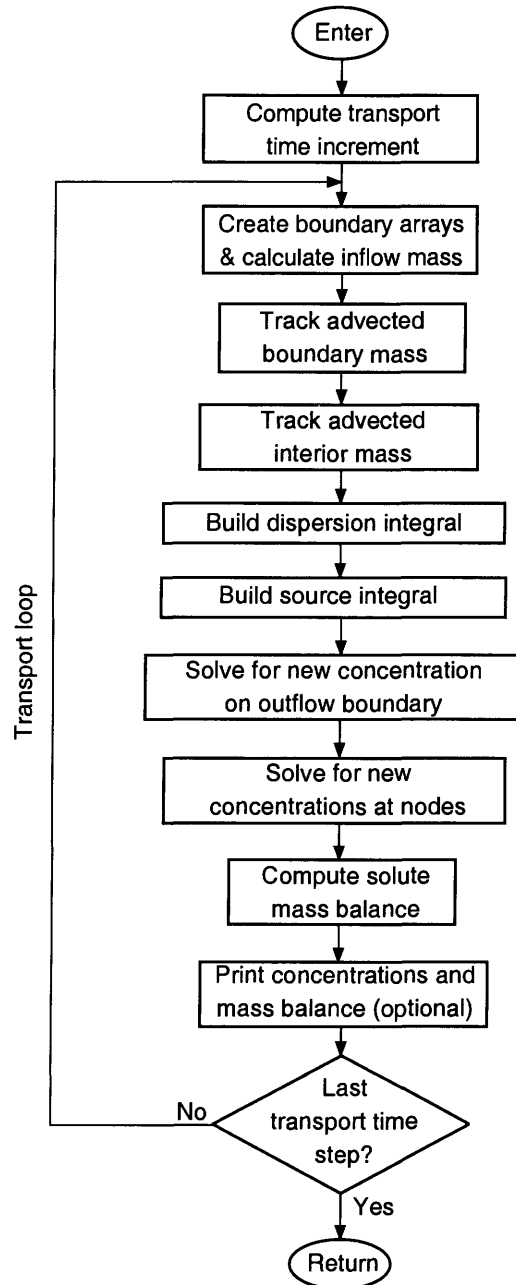


Figure 8. Simplified flow chart for the transport loop of the *ELLAM* calculation process.

Guidance on Input Parameter Values

Discretization parameters NSC, NSR, NSL, and NT must all be powers of two. In each case, the input parameter specified by the user is the exponent: NSCEXP, NSREXP,

NSLEXP, and NTEXP, respectively.

In general, use NSC = NSR = NSL = 4 (and therefore NSCEXP = NSREXP = NSLEXP = 2) except if modeling a one- or

two-dimensional problem. Here, a value of 4 is only needed in the dimension(s) of the problem, with NS values of 2 adequate in the missing direction(s). NS values greater than four may be useful when modeling with a complicated velocity field, or with numerous transport time steps, or to improve accuracy near a boundary. Computational efficiency is strongly related to the values of the NS parameters, but the impact of changing NS values is highly problem dependent.

The number of discrete mass-bearing volumes entering the transport subdomain during a time step is $NT+1$. This number must be large enough so that each grid cell, from the one at the first boundary to the one where mass entering at the beginning of that time step is advected, can receive a portion of the discretized inflow mass. If the solution shows mass becoming distributed down the flow path in clumps (this is illustrated schematically in figure 9 for a case in which two sub-time steps are used), increase NTEXP (thus increasing NT). Increasing NT will smooth the front to the point of ensuring an even distribution of mass among destination cells. NT only comes into effect

for problems with concentrations entering via fluid sources or inflow across boundaries of the transport subgrid.

For example, consider the case of one-dimensional flow discussed below as the first test problem. Here, $V = 0.1$ cm/s and each cell has a width of 0.1 cm. $NTEXP = 1$ (thus $NT = 2$) will result in a discretization of inflow mass such that one fourth enters the system at the beginning of the time step, one half in the middle, and one fourth at the end. Thus, the initial quarter of the mass will be advected the farthest and will be distributed among neighboring cells as mandated by the test functions determined by NSC. The mass entering at the end of the time step remains in the the first grid cell. The point representing half of the mass will be advected a number of cells equal to the magnitude of $(TIMV/2)$. For a time step (TIMV) of 5 s or longer, $TIMV/2 > 2.5$, so the point will move past the middle of the third cell. No advected mass will end up in the second grid cell, for any value of NSC. Increasing NTEXP will eliminate this problem.

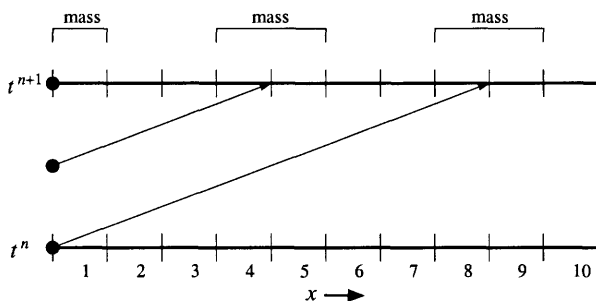


Figure 9. Schematic representation of tracking mass from an inflow boundary. For a Courant number of 8 and $NT = 2$, not every cell along the inflow path receives mass.

MODEL TESTING AND EVALUATION

The *ELLAM* simulator was tested and evaluated by running the same suite of test cases as was applied to *MOC3D* Version 1 by Konikow and others (1996) and *MOC3D* Version 2 by Kipp and others (1998). This suite includes results generated by analytical

solutions and by other numerical models. It spans a range of conditions and problem types so that the user will gain an appreciation for both the strengths and weaknesses of this particular code. It should be noted that all test cases involve steady flow conditions.

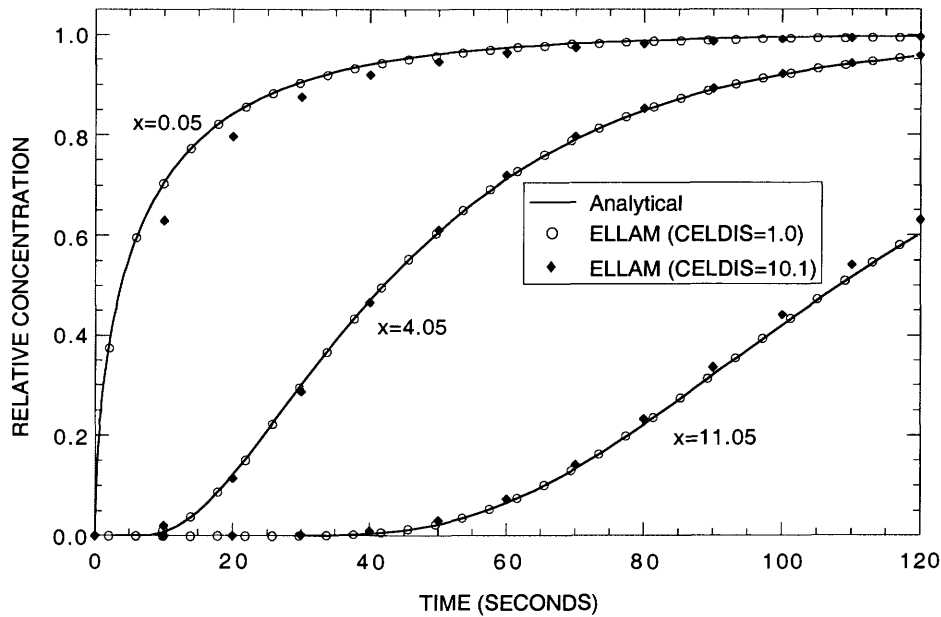


Figure 11. Numerical and analytical solutions for the case of increased dispersivity ($\alpha_L = 1.0$ cm, $D_{XX} = 0.1$ cm²/s, and other parameters as defined in table 3).

Konikow and others (1996) also present the results of these tests in the form of concentration profiles in space at various times and for various retardation factors (see their figures 22 and 23). Replication of these tests using the *ELLAM* formulation yields results comparable to those just described, as seen in figure 12 for CELDIS = 10.1 and CELDIS = 1 in the nonreactive case. To test the limits of the *ELLAM* method, we also solved this problem using CELDIS = 61 (2 time increments), NSC = 4, NSR = NSL = 2, and NT = 128. Although these numerical parameters yield too few time increments to even expect an accurate or precise match to the breakthrough curves, figure 12 indicates that even in this extreme case, a qualitatively good match for most of the breakthrough was calculated, except notably near the outflow face. Although such large values of CELDIS are not recommended, the results for CELDIS = 61 demonstrate the apparent robustness of the method.

The accuracy of the numerical method for problems in which decay is occurring was

evaluated by specifying the decay rate as $\lambda = 0.01$ s⁻¹ for the same low-dispersion, no sorption, problem as defined for figure 10. The results for CELDIS = 1, NSC = 32, NSR = NSL = 2, and NT = 128 are presented in figure 13, which shows excellent agreement between the analytical and numerical solutions. For clarity, only every fourth data point of the numerical solution is shown. As in the case of no decay, NSC = 4 (not plotted) produces a slightly low concentration at short distance.

In all cases described above, the mass-balance error was less than 0.001 percent. In contrast, the mass-balance errors for these problems using the explicit and implicit versions of the method-of-characteristics code yielded mass-balance errors of up to a few percent in some cases. *ELLAM* is mass conservative whereas *MOC* and *MOCIMP* are not. Also, as illustrated by the results for CELDIS = 61 in figure 12, an accurate mass balance does not prove that you have an accurate solution.

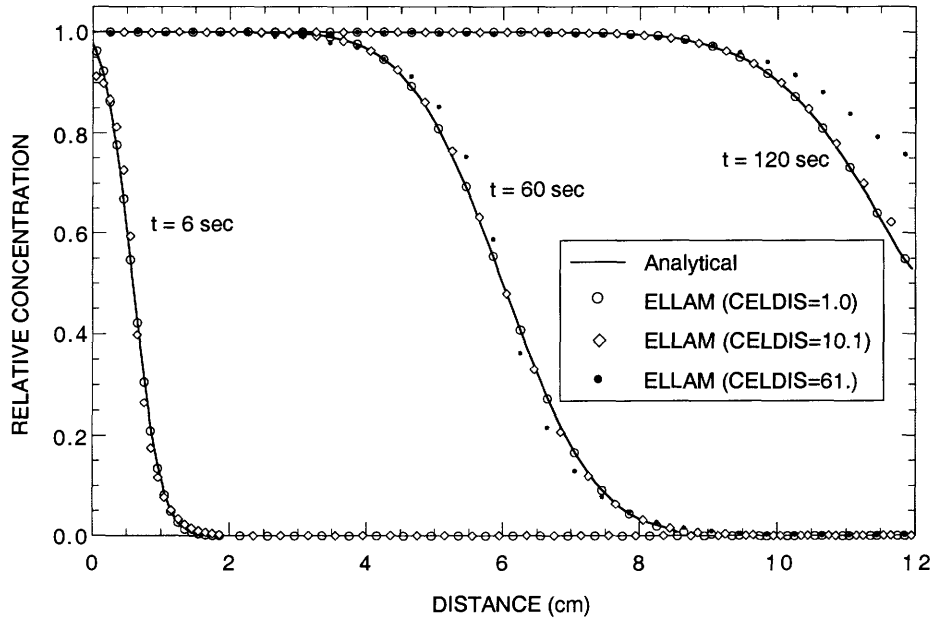


Figure 12. Numerical and analytical solutions for three different times for same one-dimensional, steady flow, solute-transport problem shown in figure 10.

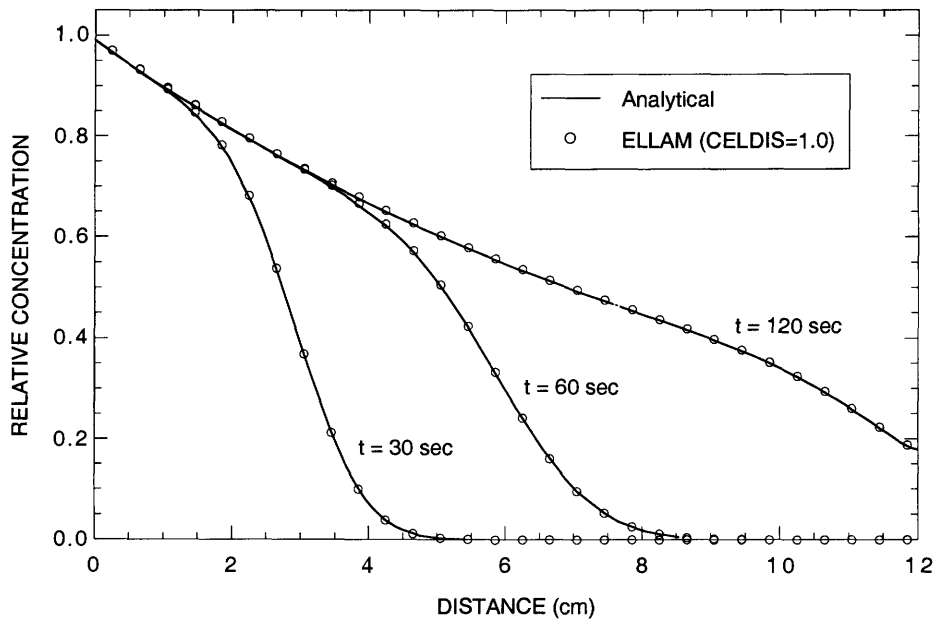


Figure 13. Numerical and analytical solutions for three different times for case in which solute is subject to decay at rate of $\lambda = 0.01 \text{ s}^{-1}$.

The sensitivity of the results to variations in the value of NT was evaluated for the case of CELDIS = 1.0, shown in figure 10. Values of NTEXP were varied from 2 to

9 (figure 10 represents NTEXP = 7). This parameter only had a small effect on the accuracy of results at the node closest to the source; there were no discernible differences

elsewhere. It also had only a minor impact on the efficiency of the solution. At the first node, the breakthrough curves were essentially identical for values of NTEXP ranging from 5 to 9, and the running time increased by less than 13 percent (from 203 s to 229 s). However, decreasing the value of NTEXP from 5 to 2 caused increasingly larger (but still minor) deviations from the

analytical solution, while decreasing the running time only by 1 s (from 203 s to 202 s). For NTEXP = 2, the solution was stable but too low by about 2 percent at most times. For this case, it appears that a value of NTEXP = 5 or 6 would be optimal, but the results were relatively insensitive to variations in NTEXP.

Uniform Flow, Three-Dimensional Transport

To evaluate and test *ELLAM* for three-dimensional cases, we compared numerical results with those of the analytical solution developed by Wexler (1992) for the case of three-dimensional solute transport from a continuous point source in a steady, uniform flow field in a homogeneous aquifer of infinite extent. Konikow and others (1996) note that this evaluation primarily is a test of the accuracy of the calculated dispersive flux in three directions because the flow field is aligned with the grid. The problem and analytical solution are described in detail by Konikow and others (1996, p. 45-48); the parameters and boundary conditions for this test case are summarized in table 4. This case also represents a test of the ability of the algorithm to represent the effects of a solute source at a specified flux boundary condition.

The results of *ELLAM* are compared graphically in figure 14 with those of the analytical solution for the x - y plane passing through the point source. Figure 14a shows the concentrations in this plane at $t = 400$ days as calculated using the analytical solution. Also shown, in figures 14b-d, are the *ELLAM* solutions using CELDIS = 7 (two transport time increments), NSC = NSR = NSL = 4, and NT = 16 (figure 14b); CELDIS = 1 (14 time increments), NSC = NSR = NSL = 4, and NT = 4 (figure 14c); and CELDIS = 0.1 (134 time increments), NSC = NSL = 4, NSR = 8, and NT = 16 (figure 14d).

Table 4. Base-case parameters used in *ELLAM* simulation of transport from a continuous point source in a three-dimensional, uniform, steady-state flow system

Parameter	Value
$T_{xx} = T_{yy}$	0.0125 m ² /day
ε	0.25
α_L	0.6 m
α_{TH}	0.03 m
α_{TV}	0.006 m
PERLEN (length of stress period)	400 days
V_y	0.1 m/day
$V_x = V_z$	0.0 m/day
Initial concentration (C_0)	0.0
Source concentration (C')	2.5×10^6 g/m ³
Q (at well)	1.0×10^{-6} m ³ /d
Source location	Column = 1, Row = 8, Layer = 1
Number of rows	30
Number of columns	12
Number of layers	40
DELR (Δx)	0.5 m
DELC (Δy)	3.0 m
Layer thickness (Δz)	0.05 m
CELDIS	1.0
NSCEXP	2
NSREXP	2
NSLEXP	2
NTEXP	2

As noted in previous *MOC3D* reports, a slightly greater spreading is evident in the numerical model results than in the analytical solution, both upstream as well as downstream of the source. Part of this difference, however, is explained by the fact that the numerical source is applied over a finite area in the horizontal plane of the model, in which the length of the source cell is 3 m in the direction parallel to flow, whereas the source is represented as a true point in the analytical solution.

The *ELLAM* results using two transport time increments (figure 14b) indicate that more time steps are needed in order to accurately simulate dispersion. The

ELLAM results for 14 time steps (figure 14c) accurately characterize the dispersive flux without the spreading upstream from the source that is produced by *MOC3D*. The *ELLAM* results for 134 time steps (figure 14d) yield even less spreading upstream of the source, but do exhibit numerical oscillations produced because the concentration gradient is too steep relative to the grid spacing.

Konikow and others (1996) also present comparisons for this case for vertical planes parallel and perpendicular to the flow direction. These same comparisons between the analytical and *ELLAM* results are as close as between figures 14a and 14c, and are not reproduced here.

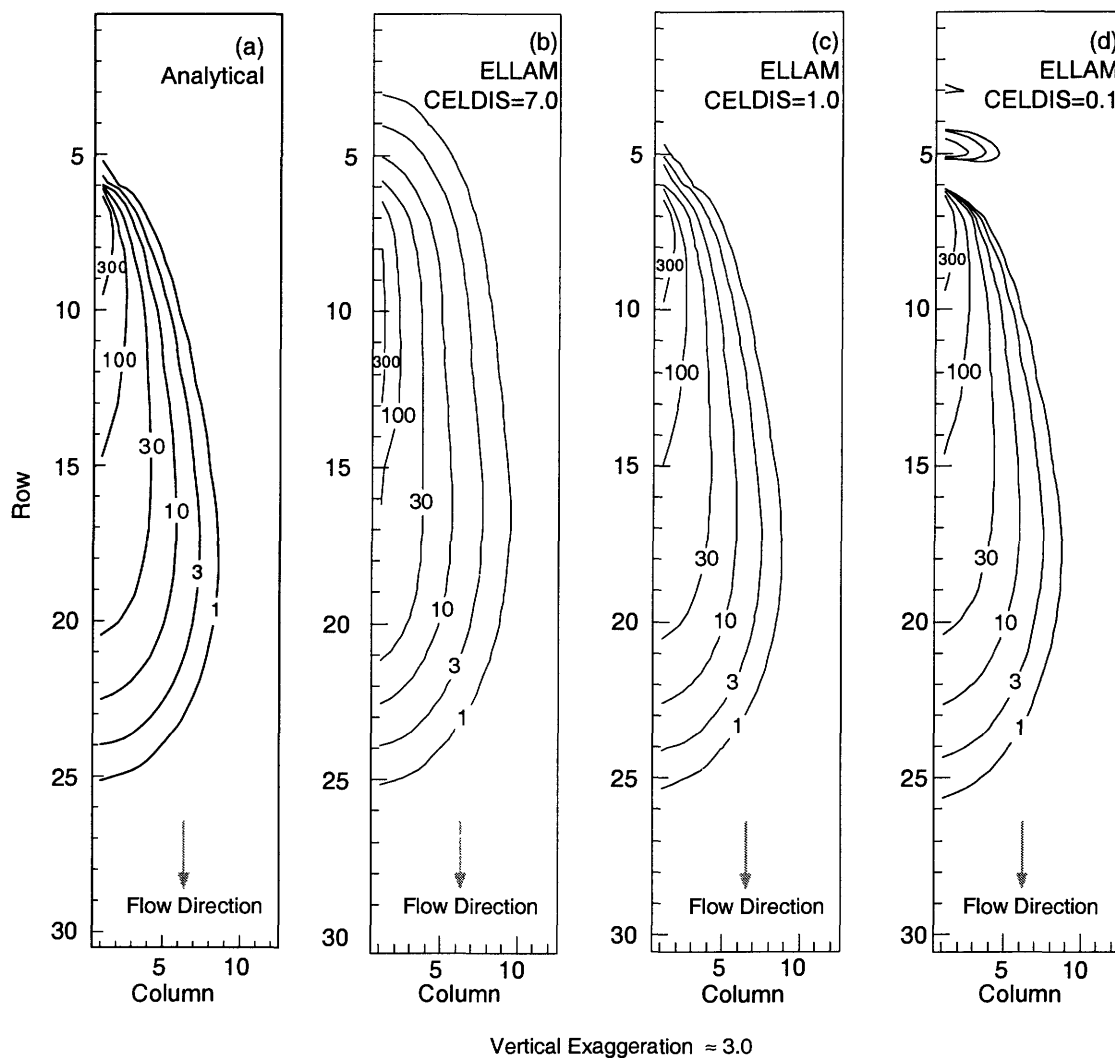


Figure 14. Concentration contours for (a) analytical and (b-d) *ELLAM* numerical solutions in the horizontal plane containing the solute source (layer 1) for three-dimensional solute transport in a uniform steady flow field at $t = 400$ days. Parameters are defined in table 4.

Table 6. Parameters used in *ELLAM* simulation of three-dimensional transport from a point source with flow in the x -direction and flow at 45 degrees to x - and y -axes

Parameter	Value
$T_{xx} = T_{yy}$	10.0 m ² /day
ε	0.1
α_L	1.0 m
$\alpha_{TH} = \alpha_{TV}$	0.1 m
PERLEN (length of stress period)	90 days
V_x	1.0275 m/day
$V_y = V_z$	0.0 m/day*
Initial concentration at source	1×10^6
Source location in transport grid	Column = 11, Row = 36, Layer = 4
Number of rows	72
Number of columns	72
Number of layers	24
DEL _R (Δx)	3.33 m
DEL _C (Δy)	3.33 m
Layer thickness ($b = \Delta z$)	10.0 m
CELDIS	5.0
NSCEXP	2
NSREXP	2
NSLEXP	2
NTEXP	1

* For flow at 45 degrees to x - and y -axes, $V_y = 1.0275$ m/day

We specified boundary conditions for the test case of flow in the x -direction such that $V_x = 1.0275$ m/d, and $V_y = V_z = 0.0$ m/d. For flow at 45 degrees to x and y , we specified boundary conditions such that $V_x = V_y = 1.0275$ m/d, and $V_z = 0.0$ m/d. For both cases, the distance the center of mass of the plume travels in the x -direction is the same for equal simulation times. Note, however, that the magnitude of velocity is higher in the latter case; therefore, there will be more

dispersion in that problem during an equal time interval.

The results for both the analytical and numerical solutions for the case in which flow occurs in the x -direction are shown in figure 16, where values of CELDIS = 5 (yielding six transport time increments), NSC = NSR = NSL = 4, and NT = 2 were used. These results represent the concentrations in the plane of the initial source of solute. The *ELLAM* transport algorithm gives results (figure 16b) for a 72 by 72 grid that are close to those of the analytical solution (figure 16a). The numerical results, however, do show some slight spreading (or numerical dispersion) relative to the analytical solution in both the transverse and longitudinal directions. Increasing the number of time increments does not completely eliminate the spreading and causes some loss of peak concentrations, even with increased NS values. In contrast to the previous *MOC3D* solutions, *ELLAM* results retain the symmetry of the analytical solution. Part of the discrepancy is attributable to the need in *ELLAM* to use four grid points to discretize a front. This precludes the possibility of modeling with high accuracy the migration of an instantaneous point source placed in a single grid cell. Therefore, we modified this test problem for *ELLAM* by using a dispersed solute mass as an initial condition. The initial condition for this *ELLAM* test is the analytical solution to the original point source problem at $t = 90$ days, and the *ELLAM* solution is evaluated against the Wexler analytical solution later in time at $t = 130$ days. These results are presented in figure 17, where it can be seen that the analytical solution (figure 17a) and the numerical solution (figure 17b) are very similar. The *ELLAM* solution, however, still clearly exhibits some numerical dispersion, which is most evident at the lower concentrations.

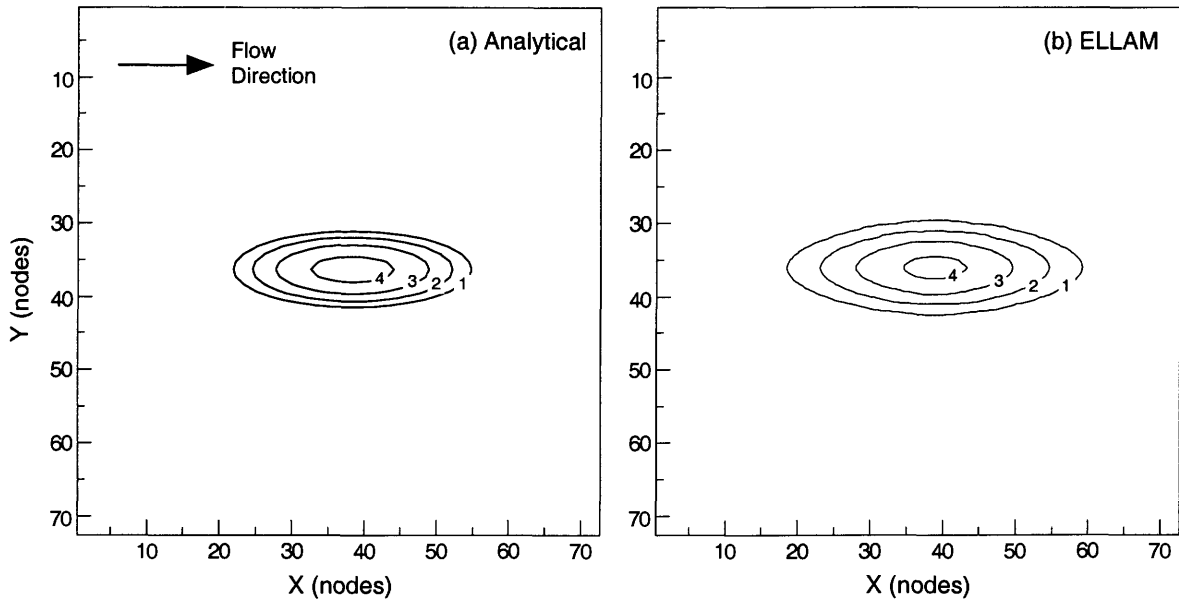


Figure 16. Concentration contours for (a) analytical and (b) numerical solutions for transport of a point initial condition in uniform flow in the x-direction at $t = 90$ days. The z-component of flow is zero, but there is dispersion in all three directions. Contour values are the log of the concentrations.

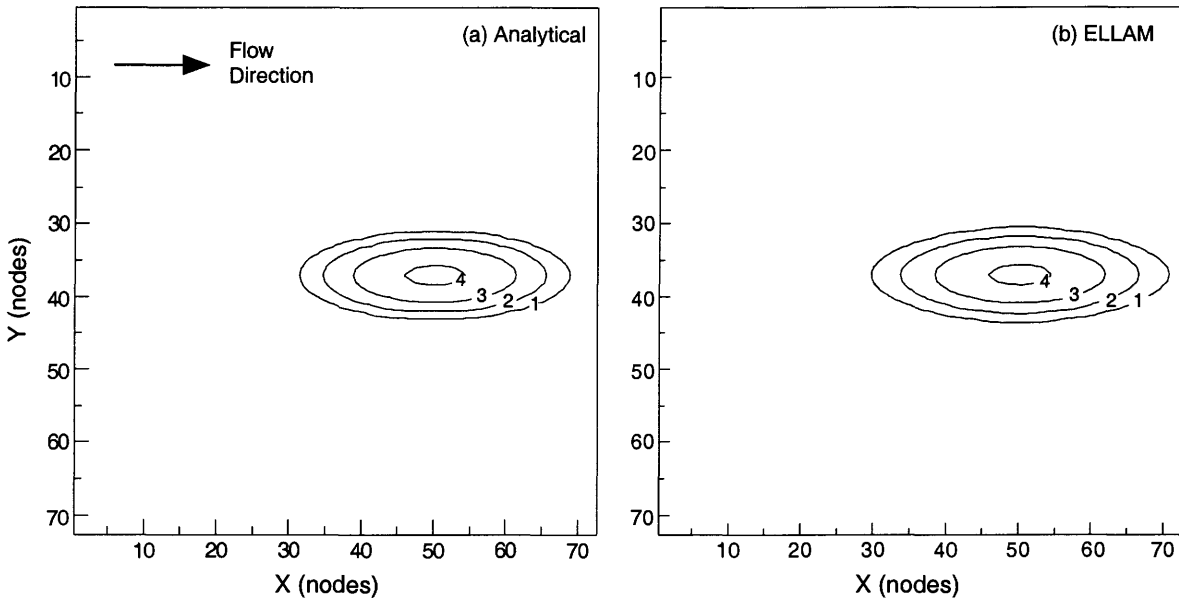


Figure 17. Concentration contours for (a) analytical and (b) *ELLAM* numerical solutions for transport of a dispersed-point initial condition in uniform flow in the x-direction at $t = 130$ days. The y- and z-components of flow are zero, but dispersion occurs in all three directions. Contour values are the log of the concentrations.

The results of the test problem for flow at 45 degrees to the grid are shown in figure 18, again using a $72 \times 72 \times 24$ grid. The analytical solution for $t = 130$ days, which

provides the basis for the evaluation, is shown in figure 18a. As was done for the previous analysis shown in figure 17, the *ELLAM* solution in this case also used the analytical

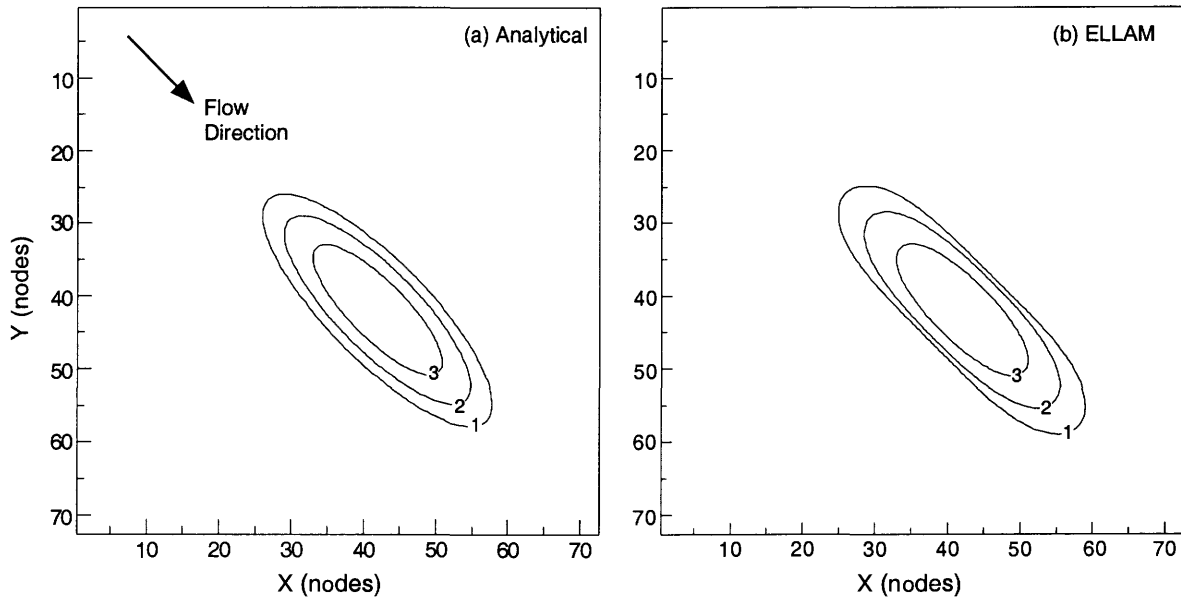


Figure 18. Concentration contours for (a) analytical and (b) *ELLAM* numerical solutions for transport of a point initial condition in uniform flow at 45 degrees to the x-direction at $t = 130$ days. Contour values are the log of the concentrations.

solution at $t = 90$ days as the initial conditions. The results using $CELDIS = 5$ (three time increments), $NSC = NSR = NSL = 4$, and $NT = 2$ are shown in figure 18b for the plane of the initial source. As in the previous case (where flow is aligned with the grid), *ELLAM* produces the symmetry characteristic of the analytical solution. There is also slight longitudinal spreading (numerical dispersion) that is not alleviated by increasing the number of time steps.

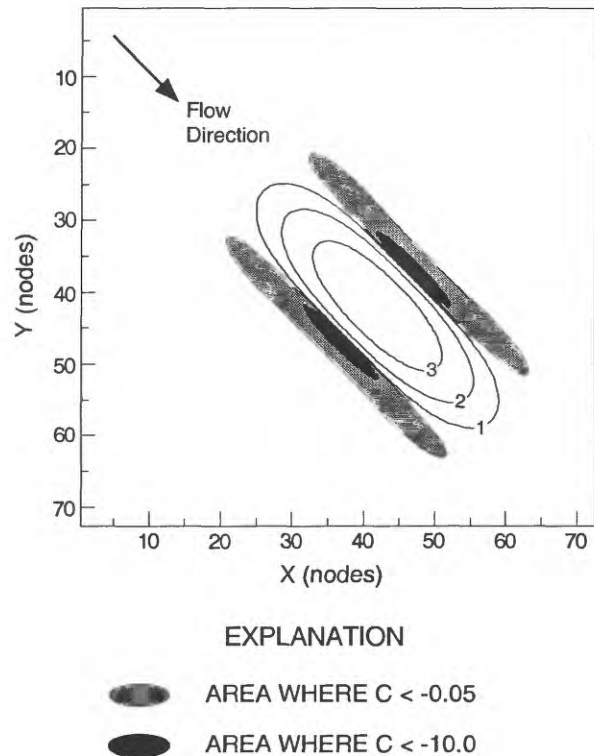
Unlike the previous case, the numerical results in figure 18b do show some distortion of the shape of the plume relative to the analytical solution. It is not as pronounced, however, as the “hourglass” shape yielded by *MOC3D* for the Dirac problem (see Kipp and others, 1998, figure 14). There is a narrowing of the plume calculated with the numerical model, which is characteristic of a grid-orientation effect and is caused primarily by the off-diagonal (cross-derivative) terms of the dispersion tensor. When flow is oriented parallel to the grid, or when longitudinal and transverse dispersivities are equal, the cross-derivative

terms of the dispersion tensor are zero. Because flow is at 45 degrees to the grid in this test problem, the cross-derivative dispersive flux terms are of maximum size and negative concentrations are most likely to occur. The calculated concentration field is less accurate in this case largely because the standard differencing scheme for the cross-derivative dispersive flux terms can cause overshoot and undershoot of concentrations. If the base (or background) is zero concentration, then undershoot will cause negative concentrations. The magnitude of this overshoot and undershoot effect can be reduced by using a finer grid.

Some small areas of negative concentrations were calculated, but they do not appear in figure 18b using logarithmic-scale contouring. To show the extent of the areas of negative concentration, we have replotted the results illustrated in figure 18b in figure 19, using two types of shading for areas where the relative concentration is less than -0.05 and less than -10.0 . We tested the sensitivity of the extent of negative concentrations to the size of the transport time

increment by reducing the value of CELDIS to 0.25. The area over which negative concentrations occurred was only slightly smaller. The increase in execution time, however, was significant, so the very small improvement does not appear to justify the extra computational costs.

Figure 19. Concentration contours for *ELLAM* numerical solution showing areas of calculated negative concentrations for problem represented in figure 18b.



Constant Source in Nonuniform Flow

Burnett and Frind (1987) used a numerical model to simulate a hypothetical problem having a constant source of solute over a finite area at the surface of an aquifer having homogeneous properties, but nonuniform boundary conditions, which result in nonuniform flow. Because an analytical solution is not available for such a complex system, we use their results for this test case as a benchmark for comparison with the results of applying the *ELLAM* algorithm in *MOC3D*, as was also done by Konikow and others (1996) and Kipp and others (1998). Burnett and Frind (1987) used an alternating-direction Galerkin finite-element technique to solve the flow and solute-transport equations in both two and three dimensions. Their model also includes the capability to vary α_T as a function of coordinate direction, thereby allowing this feature of *MOC3D* to be evaluated. A detailed description of the problem geometry and of the parameters for the numerical simulation are presented by Konikow and others (1996, p. 55-60).

Cases of both two- and three-dimensional transport were examined for this problem. The grids used in the *ELLAM* simulations were designed to match as closely as possible the finite-element mesh used by Burnett and Frind (1987). Some differences in discretization, however, could not be avoided because the finite-element method uses a point-centered grid whereas *ELLAM* uses a block-centered (or cell-centered) grid. The former allows specifications of values at nodes, which can be placed directly on boundaries of the model domain. Nodes in *ELLAM* are located at the centers of cells, and block-centered nodes are always one-half of the grid spacing away from the edge of the model domain. Among the small differences arising from the alternative discretization schemes are that, in the *ELLAM* grid, (1) the modeled location of the 14.25 m long source area is offset by 0.225 m towards the right, and (2) the total length of the domain is 199.5 m.

The first simulation of this test problem was for the case of a two-dimensional model. The input data values for this analysis are listed in table 7. The top discretization layer consisted of constant-head nodes and the solute source.

Results for the two-dimensional case from the *ELLAM* simulation closely match those of Burnett and Frind (1987) (see figure 20). The results using $CELDIS = 30$ (seven time increments), $NSC = NSR = NSL = 4$, and $NT = 32$ are shown. The shape of the plume is almost exactly the same for both models. In the *ELLAM* results, however, the highest concentration contour (0.9) does not extend as far downgradient as that of Burnett and Frind (1987), while the low concentration contour (0.3) from *ELLAM* extends slightly farther downgradient. Overall, the *ELLAM* results provide a closer match to the contours of Burnett and Frind than do the *MOC3D* contours using 381, 1901, or 4218 time increments (see Kipp and others, 1998). The *ELLAM* contours (for all NS values tested) are free of “wiggles” in the *MOC3D* solution discussed by Kipp and others (1998). Increasing the number of transport time increments produced a solution having a slightly greater downgradient extent, but still short of *MOC3D* results.

Table 7. Parameters used for *ELLAM* simulation of transport in a vertical plane from a continuous point source in a nonuniform, steady-state, two-dimensional flow system (described by Burnett and Frind, 1987)

Parameter	Value
K	1.0 m/day
ε	0.35
α_L	3.0 m
α_{TH}	0.10 m
α_{TV}	0.01 m
<i>PERLEN</i> (length of stress period)	12,000 days
Source concentration (C')	1.0
Number of rows	1
Number of columns ¹	141
Number of layers ¹	91
<i>DELR</i> (Δx)	1.425 m
<i>DELC</i> (Δy)	1.0 m
Layer thickness ($b = \Delta z$)	0.2222-0.2333 m
<i>CELDIS</i>	30.0
<i>NSCEXP</i>	2
<i>NSREXP</i>	2
<i>NSLEXP</i>	2
<i>NTEXP</i>	5

¹ One row and layer were allocated to defining boundary conditions, so concentrations calculated in only 140 columns and 90 layers were used for comparison.

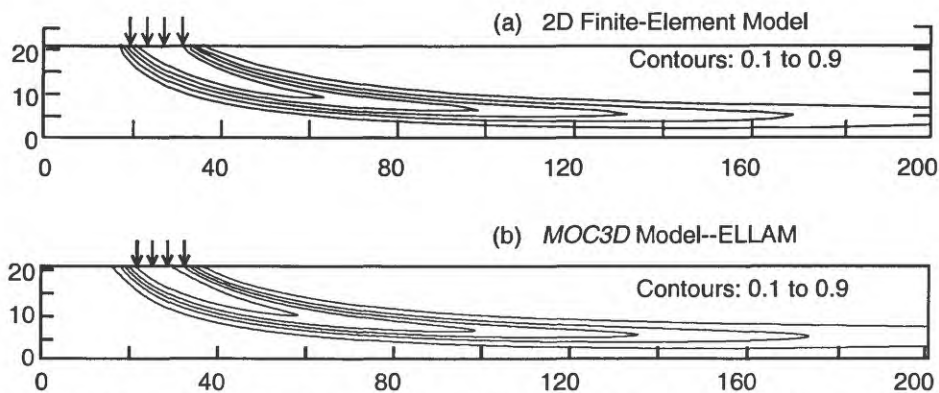


Figure 20. Two-dimensional simulation results for nonuniform-flow test case showing plume positions as contours of relative concentration; (a) finite-element model (modified from Burnett and Frind, 1987, figure 8a), and (b) *ELLAM* solution using $CELDIS = 30$. Contour interval is 0.2 relative concentration.

As was done for the *MOC3D* tests (Konikow and others, 1996; Kipp and others, 1998), the *ELLAM* grid was expanded laterally to 15 rows having Δy of 1.0 m for the three-dimensional version of this case. Figure 21 shows the transport results in a vertical plane at the middle of the plume for both models for the case in which $\alpha_{TV} = 0.01$ m and $\alpha_{TH} = 0.1$ m. The *ELLAM* results for the vertical plane in the first row are contoured in figure 21b (because of symmetry, we only simulate half of the plume, as explained by Konikow and others, 1996). The *ELLAM* plume closely matches that calculated by the finite-element model (figure 21a), although the former shows slightly farther downstream migration of low concentrations of solute. As

in the two-dimensional case, the *ELLAM* solution provides a closer match to the Burnett and Frind (1987) solution than do the previous *MOC3D* results.

Figure 22 shows the results for the case in which the vertical transverse dispersivity is increased by a factor of ten, so that $\alpha_{TH} = \alpha_{TV} = 0.1$ m. The *ELLAM* results for CELDIS = 30 yielded concentrations that were noticeably low near the source (near the upgradient end of the plume), so the simulation was repeated using CELDIS = 21 (10 time increments). These *ELLAM* results are illustrated in figure 22b and appear to agree very closely with the results of Burnett and Frind (1987) (figure 22a).

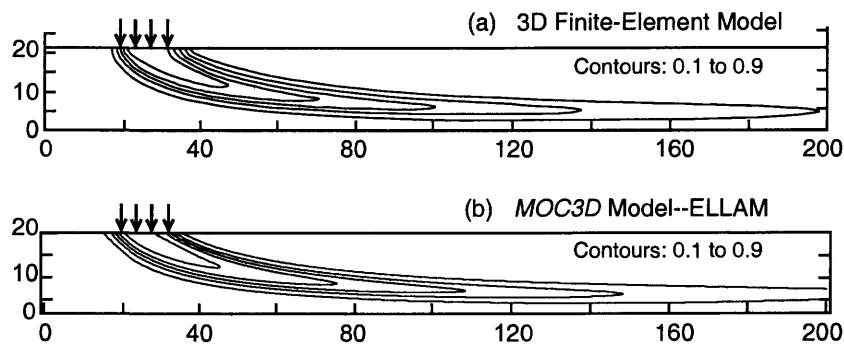


Figure 21. Three-dimensional simulation results for nonuniform-flow test case in which $\alpha_{TH} = 0.1$ m and $\alpha_{TV} = 0.01$ m: (a) finite-element model (modified from Burnett and Frind, 1987, figure 8c), and (b) numerical *ELLAM* solution using CELDIS = 30. Plume positions are represented by contours of relative concentration; contour interval is 0.2 relative concentration.

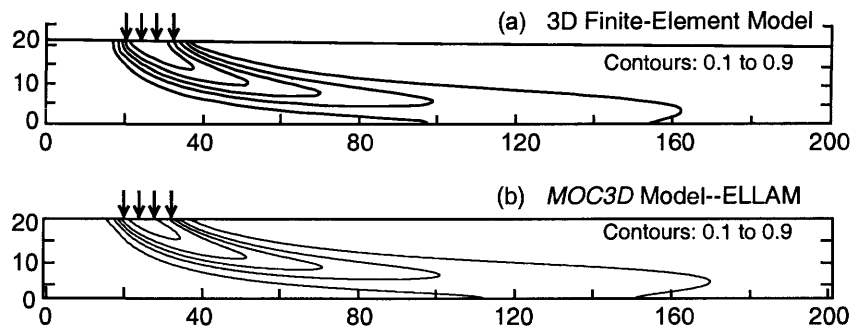


Figure 22. Three-dimensional simulation results for nonuniform-flow test case in which $\alpha_{TH} = \alpha_{TV} = 0.1$ m: (a) finite-element model (modified from Burnett and Frind, 1987, figure 9b), and (b) numerical *ELLAM* solution using CELDIS = 21. Plume positions are represented by contours of relative concentration; contour interval is 0.2 relative concentration.

Relative Computational and Storage Efficiency

Computer-memory requirements for *ELLAM* are greater than those for the explicit or implicit *MOC3D* dispersive transport algorithm. The additional arrays required can increase the memory size requirement by as much as a factor of three (see table 8).

The computational effort required by the *ELLAM* simulator is strongly dependent on the size of the problem being solved, as determined by the total number of nodes, the NS and NT values, and the total number of time increments (controlled by CELDIS). The user is cautioned that using values for NS and NT parameters that are too small for a given problem may lead to inaccurate solutions. Sensitivity testing will help the user determine appropriate values to specify. Analyses indicate that the greatest computational effort, as measured by CPU

time, is typically expended in the mass tracking routines. For a given problem, computational time may vary significantly as a function of the characteristics of the particular computer on which the simulation is performed, and on which FORTRAN compiler and options were used to generate the executable code.

For a given problem, the *ELLAM* algorithm can often yield an accurate solution more efficiently than the previously documented explicit or implicit MOC options. However, this will typically require the use of a CELDIS value of 5 or more; the explicit and implicit versions of MOC require that CELDIS be less than or equal to 1.0. Table 8 shows that *ELLAM* was more efficient for three of the six test problems evaluated.

Table 8. Execution times and storage requirements for *MOC3D* and *ELLAM* for selected test cases

Problem Description	Run Time in CPU-seconds			Array Elements Used ¹		
	Explicit	Implicit	ELLAM	Explicit	Implicit	ELLAM
One-Dimensional Steady Flow ²	7	10	9 CELDIS=10.1	11,457	17,400	33,489
Three-Dimensional Steady Flow ²	404	175	1,366 CELDIS=1	897,331	1,602,994	3,344,624
Two-Dimensional Radial Flow and Dispersion ²	930	445	138 CELDIS=5	455,737	499,900	233,564
Point Initial Condition in Uniform Flow ²	210	310	2,721 CELDIS=5	1,728,673	2,406,112	3,384,524
Constant Source in Nonuniform Flow (Two-Dimensional) ³	13,360	2,450	2,245 CELDIS=30	868,951	1,457,602	2,850,056
Constant Source in Nonuniform Flow (Three-Dimensional) ³	38,117	12,026	4,400 CELDIS=30	12,823,151	21,652,034	41,206,836

¹ Data arrays and lists for *MODFLOW* and explicit *MOC3D* are allocated space in one array, the *MODFLOW* "X" array. *ELLAM* also uses an "MX" array for integer arrays.

² Data General server with a Motorola 88110 chip running DG Unix 5.4R3.10 with 256MB RAM and a 45 MHz processor was used for this problem. Green Hills Software FORTRAN-88000 was used to compile *MOC3D*.

³ Silicon Graphics server with an R8000 chip running Irix 6.0.1 with 576MB RAM and a 90 MHz processor was used for this problem. MIPSpro F77 was used to compile *MOC3D*.

CONCLUSIONS

The *ELLAM* advective-dispersive transport algorithm presented as an alternative solution method within the *MOC3D* simulator can model the transient, three-dimensional, transport of a solute subject to decay and retardation. The numerical methods used to solve the governing equations have broad general capability and flexibility for application to a wide range of hydrogeological problems.

The accuracy and precision of the numerical results of the implicit *ELLAM* simulator were evaluated by comparison to analytical and numerical solutions for the same set of test problems as reported for *MOC3D* (Versions 1 and 2), with the instantaneous point source problem modified slightly. These evaluation tests indicate that the solution algorithms in the *ELLAM* model can successfully and accurately simulate three-dimensional transport and dispersion of a solute in flowing ground water. To avoid non-physical oscillations and loss of peak concentrations, care must be taken to use a

grid having sufficient mesh density to adequately resolve sharp fronts. The primary advantages of the *ELLAM* code are that fewer transport time steps need be used and that mass is conserved globally. Using *ELLAM* with few time steps can provide an accurate and cost-effective way of discerning salient features of a solute-transport process under a complex set of boundary conditions. Furthermore, the *ELLAM* algorithm eliminates the previous restriction in *MOC3D* that the transport grid be uniformly spaced. The computational effort required by the *ELLAM* simulator is strongly dependent on the size of the problem being solved, as determined by the total number of nodes, the NS and NT values, and the total number of time increments (controlled by CELDIS, a model parameter that is analogous to the Courant number). For test cases in which *ELLAM* was more efficient than the explicit or implicit MOC options, use of CELDIS values equal to or greater than 5.0 were required.

REFERENCES

- Burnett, R.D., and Frind, E.O., 1987, Simulation of contaminant transport in three dimensions, 2. Dimensionality effects: *Water Resources Research*, v. 23, no. 4, p. 695-705.
- Celia, M.A., Russell, T.F., Herrera, I., and Ewing, R.E., 1990, An Eulerian-Lagrangian localized adjoint method for the advection-diffusion equation: *Advances in Water Resources*, v. 13, no. 4, p. 187-206.
- Gardner, A.O., Peaceman, D.W., and Pozzi, A.L., 1964, Numerical calculation of multidimensional miscible displacement by the method of characteristics: *Soc. Petroleum Eng. Jour.*, v. 4, no.1, p. 26-36.
- Goode, D.J., 1999, Age, Double porosity, and simple reaction modifications for the MOC3D ground-water transport model: U.S. Geological Survey Water-Resources Investigations Report 99-4041, 34 p.
- Harbaugh, A.W., and McDonald, M.G., 1996a, User's documentation for MODFLOW-96, an update to the U.S. Geological Survey modular finite-difference ground-water flow model: U.S. Geological Survey Open-File Report 96-485, 56 p.
- Harbaugh, A.W., and McDonald, M.G., 1996b, Programmer's documentation for MODFLOW-96, an update to the U.S. Geological Survey modular finite-difference ground-water flow model: U.S. Geological Survey Open-File Report 96-486, 220 p.
- Healy, R.W., and Russell, T.F., 1993, A finite-volume Eulerian-Lagrangian localized adjoint method for solution of the advection-dispersion equation: *Water Resources Research*, v. 29, no. 7, p. 2399-2413.
- Hsieh, P. A., 1986, A new formula for the analytical solution of the radial dispersion problem: *Water Resources Research*, v. 22, no. 11, p. 1597-1605.
- Kipp, K.L., Konikow, L.F., and Hornberger, G.Z., 1998, An implicit dispersive transport algorithm for the U.S. Geological Survey MOC3D solute-transport model: U.S. Geological Survey Water-Resources Investigations Report 98-4234, 54 p.
- Konikow, L.F., and Bredehoeft, J.D., 1978, Computer model of two-dimensional solute transport and dispersion in ground water: U.S. Geological Survey Techniques of Water-Resources Investigations, Book 7, Chapter C2, 90 p.
- Konikow, L.F., Goode, D.J., and Hornberger, G.Z., 1996, A three-dimensional method-of-characteristics solute-transport model (MOC3D): U.S. Geological Survey Water-Resources Investigations Report 96-4267, 87 p.
- McDonald, M.G., and Harbaugh, A.W., 1988, A modular three-dimensional finite-difference ground-water flow model: U.S. Geological Survey Techniques of Water-Resources Investigations, Book 6, Chapter A1, 586 p.
- Wexler, E.J., 1992, Analytical solutions for one-, two-, and three-dimensional solute transport in ground-water systems with uniform flow: U.S. Geological Survey Techniques of Water-Resources Investigations, Book 3, Chapter B7, 190 p.

APPENDIX A: DATA INPUT INSTRUCTIONS FOR *MOC3D* (Version 3.5)

This Appendix includes a complete set of instructions for preparing a data set for the *MOC3D* model. For more comprehensive descriptions of input parameters, options, and underlying assumptions, however, the user should also refer to Konikow and others (1996), Kipp and others (1998), and Goode (1999). Major changes that have been implemented since the release of Version 3.0 are shaded to highlight the new instructions.

***MODFLOW* Name File**

Transport simulation is activated by including a record in the *MODFLOW* name file using the file type (Ftype) “CONC” to link to the transport name file. The transport name file specifies the files to be used when simulating solute transport in conjunction with a simulation of ground-water flow using *MODFLOW*. The transport name file works in the same way as the *MODFLOW* name file.

***MODFLOW* Source and Sink Packages**

Except for recharge, concentrations associated with fluid sources (C') are read as auxiliary parameters in the *MODFLOW* source package. The source concentration is read from a new column appended to the end of each line of the data file describing a fluid sink/source (see documentation for revised *MODFLOW* model; Harbaugh and McDonald, 1996a and 1996b). For example, concentrations associated with well nodes should be appended to the line in the WEL Package where the well's location and pumping rate are defined. These concentrations will be read if the auxiliary parameter “CONCENTRATION” (or “CONC”) appears on the first line of the well input data file. The concentration in recharge is defined separately, as described in following section “Source Concentration in Recharge File.”

To simulate solute transport the *MODFLOW* option enabling storage of cell-by-cell flow rates for each fluid source or sink is required in all fluid packages except recharge. The key word “CBCALLOCATE” (or “CBC”) must appear on the first line of each input data file for a fluid package (see Harbaugh and McDonald, 1996a and 1996b).

***MOC3D* Input Data Files**

All input variables are read using free formats, except as specifically indicated. In free format, variables are separated by one or more spaces or by a comma and optionally one or more spaces. Blank spaces are not read as zeros. Variables that are optional are enclosed in brackets, as in {option}.

MOC3D Transport Name File (CONC)

FOR EACH SIMULATION:

1. Data: FTYPE NUNIT FNAME

The name file consists of records defining the names and unit numbers of the files. Each “record” consists of a separate line of data. There must be a record for the listing file and for the main MOC3D input file.

The listing (or output) file (“CLST”) must be the first record. The other files may be in any order. Each record can be no more than 79 characters.

FTYPE The file type, which may be one of the following character strings:

CLST	MOC3D listing file (separate from the MODFLOW listing file) [required].
MOC or MOCIMP or ELLAM	Main MOC3D input data file [required]. Specifying MOC indicates dispersion calculations will be explicit (as described by Konikow and others, 1996); specifying MOCIMP indicates dispersion calculations will be implicit (as described by Kipp and others, 1998); and specifying ELLAM indicates that the solute-transport equation will be solved using the <i>ELLAM</i> method (as described in this report).
CRCH	Concentrations in recharge [optional].
CNCA	Separate output file containing concentration data in ASCII (text-only) format [optional]. Frequency and format of printing controlled by NPNTCL and ICONFM. If concentrations are written to a separate output file, they will not be written to the main output file.
CNCB	Separate output file containing concentration data in binary format [optional].
VELA	Separate output file with velocity data in ASCII format [optional]. Frequency and format of printing controlled by NPNTVL and IVELFM.
VELB	Separate output file with velocity data in binary format [optional].
PRTA	Separate output file with particle locations printed in ASCII format [optional]. Frequency and format of printing controlled by NPNTPL.
PRTB	Separate output file with particle locations printed in binary format [optional].
OBS	Observation wells input file [optional].
DATA	For formatted files such as those required by the OBS package and for array data separate from the main MOC3D input data file [optional].
DATA (BINARY)	For formatted input/output files [optional].
AGE	Ground-water age simulation input file [optional]. (Not compatible with <i>ELLAM</i> option.)
DP	Double porosity input file [optional]. (Not compatible with <i>ELLAM</i> option.)

DK Simple reactions(decay, zero-order growth, retardation) input file [*optional*].
(Not compatible with *ELLAM* option.)

NUNIT The FORTRAN unit number used to read from and write to files. Any legal unit number other than 97, 98, and 99 (which are reserved by *MODFLOW*) can be used provided that it is not previously specified in the *MODFLOW* name file.

FNAME The name of the file.

Note: AGE, DP, and DK file types are described by Goode (1999).

Main *MOC3D* Package Input (*MOC*, *MOCIMP*, or *ELLAM*)

Input for the method-of-characteristics (*MOC3D*) solute-transport package is read from the unit specified in the transport name file. The input consists of up to 19 separate records or data sets, as described in detail below (note that data set numbers do not necessarily correspond with line numbers in the file). These data are used to specify information about the transport subgrid, physical and chemical transport parameters, numerical solution variables, and output formats. Output file controls for the *MOC3D* package are specified in the transport name file, described previously. Compared to previous versions of *MOC3D*, use of the *ELLAM* option requires definition of several alternative input parameters and deletes two previously required data sets (6 and 13), which are used only if *MOC* or *MOCIMP* is selected.

FOR EACH SIMULATION:

1. Data: HEDMOC A two-line character-string title describing the simulation (80 text characters per line).

2. Data: HEDMOC (continued)

3. Data: ISLAY1 ISLAY2 ISROW1 ISROW2 ISCOL1 ISCOL2

ISLAY1 Number of first (uppermost) layer for transport.

ISLAY2 Last layer for transport.

ISROW1 First row for transport.

ISROW2 Last row for transport.

ISCOL1 First column for transport.

ISCOL2 Last column for transport.

Notes:

Transport may be simulated within a subgrid, which is a “window” within the primary *MODFLOW* grid used to simulate flow. Within the subgrid, the row and column spacing must be uniform if *FTYPE* *MOC* or *MOCIMP* are specified in the transport name file, but subgrid spacing can vary as in *MODFLOW*

if *ELLAM* is specified. The thickness can vary from cell to cell and layer to layer. However, the range in thickness values (or product of thickness and porosity) should be as small as possible.

4. Data: NODISP DECAY DIFFUS

NODISP	Flag for no dispersion (set NODISP = 1 if no dispersion in problem; this will reduce storage allocation).
DECAY	First-order decay rate [1/T] (DECAY = 0.0 indicates no decay occurs).
DIFFUS	Effective molecular diffusion coefficient [L^2/T].

Notes:

The decay rate (λ) is related to the half life ($t_{1/2}$) of a constituent by $\lambda = (\ln 2)/t_{1/2}$.

The effective molecular diffusion coefficient (D_m) includes the effect of tortuosity.

IF Ftype *MOC* OR *MOCIMP* IS ACTIVE:

5a. Data: NPMAX NPTPND

NPMAX	Maximum number of particles available for particle tracking of advective transport in <i>MOC3D</i> . If set to zero, the model will calculate NPMAX according to the following equation:
-------	--

$$NPMAX = 2 \times NPTPND \times NSROW \times NSCOL \times NSLAY.$$

NPTPND	Initial number of particles per cell in transport simulation (that is, at $t = 0.0$). Valid options for default geometry of particle placement include 1, 2, 3, or 4 for one-dimensional transport simulation; 1, 4, 9, or 16 for two-dimensional transport simulation; and 1, 8, or 27 for three-dimensional transport simulation. The user can also customize initial placement of particles by specifying NPTPND as a negative number, in which case the minus sign is recognized as a flag to indicate custom placement is desired. In this case, the user must input local particle coordinates as described below.
--------	---

IF Ftype *ELLAM* IS ACTIVE:

5b. Data: NSCEXP NSREXP NSLEXP NTEXP

NSCEXP	Exponent used to calculate the number of subcells in the column direction (NSC, where $NSC = 2^{*}NSCEXP$).
NSREXP	Exponent used to calculate the number of subcells in the row direction (NSR).
NSLEXP	Exponent used to calculate the number of subcells in the layer direction (NSL).
NTEXP	Exponent used to calculate the number of sub-time steps per transport time increment (NT).

Notes:

In general, numerical accuracy will be increased by increasing the value of these parameters. This will also, however, increase computational costs. For each of the four parameters above, the value represents the exponent y in the expression 2^y .

Entering a zero or negative value for any of the above variables will cause the code to use default values. Default values for NSCEXP, NSREXP, and NSLEXP are 2 in active dimensions and 1 in inactive dimensions (for example, if a simulation represented a two-dimensional areal problem in which the number of rows and columns were greater than one and the number of layers equals one, then default settings would be NSCEXP = 2, NSREXP = 2, and NSLEXP = 1, and the number of subcells in each direction would be 4, 4, and 2, respectively). The default value of NTEXP is 2.

IF *MOC* OR *MOCIMP* IS ACTIVE AND IF *NPTPND* IS NEGATIVE IN SIGN:

6. Data: PNEWL PNEWR PNEWC

PNEWL Relative position in the layer (z) direction for initial placement of particle within any finite-difference cell.

PNEWR Relative position in the row (y) direction for initial placement of particle.

PNEWC Relative position in the column (x) direction for initial placement of particle.

Notes:

The three new (or initial) particle coordinates are entered sequentially for each of the NPTPND particles. Each line contains the three relative local coordinates for the new particles, in order of layer, row, and column. There must be NPTPND lines of data, one for each particle. The local coordinate system range is from -0.5 to 0.5, and represents the relative distance within the cell about the node location at the center of the cell, so that the node is located at 0.0 in each direction.

FOR EACH SIMULATION:

7. Data: CELDIS {FZERO} {INTRPL}

CELDIS Maximum fraction of cell dimension that particle may move in one step (typically, $0.5 \leq \text{CELDIS} \leq 1.0$). For *ELLAM*, CELDIS can be greater than 1.0, and specifying CELDIS = 0.0 will result in one transport time step being used (which is not generally recommended).

FZERO If the fraction of active cells having no particles exceeds FZERO, the program will automatically regenerate an initial particle distribution before continuing the simulation (typically, $0.01 \leq \text{FZERO} \leq 0.05$). Only specify if *MOC* or *MOCIMP* is active.

INTRPL Flag for interpolation scheme used to estimate velocity of particles. The default (INTRPL = 1) will use a linear interpolation routine; if INTRPL = 2, a scheme will be implemented that uses bilinear interpolation in the row and column (j and i) directions only (linear interpolation will still be applied in the k , or layer, direction). (See section "Discussion—Choosing appropriate interpolation scheme.") Only specify if *MOC* or *MOCIMP* is active. If *ELLAM* is specified, the code will automatically set INTRPL = 1.

IF *MOCIMP* IS ACTIVE:

7.1 Data: FDTMTH NCXIT IDIREC EPSSLV MAXIT

- FDTMTH Weighting factor for temporal differencing of dispersion equation ($0.0 \leq \text{FDTMTH} \leq 1.0$). We suggest using either a value of $\text{FDTMTH} = 0.5$, a centered-in-time (or Crank-Nicolson) approximation, or $\text{FDTMTH} = 1.0$, a backward-in-time (or fully implicit) approximation. [Default value = 1]
- NCXIT Number of iterations for the explicitly-lagged cross-dispersive flux terms ($\text{NCXIT} \geq 1$). We suggest that the user initially specify a value of 2, but if the solution exhibits significant areas of negative concentrations, then the value of NCXIT should be increased to require more iterations, which typically will reduce the extent and magnitude of negative concentrations (at the cost of increased computational time). [Default value = 2]
- IDIREC Direction index for permutation of the red-black node renumbering scheme. The order is as follows: 1: x,y,z; 2: x,z,y; 3: y,x,z; 4: y,z,x; 5: z,x,y; and 6: z,y,x. The first direction index is advanced most rapidly and the last direction index is advanced least rapidly. In some cases, there can be a significant variation in the number of iterations needed to achieve convergence, depending on the order of the directions for the red-black renumbering. We suggest that the user initially specify $\text{IDIREC} = 1$. If this leads to a relatively large number of iterations (more than 10), then the user should experiment with alternate choices to determine the one requiring the fewest number of iterations for their particular problem. [Default value = 1]
- EPSSLV Tolerance on the relative residual for the conjugate-gradient solution of the matrix of the difference equations. We suggest that the user initially specify $\text{EPSSLV} \leq 10^{-5}$. An adequately small value of EPSSLV has the property that a smaller value does not change the numerical solution within the number of significant digits desired by the user. In the single-precision code implemented here, EPSSLV should not be less than 10^{-7} . [Default value = 10^{-5}]
- MAXIT Maximum number of iterations allowed for the iterative solution to the difference equations for dispersive transport. In most cases, $\text{MAXIT} = 100$ is satisfactory. [Default value = 100]

Notes:

Entering a zero or out-of-range value for any of these five variables will cause the code to use the indicated default value.

FOR EACH SIMULATION:

8. Data: NPNTCL ICONFM NPNTVL IVELFM NPNTDL IDSPFM {NPNTPL}

- NPNTCL** Flag for frequency of printing concentration data. If NPNTCL = -2, concentration data will be printed at the end of every stress period; if NPNTCL = -1, data will be printed at the end of every flow time step; if NPNTCL = 0, data will be printed at the end of the simulation; if NPNTCL = N > 0, data will be printed every Nth particle moves, and at the end of the simulation. Initial concentrations are always printed. Solute budget and mass balance information are only printed every time concentration data are saved.
- ICONFM** Flag for output format control for printing concentration data. If concentration data are written to main output file (file type CNCA is not used), ICONFM represents a code indicating the format style (table 9, also see Harbaugh and McDonald, 1996a, p. 19). If concentration data are written to a separate output file (file type CNCA exists), specifying ICONFM ≥ 0 will indicate that concentration data are to be written as a matrix of values for each layer of the subgrid, whereas specifying ICONFM < 0 will indicate that concentration data are to be written as a table of values having one row for each node in the subgrid and four columns (x, y, z, and concentration), where x, y, and z are the actual nodal coordinates in the length units of the model simulation. Note that we follow the *MODFLOW* convention in that y increases from top to bottom row, and z increases from top layer to bottom layer. Also note that the x and y values are given with respect to the entire MODFLOW grid, but the z location is calculated only for vertical distances within the layers of the transport subgrid. If data are written in matrix style, one header line precedes and identifies the data for each layer. If data are written as a table of values, one header line is written each time that concentration data are saved.
- NPNTVL** Flag for printing velocity data. If NPNTVL = -1, velocity data will be printed at the end of every stress period; if NPNTVL = 0, data will be printed at the end of the simulation; if NPNTVL = N > 0, data will be printed every Nth flow time steps, and at the end of the simulation.
- IVELFM** Specification for format of velocity data, if being printed in main output file (see table 9).
- NPNTDL** Flag for printing dispersion equation coefficients that include cell dimension factors (see section “Program Segments”). If NPNTDL = -2, coefficients will be printed at the end of every stress period; if NPNTDL = -1, coefficients will be printed at the end of the simulation; if NPNTDL = 0, coefficients will not be printed; if NPNTDL = N > 0, coefficients will be printed every Nth flow time step.
- IDSPFM** Specification for format of dispersion equation coefficients (see table 9).
- NPNTPL** Flag for printing particle locations in a separate output file (only used if file types “PRTA” or “PRTB” appear in the *MOC3D* name file). If neither “PRTA” or “PRTB” is entered in the name file, NPNTPL will be read but ignored (so you must always have some value specified here). If either “PRTA” or “PRTB” is entered in the name file, initial particle locations will be printed to the separate file first,

followed by particle data at intervals determined by the value of NPNTPL. If NPNTPL = -2, particle data will be printed at the end of every stress period; if NPNTPL = -1, data will be printed at the end of every flow time step; if NPNTPL = 0, data will be printed at the end of the simulation; if NPNTPL = N > 0, data will be printed every Nth particle moves, and at the end of the simulation. Only specify if *MOC* or *MOCIMP* is active.

Table 9. Formats associated with *MOC3D* print flags. (Positive values for wrap format; negative values for strip format. Also see Harbaugh and McDonald, 1996a, p. 19.)

Print flag	Format	Print flag	Format	Print flag	Format
0	10G11.4	7	20F5.0	14	10F6.1
1	11G10.3	8	20F5.1	15	10F6.2
2	9G13.6	9	20F5.2	16	10F6.3
3	15F7.1	10	20F5.3	17	10F6.4
4	15F7.2	11	20F5.4	18	10F6.5
5	15F7.3	12	10G11.4		
6	15F7.4	13	10F6.0		

FOR EACH SIMULATION:

9. Data: CNOFLO Concentration associated with inactive cells of subgrid (used for output purposes only).

FOR EACH LAYER OF THE TRANSPORT SUBGRID:

10. Data: CINT (NSCOL, NSROW) Initial concentration.
Module: U2DREL*

FOR EACH SIMULATION, ONLY IF TRANSPORT SUBGRID DIMENSIONS ARE SMALLER THAN FLOW GRID DIMENSIONS:

11. Data: CINFL (ICINFL) C' to be associated with fluid inflow across the boundary of the subgrid.
Module: U1DREL*

Notes:

The model assumes that the concentration outside of the subgrid is the same within each layer, so only one value of CINFL is specified for each layer within and adjacent to the subgrid. That is, the size of the array (ICINFL) is determined by the position of the subgrid with respect to the entire (primary)

* Module is a standard *MODFLOW* input/output module.

MODFLOW grid. If the transport subgrid has the same dimensions as the flow grid, this parameter should not be included in the input data set. If the subgrid and flow grid have the same number of layers, but the subgrid has fewer rows or fewer columns, $ICINFL = NSLAY$. Values are also required if there is a flow layer above the subgrid and/or below the subgrid. The order of input is: C' for first (uppermost) transport layer (if required); C' for each successive (deeper) transport layer (if required); C' for layer above subgrid (if required); and C' for layer below subgrid (if required).

FOR EACH SIMULATION:

12. Data: NZONES Number of zone codes among fixed-head nodes in transport subgrid.

IF *NZONES* > 0:

Data: IZONE ZONCON

IZONE Value identifying a particular zone.

ZONCON Source concentration associated with nodes in the zone defined by IZONE above.

Notes:

Zones are defined within the IBOUND array in the BAS Package of *MODFLOW* by specifying unique negative values for fixed-head nodes to be associated with separate fluid source concentrations. Each zone is defined by a unique value of IZONE and a concentration associated with it (ZONCON). There must be NZONES lines of data, one for each zone. Note that values of IZONE in this list must be negative for consistency with the definitions of fixed-head nodes in the IBOUND array in the BAS Package. If a negative value of IBOUND is defined in the BAS package but is not assigned a concentration value here, *MOC3D* will assume that the source concentrations associated with those nodes equal 0.0.

FOR EACH LAYER OF THE TRANSPORT SUBGRID IF *MOC* OR *MOCIMP* IS ACTIVE:

13. Data: IGENPT (NSCOL, NSROW) Flag to treat fluid sources and sinks as either "strong" or "weak."

Module: U2DINT*

Notes:

Where fluid source is "strong," new particles are added to replace old particles as they are advected out of that cell. Where a fluid sink is "strong," particles are removed after they enter that cell and their effect accounted for. Where sources or sinks are weak, particles are neither added nor removed, and the source/sink effects are incorporated directly into appropriate changes in particle positions and concentrations. If IGENPT = 0, the node will be considered a weak source or sink; if IGENPT = 1, it will be a strong source or sink. See section on "Special Problems" and discussion by Konikow and Bredehoeft (1978).

* Module is a standard *MODFLOW* input/output module.

IF *NODISP* \neq 1 (If dispersion is included in simulation):

- | | | | |
|------------|---------|---------------|-------------------------------------|
| 14. | Data: | ALONG (NSLAY) | Longitudinal dispersivity. |
| | Module: | U1DREL* | |
| | | | |
| 15. | Data: | ATANH (NSLAY) | Horizontal transverse dispersivity. |
| | Module: | U1DREL* | |
| | | | |
| 16. | Data: | ATRV (NSLAY) | Vertical transverse dispersivity. |
| | Module: | U1DREL* | |

Notes:

Data sets 14-16 should include one value for each layer in subgrid.

FOR EACH SIMULATION:

- | | | | |
|------------|---------|------------|---|
| 17. | Data: | RF (NSLAY) | Retardation factor (RF = 1 indicates no retardation). |
| | Module: | U1DREL* | |

Notes:

If RF = 0.0 in input, the code automatically resets it as RF = 1.0 to indicate no retardation.

FOR EACH LAYER OF TRANSPORT SUBGRID:

- | | | | |
|-------------|---------|---------------------|-----------------|
| 18a. | Data: | THCK (NSCOL, NSROW) | Cell thickness. |
| | Module: | U2DREL* | |
| | | | |
| 18b. | Data: | POR (NSCOL, NSROW) | Cell porosity. |
| | Module: | U2DREL* | |

Notes:

The thickness and porosity are input as separate arrays for each layer of the transport subgrid. The sequence used in data set 18 is to first define the thickness of the first layer of the transport subgrid, and then define the porosity of that same layer. Next, that sequence is repeated for all succeeding layers. The product of thickness and porosity should not be allowed to vary greatly among cells in the transport subgrid. If the *ELLAM* option is being used, the variation in thickness (not thickness times porosity) between neighboring cells within a layer should be small.

* Module is a standard *MODFLOW* input/output module.

Source Concentration in Recharge File (CRCH)

Concentrations in recharge, if the recharge package is used, are read from a separate unit specified in the *MOC3D* name file. This is defined using the file type (Ftype) "CRCH."

FOR EACH STRESS PERIOD, IF RECHARGE PACKAGE USED:

1. Data: INCRCH Flag to reuse or read new recharge concentrations.

Notes:

Read new recharge concentrations if $\text{INCRCH} \geq 0$. Reuse recharge concentrations from the last stress period if $\text{INCRCH} < 0$.

2. Data: CRECH (NSCOL, NSROW) Source concentration associated with fluid entering the aquifer in recharge.
Module: U2DREL*

Observation Well File (OBS)

Nodes of the transport subgrid can be designated as "observation wells." At each such node, the time, head, and concentration after each move increment will be written to a separate output file to facilitate graphical postprocessing of the calculated data. The input file for specifying observation wells is read if the file type (Ftype) "OBS" is included in the *MOC3D* name file.

FOR EACH SIMULATION, IF OBS PACKAGE USED:

1. Data: NUMOBS IOBSFL

NUMOBS Number of observation wells.

IOBSFL If $\text{IOBSFL} = 0$, well data are saved in NUMOBS separate files. If $\text{IOBSFL} > 0$, all observation well data will be written to one file, and the file name and unit number used for this file will be that of the first observation well in the list.

FOR EACH OBSERVATION WELL:

2. Data: LAYER ROW COLUMN UNIT

LAYER Layer of observation well node.

ROW Row of observation well node.

COLUMN Column of observation well node.

UNIT Unit number for output file.

Notes:

If $\text{NUMOBS} > 1$ and $\text{IOBSFL} = 0$, you must specify a unique unit number for each observation well and match those unit numbers to DATA file types and file names in the *MOC3D* name file. If $\text{IOBSFL} > 0$, you must specify a unique unit number for the first observation well and match that unit number to a DATA file type and file name in the *MOC3D* name file.

* Module is a standard *MODFLOW* input/output module.

File name: *finite.sip*

500	5	; MXITER,NPARM			SIP Input	← 1
1.	0.0000001	0	0.001	0	; ACCL,HCLOSE,IPCALC,WSEED,IPRSIP	← 2

¹ Maximum iterations, number of iteration parameters

² Acceleration parameter, head change criterion, flag for seed, seed, printout interval for SIP

Following (enclosed in a border) are the contents of the *MOC3D* name file for the sample problem; explanations are noted outside of border:

File name: *fint_moc.nam*

clst	94	finite.out	← Designates main output file for <i>MOC3D</i>
ellam	96	finite.ell	← Main input data file for <i>ELLAM</i>
obs	44	finite.obs	← Input data file for observation wells
data	45	finite.oba	← Output file for observation well data
cnca	22	finite.cna	← Separate output file for concentration data (ASCII)

↑	↑	↑
1	2	3

¹ Ftype

² Unit number

³ File name

Following (enclosed in a border) are the contents of the main input data file for the *MOC3D* simulation for the sample problem; selected explanations are noted outside of border:

File name: *finite.ell*

One-dimensional, Steady Flow, No Decay, Low Dispersion: MOC3D (ELLAM)						← 1
ISLAY1	ISLAY2	ISROW1	ISROW2	ISCOL1	ISCOL2	← 1
1	1	1	1	2	121	← 2
0	0.00	0.0	; NODISP, DECAY, DIFFUS			← 3
5	1	1	7	; NSCEXP, NSREXP, NSLEXP, NTEXP		← 4
1.0	1	; CELDIS, INTRPL				← 5
0	-1	0	0	0	0	← 6
; NPNTCL, ICONFM, NPNTVL, IVELFM, NPNTDL, IDSPFM, NPRTPL						← 7
0.0	; CNOFLO					
0	0.0	(122F3.0)				
0	; initial concentration					
2	1.	; C' inflow				
-1	1.0	; NZONES to follow				← 8
-2	0.0	; IZONE, ZONCON				← 8
0	0.1	; IZONE, ZONCON				← 8
0	0.1	; longitudinal disp.				
0	0.1	; transverse disp. horiz.				
0	0.1	; transverse disp. vert.				
0	1.0	; retardation factor				
0	1.0	; thickness				
0	0.1	; porosity				

- ¹ Two header lines of comments. For convenience and clarity, the second line is used to label names of parameters on subsequent line of file.
- ² Indices for transport subgrid
- ³ Flag for no dispersion, decay rate, diffusion coefficient
- ⁴ *ELLAM* parameters (exponents for NSC, NSR, NSL, and NT)
- ⁵ Courant number and interpolation method flag
- ⁶ Print flags
- ⁷ Value of concentration associated with inactive cells
- ⁸ Concentrations associated with fixed-head nodes (fixed head nodes are defined in the IBOUND array in the *MODFLOW* BAS package)

Following (enclosed in a border) are the contents of the observation well input data file for the sample problem; explanations are noted outside of border:

File name: *finite.obs*

3	1	; NUMOBS IOBSFL		Observation well data	← 1
1	1	2	45	; layer, row, column, unit number	← 2
1	1	42	; layer, row, column		← 2
1	1	112	; layer, row, column		← 2

- ¹ Number of observation wells, flag to print to one file or separate files
- ² Node location and unit number for output file (linked to the Ftype DATA in *MOC3D* name file)

APPENDIX C: SELECTED OUTPUT FOR EXAMPLE PROBLEM

This example output was generated from the input data sets listed in Appendix B for a case of the one-dimensional steady-state flow problem. We do not include the main *MODFLOW* listing (output) file. The line spacing and font sizes of the output files have been modified in places to enhance the clarity of reproduction in this report. Some repetitive lines of output have been deleted where indicated by an ellipsis (...). Output related specifically to *ELLAM* is highlighted by shading.

Some brief annotations were added to this sample output listing to help the reader understand the purpose of various sections of output. These annotations are written in bold italics.

Following are the contents of the *MOC3D* main output file (*finite.out*) for the sample problem.

U.S. GEOLOGICAL SURVEY
METHOD-OF-CHARACTERISTICS SOLUTE-TRANSPORT MODEL
MOC3D (Version 3.5) July 2000

MOC BASIC INPUT READ FROM UNIT
LISTING FILE: finite.out UNIT 94

OPENING finite.ell
FILE TYPE: ELLAM UNIT 96

OPENING finite.obs
FILE TYPE: OBS UNIT 44

OPENING finite.oba
FILE TYPE: DATA UNIT 45

OPENING finite.cna
FILE TYPE: CNCA UNIT 22

MOC BASIC INPUT READ FROM UNIT 96

FILE INFORMATION

2 TITLE LINES:

One-dimensional, Steady Flow, No Decay, Low Dispersion: ELLAM
ISLAY1 ISLAY2 ISROW1 ISROW2 ISCOL1 ISCOL2

PROBLEM DESCRIPTORS, INCLUDING GRID CHARACTERISTICS AND ELLAM INPUT INFORMATION:

MAPPING OF SOLUTE TRANSPORT SUBGRID IN FLOW GRID:
FIRST LAYER FOR SOLUTE TRANSPORT = 1 LAST LAYER FOR SOLUTE TRANSPORT = 1
FIRST ROW FOR SOLUTE TRANSPORT = 1 LAST ROW FOR SOLUTE TRANSPORT = 1
FIRST COLUMN FOR SOLUTE TRANSPORT = 2 LAST COLUMN FOR SOLUTE TRANSPORT = 121

NONUNIFORM DELCOL AND DELROW IN SUBGRID FOR SOLUTE TRANSPORT
NO. OF LAYERS = 1 NO. OF ROWS = 1 NO. OF COLUMNS = 120

NO SOLUTE DECAY
NO MOLECULAR DIFFUSION

ELLAM INPUT PARAMETERS:

NSCEXP, NSREXP, NSLEXP, NTEXP

5 1 1 7

NSC, NSR, NSL, NT (CALCULATED)

32 2 2 128

18288 ELEMENTS IN X ARRAY ARE USED BY MOC

CELDIS= 1.0

NPNTCL= 0: CONCENTRATIONS WILL BE WRITTEN AT THE END OF THE SIMULATION
MODFLOW FORMAT SPECIFIER FOR CONCENTRATION DATA: ICONFM= -1

NPNTVL= 0: VELOCITIES WILL BE WRITTEN AT THE END OF THE SIMULATION
MODFLOW FORMAT SPECIFIER FOR VELOCITY DATA: IVELFM= 0

NPNTDL= 0: DISP. COEFFICIENTS WILL NOT BE WRITTEN

CONCENTRATION WILL BE SET TO 0.00000E+00 AT ALL NO-FLOW NODES (IBOUND=0).

INITIAL CONCENTRATION = 0.0000000E+00 FOR LAYER 1

VALUES OF C' REQUIRED FOR SUBGRID BOUNDARY ARRAY = 1
ONE FOR EACH LAYER IN TRANSPORT SUBGRID

ORDER OF C' VALUES: FIRST LAYER IN SUBGRID, EACH SUBSEQUENT LAYER,
LAYER ABOVE SUBGRID, LAYER BELOW SUBGRID:

SUBGRID BOUNDARY ARRAY = 1.000000

NUMBER OF ZONES FOR CONCENTRATIONS AT FIXED HEAD CELLS = 2

ZONE FLAG = -1 INFLOW CONCENTRATION = 1.0000E+00

ZONE FLAG = -2 INFLOW CONCENTRATION = 0.0000E+00

SINK-SOURCE FLAG = 0 FOR LAYER 1

LONGITUDNL. DISPERSIVITY = 0.1000000

HORIZ. TRANSVERSE DISP. = 0.1000000

VERT. TRANSVERSE DISP. = 0.1000000

RETARDATION FACTOR = 1.000000

INITIAL THICKNESS = 1.000000 FOR LAYER 1

INITIAL POROSITY = 0.1000000 FOR LAYER 1

COORDINATES FOR 3 OBSERVATION WELLS:

WELL #	LAYER	ROW	COLUMN	UNIT
1	1	1	2	45
2	1	1	42	45
3	1	1	112	45

OUTPUT
CONTROL

INITIAL AND
BOUNDARY
CONDITIONS
FOR SOLUTE

ALL OBSERVATION WELL DATA WILL BE WRITTEN ON UNIT 45

CONCENTRATION DATA WILL BE SAVED ON UNIT 22 IN X,Y,Z,CONC FORMAT

CALCULATED VELOCITIES (INCLUDING EFFECTS OF RETARDATION, IF PRESENT):

EFFECTIVE MEAN SOLUTE VELOCITIES IN COLUMN DIRECTION
AT NODES

```
1
  VELOCITY (COL)   IN LAYER 1 AT END OF TIME STEP 1 IN STRESS PERIOD 1
  -----
      1         2         3         4         5         6         7...
...
.....
1  0.1000      0.1000      0.1000      0.1000      0.1000      0.1000      0.1000...
...
.....
```

EFFECTIVE MEAN SOLUTE VELOCITIES IN ROW DIRECTION
AT NODES

```
1
  VELOCITY (ROW)   IN LAYER 1 AT END OF TIME STEP 1 IN STRESS PERIOD 1
  -----
      1         2         3         4         5         6         7         8         9         ...
...
.....
1  0.000E+00  0.000E+00  0.000E+00  0.000E+00  0.000E+00  0.000E+00  0.000E+00  0.000E+00  0.000E+00  ...
...
.....
```

EFFECTIVE MEAN SOLUTE VELOCITIES IN LAYER DIRECTION
AT NODES

```
1
  VELOCITY (LAYER) IN LAYER 1 AT END OF TIME STEP 1 IN STRESS PERIOD 1
  -----
      1         2         3         4         5         6         7         8         9         ...
...
.....
1  0.000E+00  0.000E+00  0.000E+00  0.000E+00  0.000E+00  0.000E+00  0.000E+00  0.000E+00  0.000E+00  ...
...
.....
```

NUMBER OF MOVES FOR CELDIS CRITERIA:
120

TIME STEP 1 IN STRESS PERIOD 1
NO. OF PARTICLE MOVES REQUIRED TO COMPLETE THIS TIME STEP = 120
MOVE TIME STEP (TIMV) = 1.000000000000E+00

SOLUTE BUDGET AND MASS BALANCE FOR TRANSPORT SUBGRID

VALUES CALCULATED AT END OF:
 STRESS PERIOD 1 OUT OF 1
 FLOW TIME STEP 1 OUT OF 1
 TRANSPORT TIME INCREMENT 120 OUT OF 120

ELAPSED TIME = 1.2000E+02

CHEMICAL MASS IN STORAGE:

INITIAL: MASS DISSOLVED = 0.0000E+00 MASS SORBED = 0.0000E+00
 PRESENT: MASS DISSOLVED = 1.0203E-01 MASS SORBED = 0.0000E+00

CHANGE IN MASS STORED = -1.0203E-01

CUMULATIVE SOLUTE MASS (L**3) (M/VOL)

IN:

DECAY = 0.0000E+00
 CONSTANT HEAD = 0.0000E+00
 SUBGRID BOUNDARY = 1.2000E-01
 RECHARGE = 0.0000E+00
 WELLS = 0.0000E+00
 RIVERS = 0.0000E+00
 DRAINS = 0.0000E+00
 GENL. HEAD-DEP. BDYS. = 0.0000E+00
 EVAPOTRANSPIRATION = 0.0000E+00
 SPECIFIED FLOW (FHB) = 0.0000E+00

TOTAL IN = 1.2000E-01

OUT:

DECAY = 0.0000E+00
 CONSTANT HEAD = 0.0000E+00
 SUBGRID BOUNDARY = -1.7972E-02
 RECHARGE = 0.0000E+00
 WELLS = 0.0000E+00
 RIVERS = 0.0000E+00
 DRAINS = 0.0000E+00
 GENL. HEAD-DEP. BDYS. = 0.0000E+00
 EVAPOTRANSPIRATION = 0.0000E+00
 SPECIFIED FLOW (FHB) = 0.0000E+00

TOTAL OUT = -1.7972E-03

SOURCE-TERM DECAY = 0.0000E+00

RESIDUAL = 2.5183E-06

PERCENT DISCREPANCY = 2.0986E-03 RELATIVE TO MASS FLUX IN

ITEMIZED BUDGETS

FOR SOLUTE

FLUXES

Following (enclosed in a border) are the abridged contents of the observation well output file for the sample problem. This output file was generated using the option to write all observation well data to a single file (IOBSFL = 1).

File name: *finite.oba*

"OBSERVATION WELL DATA"						
"TIME, THEN HEAD AND CONC. FOR EACH OBS. WELL AT NODE (K,I,J) "						
" TIME:	H & C AT	1, 1, 2	H & C AT	1, 1, 42	H & C AT	1, 1, 112 "
0.0000E+00	0.000E+00	0.000E+00	0.000E+00	0.000E+00	0.000E+00	0.000E+00
1.0000E+00	1.200E+01	2.587E-01	8.000E+00	8.100E-07	1.000E+00	1.887E-16
2.0000E+00	1.200E+01	3.780E-01	8.000E+00	8.271E-06	1.000E+00	2.499E-15
3.0000E+00	1.200E+01	4.558E-01	8.000E+00	4.364E-05	1.000E+00	2.128E-14
4.0000E+00	1.200E+01	5.140E-01	8.000E+00	1.591E-04	1.000E+00	1.447E-13
5.0000E+00	1.200E+01	5.606E-01	8.000E+00	4.519E-04	1.000E+00	8.378E-13
6.0000E+00	1.200E+01	5.993E-01	8.000E+00	1.070E-03	1.000E+00	4.233E-12
7.0000E+00	1.200E+01	6.322E-01	8.000E+00	2.202E-03	1.000E+00	1.971E-11
8.0000E+00	1.200E+01	6.606E-01	8.000E+00	4.060E-03	1.000E+00	8.664E-11
9.0000E+00	1.200E+01	6.856E-01	8.000E+00	6.853E-03	1.000E+00	3.646E-10
1.0000E+01	1.200E+01	7.078E-01	8.000E+00	1.076E-02	1.000E+00	1.404E-09
...						
...						
1.1100E+02	1.200E+01	9.955E-01	8.000E+00	9.428E-01	1.000E+00	5.284E-01
1.1200E+02	1.200E+01	9.957E-01	8.000E+00	9.446E-01	1.000E+00	5.374E-01
1.1300E+02	1.200E+01	9.958E-01	8.000E+00	9.463E-01	1.000E+00	5.464E-01
1.1400E+02	1.200E+01	9.959E-01	8.000E+00	9.479E-01	1.000E+00	5.553E-01
1.1500E+02	1.200E+01	9.960E-01	8.000E+00	9.495E-01	1.000E+00	5.640E-01
1.1600E+02	1.200E+01	9.962E-01	8.000E+00	9.511E-01	1.000E+00	5.727E-01
1.1700E+02	1.200E+01	9.963E-01	8.000E+00	9.526E-01	1.000E+00	5.813E-01
1.1800E+02	1.200E+01	9.964E-01	8.000E+00	9.540E-01	1.000E+00	5.897E-01
1.1900E+02	1.200E+01	9.965E-01	8.000E+00	9.554E-01	1.000E+00	5.980E-01
1.2000E+02	1.200E+01	9.966E-01	8.000E+00	9.568E-01	1.000E+00	6.063E-01

Following (enclosed in a border) are the partial contents of the separate ASCII output file for concentration in a table format style. Initial concentrations follow the first header line; final concentrations follow the second (internal) header line.

File name: *finite.cna*

```

CONCENTRATIONS AT NODES (X,Y,Z,CONC): IMOV= 0, KSTP= 0, KPER= 0, SUMTCH=0.0000E+00
 1.5000E-01  5.0000E-02  5.0000E-01  0.0000E+00
 2.5000E-01  5.0000E-02  5.0000E-01  0.0000E+00
 3.5000E-01  5.0000E-02  5.0000E-01  0.0000E+00
 4.5000E-01  5.0000E-02  5.0000E-01  0.0000E+00
 5.5000E-01  5.0000E-02  5.0000E-01  0.0000E+00
 6.5000E-01  5.0000E-02  5.0000E-01  0.0000E+00
 7.5000E-01  5.0000E-02  5.0000E-01  0.0000E+00
 8.5000E-01  5.0000E-02  5.0000E-01  0.0000E+00
 9.5000E-01  5.0000E-02  5.0000E-01  0.0000E+00
...
 1.1650E+01  5.0000E-02  5.0000E-01  0.0000E+00
 1.1750E+01  5.0000E-02  5.0000E-01  0.0000E+00
 1.1850E+01  5.0000E-02  5.0000E-01  0.0000E+00
 1.1950E+01  5.0000E-02  5.0000E-01  0.0000E+00
 1.2050E+01  5.0000E-02  5.0000E-01  0.0000E+00
CONCENTRATIONS AT NODES (X,Y,Z,CONC): IMOV= 120, KSTP= 1, KPER= 1, SUMTCH=1.2000E+02
 1.5000E-01  5.0000E-02  5.0000E-01  9.9660E-01
 2.5000E-01  5.0000E-02  5.0000E-01  9.9667E-01
 3.5000E-01  5.0000E-02  5.0000E-01  9.9634E-01
 4.5000E-01  5.0000E-02  5.0000E-01  9.9600E-01
 5.5000E-01  5.0000E-02  5.0000E-01  9.9563E-01
 6.5000E-01  5.0000E-02  5.0000E-01  9.9524E-01
 7.5000E-01  5.0000E-02  5.0000E-01  9.9482E-01
 8.5000E-01  5.0000E-02  5.0000E-01  9.9438E-01
 9.5000E-01  5.0000E-02  5.0000E-01  9.9392E-01
...
 1.1650E+01  5.0000E-02  5.0000E-01  5.8604E-01
 1.1750E+01  5.0000E-02  5.0000E-01  5.8337E-01
 1.1850E+01  5.0000E-02  5.0000E-01  5.8129E-01
 1.1950E+01  5.0000E-02  5.0000E-01  5.7985E-01
 1.2050E+01  5.0000E-02  5.0000E-01  5.7913E-01

```

