Finite Element Method for Subsurface Hydrology
Using a Mixed Explicit-Implicit Scheme

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The mixed explicit-implicit Galerkin finite element method developed previously by the authors is shown to be ideally suited for a wide class of problems arising in subsurface hydrology. These problems include confined saturated flow, unconfined flow under free surface conditions subject to the Dupuit assumption, flow in aquifers which are partly confined and partly unconfined, axisymmetric flow to a well with storage, and flow in saturated-unsaturated soils. A single computer program, entitled Flump, can now handle all of these problems. The mixed explicit-implicit solution strategy employed in the program insures a high level of accuracy and computation efficiency in most cases. It eliminates many of the difficulties that groundwater hydrologists have been encountering in trying to simulate extensive aquifer systems by finite elements. Some of the outstanding features of this solution strategy include an automatic control of time step size, reclassification of nodes from explicit to implicit during execution, automatic adjustment of the implicit time-weighting factor, and the treatment of boundary conditions and source terms as arbitrary functions of time of the state of the system. Five examples are presented to demonstrate the versatility and power of this new approach. A purely physical derivation of the finite element equations which does not rely on the Galerkin formalism is also included in one of the appendices.

INTRODUCTION

A large class of problems in subsurface hydrology are governed by linear or quasi-linear parabolic partial differential equations of the type

\[ \nabla \cdot (K \nabla h) - q = C(\partial h/\partial t) \]  

where \( h \) is the dependent variable, \( K \) is the second-rank symmetric positive definite tensor, \( q \) is the source term, and \( C \) is a fluid mass capacity coefficient. For example, in the case of confined saturated flow, \( h \) represents the hydraulic head, \( K \) the hydraulic conductivity (or transmissivity), and \( C \) the specific storage (or storage coefficient). In the case of flow in an unconfined aquifer, \( h \) represents the hydraulic head, \( K \) the transmissivity (i.e., the product of hydraulic conductivity and saturated thickness), and \( C \) the specific yield. The resulting expression is known as the Boussinesq equation. If \( h \) is replaced by the volumetric water content \( \psi \), \( K \) is replaced by the diffusion coefficient \( D(\psi) \), and \( C \) is set equal to unity, then (1) becomes the well-known diffusivity equation for unsaturated flow in a homogeneous soil. Richards's equation for saturated-unsaturated flow is obtained from (1) simply by writing \( h = \psi + Z \), where \( \psi \) is pressure head and \( Z \) is elevation above an arbitrary datum, by allowing \( K \) to vary in a prescribed fashion with \( \psi \) while at the same time treating \( \psi \) as a prescribed function of \( \psi \), and by setting \( C = \partial^2 \psi/\partial \psi^2 \). In the case of flow in a confined saturated medium the coefficients \( K \) and \( C \) are prescribed at each point in the medium independently of \( h \), and therefore (1) is generally linear in the dependent variable. On the other hand, Boussinesq, diffusion, and Richards's equations are nonlinear because at least one of the coefficients \( K \) and \( C \) varies with \( h \).

When (1) is discretized by the Galerkin finite element method, the result is a system of first-order linear or quasi-linear differential equations of the form [cf. Neuman, 1975b]

\[ A h + D h = Q \]  

where \( A \) is the conductance (or stiffness) matrix \( D \) is the capacity matrix, \( h \) is the dependent variable vector (e.g., hydraulic head or water content), and \( Q \) is a vector representing sources or sinks. The term \( D h \) represents time derivatives of \( h \).

In most finite element schemes [e.g., Javandel and Witherspoon, 1968; Pinder and Frind, 1972] the capacity matrix \( D \) includes nonzero off-diagonal terms (i.e., it is non-diagonal), and there is growing evidence in the literature suggesting that this may lead to conceptual as well as numerical difficulties. For example, a recent analysis by Narasimhan [1978] indicates that a nondiagonal \( D \) matrix may upset the maintenance of local mass balance in the vicinity of each node, although overall balance over the entire finite element grid may still be preserved. Fujii [1973] showed that when \( D \) is nondiagonal, the time increment \( \Delta t \) in the numerical scheme must not be too large or too small if the maximum principle is to be preserved (the maximum principle states that in the absence of internal sources \( q \) the maximum value of \( h \) must occur at the initial time or at the boundary). When \( D \) is diagonal, the permissible value of \( \Delta t \) is larger, and there is no lower limit. This may perhaps explain why, as our experience indicates, (2) sometimes yields physically unrealistic values of \( h \) when there is a sudden and drastic change in \( Q \) (e.g., the rate of pumpage from certain wells in an aquifer) and why this can be remedied by diagonalizing the \( D \) matrix, as has been done by Wilson [1968],
Emery and Carson [1971], Neuman [1973, 1975], Neuman et al. [1977], and others. In fact, Neuman [1975] found by experience that in solving the highly nonlinear problem of saturated-unsaturated groundwater flow with the aid of finite elements the rate of convergence can be improved dramatically by diagonalizing the D matrix. Neuman used triangular elements, but the same happens when one uses isoparametric elements (G. Segol, personal communication, 1976). Similar conclusions have also been reached by Mercer and Faust [1977] in connection with two-phase immiscible flow in porous media.

Neuman and Narasimhan [1977] described an alternative form for (2) with a diagonal capacity matrix which, they showed, has many advantages over the traditional approach. In this new approach the system of differential equations (2) is discretized in time by finite differences, a procedure which enables one to treat the equations either explicitly, implicitly, or by an optimum combination of both schemes. The implicit equations are solved by a point iterative technique rather than by a direct method such as Gaussian elimination or Cholesky decomposition. This eliminates one of the major difficulties that groundwater hydrologists have faced in trying to utilize extensive aquifers. Neuman and Narasimhan [1977] derived local stability and convergence criteria for the explicit-implicit scheme, as well as convergence criteria for the proposed point iterative technique. Their theory assumes that the matrix A is diagonally dominant, and it is shown that one can always construct a finite element mesh satisfying this condition.

Narasimhan et al. [1977] have incorporated these new ideas into a powerful and versatile computer program entitled Flump. They applied it to various linear problems and concluded that the mixed explicit-implicit approach is highly efficient for problems that might otherwise involve matrices with large bandwidths (as they do in the case of extensive aquifers), problems in which the boundary conditions or source terms vary often and rapidly with time, as they do in the case of variable pumpage rates, and problems characterized by a significant spatial variability of element sizes and material properties. They showed that the mixed scheme is capable of yielding highly accurate results to such linear problems and that it often requires a lesser amount of computer time than other explicit or iterative implicit methods. Neuman et al. [1977] applied Flump to various quasi-linear problems involving unconfined flow under free surface and saturated-unsaturated conditions. Their results indicated that the mixed approach is especially well suited for quasi-linear problems in which the coefficients, the source terms, and the boundary conditions vary with the dependent variable.

The purpose of this paper is to review briefly the theory of the mixed explicit-implicit iterative finite element method and to demonstrate its flexibility and reliability in dealing with a variety of problems arising in subsurface hydrology. The power of the new method is demonstrated by five examples, including the buildup of a groundwater ridge, infiltration into an unsaturated soil, drainage of a saturated-unsaturated system, radial flow to a well with well bore storage, and areal flow in an extensive stream-aquifer system. All of these examples were computed with the aid of a single computer program, Flump. A purely physical derivation of the finite element equations which does not rely on the Galerkin formalism is also presented in Appendix B.

**Explicit-Implicit Formulation**

In order to discretize (1) in space we adopt a network of triangular elements for plane flow and a network of concentric rings of constant triangular cross section for axisymmetric problems (Figure 1). Each triangle is further subdivided along its medians into three subdomains of equal areas, each of which is associated with the nearest nodal point. The collection of all such subdomains associated with any given node n (see the shaded area in Figure 1) is referred to as the exclusive subregion $R_n$ of n.

In each individual element e the dependent variable is described approximately by

$$h(x, t) = \sum_n h_n(t) \xi_e(x)$$

where $h_n$ are the values of $h$ at the nodal coordinates $x_n$ and $\xi_e(x)$ are linear shape functions of the space coordinates defined in Appendix A and satisfying $\xi_e(x_n) = \delta_{en}$, $\delta_{en}$ being Kronecker delta. With this we are ready to apply the Galerkin method to (1). However, since this method is applicable only at a given instant of time, the time derivative $\partial h/\partial t$ must be determined independently of the Galerkin orthogonalization process, as was pointed out earlier by Neuman [1975a]. Thus instead of replacing $h$ in the time derivative by (3), as is usually done in the conventional finite element approach, one is justified in defining the nodal values of $\partial h/\partial t$ as averages over the exclusive subregions $R_n$ associated with each node. This leads to a system of first-order linear or quasi-linear differential equations of the form given in (2) but with a diagonal capacity matrix. The individual terms of the matrices A and D are given by [Neuman, 1975a]

$$A_{nm} = \sum_e \frac{\alpha}{4\Delta} [K_{1e} b_e b_m + K_{2e} b_e c_m + b_m c_e + K_{3e} c_e c_m]$$

$$D_{nm} = \delta_{nm} \sum_e \frac{\alpha \Delta}{3} C$$

**Fig. 1. Exclusive subdomain $R_n$ associated with node n.**

A purely physical derivation of the finite element equations which does not rely on the Galerkin formalism is also presented in Appendix B.
where $A$ is the area of a triangle, $\alpha = 1$ for plane flow and $\alpha = 2\pi r$ for axisymmetric flow ($r$ being the average radius to the centroid of a triangle), $a$ and $c$ are geometric coefficients defined in Appendix A, and the summation sign applies to all elements adjacent to nodal point $n$.

The finite element differential equations (2), $A$ and $D$ being defined as they are in (4) and (5), can also be derived from purely physical considerations (that is, without using the Galerkin formalism). Such a derivation is given in Appendix B [see also Narasimhan, 1978]. It is clear from the physical derivations that $-A_{nm}$ represents the rate of fluid transfer into $R_n$, the exclusive subregion of node $n$, due to a unit difference in head between nodes $m$ and $n$. On the other hand, $D_{nn}$ is the fluid mass capacity [Narasimhan and Witherspoon, 1977] of $R_n$, denoting the ability of $R_n$ to take fluid into storage because of a unit change in head. The fact that $D_{nn}$ is the diagonal matrix simply means that the amount of fluid in $R_n$ is a function only of $h_n$.

If we replace the time derivatives in (2) by finite differences and introduce a weighting factor $\theta$, we obtain a system of simultaneous linear or quasi-linear algebraic equations of the form

$$A[\theta h^{t+1} + (1 - \theta)h^t] + D^{t+1}_n = Q$$

(6)

where $0 \leq \theta \leq 1$, $\Delta t$ is the time increment, and $k$ indicates the number of time steps. Defining a new term

$$\lambda_{nm} = A_{nm} \Delta t / D_{nn}$$

(7)

and recognizing from (A6) in Appendix A that

$$\lambda_{nn} = -\sum_{m=1}^{N} \lambda_{nm}$$

(8)

we can rewrite (6) as

$$h^{t+1} - h^t = \theta \sum_{m=1}^{N} \lambda_{nm} (h^{t+1}_m - h^t_m) + (1 - \theta)$$

$$\sum_{m=1}^{N} \lambda_{nm} (h^t_m - h^t_n) + \frac{Q \Delta t}{D_{nn}}$$

(9)

where $N$ is the total number of nodes and the summation is taken over all nodes $m$ in the immediate neighborhood of $N$. When $\theta = 0$, all the values of $h^{t+1}_n$ can be calculated explicitly from the system of equations (9), which now corresponds to a forward difference scheme in time. When $\theta = 1$, the result is a fully implicit backward difference scheme, whereas $\theta = 1/2$ corresponds to a time-centered or Crank-Nicolson scheme. It is important to recognize that conventional finite element schemes with a nondiagonal $D$ matrix are inherently implicit [Narasimhan and Witherspoon, 1976] in that $h^{t+1}_n$ can never be computed explicitly from the time discrete form of (2). Furthermore, our formulation in (9) does not include any diagonal terms of the matrix $A$, a feature which leads to a considerable saving in computer time and storage, especially in dealing with nonlinear problems in which the matrix must be recomputed at each time step.

**Stability and Convergence**

For the purpose of defining stability and convergence criteria, let us assume that $\lambda_{nm} \leq 0$ for all $m \neq n$. Since $\lambda_{nn} = -\sum_{m=1}^{N} \lambda_{nm}$, this implies that $\lambda_{nn}$ is diagonally dominant, i.e.,

$$\lambda_{nn} = \sum_{m=1}^{N} |\lambda_{nm}|$$

(10)

Later in the text we will show that one can always construct a finite element mesh which satisfies this requirement at all nodes. Neuman and Narasimhan [1977] showed that when (10) is satisfied, the finite element scheme in (9) is unconditionally stable at node $n$ for all values of $\theta$ which are not less than 0.5. When $\theta < 0.5$, stability is conditioned upon the criteria

$$\lambda_{nn} \leq 1/(1 - 2\theta)$$

(11a)

or

$$\Delta t \leq D_{nn}/(1 - 2\theta)A_{nn}$$

(11b)

For example, the explicit version of (9) is stable at any node at which the ratio between capacity and conductance is large enough that $\Delta t \leq D_{nn}/A_{nn}$ for any given $\Delta t$. Conversely, the explicit scheme can be made stable at all nodes by choosing a sufficiently small $\Delta t$.

The stability criterion $\Delta t \leq D_{nn}/A_{nn}$ for the explicit scheme has a simple physical interpretation. According to Appendix B, $\Delta t/\beta_{nn}$ is the amount of fluid entering in the exclusive subregion of node $n$, $R_n$, when heads at all adjacent nodes, $h_m$, exceed $h_n$ by unity. On the other hand, $D_{nn}$ is the capacity of the subregion to absorb fluid when $h_n$ changes by one unit. Thus the above stability criterion merely states that the amount of fluid entering $R_n$ must not exceed the capacity of $R_n$ to absorb fluid. A value of $\Delta t$ in excess of what is prescribed by the stability criterion would imply that $h_n$ must change by more than unity, which is contrary to the maximum principle (recall that in the explicit scheme the values of $h_n$ remain fixed during a time step). Note that the identification of the stability criterion based on physical considerations focuses attention on the invariant nature of stability.

We mention that (11a) and (11b) include the term $(1 - 2\theta)$ in the denominator; Fujii [1973] derived a global criterion for the maximum principle associated with (9) which includes the term $(1 - \theta)$ in the denominator. This implies that our local stability criterion is less restrictive than the global criterion of Fujii.

Neuman and Narasimhan [1977] have also investigated an important question: Under what conditions does the numerical scheme in (9) converge to the exact solution of (1) as the mesh is made finer and finer in space and time? Many conventional finite element formulations are known to converge in the mean; however, they seldom guarantee convergence at each point, as is the case with many finite difference schemes. Neuman and Narasimhan [1977] were able to demonstrate that in the linear case the explicit-implicit scheme in (9) converges to the exact solution of (1) at each node at which stability and certain local symmetry conditions are satisfied.

**Point Iterative Method**

The particular iterative scheme adopted in this work is known as the point acceleration method and was developed originally by Evans et al. [1954]. As will be seen below, it differs from the more familiar point successive overrelaxation technique [Gambolati, 1975]; the latter can be viewed as an extension of the Gauss-Seidel method, whereas the former is more closely related to the point Jacobi method. The acceleration method is readily amenable to an analysis of pointwise convergence and is therefore well suited for the mixed explicit-implicit scheme proposed in this work. This by no means implies that one would not be able to achieve an equally fast, perhaps even faster, rate of convergence with other iterative
techniques, such as the point successive overrelaxation method.

The system of equations (9) can be rewritten as

$$\Delta h_n = \sum_{m \neq n} \lambda_{nm}[h_n^* - h_m^*] + Q_n \Delta t / D_n$$

$$n = 1, 2, \ldots N$$ (12)

where $\lambda_{nm}(\Delta h_n - \Delta h_m)$ denotes the explicit part of the potential change, while $\lambda_{nm}(h_n^* - h_m^*)$ denotes the implicit part. The acceleration method consists of introducing the following substitutions into (12):

$$\Delta h_n \text{ (left side)} - \Delta h_n^{j+1}$$
$$\Delta h_n \text{ (right side)} - (1 + g)\Delta h_n^{j+1} - g\Delta h_n^j$$
$$\Delta h_m \text{ (right side)} - \Delta h_m^j$$

where $j$ is the number of iterations and $g$ is the acceleration factor. When we solve for $\Delta h_n^{j+1}$, the acceleration algorithm takes the form

$$\Delta h_n^{j+1} = \left[ \sum_{m \neq n} \lambda_{nm}(h_n^* - h_m^*) - \theta \sum_{m, n} \lambda_{nm}(\Delta h_n^* + \Delta h_m^*) \right]$$
$$+ Q_n \Delta t / D_n \left[ 1 - \theta(1 + g) \sum_{m \neq n} \lambda_{nm} \right]^{-1}$$ (13)

The same algorithm can also be written in terms of the residuals $\epsilon$ as

$$\epsilon_n^{j+1} = \left[ \theta \sum_{m \neq n} \lambda_{nm}(g^* + \epsilon_m^*) \right]$$
$$- (1 - \theta(1 + g)) \sum_{m \neq n} \lambda_{nm} \epsilon_n^{j+1}$$ (14)

where $\epsilon_n^{j+1} = \Delta h_n^{j+1} - \Delta h_n^j$.

The reader may easily recognize the fact that when $g$ is set equal to zero, (13) and (14) reduce to the point Jacobi algorithm. As a matter of contrast, when the relaxation factor in the point successive overrelaxation algorithm is set equal to unity, it reduces to the Gauss-Seidel algorithm.

Neuman and Narasimhan [1977] showed that for linear problems the acceleration method converges at any node at which $\lambda_{nm}$ is locally diagonally dominant (we mentioned earlier that it is always possible to construct a mesh which satisfies this requirement), provided that $g \geq 0$. For optimum results, $g$ should not exceed 1 and should usually be less than 0.5. Experience indicates that near-zero values of $g$ may cause difficulties and the optimum value tends to be in the vicinity of 0.2. The optimum value of $g$ appears to be more stable (i.e., to have a narrower range of values) than the optimum value of the relaxation factor $\omega$ in the point successive overrelaxation method.

**Local Diagonal Dominance**

The purpose of this section is to show that one can always construct a finite element network of triangles so as to guarantee that $\lambda_{nm}$ will be diagonally dominant. To do so for isotropic domains, we will follow an approach suggested earlier by Gambolati [1973]. However, we will follow a simpler and more practical line of reasoning for anisotropic domains.

Consider a triangular element in a plane described by the coordinate system $x$, $y$ as shown in Figure 2. Then it is easy to show that

$$b_1 = -a_1 \sin \gamma \quad c_1 = a_1 \cos \gamma$$
$$b_2 = a_2 \sin (\gamma - \beta_3) \quad c_2 = -a_2 \cos (\gamma - \beta_3)$$
$$b_3 = a_3 \sin (\gamma + \beta_3) \quad c_3 = -a_3 \cos (\gamma + \beta_3)$$ (15)

where $b$ and $c$ are geometric coefficients defined in Appendix A and all other terms are defined in Figure 2. If $A'$ is the contribution of this triangle to the global matrix $A$, then (4) and (15) imply that in the isotropic domain,

$$A' = \frac{K}{4\Delta} \begin{bmatrix} a_1^2 & -a_1a_2 \cos \beta_1 & -a_1a_3 \cos \beta_1 \\ -a_1a_2 \cos \beta_1 & a_2^2 & -a_2a_3 \cos \beta_1 \\ -a_1a_3 \cos \beta_1 & -a_2a_3 \cos \beta_1 & a_3^2 \end{bmatrix}$$ (16)

where $\Delta$ is the area of the triangle and $K$ is the scalar equivalent of $K$. From Appendix A we know that

$$A_{nm}^x = -\sum_{m \neq n} A_{nm}$$ (17)

indicating that $A^x$ is diagonally dominant if and only if all the off-diagonal terms in (16) are negative. This means that none of the angles $\beta$ may exceed $90^\circ$, and therefore a sufficient (though not always necessary) condition for the global matrix $A$ network consists entirely of right- and/or acute-angled triangles. The same, of course, holds for $\lambda_{nm}$.

Next, consider an anisotropic domain with principal hydraulic conductivities $K_x$ and $K_y$ oriented parallel to the $x$ and $y$ coordinates, respectively. Then according to (4),

$$A_{nm}^x = \frac{1}{4\Delta} (K_x b_n b_m + K_y c_n c_m)$$ (18)

We can now define a new set of coordinates $x' = x/(K_x/K_y)^{1/2}$ and $y' = y$, so that in the transformed domain of $x'$ and $y'$, (1) will take form

$$(K_x, K_y)^{1/2} \cdot \Delta h = (K_x, K_y)^{1/2} \cdot C(\partial h/\partial t) + q$$ (19)

In other words, the original anisotropic domain in the $x$, $y$ plane has been transformed into an equivalent isotropic domain with conductivity $(K_x, K_y)^{1/2}$ in the $x'$, $y'$ plane merely by expanding or contracting one of the coordinate axes. If $b'$ and $c'$ are the equivalents of $b$ and $c$ in the transformed domain, then it is easy to verify that (18) can also be written as

$$A_{nm}^x = (K_x, K_y)^{1/2} \cdot 4\Delta (h_n^* b_m^* + c_n^* c_m^*)$$ (20)

Let $a$, $b$, and $\gamma$ in Figure 2 transform into $a'$, $b'$, and $\gamma'$ in the $x'$, $y'$ plane. Then it is immediately obvious from (15), (16), and (19) that...
\[ A^* = \frac{(K_1 K_2)^{1/3}}{4\Delta} \begin{bmatrix} a'_1\cos \beta_1' & -a'_1 a'_2 \cos \beta_1' & -a'_1 a'_3 \cos \beta_1' \\ -a'_1 a'_2 \cos \beta_1' & (a'_2)^2 & -a'_2 a'_3 \cos \beta_1' \\ -a'_1 a'_3 \cos \beta_1' & -a'_2 a'_3 \cos \beta_1' & (a'_3)^2 \end{bmatrix} \] (21)

Thus if we follow the same line of reasoning as we did before, it is evident that the global matrix \( A \) is locally diagonally dominant whenever the local network in the transformed domain consists entirely of right- and/or acute-angled triangles. Again, so is \( \lambda_{on} \).

It is easy to show that the same holds true when the principal conductivities \( K_1 \) and \( K_2 \) are not parallel to the \( x \) and \( y \) coordinates. For this purpose it is sufficient to recognize that the solutions of (1) and (9) at any given point in space are independent of the choice of coordinates. Thus \( \lambda_{om} \) must remain invariant under a rotation of coordinates, and therefore if it is diagonally dominant in a set of coordinates which is parallel to \( K_1 \) and \( K_2 \), it must also remain diagonally dominant in another set of coordinates oriented at an angle to the first one. One can therefore transform any anisotropic domain into an equivalent isotropic domain merely by expanding or contracting it parallel to \( K_1 \) by the amount \( (K_1/K_2)^{1/3} \). If all the triangles in the transformed domain are constructed without obtuse angles, \( \lambda_{om} \) will be diagonally dominant.

In a composite material consisting of several segments with different degrees and orientations of anisotropy, each segment must be transformed separately parallel to its own principal direction of conductivity. Here, in addition to ensuring that all the triangles in the transformed domain are free of obtuse angles, one must also make sure that corresponding nodal points at both sides of a material interface will coincide with each other when the meshes are transformed back into the original plane. This is usually not a very difficult task, as has been demonstrated by an example in the work of Narasimhan et al. [1977].

**Mixed Explicit-Implicit Solution Strategy**

The local nature of the stability criteria for (9), together with the use of a point iterative technique, suggests the interesting possibility of solving the finite element equations explicitly at some nodes and implicitly at some other nodes during a single time step. If \( \Delta t \) satisfies the stability conditions (11a) and (11b) for some node \( n \), then (9) can be solved explicitly for \( \Delta h_n \) at that node. At nodes which do not satisfy the stability criterion, \( \Delta h_n \) is determined iteratively by using the algorithms in (13) and (14). A final correction is then made to the \( \Delta h_n \) values calculated explicitly, when it is required, to conserve mass. We refer to this approach as a mixed explicit-implicit solution strategy.

The mixed strategy is very useful in dealing with meshes characterized by a significant spatial variability of element sizes and material properties. For example, if the region of interest consists of two materials having different conductivities and capacities, it may be possible to solve explicitly in one material and implicitly in the other. The mixed approach is also useful when there is a sudden temporal change in boundary conditions. In this case it is often desirable to use small \( \Delta t \) values for a short period of time until the system reaches a certain level of equilibrium; otherwise, there may be a loss of accuracy. The attractive possibility of using an explicit solution procedure during this period may lead to significant savings in computer time.

The idea of combining explicit and implicit calculations in a single time step was previously used in conjunction with an integrated finite difference scheme by Edwards [1972]. The procedure has been incorporated by Edwards into a powerful computer program, called Trump, which can handle multidimensional steady state and transient temperature distributions in complex, nonuniform, and isotropic systems, taking into account conduction, convection, and radiation. The program has also been applied by Edwards [1969] to Darcian fluid flow in porous media. The conduction aspects of Trump are based on a set of algebraic equations which have the same general form as (9). This made it possible for us to develop a computer program which combines the advantages of the finite element method (such as the ability to treat anisotropic regions with complex geometry) with the remarkable logic and facilities of Trump. The new program is called Flump, as a mnemonic for finite element and Trump.

In addition to the mixed explicit-implicit solution strategy the user of Flump has the option of using a fully explicit forward difference scheme (\( \theta = 0 \)), a time-centered Crank-Nicolson scheme (\( \theta = 0.5 \)), or a fully implicit backward difference scheme (\( \theta = 1.0 \)), throughout any part of the solution process. However, in practice these options are seldom used because Flump has the facility to adjust the weight factor \( \theta \) automatically during execution in a manner that ensures a high level of accuracy at each time step. This, as well as other special features of Flump, is described briefly below.

**Iterative Approach**

The iterative approach is based on (13) and (14). During a given time step \( \Delta t \) the algorithm is applied only to a selected number of nodes (called implicit nodes) which cannot be treated explicitly without endangering stability. For the first iteration (\( j = 0 \)) the initial guesses used are \( \Delta h_n^0 = \hat{h}_n \Delta t \) and \( \Delta h_n^0 = \hat{h}_n \Delta t \), where \( \hat{h}_n \) and \( \hat{h}_n \) are estimated time derivatives (a method for obtaining these derivatives is described later in the text). The values of \( \Delta h_n^1 \) are then calculated by using (13), and the first set of residuals is determined from the relation

\[ e_0 = \Delta h_n^1 - \hat{h}_n \Delta t. \]

This procedure is continued until convergence is achieved or until 80 iterations have been completed. In the latter case the calculations are repeated with half the original value of \( \Delta t \). Thus if we follow the same line of reasoning as we did before, the mixed strategy is very useful in dealing with meshes characterized by a significant spatial variability of element sizes and material properties. For example, if the region of interest consists of two materials having different conductivities and capacities, it may be possible to solve explicitly in one material and implicitly in the other. The mixed approach is also useful when there is a sudden temporal change in boundary conditions. In this case it is often desirable to use small \( \Delta t \) values for a short period of time until the system reaches a certain level of equilibrium; otherwise, there may be a loss of accuracy. The attractive possibility of using an explicit solution procedure during this period may lead to significant savings in computer time.

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Two convergence criteria must be satisfied simultaneously before terminating the iterations. The first criterion is

\[ \max_n |e_n| \leq 10^{-4} \Delta h_{max} \] (22)

where \( \Delta h_{max} \) is the desired maximum change in \( h \) at any node during a time step (as will be seen later, \( \Delta t \) is adjusted during execution to maintain the maximum change in \( h \) near the value of \( \Delta h_{max} \) and less than \( 2\Delta h_{max} \)). The second criterion is based on the net correction to the fluid content,

\[ \Delta H_{net} = \sum_n D_{an} e_n \]

and the fluid capacity,

\[ D_{net} = \sum_n D_{nn} \]
of all implicit nodes (the summation is taken over all implicit nodes). The iterative procedure is stopped when (22) is satisfied together with (23) below:

$$\Delta H_{\text{new}} < 10^{-4} \Delta h_{\text{old}} \Delta h_{\text{max}}$$

(23)

i.e., the net error in fluid content is less than $10^{-5}$ of the amount of fluid required to change $h$ at each implicit node by the amount $\Delta H_{\text{new}}$. Experiments by Edwards [1972] on a large number of sample problems using Trump indicate that the net cumulative error in the average value of $h$ tends to be no more than $0.01 \Delta h_{\text{des}}$ after several hundred time steps; the cumulative error at individual nodes does not usually exceed $0.1 \Delta h_{\text{des}}$ and is much less if some values of $h$ are fixed at the boundary of the system.

After having completed the iterative procedure for all implicit nodes, one must now correct the values of $h$ at all explicit nodes connected to implicit nodes, according to (see (12))

$$\Delta h_{\text{new}} \text{ corrected} = \Delta h_{\text{new}} \text{ explicit} + \theta \sum_{m,n} \lambda_{mn} (\Delta h_{\text{m}} - \Delta h_{\text{n}})$$

(24)

where the summation is taken only over implicit nodes. This correction is necessary for a correct material balance. Since internal fluxes between adjacent subregions are calculated simultaneously with the $h$ values, these fluxes must also be corrected in a similar manner.

**Automatic Control of Time Step Size**

The size of $\Delta t$ is controlled automatically by several factors such as the lower and upper limits specified by the user ($\Delta t_{\text{min}}$ and $\Delta t_{\text{max}}$, respectively), the desired maximum change in $h$ at any node during a time step ($\Delta h_{\text{max}}$), the smallest time step allowed at any explicit node by the stability criterion ($\Delta t_{\text{stab}}$), the average number of iterations required for convergence, and the desired interval between printed outputs.

The first time step is always $10^{-19}$ and is used primarily for checking the input data, establishing time derivatives, and determining $\Delta t_{\text{stab}}$ (the latter is recalculated whenever the conductance or capacitance matrices change). The maximum allowed time step $\Delta t_{\text{max}}$ is then set equal to $\frac{1}{3}$ of $\Delta t_{\text{stab}}$ or $\Delta t_{\text{high}}$, whichever is smallest (the use of $\frac{1}{3}$ of $\Delta t_{\text{stab}}$ instead of $\Delta t_{\text{high}}$ greatly increases the accuracy in coarse meshes). The minimum allowed time step $\Delta t_{\text{min}}$ is set equal to $\Delta t_{\text{stab}}$ or $10^{-20}$, whichever is greater. If $\Delta t_{\text{new}} > \Delta t_{\text{max}}$, the value of $\Delta t_{\text{new}}$ is reduced to slightly less than $\Delta t_{\text{max}}$ to prevent the input value from causing instabilities. The default value for $\Delta t_{\text{new}}$ is taken to be $\Delta t_{\text{max}}/100$.

During the subsequent calculations the size of $\Delta t$ is gradually adjusted to obtain a maximum change in $h$ close to $\Delta h_{\text{des}}$ and not exceeding $2 \Delta h_{\text{des}}$, to maintain the maximum change in any tabulated material property in nonlinear problems near 1% or less and not exceeding 2%, and to prevent the number of iterations from averaging more than 40, the maximum allowed being 80. The technique for doing this has been designed by Edwards [1972] to cause a rapid decrease in $\Delta t$ when the above limits are exceeded and a more gradual increase in $\Delta t$ when changes are relatively slow. For this purpose, let $\theta$ be either the largest percentage change property or $\Delta h_{\text{des}}$ of the number of iterations required for convergence, whichever is larger. We then calculate the ratio

$$R = \Delta h_{\text{des}} / \max_n |\Delta h_n|, \delta \Delta h_{\text{des}}$$

(25)

and if $R \leq 0.5$ and $\Delta t \geq 1.0 \Delta t_{\text{min}}$, the entire computation for the recent time step is repeated with a modified value of $\Delta t$. If all the nodes in the mesh are set to be implicit, $\Delta t$ is reduced by a factor of 100 for the first time step to start the calculation out smoothly. If $R \leq 1.0$, the new time step is calculated according to

$$\Delta t_{\text{new}} = R \Delta t_{\text{old}}$$

(26)

whereas if $R > 1$, the formula is

$$\Delta t_{\text{new}} = 0.5(1 + R) \Delta t_{\text{old}}$$

(27)

In both cases the adjustments are subject to the constraints $\Delta t_{\text{min}} \leq \Delta t_{\text{new}} \leq \Delta t_{\text{max}}$ and $0.5 \Delta t_{\text{old}} \leq \Delta t_{\text{new}} \leq 2.0 \Delta t_{\text{old}}$. An additional adjustment in the size of $\Delta t$ may be required in order to meet a desired interval between printed outputs.

**Reclassification of Nodes from Explicit to Implicit**

If the recent time step was equal to $\Delta t_{\text{new}}$ and less than $\Delta t_{\text{stab}}$, the stability limits of all explicit nodes are tested. All explicit nodes with stability limits equal to or less than 1.8 $\Delta t_{\text{max}}$ are then reclassified as implicit nodes. Since $\Delta t_{\text{stab}}$ is $\Delta t_{\text{stab}}$, the reclassification affects all explicit nodes having stability limits from 1.0 to 1.2 times $\Delta t_{\text{stab}}$. This range was chosen empirically by Edwards [1972] in an effort to minimize the required computation time for a large group of test problems using Trump.

The rate at which the nodes are reclassified from explicit to implicit depends on the input parameter $\Delta h_{\text{des}}$. The larger this parameter, the faster the increase in the size of $\Delta t$, and therefore the earlier the stability limits of most nodes are reached.

**Estimation of Time Derivatives**

The initial guess of $h$ for the iterative procedure requires a preliminary estimate of the time derivatives $h$. In nonlinear problems the time derivatives are also used to estimate the average values of $h$ to be used in evaluating $h$-dependent parameters. Instead of saving $h_n$ values from several preceding time steps, which could be used to calculate more accurate time derivatives, a simpler method is used which requires less memory space and machine time and is sufficiently accurate for most problems.

The time derivatives for any time step $t_{n+1} - t_n$ are estimated from the maximum rates of change in $h$ occurring during the two preceding time steps, $\Delta t_n = t_{n+1} - t_n$ and $\Delta t_{n-1} = t_{n} - t_{n-1}$. For this purpose, let us define the two ratios

$$R_k = \frac{\max_n \left| (h_n^k - h_n^{k-1}) / \Delta t_k \right|}{\max_n \left| (h_n^{k-1} - h_n^{k-2}) / \Delta t_{k-1} \right|}$$

(28)

$$R_t = (\Delta t_k + \Delta t_{k+1}) / (\Delta t_{k-1} + \Delta t_k)$$

(29)

where $0.5 \leq R_t \leq 2.0$ because, as will be recalled, $\Delta t$ is not allowed to vary from one time step to another by more than a factor of 2. If $R_t < 1.0$, the maximum rate of change in $h$ is decreasing with time, and the estimate is based on the assumption that all $h$ values are approaching equilibrium exponentially according to the formula $h(t) = h(0)e^{-\alpha t}$. Since $e^{-\alpha t} = [h(t)/h(0)]^{1/\alpha}$, it follows that

$$R_{k+1}^{\text{est}} = R_k R_t$$

(30)
where $R_{n+1}^{est} \leq 1$ is the estimated value of $R_{n+1}$. If $R > 1$, the maximum rate of change in $h$ is increasing, and the estimate is based on the assumption that all $h$ values vary quadratically according to the formula $h(t) = h(0) + h(0)\Delta t + a\Delta t^2$. Since $h(t)/h(0) = 1 - 2a/\Delta h(0)$, it follows that

$$R_{n+1}^{est} = 1 + (1 - R^{-1})R$$

(31)

where $1 \leq R_{n+1}^{est} < 3$ because of the limits imposed on $R$. Equation (31) gives a more conservative estimate of the maximum rate of change in $h$ than (30) does. The estimated time derivative at each node is calculated as the product of the actual derivative during the previous time step and $R_{n+1}^{est}$:

$$h_n = R_{n+1}^{est}[(h_n^k - h_n^{k-1})/\Delta t_k]$$

(32)

Numerical experiments with Trump led Edwards [1972] to conclude that it is advisable to keep $R_{n+1}^{est} = 1.0$ during the first two time steps (1) at the beginning of each problem, (2) after repeating a time step with a modified $\Delta t$, and (3) after a node has been reclassified from explicit to implicit. Edwards further concluded that the time derivatives should be set equal to zero or a very small number during the initial time step $(\Delta t = 10^{-14})$ as well as when they change sign. It was also found that more accurate results can be obtained for implicit nodes having stability limits smaller than $\Delta t$ by estimating their time derivatives during the first two time steps according to

$$h_n = (h_n^k - h_n^{k-1})/\Delta t_k \left(1 - \sum_{m \neq n} \lambda_{nm}\right)$$

(33)

where the values of $\lambda_{nm}$ correspond to the time step just completed, $\Delta t_k$.

**Estimation of Implicit Weight Factor**

In most implicit procedures it is customary to employ either a time-centered scheme with $\theta = 0.5$ or a backward difference scheme with $\theta = 1.0$. In Flump, $\theta$ is allowed to be zero for explicit nodes or to vary between 0.57 and 1.0 for implicit nodes. Experience indicates that small oscillations caused by rapid changes in boundary conditions or variable parameters tend to persist when $\theta$ is close to 0.5. The lower limit of 0.57 was chosen empirically by Edwards [1972] to eliminate persistent oscillations and to optimize the stability and accuracy of a large number of test problems using Trump.

The average value of $h$ at any node during a time step is calculated in the program as

$$\bar{h}_n = h_n^k + \theta(h_n^{k+1} - h_n^k)$$

Let us assume that $h$ approaches equilibrium exponentially. Then for small time steps and for time steps during which the slope of $h$ remains nearly constant the correct average value is obtained with $\theta = 0.5$. On the other hand, for large time steps near equilibrium the correct average value is obtained with $\theta = 1.0$. Thus $\theta$ should be in the vicinity of 0.57 during the period when rapid changes in $h$ take place and should gradually shift toward 1.0 as equilibrium is approached; otherwise, there may be a loss of accuracy. One way to accomplish this is by using the empirical formula

$$\theta = \max [0.57, \max (1.0, R_{n+1}^{est})/(1.0 + R_{n+1}^{est})]$$

(34)

Experiments conducted by Edwards [1972] on a large number of problems using Trump have shown that the approach to equilibrium is usually too rapid when a forward difference scheme or a time-centered scheme is used and much more accurate results can be obtained with a variable $\theta$. His experiments also showed that $\theta$ should be set equal to 1.0 during the initial time step $(\Delta t = 10^{-4})$ as well as during any time step following a rejected time step. This enables nodes with small stability limits to reach equilibrium with their neighbors when there is a rapid change in a boundary condition or a variable parameter without overshoot which may lead to damped oscillations.

As was mentioned earlier, the computer program also provides an option to fix the value of $\theta$ at 0, 0.5, or 1.0 for the entire period of computation, corresponding to explicit forward difference, time-centered, or backward difference schemes, respectively. However, this tends to reduce accuracy and increase computer time and is therefore not advisable. The purpose of including these options is to allow the calculational results and machine time to be compared with those of other methods using a fixed value of $\theta$.

**Additional Features of Flump**

The printed output of Flump provides information on the nodal values of $h$ and $\Delta h$ and on the estimated value of $h$ at discrete time intervals specified by the user. Additional information includes the amount of fluid contained in the exclusive subdomain of each node, change in the amount of fluid in each subdomain during $\Delta t$ as well as from the start of the problem, total fluid content in the system, flux across the boundary of the system, and net flux into or out of the exclusive subdomain of each node. This information makes it possible to maintain a continuous check on material balance in the subdomain of each node as well as in the system as a whole.

The program also includes a built-in safety feature to warn the user about nodes at which the matrix $\lambda_{nn}$ is not diagonally dominant. If the degree of deviation from local diagonal dominance is significant, there are risks that the solution may be locally unstable (if the node is explicit) and inaccurate and that convergence will be relatively slow. The problem can always be remedied by locally redesigning the finite element mesh according to the guidelines given earlier in the text. Since the numbering of nodes and elements is completely arbitrary (as opposed to direct methods, such as Gaussian elimination or Cholesky decomposition, in which numbering has an effect on the bandwidth), local modifications of the mesh can be easily introduced merely by changing a few cards in the data deck.

![Fig. 4. Finite element mesh used to simulate Marino's [1967] experiment.](image-url)
TABLE 1. D-θ Relationship for Adsorption Problem

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<tr>
<th>θ</th>
<th>D, cm²/s</th>
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<td>0.2376</td>
<td>0</td>
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<tr>
<td>0.2440</td>
<td>1.59 × 10⁻⁴</td>
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APPLICATION TO SUBSURFACE HYDROLOGY PROBLEMS

The nature of our solution process makes it ideally suited for problems encountered under confined, unconfined, and saturated-unsaturated subsurface flow conditions. In the program, such problems are treated simply by specifying selected values of nodal conductivities, capacities, and sources or sinks as tabulated functions either of time or of the dependent variable h. The boundary conditions can also be controlled in a similar manner by tabulating them as functions of time or h.

For saturated-unsaturated flow problems, one has the option of tabulating the variables not as a function of total hydraulic head h but as a function of pressure head \( \psi = h - z \), where z is elevation. These features of Flump are demonstrated below by five examples involving various degrees of complexity and nonlinearity.

BUILDUP OF A GROUNDWATER RIDGE UNDER FREE SURFACE CONDITIONS

Our first example concerns the growth of a groundwater ridge under free surface conditions and the influence of uniform recharge from an infinite strip. This problem has been studied experimentally by Marino [1967] with the aid of a vertical viscous analogue (Hele-Shaw) model and has been treated analytically by Hantush [1967]. Figure 3 is a schematic diagram of the hydraulic conditions at the bottom and on both sides. It represents a fictitious porous medium having a hydraulic conductivity \( K = 0.42 \text{ cm/s} \) and a specific yield \( S_y = 1.00 \). Initially, the water table in the model was at an elevation of 11.25 cm. At time \( t = 0 \) a uniform recharge rate of 0.056 cm/s was inducted over a segment of the model extending from \( x = 0 \) to \( x = 23.8 \text{ cm} \), and this situation was maintained for a prolonged period. This caused the water table to rise at the rate indicated by the solid curves in Figure 3.

If one assumes that flow underneath the water table is essentially horizontal and that the recharging water reaches the water table instantaneously at \( t = 0 \), the flow in the model is governed by the one-dimensional Boussinesq equation

\[
\frac{\partial}{\partial x} \left( T(h) \frac{\partial h}{\partial x} \right) - q = S_y \frac{\partial h}{\partial t} \tag{35}
\]

Here, \( h \) is the height of the water table above the bottom of the model, \( q \) is the specific rate of recharge, and \( T(h) = Kh \) is the transmissivity. An analytical solution for a linearized version of this problem has been published by Hantush [1967], and some of his results are shown by triangles in Figure 3.

To solve (35) with Flump, we used a finite element mesh of quadrilaterals as shown in Figure 4. In the program each quadrilateral is automatically divided into two triangles, as is indicated by the diagonal dashed line in Figure 4. The numerical results for \( t = 60, 300, \) and \( 540 \) are shown by circles in Figure 5.

Fig. 5. Finite element mesh for adsorption problem.
Figure 3, and it is seen that they compare favorably with Hantush's analytical solution. The theoretical results also show a good fit with Marino's experimental data at early time. At later values of time the fit is good at distances exceeding 50 cm from the crest of the ridge but is somewhat less satisfactory near the crest. This may be due to the development of significant vertical hydraulic gradients under the recharging area; such gradients are not taken into account in the Boussinesq equation.

INfiltration INTO AN UnSATURATED SOIL COLUMN

The second example deals with the early adsorption stage of infiltration into a vertical unsaturated soil column. Initially, the volumetric water content in the soil is \( \theta = 0.2376 \). At time \( t = 0 \) the top of the column is brought to a state of saturation at \( \theta_{sat} = 0.4950 \), and this situation is maintained indefinitely. If one is interested only in the early adsorption stage of the resulting infiltration process when gravity is not important (or if the soil column is horizontal), then the flow in the soil is governed by the diffusion equation

\[
\frac{\partial \theta}{\partial z} D(\theta) \frac{\partial \theta}{\partial z} = \frac{\partial \theta}{\partial t} \tag{36}
\]

Here, \( z \) is depth below the soil surface, and \( D(\theta) \) is hydraulic diffusivity.

The highly nonlinear relationship between \( D \) and \( \theta \) is given in Table 1. Figure 5 shows the finite element mesh employed in solving this problem with Flump. The resulting water content profile at \( t = 10^6 \) s is shown in Figure 6. The numerical results are seen to be in excellent agreement with an analytical solution derived for this problem by Philip [1969] and with numerical results obtained by Van der Ploeg and Benecke [1974].

DRAINAGE FROM A SATURATED-UNSATURATED SYSTEM

The third example is devoted to two-dimensional flow in a vertical plane in which part of the porous medium is saturated and part is unsaturated. Vauclin [1975] and Vauclin et al. [1975] described a sophisticated and highly accurate drainage experiment in a sandbox shown schematically in Figure 7. Initially, the water table in the box was at an elevation of 143 cm. At time \( t = 0 \) the water level in one of the reservoirs attached to the sandbox was lowered to an elevation of 80 cm, and this situation was maintained indefinitely. The resulting hydraulic gradient caused the water table to drop gradually at

<table>
<thead>
<tr>
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the rate indicated by the solid curves in Figure 7. During the drainage process the sand remained saturated below the water table; the region above the water table had varying degrees of saturation.

Under this condition, flow in the model can be described by Richards's equation

$$\frac{\partial}{\partial x} \left( K(\theta) \frac{\partial \psi}{\partial x} \right) + \frac{\partial}{\partial z} \left( K(\theta) \frac{\partial \psi}{\partial z} \right) = C(\theta) \frac{\partial \psi}{\partial t}$$

Here, $x$ and $z$ are horizontal and vertical coordinates, respectively; $\psi$ is pressure head, defined as $\psi = h - z$, where $h$ is total hydraulic head; $K$ is hydraulic conductivity; and $C$ is specific moisture capacity, defined as $C = \frac{d\theta}{d\psi}$. The highly nonlinear relationships between $\psi$, $\theta$, $K$, and $C$ are given in Table 2.

Figure 8 shows the finite element network used in simulating the experiment with Flump. As was seen in Figure 7, the numerical results for the position of the water table are in excellent agreement with the corresponding experimental data. Figure 9 compares the cumulative depletion from storage as obtained by direct measurement, as computed with Flump, and as obtained by an alternating direction implicit iterative technique approach by Vauclin et al. [1975]; here again, the numerical results are in fairly good agreement with each other as well as with the experimental data.

**Confined Axisymmetric Flow to a Well with Well Bore Storage**

In groundwater hydrology, wells are often treated as line sources having negligible radii. However, when the aquifer transmissivity is small, the water stored in a well may have a considerable effect on discharge and on the hydraulic head in the aquifer [Papadopulos and Cooper, 1967; Narasimhan, 1968; Kipp, 1973; Neuman, 1973]. The problem of simulating well bore storage under saturated-unsaturated conditions by finite elements was previously discussed by Neuman [1975b] and Neuman et al. [1975]. In the present paper we consider an alternative approach based on the explicit-implicit formulation in (9).

Consider a partially penetrating well together with a superimposed finite element network as shown in Figure 10. The well has a radius $r_w$ and discharges at a rate $Q_w$ which may vary with time in a prescribed fashion. If the aquifer is confined, the head $h_w$ in the well is the same at all the nodal points $n = 1, 2, \ldots, 6$. Neglecting the storage capacity of the porous medium near the well compared with that of the well itself, we obtain

$$h_w^{k+1} - h_w^k = \frac{\Delta t}{D_w} \left( \sum \sum A_{nm} (h_w^{k+1} - h_m^{k+1}) \right) + (1 - \theta) \sum \sum A_{nm} (h_w^k - h_m^k) + Q_w$$

where $D_w$, the capacity of the well, is equal to $\pi r_w^2$; the first summation is taken over all nodes $n = 1, 2, \ldots, 6$ along the well bore.

Consider now an example of a fully penetrating well. The aquifer has a transmissivity of $9.29 \times 10^{-4}$ m$^2$/s and a storage...
coefficient of $10^{-8}$. The well is located at the center of an impermeable circular boundary having a radius of 400 m. It discharges at a constant rate of $1.42 \times 10^{-8}$ m$^3$/s. To solve this problem with the mixed explicit-implicit scheme, the flow region was discretized into 130 nodes and 100 elements. The radial spacing of the nodes increased logarithmically from the center.

Figure 11 shows a comparison between the numerical results and two analytical solutions, one developed by Papadopulos and Cooper [1967] for drawdown in a large-diameter well and the other developed by Muskat [1946] for an impermeable circular boundary. Their solutions are included for reference purposes. It is seen that the numerical results compare favorably with the analytical solution of Papadopulos and Cooper. The agreement with Muskat's solution is slightly less favorable, possibly because the analytical solution disregards well storage.

Recently, Jargon [1976] pointed out that well bore storage may also affect the drawdowns at observation wells located at a distance from the pumping well. Figure 12 shows the variation of drawdown with time at a distance of $r = 12.5$ m from the center of the pumping well.

**FLOW IN AN EXTENSIVE STREAM-AQUIFER SYSTEM**

As was mentioned earlier, our new approach eliminates many of the difficulties previously encountered in trying to simulate extensive aquifer systems by finite elements. To illustrate the capabilities of Flump in treating such systems, we have considered a hypothetical basin as shown in Figure 13. The basin is 20 km in length and 10 km in width, and it consists of an unconfined aquifer in contact with a stream and its tributary. The aquifer material consists of four different sedimentary units having distinct permeabilities. The thickness of the sediments varies in a steeple fashion from 20 m near the margins to 100 m below the streams. The transmissivity is assumed to vary linearly with the saturated thickness, as is shown in Figure 14. Water levels in the streams are taken to be constant with time, decreasing monotonically at a slow rate in the downstream direction. The hydraulic contact between the aquifer and the streams is such that water can flow either from the streams into the aquifer or vice versa, depending on the difference in head between the aquifer and the stream, as is
shown in Figure 15. The aquifer is tapped by 21 wells which are indicated by circles in Figure 13.

The system was discretized into 806 elements with 770 nodal points as illustrated in Figure 16. The first phase of the simulation consisted of determining the steady state distribution of hydraulic heads in the aquifer prior to pumpage, due to constant head along the streams and along segments of the aquifer boundary. The resulting piezometric surface is depicted in Figure 17. It can be seen that under steady state conditions, water flows from the stream into the aquifer east of the 106-m contour, whereas the reverse is true in the downstream portion of the basin west of this contour.

The second phase was to investigate what happens if all the 21 wells are turned on simultaneously, each pumping at a rate of 5451 m³/d (1000 gpm), while head along the streams and the boundaries remains the same as before. Figure 18 shows the distribution of hydraulic heads in the aquifer after 1 year of pumping. It is seen that steep cones of drawdown have developed around all major pumping centers. In the left central part of the basin the mutual interference of wells has caused a regional decline in water levels over a significant portion of the basin. Moreover, the pumping wells have captured much of the flow which was originally directed into the streams, so that in the downstream part of the basin the streams are now recharging the aquifer.

The problem treated in this example is nonlinear owing to the dependence of transmissivities and source terms along the streams on the hydraulic head in the aquifer. For this reason we found it necessary to restrict the size of each time step to approximately 1 day. This time step was less than the stability limit of all but 32 nodes in the mesh, so that most of the nodes were treated explicitly during the entire calculation. Each time step required an average of 1.45 s of computer time on the CDC 7600. We also ran the same problem with the Crank-Nicolson scheme, in which all the nodes are treated implicitly. Here, the results were similar to those of the mixed explicit-implicit scheme. However, each time step required on the average 2.89 s, i.e., twice as much computer time as the mixed scheme.

**APPENDIX A**

Consider the matrix $A$ as defined in (4) for plane flow (i.e., $\alpha = 1$). A typical term contributed by a single triangle such as that shown in Figure 2 has the form

$$A_{nm} = \frac{1}{4\Delta} [K_{xx}b_n b_m + K_{xy}(b_n c_m + b_m c_n) + K_{yy}c_n c_m]$$  (A1)

where

$$b_1 = y_s - y_s \quad c_1 = x_s - x_s$$
$$b_2 = y_s - y_1 \quad c_2 = x_1 - x_s$$
$$b_3 = y_1 - y_s \quad c_3 = x_s - x_1$$  (A2)

If $K_1$ and $K_2$ are the two principal conductivities and $\sigma$ is the angle between $K_1$ and the $x$ coordinate, then it can be shown that

$$K_{xx} = K_1 \cos^2 \sigma + K_2 \sin^2 \sigma$$
$$K_{yy} = K_1 \sin^2 \sigma + K_2 \cos^2 \sigma$$
$$K_{xy} = (K_1 - K_2) \sin \sigma \cos \sigma$$  (A3)

Substituting (A3) into (A1) and rearranging, we obtain

$$A_{nm} = \frac{1}{4\Delta} [K_{1}(b_n \cos \sigma + c_n \sin \sigma)(b_m \cos \sigma + c_m \sin \sigma) + K_2(b_n \sin \sigma - c_n \cos \sigma)(b_m \sin \sigma - c_m \cos \sigma)]$$  (A4)
indicating that \( A_{nn}^e \geq 0 \), i.e., the diagonal terms of \( A^e \) and \( A \) are always nonnegative. Furthermore, recognizing that \( b_1 + b_2 + b_3 = 0 \) and \( c_1 + c_2 + c_3 = 0 \), we find that (A4) can also be written for \( n = m = 1 \) as

\[
A_{11}^e = -(K_1/4S)(b_1 \cos \sigma + c_1 \sin \sigma)(b_1 + b_3)\cos \sigma + (c_2 + c_3) \sin \sigma - (K_2/4S)(b_1 \sin \sigma - c_1 \cos \sigma)(b_2 + b_3) 
\]

\[
- \cos \sigma - (c_2 + c_3) \sin \sigma = -A_{13}^e - A_{13}^e
\]

Thus in general we have

\[
A_{nn}^e = \sum_{m \neq n} A_{nm}^e, \quad A_{nn} = \sum_{m \neq n} A_{nm}
\]

where the summation is taken over all nodes other than \( n \).

The linear shape functions \( \xi_n^e(x) \) in (3) are given by

\[
\xi_n^e = \frac{1}{2A} (a_n x + b_n y + c_n) 
\]

where \( b_n \) and \( c_n \) are as they were in (A2) and

\[
a_1 = x_{13} - x_{23}
\]

\[
a_2 = x_{21} - x_{12}
\]

\[
a_3 = x_{12} - x_{23}
\]

Note that

\[
\frac{\partial \xi_n^e}{\partial x} = \frac{b_n}{2A} \quad \frac{\partial \xi_n^e}{\partial y} = \frac{c_n}{2A}
\]

**APPENDIX B**

To derive the finite element equations (2) on the basis of purely physical considerations (i.e., without the Galerkin formalism), let us confine our attention to the exclusive subregion \( R_n \) associated with a given node \( n \) (Figure 19). According to (3), \( h \) inside the particular triangle \( (n_{12}) \) is given by the linear interpolation formula

\[
h(x, t) = h_n(t)\xi_n^e(x) + h_1(t)\xi_1^e(x) + h_2(t)\xi_2^e(x)
\]

Since the functions \( \xi^e(x) \) are linear in the coordinates \( X \), (B1) implies that the gradient of \( h, \nabla h \), is constant inside the triangle. Furthermore, \( K \) is assumed to be constant in each element, and therefore the flux \( q = -K \nabla h \) must be uniform inside the triangle. Thus the sum of the flow rates across the line segments AG and BG must be equal to the flow rate across AB. In other words, the flow rate \( q_{12} \) into \( R_n \), contributed by the triangle \( (n_{12}) \), is given by

\[
q_{12} = K \nabla h \cdot n_{12} AB
\]

where \( n_{12} \) is a unit normal pointing out of the triangle as is shown in Figure 19.

Now \( \xi_n^e(x) \) is by definition unity at node \( n \) and zero at nodes 1 and 2. Therefore the derivative of \( \xi_n^e(x) \) parallel to the vector \( n_{12} \) is \(-1/L\), where \( L \) is the distance between node \( n \) and the line segment joining nodes 1 and 2. The gradient of \( \xi_n^e(x) \) is therefore given by

\[
\nabla \xi_n^e = -(n_{12}/L)
\]

Solving (B3) for \( n_{12} \) and substituting into (B2), we obtain

\[
q_{12} = -K \nabla h \cdot \nabla \xi_n^e AB L
\]

However, \( ABL \) is equal to the area of the triangle, \( \Delta \), and from (B1) we see that 
\( \nabla h = h_n \nabla \xi_n^e + h_1 \nabla \xi_1^e + h_2 \nabla \xi_2^e \). Therefore (B4) can be rewritten as

\[
q_{12} = -\Delta K(h_n \nabla \xi_n^e + h_1 \nabla \xi_1^e + h_2 \nabla \xi_2^e) \cdot \nabla \xi_n^e
\]
Similar expressions can be developed for \( q_{n3n} \), \( q_{n4n} \), \( q_{n45} \), and \( q_{a3} \). The sum of all of these terms gives the net flow rate into \( R_n \), \( q_n \), according to

\[
q_n = - \sum_m A_{nm}(h_m - h_n)
\]

where

\[
A_{nm} = \sum_e \Delta K \nabla \xi_n \cdot \nabla \xi_m
\]

and \( m \) takes on the values \( n, 1, 2, \ldots, 5 \).

Substituting (A9) into (B7) leads immediately to the final expression for \( A_{nm} \) as given in (4) for \( \alpha = 1 \). According to (A6), \( q_n \) can also be rewritten as

\[
q_n = \sum_{m \neq n} A_{nm}(h_m - h_n)
\]

This indicates that \(-A_{nm}\) represents the new flow rate into \( R_n \) due to unit difference in head between nodes \( m \) and \( n \). The same result was derived by Narasimhan [1975] by evaluating the Galerkin spatial integral over the triangular element \( e \).

The fluid capacitance of the segment \( nAGB \) of \( R_n \) is equal to

\[
C\Delta/3,
\]

where \( C \) is the capacitance of a unit area. Therefore the total capacitance \( D_{nn} \) of \( R_n \) is given by

\[
D_{nn} = \sum_e \frac{\Delta}{3} C
\]

which corresponds to the diagonal terms of (5) when \( \alpha = 1 \). Therefore a complete expression of inartential balance for \( R_n \) can be written as

\[
D_{nn}(dh_n/dt) = q_n + Q_n
\]

where \( Q_n \) is the net rate at which fluid is generated inside \( R_n \) by internal sources. When (B6) is substituted into (B10), we finally obtain

\[
\sum_{m \neq n} A_{nm}(h_m - h_n) + D_{nn}(dh_n/dt) = Q_n
\]

which is the \( n \)th equation in the set of finite element equations (2). A similar derivation applies to the axisymmetric case where \( \alpha \neq 1 \).

Acknowledgment. This work was partly supported by the U.S. Department of Energy.

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(Received October 5, 1977; accepted March 8, 1978.)