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A Two-Scale Numerical Subgrid Technique for Waterflood Simulations

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Summary
We present a two-scale numerical subgrid technique for simulating waterflooding. Local subgrid computations are combined with a coarse grid computation to provide a fine scale resolution of the solution. We use on the fine scale porosity, relative and absolute permeabilities, the location of wells, and capillary pressure curves. No explicit macroscopic coefficients or pseudo-functions result. The method is several times faster than solving the fine scale problem directly, generally more robust, and yet achieves good results as it requires no ad hoc assumptions at the coarse scale and retains all the physics of the original multiphase flow equations.

Introduction
One of the more challenging problems in reservoir simulation is to resolve all the pertinent scales in both the data and the solution. Computational power limitations generally prevent one from using as fine a grid as would be desired, especially when multiple simulations for Monte-Carlo or other statistical analyses are required. There are two natural approaches to address this limitation. One is to change the scale on which the data are represented by some kind of averaging procedure, and then to solve the problem on a coarse grid.

The other natural approach is to improve the resolution of the numerical simulation on a coarse grid through some type of subgrid technique. In either case, the desire is to change from the fine scale on which the problem is defined to a coarser scale by defining an appropriate coarse scale problem that captures in some way the fine scale details. Broadly speaking, we might call either approach upscaling; however, as is common in the petroleum industry, we will use the term “upsampling” to refer to the former approach of finding averaged parameters (such as permeability) suitable for use on a coarse scale. In this paper we present an approach of the latter, subgrid type.

There is a large and growing literature on upscaling techniques. We will not attempt a literature review here (see, e.g., the excellent review by Renard and de Marsily); we merely mention a few of the main techniques. Early techniques involved in an essential way averaging or homogenization of physical parameters such as permeability. While such upscaling techniques can be very effective for purely linear problems, they are less satisfactory for nonlinear problems. They suffer from the elementary observation that a nonlinear function of an average is not the average of the nonlinear function. For example, over a coarse grid-block, the value of capillary pressure evaluated at the average saturation is not at all the same as the average over the grid-block of the capillary pressure.

More sophisticated upscaling techniques have been developed to circumvent the inadequacies of simple averaging, including the development of renormalization techniques to successively upscale to coarse levels, pseudo-functions, and stochastic methods. Numerical subgrid techniques have also been developed, including using modified finite element basis functions and explicit subgrid techniques that seek to improve the resolution of the coarse solution after it has been computed.

These techniques all attempt in some way to represent fine scale information on coarse scales in an indirect way, and sometimes require at least some information about the nature of the flow that is expected under field management conditions. Although most upscaling and subgridding techniques are dynamic in that they respond to the changing state of the reservoir, many do so through anticipation of the possibilities. Often one needs some kind of closure assumption such as the imposition of local boundary conditions, the expected primary flow direction, or expected limits on certain parameters such as flow rates.

To handle the dynamic and sometimes unanticipated nature of reservoir conditions, we present in this paper an implicit numerical subgrid technique that allows us to finely resolve the pressure equation even though we end up solving the problem on a coarse grid. This technique is a locally conservative variational multiscale method. We also discuss its implementation in a sequential two-phase waterflood research simulator. We maintain a fully implicit (as opposed to an explicit) coupling between the coarse and fine or subgrid scales, and we obtain a fine scale representation of the reservoir state. As a consequence, we make use of the capillary pressure and relative permeability curves directly and accurately on the fine scale on which they are defined. No pseudo-functions are needed, nor do any arise in our technique. Our technique allows us to handle fine scales in the heterogeneous absolute permeability and porosity, the non-
linear functions relative permeability and capillary pressure, and even the fine scale position of wells.

The idea is to consider the simulation as defined on a fine grid, and to re-fine this grid to form a reasonable coarse grid over which we can compute the solution. We break the solution into two parts, the coarse scale representation of the solution plus the subgrid part. The subgrid part is defined inside the coarse grid blocks. In order to be able to compute it efficiently, it must involve only the coarse solution itself and local information. Because of this computational restriction and the need to maintain accuracy, we compute the coarse scale Darcy velocity using a second order accurate method. Thus, even though the coarse part of the velocity has accuracy based on the coarse scale resolution \( H \) (the diameter of a coarse grid block), the expected accuracy is actually proportional to \( H^2 \). Thus the coarse scale velocity is accurate from the point of view of the subgrid-scale, for which we use a more standard first order accurate method such as cell-centered finite differences. The use of a low order method for the solution on the subgrid scale is natural, since, e.g., heterogeneities in the permeability are likely to produce solutions with large spatial gradients. It is well-known that higher order methods do not in general improve such solutions much, and certainly not enough to justify the added cost.

Because we insist on an implicit coupling between the coarse and subgrid scales, there will be a mixture of these two parts of our solution in the equations. Some kind of static condensation or Schur complement technique is needed to eliminate the subgrid unknowns from the equations. We do this using a technique involving numerical Greens functions, also called influence functions. This technique allows us to treat the subgrid and coarse scales in completely separate parts of the computer code. It is also relatively memory efficient.

To maintain local mass conservation, our procedure is based on mixed finite element methods.\(^{9,10}\) It is known that the lowest order Raviart-Thomas method,\(^11\) RT0, when combined with numerical quadrature to evaluate some of the integrals that arise, is the same as cell-centered finite differences. A similar method that gives higher order velocities uses the finite element spaces defined in 2-D by Brezzi, Douglas, and Marini,\(^12\) BDM1, and generalized in 3-D by Brezzi, Douglas, Durán and Fortin,\(^13\) BDDF1.

The Numerical Subgrid Technique

In this section we present in some detail our two-scale numerical subgrid technique, i.e., our locally conservative variational multiscale method, as applied to the pressure equation. Because of the need for a higher order accurate coarse solution, we must present the technique in a finite element context rather than a cell-centered finite difference context.

**The pressure equation.** In order to simplify the presentation, we illustrate our ideas on the differential equations

\[
\begin{align*}
\n a P + \nabla \cdot \mathbf{u} &= b, \\
\n \mathbf{u} &= -d(\nabla P - c),
\end{align*}
\]

where \( a, b, c \) and \( d \) are constants. The pressure equation is an example of such a system, as we will see later.

**The finite element basis functions.** In a mixed finite element method, finite element basis functions are used to approximate both the pressure \( P \) and the Darcy velocity \( \mathbf{u} \). The Darcy velocity is not defined from the pressure as a simple finite difference involving nearby pressures; in fact, it is defined from all the pressures in the reservoir, although the nearby ones are the most important. That is, \( \mathbf{u} \) is not defined from \( P \) through a finite stencil or molecule, and so the method does not reduce to finite differences. (However, for RT0, we can approximate \( \mathbf{u} \) as a finite stencil of the pressures, and recover the usual 7 point stencil,\(^14,15\) or even a 19-point stencil if the permeability is a full tensor.\(^16\))

**Pressure basis functions.** In both the RT0 and BDDF1 (or BDM1) spaces, the pressure is approximated by a piecewise discontinuous constant function. That is, the pressure is approximated on each grid block by a constant value, just like it is in cell-centered finite differences. Such functions are linear combinations of simpler basis functions.

Let us number the coarse grid blocks \( 1, 2, \ldots, N \). For the \( i \)th coarse grid block, there is a pressure basis function, call it \( \tilde{w}_i \), which takes the value 1 on the block and 0 everywhere else. Within each coarse grid block are similar subgrid-scale basis functions. Suppose that there are \( M_i \) fine grid blocks in the \( i \)th coarse grid block. For each \( k = 1, 2, \ldots, M_i \), we let \( w_k^i \) be 1 on the \( k \)th subgrid block and 0 everywhere else.

In a finite element method, we approximate the pressure in the finite element space; thus, defining both a coarse and subgrid pressure, we have that for any point \( x \) in the reservoir,

\[
\begin{align*}
\tilde{P}(x) &= \sum_{i=1}^{N} a_i \tilde{w}_i(x), \\
\tilde{\tilde{P}}(x) &= \sum_{i=1}^{N} M_i \sum_{k=1}^{M_i} \beta_k^i w_k^i(x), \\
P(x) &= \tilde{P}(x) + \tilde{\tilde{P}}(x),
\end{align*}
\]

where the coefficients \( a_i \) and \( \beta_k^i \) are to be determined. Even though our intent is to scale up to the coarse grid, we have in fact a pressure value for each fine grid block, as illustrated in Fig. 1.

**Velocity basis functions.** The velocity is a vector, so the velocity basis functions are also vector functions. We describe the basis functions as if the grid block was simply the unit cube \( 0 \leq x \leq 1, \ 0 \leq y \leq 1, \) and \( 0 \leq z \leq 1 \). There are 6 basis functions common to RT0 and BDDF1, which
have a nonzero divergence. These are given by
\[ \begin{align*}
\mathbf{v}_1 &= (1 - x)e_1, \\
\mathbf{v}_2 &= xe_1, \\
\mathbf{v}_3 &= (1 - y)e_2, \\
\mathbf{v}_4 &= ye_2, \\
\mathbf{v}_5 &= (1 - z)e_3, \\
\mathbf{v}_6 &= ze_3.
\end{align*} \]

There is one function for each face of the grid block, and each represents a unit flow either into or out of the block across that face, and no flow across the other faces. For example, \( \mathbf{v}_1 \) gives a unit flow into the block across the face \( x = 0 \), while \( \mathbf{v}_2 \) gives a unit flow out of the block across \( x = 1 \).

The BDDF1 spaces have the 12 additional basis functions
\[ \begin{align*}
\mathbf{v}_7 &= (1 - x)(2y - 1)e_1 + (y^2 - y)e_2, \\
\mathbf{v}_8 &= x(2y - 1)e_1 + (y - y^2)e_2, \\
\mathbf{v}_9 &= (x^2 - x)e_1 + (1 - y)(2x - 1)e_2, \\
\mathbf{v}_{10} &= (x - x^2)e_1 + y(2x - 1)e_2, \\
\mathbf{v}_{11} &= (x - x^2)e_1 + (1 - z)(2x - 1)e_3, \\
\mathbf{v}_{12} &= (x - x^2)e_1 + z(2x - 1)e_3, \\
\mathbf{v}_{13} &= (1 - x)(2z - 1)e_1 + (z^2 - z)e_3, \\
\mathbf{v}_{14} &= x(2z - 1)e_1 + (z - z^2)e_3, \\
\mathbf{v}_{15} &= (1 - y)(2z - 1)e_2 + (z^2 - z)e_3, \\
\mathbf{v}_{16} &= y(2z - 1)e_2 + (z - z^2)e_3, \\
\mathbf{v}_{17} &= (y^2 - y)e_2 + (1 - z)(2y - 1)e_3, \\
\mathbf{v}_{18} &= (y - y^2)e_2 + z(2y - 1)e_3.
\end{align*} \]

Each has vanishing divergence, so these basis functions represent no net flow. They merely redistribute flow across the face from one side to the other. For example, the \( \mathbf{v}_7 \) represents a redistribution of flow on the \( x = 0 \) face, by allowing flow into the block on the \( y = 0 \) side and taking it back out on the \( y = 1 \) side. The function \( \mathbf{v}_{13} \) is similar, except that flow is redistributed in the \( z \) direction.

For an arbitrary rectangular grid block, we translate and scale these functions. It should be noted that \( \mathbf{v}_7 \) to \( \mathbf{v}_{18} \) must be scaled so that the divergence remains zero. For example, \( \mathbf{v}_7 \) becomes
\[ \mathbf{v}_7 = (1 - x/h_x)(2y/h_y - 1)e_1 + (y^2/h_y - y)/h_x e_2. \]

For consistency of flow, these basis functions are paired across grid blocks. For example, two blocks that share an \( x \) face would have \( \mathbf{v}_2 \) on the left and \( \mathbf{v}_1 \) on the right, or \( \mathbf{v}_8 \) on the left and \( \mathbf{v}_7 \) on the right. Thus there are 3 basis functions per grid block face for BDDF1, and 1 for RT0. (In 2-D, the RT0 and BDM1 spaces are similar, except that only \( \mathbf{v}_1 \) to \( \mathbf{v}_4 \) and \( \mathbf{v}_7 \) to \( \mathbf{v}_{10} \) arise.)

In our subgrid approach, we need 3 BDDF1 basis functions defined for each coarse grid block face. Call these basis functions \( \mathbf{v}_j \), \( j = 1, 2, ..., N' \). Within coarse grid block \( i \), we need RT0 basis functions defined for each fine grid block face, which we denote by \( \mathbf{v}_j \), \( j = 1, 2, ..., M_i \). Finally, defining both a coarse and subgrid velocity, we have that
\[ \begin{align*}
\mathbf{u}(x) &= \sum_{j=1}^{N'} \gamma_j \mathbf{v}_j(x), \\
\mathbf{u}(x) &= \sum_{i=1}^{N} \sum_{j=1}^{M_i} \delta_{ij} \mathbf{v}_j(x), \\
\mathbf{u}(x) &= \bar{u}(x) + \mathbf{u}(x),
\end{align*} \]
where the coefficients \( \gamma_j \) and \( \delta_{ij} \) are to be determined.

As illustrated in Fig. 1, the velocity has 3 fluxes (2 in 2-D) on each coarse grid block face representing flow between the grid blocks that varies linearly over the face, and a single constant flux across fine grid block faces that lie within the coarse grid blocks. Thus we have good resolution. It is the special placement of these fluxes, i.e., these finite element nodal values, that allows us to scale up to the coarse grid.

The direct finite element formulation. Multiply each equation (1)-(2) by a finite element basis function, which is called a test function, and integrate. Let \( \mathbf{w} \) be any of the \( \mathbf{w}_i \) or \( \mathbf{w}_f \), and \( \mathbf{v} \) be any of the \( \mathbf{v}_j \) or \( \mathbf{v}_f \). This results in the equations
\[ \begin{align*}
\int a \mathbf{P} \mathbf{w} dV &= \int \nabla \cdot \mathbf{u} \mathbf{w} dV = \int \mathbf{b} \mathbf{w} dV, \\
\int d^{-1} \mathbf{u} \cdot \mathbf{v} dV &= -\int \nabla \cdot \mathbf{P} \cdot \mathbf{v} dV + \int \mathbf{c} \cdot \mathbf{v} dV \\
&= \int \nabla \cdot \mathbf{v} dV + \int \mathbf{c} \cdot \mathbf{v} dV,
\end{align*} \]
where we have made use of the divergence theorem to replace \( -\int \nabla \cdot \mathbf{P} \cdot \mathbf{v} dV \) by \( \int \mathbf{P} \nabla \cdot \mathbf{v} dV \), since there is no flow on the external boundary of the reservoir. (If other boundary conditions are used, certain boundary terms appear at this stage, but otherwise no particular complications arise later.)

Combining (9)-(10) with (3)-(5) and (6)-(8) results in a fully implicit system of linear equations, a square matrix problem of size \( N + \sum_i M_i + N' + \sum_i M_i' \), for the unknowns \( \alpha_i, \beta_k, \gamma_j, \) and \( \delta_{ij} \). This system is too large to solve as formulated, so we will not discuss its implementation. It has nearly as many unknowns as the fine scale solution itself: the same number of pressures, and as many velocity fluxes within the coarse blocks. On the faces of the coarse blocks, we have fewer velocity fluxes than one would have in a fine scale solution, but these fluxes are higher-order accurate and so give a good approximation of the velocity.

The system is actually underdetermined, since the pressure basis functions sum to 1, i.e., \( \sum_{i=1}^N \mathbf{w}_i = \mathbf{w} \). That is, the decomposition of \( \mathbf{P} \) into \( \bar{P} \) and \( \bar{P} \) is not unique. The removal of this ambiguity depends on the nature of \( a \). For
each coarse block \(i\), if \(a = 0\) on the block, we cannot solve our subgrid problems below as they are written. Instead, we remove, say, the test function \(w = \tilde{w}_i^j\) and replace that equation by the requirement that

\[
\int \tilde{P} \tilde{w}_i \, dV = 0 .
\]

If, however, \(a \neq 0\) on the block, then we can solve our subgrid problems below as written (i.e., we do not replace the \(w = \tilde{w}_i^j\) equation as above), and we require either (11) or instead

\[
\int a \tilde{P} \tilde{w}_i \, dV = 0 ,
\]

as we will make clear later. In both cases, however, we can solve our equations, and the decomposition \(P = \tilde{P} + \bar{P}\) is unique.

Our goal now is to exploit our choice of basis functions so that the system can be solved in smaller steps, with the last step being the solution of a substantially reduced matrix problem of size \((N + N')\times(N + N')\) involving only the coarse scale unknowns \(a_i\) and \(\gamma_j\). That is, we scale up to the coarse level. We emphasize that we are not changing the method below, but merely describing how to obtain the solution. Our solution technique requires 3 steps; no iteration is performed.

**Numerical Greens Functions.** The key is to use numerical Greens functions, also known as influence functions. The idea can be illustrated in a trivial example. Suppose we want to solve \(ax + by + cz = d\) for \(x\) in terms of \(y\) and \(z\), which are not known. The solution is obviously \(x = (d - by - cz)/a\). We can arrive at this same result by solving three problems that involve neither \(y\) nor \(z\) as follows. First set \(y = z = 0\) and solve \(ax_0 = d\). Next set the nonhomogeneous terms to 0 (i.e., \(d = 0\)), set \(y = 1\) and \(z = 0\), and solve \(ax_1 = -b\). Finally, set \(d = 0\), \(y = 0\), and \(z = 1\) and solve \(ax_2 = -c\). Then the solution is the combination of these simple solutions: \(x = x_0 + x_1y + x_2z\).

Note that \(x_1\) gives the influence of \(y\) on \(x\) \((x_1 = \partial x/\partial y)\) and \(x_2\) gives the influence of \(z\) on \(x\) \((x_2 = \partial x/\partial z)\). Alternatively, one can view \(x_1\) as the response of the system to a unit stimulus in \(y\), that is, a Greens function, and similarly for \(x_2\) and \(z\).

This technique works even when \(x\) is a vector, \(a\) is a matrix, and \(b, c,\) and \(d\) are vectors.

In our case, we apply the idea to each coarse grid block. On the \(i\)th block, \(x\) refers to the subgrid unknowns \(\beta^i_k\) and \(\delta^i_{\ell}\), and \(y\) and \(z\) refer to the coarse grid unknowns \(a_i\) and all \(\gamma_j\) for which \(\tilde{v}_j\) lives (i.e., is not zero) on the \(i\)th block. We have arranged that there are very few such coarse block unknowns.

**The two-scale finite element formulation.** We begin by solving the subset of equations (9)–(10) for which the test functions live on the subgrid scale. These subgrid equations are coupled to the coarse scale. On coarse block \(i\), the subgrid is affected only by the coarse coefficients \(a_i\) and the 18 \(\gamma_j\) for which \(\tilde{v}_j\) lives on the block. Because of (11)–(12), we have arranged the computation so that \(a_i\), i.e., the coarse pressure \(\tilde{P}\), does not affect the subgrid scale. Thus only 18 parameters affect the subgrid scale, and we can determine the influence of each through the numerical Greens function approach.

**Step 1: The subgrid problems.** For coarse block \(i\), consider (9)–(10) with test functions \(w = \tilde{w}_k^i\) and \(v = \tilde{v}_\ell^i\). In the numerical Greens function approach, we first solve assuming the coarse scale information is set to zero. Solve for

\[
\tilde{P}_0^i = \sum_{k=1}^{M_i} \tilde{\tilde{\beta}}_{0,k}^i \tilde{w}_k^i ,
\]

\[
\tilde{u}_0^i = \sum_{\ell=1}^{M'_i} \tilde{\tilde{\delta}}_{0,\ell}^i \tilde{v}_\ell^i ,
\]

the equations represented by

\[
\int a \tilde{P}_0^i \tilde{w}_k^i \, dV + \int \nabla \cdot \tilde{u}_0^i \tilde{w}_k^i \, dV = \int b \tilde{w}_k^i \, dV ,
\]

\[
\int d^{-1} \tilde{u}_0^i \cdot \tilde{v}_\ell^i \, dV
\]

\[
= \int \tilde{P}_0^i \nabla \cdot \tilde{v}_\ell^i \, dV + \int c \cdot \tilde{v}_\ell^i \, dV .
\]

In matrix form, these equations are

\[
\sum_{k=1}^{M_i} \tilde{\tilde{\beta}}_{0,k}^i \tilde{w}_k^i \, dV + \sum_{\ell=1}^{M'_i} \tilde{\tilde{\delta}}_{0,\ell}^i \tilde{v}_\ell^i \, dV
\]

\[
= \int b \tilde{w}_k^i \, dV ,
\]

\[
\sum_{\ell=1}^{M'_i} \tilde{\tilde{\delta}}_{0,\ell}^i \tilde{v}_\ell^i \, dV
\]

\[
= \sum_{k=1}^{M_i} \tilde{\tilde{\beta}}_{0,k}^i \tilde{w}_k^i \nabla \cdot \tilde{v}_\ell^i \, dV + \int c \cdot \tilde{v}_\ell^i \, dV .
\]

We then solve for the influence of each \(\tilde{v}_j\) living on the \(i\)th block. There are 18 such \(j\). We solve for

\[
\tilde{P}_j^i = \sum_{k=1}^{M_i} \tilde{\tilde{\beta}}_{j,k}^i \tilde{w}_k^i ,
\]

\[
\tilde{u}_j^i = \sum_{\ell=1}^{M'_i} \tilde{\tilde{\delta}}_{j,\ell}^i \tilde{v}_\ell^i ,
\]

the equations represented by

\[
\int a \tilde{P}_j^i \tilde{w}_k^i \, dV + \int \nabla \cdot (\tilde{v}_j^i + \tilde{u}_j^i) \tilde{w}_k^i \, dV = 0 ,
\]

\[
\int d^{-1}(\tilde{v}_j^i + \tilde{u}_j^i) \cdot \tilde{v}_\ell^i \, dV = \int \tilde{P}_j^i \nabla \cdot \tilde{v}_\ell^i \, dV .
\]
In matrix form, we have
\[
\sum_{k=1}^{M_i} \beta_{j,k}^i \int a \, \hat{w}_k \, \hat{w}_k^i \, dV + \sum_{\ell=1}^{M_i} \delta_{j,\ell}^i \int \nabla \cdot \hat{v}_\ell \, \hat{w}_k^i \, dV
\]
\[= - \int \nabla \cdot \hat{v}_j \, \hat{w}_k^i \, dV, \quad (23)\]
\[
\sum_{\ell=1}^{M_i} \delta_{j,\ell}^i \int d^{-1} \hat{v}_\ell \cdot \hat{v}_\ell^i \, dV
\]
\[= \sum_{k=1}^{M_i} \beta_{j,k}^i \int \hat{w}_k \, \nabla \cdot \hat{v}_\ell \, dV - \int d^{-1} \hat{v}_j \cdot \hat{v}_\ell^i \, dV. \quad (24)\]

We must insist that the normalization (11) applies to \(\bar{P}_0^i\) and that (11) or (12) apply to each \(\bar{P}_j\), depending on whether \(a = 0\). This normalization is done before trying to solve the equations if \(a = 0\) on the block; otherwise, it is done after obtaining the solution.

We use a direct solver in this step, since there are multiple right-hand sides (19 in all), and these problems are generally small (at most the subgrid is perhaps \(10 \times 10 \times 10\)). These problems also parallelize trivially.

Once this step is complete, we then have the formal representation of the subgrid solution on coarse block \(i\) as
\[
\bar{P} = \bar{P}_0^i + \sum_j \gamma_j \bar{P}_j^i = \sum_{k=1}^{M_i} \left( \sum_j \gamma_j \beta_{j,k}^i \right) \hat{w}_k^i, \quad (25)\]
\[
\bar{u} = \bar{u}_0^i + \sum_j \gamma_j \bar{u}_j^i = \sum_{\ell=1}^{M_i} \left( \sum_j \gamma_j \delta_{j,\ell}^i \right) \hat{v}_\ell^i. \quad (26)\]

Thus
\[
\beta_{j,k}^i = \gamma_j \beta_{j,k}^i, \quad (27)\]
\[
\delta_{j,\ell}^i = \gamma_j \delta_{j,\ell}^i. \quad (28)\]

We know every quantity above except the \(\gamma_j\) so these are not yet computable. However, these formulas can be used in the coarse scale equations to obtain a linear system for the \(\alpha_i\) and \(\gamma_j\) only.

**Step 2: The coarse problem.** In the second step, we solve the coarse problem, which is given by (9)–(10) with the test functions which live on the coarse scale \(w = \bar{w}\) and \(v = \hat{v}_j\). This system, when combined with (25)–(26), gives a matrix problem for the \(\alpha_i\) and \(\gamma_j\). It is derived in detail in the Appendix. The system is
\[
\sum_{i=1}^{N} \alpha_i \int a \, \bar{w}_i \, dV + \sum_{j=1}^{N} \gamma_j \int \nabla \cdot \bar{v}_j \, \bar{w}_i \, dV
\]
\[= \int (b - a \bar{P}_0) \bar{w}_i \, dV, \quad (29)\]
\[
\sum_{j=1}^{N} \gamma_j \left\{ \int d^{-1} (\bar{v}_j + \bar{u}_j) \cdot (\bar{v}_j + \bar{u}_j) \, dV \right\}
\]
\[+ \int a \bar{P}_j \bar{P}_j \, dV \right\}
\[= \sum_{i=1}^{N} \alpha_i \int \bar{w}_i \nabla \cdot \bar{v}_j \, dV + \int (c - a \bar{u}_0) \cdot \bar{v}_j \, dV. \quad (30)\]

where, of course, \(\bar{P}_j\) is \(\bar{P}_j^i\) and \(\bar{u}_j\) is \(\bar{u}_j^i\) whenever we integrate over the ith coarse grid block.

If the subgrid scale influence functions are set to zero, this is a coarse grid BDFD1 mixed finite element discretization of the pressure equations representing conservation of mass and Darcy’s law. With the influence functions, the coarse equations represent a coarse grid conservation of mass and a coarse grid Darcy law, with the matrix coefficients modified by the subgrid-scale influence functions. Thus to implement, one first develops a code to handle the BDFD1 method on the coarse grid, and then modifies the integration routines to solve (29)–(30).

**Step 3: The fine scale solution.** Finally, we construct a fine scale representation of our solution by combining the previous results to obtain
\[
P = \bar{P} + \sum_{i=1}^{N} \left( \bar{P}_0^i + \sum_j \gamma_j \bar{P}_j^i \right)
\]
\[= \sum_{i=1}^{N} \left[ \alpha_i \bar{w}_i + \sum_{k=1}^{M_i} \left( \sum_j \gamma_j \beta_{j,k}^i \right) \hat{w}_k^i \right]. \quad (31)\]
\[
\bar{u} = \bar{u} + \sum_{j=1}^{N} \left( \bar{u}_0^j + \sum_j \gamma_j \bar{u}_j^i \right)
\]
\[= \sum_{j=1}^{N} \gamma_j \bar{v}_j + \sum_{i=1}^{N} \left( \sum_{k=1}^{M_i} \left( \delta_{j,k}^i \right) \hat{v}_\ell^i \right). \quad (32)\]

In summary, in Step 1, we solve in parallel (17)–(18) and (23)–(24), possibly modified as mentioned in (11) for solvability, for each coarse grid block \(i\) for the coefficients of the numerical Greens functions (or influence functions). We then construct \(\bar{P}_0^i, \bar{u}_0^i, \bar{P}_j^i\), and \(\bar{u}_j^i\) using (13)–(14), and normalize by (11) or (12) if this was not already done for solvability. In Step 2, we solve the coarse equations (29)–(30) for the coarse coefficients \(\alpha_i\) and \(\gamma_j\). This gives part of our solution \(P\) and \(\bar{u}\). Finally, in Step 3, we construct a fine scale representation of our solution from (31)–(32). This is precisely the solution
to (9)–(10). Further details of implementation can be found in Arbogast.\textsuperscript{17}

Application to Two-Phase Incompressible, Immiscible Flow

We can easily adapt our subgrid technique to two-phase immiscible flow if the system is written as a pressure equation coupled to a saturation equation. While our technique could be adapted for a fully implicit system, we chose to develop a research simulator that used a sequential approach. Thus both the pressure and saturation equations are individually fully implicit, but they are decoupled in time. We solve for the pressure using the saturation at the previous time, update the velocities, and then advance the saturation. Extensions to three-phase black-oil simulation would be similar.

The formulation of the equations. We use the standard two-phase flow equations; however, they are formulated in a somewhat nonstandard way. This has no bearing on the numerical subgrid technique, but we mention it for completeness.

The global pressure formulation. The formulation of the pressure equation that we use is due to Chavent.\textsuperscript{18} When the fluids are incompressible, we can define a "global" pressure $P$ from one of the phase pressures and the saturation $S$ as

$$P = P_0 + \int_S \frac{\lambda_w(s)}{\lambda(s)} P'_w(s) \, ds,$$

where $\lambda_w(s) = k_{r,w}(S)/\mu_w$ and $\lambda_o(s) = k_{r,o}(S)/\mu_o$ are the relative mobilities, and $\lambda(S) = \lambda_w(S) + \lambda_o(S)$. Then

$$\lambda \nabla P = \lambda_w \nabla P_w + \lambda_o \nabla P_o,$$

and, with $u = u_w + u_o$ being the total velocity, we have the pressure equation

$$\frac{\partial \phi}{\partial t} + \nabla \cdot u = g(P),$$

$$u = -K \lambda(S) (\nabla P - \rho(S) e_2),$$

where $g(P) = q_w + q_o$ represents the total flow of the wells and the density is

$$\rho(S) = \frac{\lambda_w(S)}{\lambda(S)} \rho_w + \frac{\lambda_o(S)}{\lambda(S)} \rho_o.$$

With $S$ being fixed in the equation to its value at the previous time step, this equation is the same as (1)–(2) if the time derivative is replaced with a backward finite difference and $g(P)$ is an affine function. Generally one assumes the rock compressibility is constant, so $\phi = \phi(P)$ is itself affine. If $g(P)$ is not affine, we linearize it in a Newton-Raphson procedure to obtain an equation of the form (1)–(2).

The Kirchhoff saturation formulation. The saturation equation is formulated so as to use a standard fractional flow, but with the diffusive terms modified with a Kirchhoff transformation.\textsuperscript{19} That is, we have

$$\frac{\partial \phi}{\partial t} + \nabla \cdot u_w = q_w(S),$$

$$u_w = -K \nabla Q + c(u, S),$$

where the "complementary" potential $Q$ and $c$ are given by

$$Q(S) = - \int_0^S \frac{\lambda_w(s)}{\lambda(s)} P'_w(s) \, ds,$$

$$c(u, S) = \frac{\lambda_w(S)}{\lambda(S)} \left[ u - K \lambda_o(S)(\rho_o - \rho_w) e_2 \right].$$

In the sequential approach, we have already solved for $u$, and we use its fine scale representation here. After replacing the time derivative with a backward finite difference, and after a Newton linearization, this equation is also of the form (1)–(2).

We can adapt our subgrid idea to the saturation equation, and did so with a good deal of success.\textsuperscript{20} However, because the saturation equation is diagonally dominant, we found that the time savings were marginal when using our particular subgrid technique for the saturation part of the solution. Moreover, in certain difficult problems,\textsuperscript{21} we found that the use of a higher order method prevented us from satisfying the maximum principle that requires the saturation to lie between its residual and one minus the residual of the other fluid. It is well known that such monotone methods must necessarily be of the first order (if the method itself is linear). Thus, we chose to solve the saturation equation on the fine scale. For simplicity, we used one-point upstream weighting.

Bottomhole pressure wells. All terms associated with constant rate injection wells have been treated in our description of the subgrid technique, since $q$ is known and fixed. Bottomhole pressure wells, as modeled by Peaceman,\textsuperscript{22} present no difficulties either for the subgrid technique or for the global pressure formulation. We have along the well at depth level $k$,

$$q_{w,k} = -\kappa_{w,k}(P_{w,k} - P_{well,k}),$$

$$q_{o,k} = -\kappa_{o,k}(P_{o,k} - P_{well,k}),$$

$$P_{well,k} = P_{BH} + \rho_{well} g(z_k - z_{BH}),$$

$$\rho_{well} = \frac{\sum_k (q_{w,k} \rho_{w,k} + q_{o,k} \rho_{o,k})}{\sum_k (q_{w,k} + q_{o,k})},$$

where the $\kappa_w$ and $\kappa_o$ are given by Peaceman. Thus, using (33),

$$q_k = q_{w,k} + q_{o,k}$$

$$= -(\kappa_{w,k} + \kappa_{o,k})(P_k - P_{BH} + \rho_{well} g(z_k - z_{BH}))$$

$$- \kappa_{w,k} F_w(S_k) - \kappa_{o,k} F_o(S_k).$$
where

\[ P_a = P + P_o(S), \]
\[ P_o = P + F_o(S), \]
\[ F_o(S) = - \int_S \frac{\lambda_0(s)}{\lambda(s)} p^2(s) \, ds, \]
\[ F_w(S) = F_o(S) - P_w(S). \]

We found it convenient to use the numerical Greens function approach to solve the bottomhole pressure wells. We let \( \rho_{\text{well}} \) be our influence parameter and solve for the pressure with \( \rho_{\text{well}} \) set to 0, and then solve for the pressure again with the nonhomogeneous terms set to 0 and \( \rho_{\text{well}} \) set to 1. Inspection of (42) shows that both of these equations are linear in \( P \), so they simply result in a modification of \( a \) and \( b \) in (1)–(2). Finally a nonlinear iteration on (41) determines \( \rho_{\text{well}} \) and completes the solution.

We have one additional pressure solve for each additional bottomhole pressure well, so this technique is feasible only for a small number of wells. This implementation of the wells would not be practical in a full field simulation. In the general case, we would need to include the well terms as linearized for a Newton-Raphson procedure. Again, only \( a \) and \( b \) need to be modified.

Due to the evolution of the simulator, we found it convenient for the saturation equation to incorporate bottomhole pressure wells in a somewhat unusual way. We took the total injection rate as determined from the pressure equation and divided it into the phases according to their relative mobility.

Simulations exploring the ability of the subgrid technique to simulate near well behavior and using the Peaceman model can be found in Arbogast. Very good agreement with fine grid computations was found.

General Numerical Performance

Before presenting our reservoir simulation examples, we mention a few general features of the subgrid method, and an important modification.

Convergence rate for smooth solutions. We tested the convergence rate of the method in Arbogast by solving the pressure equation (1)–(2) in cases for which the solution is smooth and known. When the number of subgrid blocks per coarse block is held fixed, the errors in the velocities display second order convergence. This is to be expected for the BDDF1 method, however the RT0 method is only linearly convergent. Thus, the use of higher order coarse elements is indeed sufficient to maintain accuracy, and in fact this higher order is reflected in the subgrid computation.

Savings in computation time. The speed of the subgrid method is difficult to quantify, as it depends on so many factors. In our test cases, we used simply Jacobi pre-conditioned conjugate gradients. It is well known that such a solver is very easy to implement, but also that it does not perform particularly well. However, we have consistently observed a reduction in the solution times of about a factor of 2 to 10 for our subgrid method compared to solving the fine scale pressure equation with RT0 (which is essentially standard cell-centered finite differences). Since our method is second order accurate compared to first order for RT0, this is quite a savings in time. The savings in solution time compared to solving the second order accurate fine scale BDDF1 is much more than a factor of 100 in most cases.

A more nearly optimal solver should close the timing gap somewhat, so these reduction factors should not be taken too literally. What is clear is that it is easier to precondition a smaller, coarser system than a larger, finer one. This is due partly to the reduced size, but also more importantly to the reduction in condition number that an averaged or upcaled problem exhibits. The performance of Jacobi preconditioned conjugate gradients is very sensitive to the condition number, and thus we see good speedup for the two-scale system.

Even a more efficient solution strategy will benefit from the improved condition number of the coarse problem compared to the fine scale problem. We note in passing that if one wanted to solve the fine scale problem, one could also use our subgrid technique as a type of preconditioner. In that case, perhaps one would be content with RT0 on the coarse scale.

One additional remark about the computational speed should be made. The two-scale subgrid method improves data locality, so there is the potential to reuse more data within high-speed memory caches than one might find in a fine scale solution technique.

Average permeability in the coarse equation.

When solutions are not so smooth, it is possible that the coarse flow is not quite of the proper magnitude. This is due to using the fine scale permeability in the coarse equation (30), coefficient \( d \). To remedy this, we can use an averaged permeability \( \bar{d} \) in place of \( d \) on the left-hand side of the equality. That is, replace \( \int d^{-1}(\bar{v}_j + \bar{u}_j) \cdot (\bar{v}_j + \bar{u}_j) \, dV \) by \( \int (\bar{d})^{-1}(\bar{v}_j + \bar{u}_j) \cdot (\bar{v}_j + \bar{u}_j) \, dV \), but leave \( d \) alone in every other term, especially in the subgrid equations. As we will see in our two reservoir examples, this can bring about an improvement in the overall performance of the subgrid method.

A 3-D Numerical Example

In this section we present numerical results that demonstrate the application of the subgrid method described above to some simple test cases. We compare various subgrid computations to a fine scale computation which we treat as the true solution. We consider a rectangular domain 500m \( \times \) 250m \( \times \) 40m with a water injector and a single producer, discretized into 1600 fine scale cells \((20 \times 10 \times 8)\). We compute on a \( 4 \times 2 \times 2 \) coarse grid, corresponding to a 100-fold reduction in the number of cells. The porosity is assumed constant at 0.25 and the permeability field is a realization of a spatially correlated distribution derived from observations...
on an outcrop. A constant-rate injector fully penetrates the reservoir at \( x = 100m, \ y = 75m \), while a constant bottomhole pressure producer at \( x = 400m, \ y = 200m \) is completed only in the upper 30m of the reservoir. Fig. 2 shows the saturation distribution in a vertical plane containing the injector and a horizontal plane at the bottom of the producer after 0.22 PVI (pore volumes of water injected). The influence of zones of low permeability is evident in the "islands" of high and low saturation. The subgrid solution (using the fine scale permeabilities directly for coefficient \( d \) in (30)) captures many of these small scale features, as illustrated in Fig. 3. Fig. 4 compares the computed water-oil ratio (WOR) as a function of PVI for several schemes for averaging the fine scale permeabilities to obtain \( d \) in (30). The fine solution, subgrid solution, and subgrid solutions with \( d \) in (30) given by the arithmetic average, the harmonic average in the flow direction and the arithmetic average in the direction perpendicular to flow, and the one-third power average. The latter seems to do the best job of capturing the behavior, though it predicts slightly earlier water breakthrough.

### An SPE Comparative Solution Example

Our second numerical example is based on the 2-D model described in the 2001 SPE comparative solution project for upscaling. The differences are relatively minor, and consist mainly in not being able to set the bottomhole pressure properly and setting gravity to zero. The latter was done to avoid complications with the saturation solver, and not due to the pressure equation. Thus our results, though not strictly comparable, do show the potential of our numerical subgrid technique for the pressure equation.

Fig. 5 shows the fine scale solution after 1000 days of injection. The subgrid solution captures several of the water channels on the left side of the domain and several of the bypassed regions on the right side. Fig. 6. Using the root mean square averaging scheme in the coarse part of the method smear the saturation profiles, Fig. 7. The horizontal component of the total velocity at 1000 days shows good qualitative agreement between the three computations as shown in Figs. 8, 9, and 10. The straightforward subgrid approach yields a slightly wider range of velocities than the true solution, while the approach using average permeability in the coarse equation yields a somewhat narrower range.

### A 2-D Quarter Five-Spot Example

Our final example is taken from White and Horne. It is a 2-D quarter five-spot waterflood using the three value permeability field depicted in Fig. 11, reminiscent of a fluvial geological environment that is correlated over large distances. The problem is posed on a 30 \times 30 grid, and solved using a 6 \times 6 coarse grid with a 5 \times 5 subgrid in each coarse block. Note that this does a fairly good job of resolving the high permeability channel. Thus, we also solve the problem on a 3 \times 3 coarse grid with a 10 \times 10 subgrid. The injection well is at fine block (3, 3), and the production well is at fine block (28, 28). We use the simplified data of Table 1, designed so that nonuniformities in the flow are due primarily to the permeability variations and not to differences between the two fluids.

We compare the 30 \times 30 fine grid computation to the subgrid technique used with a coarse grid of size either 6 \times 6 or 3 \times 3. We also compare to 6 \times 6 coarse simulations using the harmonic-arithmetic coarse block average diagonal permeability (harmonic average in primary direction, followed by arithmetic average in the cross direction on each block). We use either standard RT0 cell-centered finite differences (used also for the fine grid computation), or the BDM1 second order accurate discretization, since the latter is more comparable to the subgrid method.

Results are summarized in Table 2. The maximum pressure errors (assuming the fine solution is true) show that the subgrid method is extremely accurate. These errors include errors due to saturation errors and their subsequent effect on the relative permeability. Breakthrough is predicted very well with the subgrid method, and not so well when the average permeability is used. We show the complete producing water cut curve in Fig. 12. The cumulative oil production, shown in Fig. 13, also shows that the best results are achieved when the subgrid technique is used. Overall, the harmonic-arithmetic coarse block average diagonal permeability on a 6 \times 6 grid compares well with the 3 \times 3 subgrid method.

We show the saturation contours at 400 days for the fine grid solution in Fig. 14, the subgrid method on both 6 \times 6 and 3 \times 3 coarse grids in Figs. 15 and 16, and a 6 \times 6 coarse simulation using the harmonic-arithmetic coarse block average diagonal permeability in Fig. 17 (the second order method is similar, but maintains somewhat sharper fronts). The subgrid method, even with a 3 \times 3 coarse grid, does a reasonably good job of resolving the key feature of the problem: preferential flow through the channel.

Timing results are summarized in Table 3. The problem is much too small to obtain representative timing results; however, a few points are clear. First, the subgrid computations are a very small fraction of the total computation time (except for the overly small 3 \times 3 case). Moreover, the subgrid part of the computation would scale linearily in parallel, and could even be omitted in coarse blocks that experience negligible saturation change, since the numerical Greens functions only change when the relative permeability changes. Second, the subgrid technique requires fine scale integration to assemble the matrix. However, in larger problems, the time needed to solve the matrix problem is much greater than the time to assemble the matrix, so this part of the computation is not generally significant. Third, the time to solve the coarse pressure problem, which normally dominates the computation time, is much better for the subgrid method as compared to the fine grid computation (by a factor of about 10 for the 6 \times 6 simulation, and by a factor of almost 100 for the 3 \times 3 case). The coarse problem using an average permeability is quite fast. For comparable size grids, the difference in speed between the subgrid and average per-
meability methods is due to the difference between using a second or first order accurate method. However, the $6 \times 6$ average permeability method compares in accuracy with the $3 \times 3$ subgrid method, which is in fact somewhat faster in solving the coarse grid problem. Fourth, the timings reflect the fact that we chose to solve the saturation equation on the fine grid for the subgrid methods.

**Application to Traditional Upscaling**
The White and Horne technique\textsuperscript{24} obtains coarse grid effective permeabilities from postprocessing a small number of fine grid single-phase simulations. We could use their technique, combined with the subgrid technique, to obtain the effective permeability on a coarse grid, for use in a traditional reservoir simulator.

**Conclusions**
We presented a two-scale numerical subgrid approach, also called a locally conservative multiscale variational method, that decomposes the governing differential equations into a coarse grid-scale operator that is coupled to a subgrid-scale operator. The subgrid-scale problems can be solved independently of the coarse grid approximation thanks to a numerical Greens function technique. The global problem is solved only at the coarse scale, and this solution is corrected at the subgrid scale to obtain the fine scale representation of the problem. In this way no explicit macroscopic coefficients (e.g., effective permeabilities or pseudo-functions) arise, and thus no assumptions about the physics or the expected flow behavior are required.

We implemented the implementation of this technique for the flow of two immiscible, incompressible phases. For simplicity the implementation is sequential, solving the flow equation fully implicitly with saturation-dependent quantities evaluated at the previous time step, then solving the saturation equation fully implicitly and on the fine scale. We observed a reduction in pressure solution times by a factor of 2 to 10 to almost 100 for small problems (larger problems should experience better speed-up). Moreover, we note that we obtain second order accuracy while the fine scale solution used for timing comparison is only first-order accurate. Good agreement between the subgrid and fine scale solutions is observed for test cases in which the coarse grid contains two orders of magnitude fewer grid blocks. Using an appropriate average of the fine scale permeabilities may improve the fidelity of the subgrid solution under some conditions.

**Nomenclature**

- **BDDF1** = first order Brezzi-Douglas-Durán-Fortin mixed finite element function space
- **BDM1** = first order Brezzi-Douglas-Marini mixed finite element function space
- $dV$ = differential of volume, $L^3$
- $e_1 = \text{unit vector in the } x\text{-direction}$
- $e_2 = \text{unit vector in the } y\text{-direction}$
- $e_3 = \text{unit vector in the } z\text{-direction}$
- $H = \text{coarse grid block diameter, } L$
- $h = \text{fine grid block diameter, } L$
- $h_x = \text{grid block length in the } x\text{-direction, } L$
- $h_y = \text{grid block length in the } y\text{-direction, } L$
- $h_z = \text{grid block length in the } z\text{-direction, } L$
- $K = \text{absolute permeability, } L^2$
- $k_r = \text{relative permeability}$
- $M_i = \text{number of fine grid blocks in coarse grid block } i$
- $M_i' = \text{number of fine grid block faces internal to coarse grid block } i$
- $N = \text{number of coarse grid blocks}$
- $N'$ = 3 times the number of coarse grid block faces
- $P = \text{pressure, } m/L^2$
- $P_c = \text{capillary pressure, } m/L^2$
- $Q = \text{Kirchhoff "complementary" potential, } 1/t$
- $q = \text{external well sources and sinks, } 1/t$
- $RT_0 = \text{lowest order Ravindran-Thomas mixed finite element function space}$
- $S = \text{water (or gas) saturation}$
- $u = \text{Darcy velocity, } L/t$
- $v = \text{velocity test function}$
- $\bar{v} = \text{velocity test function on the unit cube}$
- $w = \text{pressure test function}$
- $x = \text{spatial point, } L$
- $(x, y, z) = \text{spatial point, } L$
- $a = \text{finite element coefficients of coarse pressure}$
- $\beta = \text{finite element coefficients of subgrid pressure}$
- $\gamma = \text{finite element coefficients of coarse velocity}$
- $\delta = \text{finite element coefficients of subgrid velocity}$
- $\lambda = \text{relative mobility, } Lt/m$
- $\mu = \text{viscosity, } m/Lt$
- $\rho = \text{density, } m/L^3$
- $\phi = \text{porosity}$

**Subscripts and superscripts**

- $i = \text{coarse grid block}$
- $j = \text{coarse grid block face}$
- $k = \text{subgrid block (or the depth index)}$
- $\ell = \text{subgrid block face}$
- $a = \text{oil}$
- $w = \text{water (or gas)}$

**Acknowledgments**

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**References**


Appendix—Derivation of the Coarse Problem

To scale up the problem (9)–(10) from the fine to the coarse scale, we choose coarse scale test functions. It is easier to describe the derivation if our choice of test functions is actually \( w = \tilde{w} \) and \( v = \tilde{v} + u \tilde{v}_x \), since it more readily results in a symmetric problem with respect to the velocity unknowns. Thus, using our coarse and subgrid decomposition (5) and (8) and the numerical Greens functions as in (25)–(26), we
have that
\[
\int a \left( \bar{P} + \bar{P}_0 + \sum_j \gamma_j \bar{P}_j \right) \bar{w}_j \, dV \\
+ \int \nabla \cdot \left( \bar{u} + \bar{u}_0 + \sum_j \gamma_j \bar{u}_j \right) \bar{w}_j \, dV \\
= \int b \bar{w}_j \, dV ,
\]  
(A-1)
\[
\int d^{-1} \left( \bar{u} + \bar{u}_0 + \sum_j \gamma_j \bar{u}_j \right) \cdot \left( \bar{v}_j + \bar{u}_j \right) \, dV \\
= \int \left( P + \bar{P}_0 + \sum_j \gamma_j \bar{P}_j \right) \nabla \cdot \left( \bar{v}_j + \bar{u}_j \right) \, dV \\
+ \int c \cdot \left( \bar{v}_j + \bar{u}_j \right) \, dV .
\]  
(A-2)

Note that in all cases
\[
\int \bar{w} \nabla \cdot \bar{v} \, dV = 0 ,
\]  
(A-3)
since \( \bar{v} \) has no flux external to the coarse block. Thus several terms cancel above. For either normalization (11) or (12),
\[
\int \bar{a} \bar{P}_j \bar{w}_j \, dV = 0 ,
\]
so (A-1) combined with (3) and (6) gives (29).

Next we note that (21), with test function \( \bar{w} = \bar{P}_j \), implies that
\[
\int \bar{a} \bar{P}_j \bar{P}_j \, dV + \int \nabla \cdot \left( \bar{v}_j + \bar{u}_j \right) \, dV = 0 ,
\]
so
\[
\bar{P}_j \nabla \cdot \left( \bar{v}_j + \bar{u}_j \right) \, dV = - \int \bar{a} \bar{P}_j \bar{P}_j \, dV .
\]  
(A-4)

Similarly, (16) with test function \( \bar{v} = \bar{u}_j \) yields
\[
\int d^{-1} \bar{u}_0 \cdot \bar{u}_j \, dV \\
= \int \bar{P}_0 \nabla \cdot \bar{u}_j \, dV + \int c \cdot \bar{u}_j \, dV .
\]  
(A-5)

Finally,
\[
\int \bar{P}_0 \nabla \cdot \bar{v}_j \, dV = 0
\]  
(A-6)

by normalization (11). Combining (A-2)–(A-6) results in
\[
\int d^{-1} \left( \bar{u} + \sum_j \gamma_j \bar{u}_j \right) \cdot \left( \bar{v}_j + \bar{u}_j \right) \, dV \\
+ \sum_j \gamma_j \int \bar{a} \bar{P}_j \bar{P}_j \, dV \\
= \int \bar{P} \nabla \cdot \bar{v}_j \, dV + \int c \cdot \bar{v}_j \, dV \\
- \int d^{-1} \bar{u}_0 \cdot \bar{v}_j \, dV .
\]  
(A-7)

Combining now (A-7) with (3) and (6) gives (30).

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Fig. 1. A 2-D example of a $12 \times 12$ fine grid decomposed into a $3 \times 3$ coarse grid with $4 \times 4$ subgrids. The dots represent the pressure values, one per fine grid block. The crosses represent the grid block face velocity fluxes. The circled crosses apply to the coarse grid block faces, and represent linear flux variation across the face. The other crosses represent constant fluxes across the subgrid block faces internal to the coarse blocks.

Fig. 2. Fine scale saturation distribution at 0.22 PVI for a 3-D waterflood example solved on a $20 \times 10 \times 8$ grid, which we take to be the true solution. Spatial correlation in the permeability field gives rise to significant small scale heterogeneities in the saturation distribution.

Fig. 3. Subgrid saturation distribution at 0.22 PVI for the same example as in Fig. 2 solved on a $4 \times 2 \times 2$ grid (using a $5 \times 3 \times 4$ subgrid in each coarse grid block). The subgrid procedure captures much of the fine scale behavior.

Fig. 4. Computed ratio of water production rate to oil production rate (WOR) as a function of pore volumes water injected (PVI) for a 3-D waterflood example. The fine scale solution (heavy line) is taken to be the true solution. The subgrid technique (dashed line) uses the fine scale permeabilities directly in coefficient $d$ in (30). The other cases replace $d$ with an average $\bar{d}$ in (30); the average is the arithmetic (dash-dot), harmonic in flow direction and arithmetic in the transverse direction (light line, lowest line near 0.2 PVI), or one-third power average (medium line, highest line near 0.2 PVI).
Fig. 5. Saturation distribution after 1000 days injection in a 2-D example based on the SPE comparative solution project. The solution was obtained on a fine $100 \times 20$ grid.

Fig. 6. Solution to the 2-D example of Fig. 5 after 1000 days injection obtained with the subgrid method on a $5 \times 5$ grid.

Fig. 7. Solution to the 2-D example of Fig. 5 after 1000 days injection obtained on a $5 \times 5$ grid, using the root mean square power average permeability in the subgrid method.
Fig. 8. Fine scale computation of the horizontal component of the total velocity after 1000 days injection in the 2-D example of Fig. 5.

Fig. 9. Computation of the horizontal component of the total velocity after 1000 days injection in the 2-D example of Fig. 5, computed on a $5 \times 5$ grid with the subgrid method.

Fig. 10. Computation of the horizontal component of the total velocity after 1000 days injection in the 2-D example of Fig. 5, computed on a $5 \times 5$ grid with the root mean square power average permeability in the subgrid method.
Table 1. Data for the 2-D quarter five-spot example.

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<td>Production rate</td>
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<td>Water relative permeability, ( k_{rw} )</td>
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</tbody>
</table>

Table 2. Summary of results for the 2-D quarter five-spot example.

<table>
<thead>
<tr>
<th>Case</th>
<th>Max. ( P ) error (psi) at 400 days</th>
<th>Max. ( P ) error (psi) at 1000 days</th>
<th>Breakthrough (days)</th>
<th>Cum. BBL oil at 1000 days (in 1000's)</th>
<th>% error</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 ( \times ) 30 Fine</td>
<td>—</td>
<td>—</td>
<td>290</td>
<td>46.8</td>
<td>—</td>
</tr>
<tr>
<td>6 ( \times ) 6 Subgrid</td>
<td>3.6</td>
<td>3.9</td>
<td>290</td>
<td>49.7</td>
<td>6.2</td>
</tr>
<tr>
<td>3 ( \times ) 3 Subgrid</td>
<td>23.7</td>
<td>24.7</td>
<td>310</td>
<td>56.2</td>
<td>19.9</td>
</tr>
<tr>
<td>6 ( \times ) 6 Avg. ( K )</td>
<td>56.6</td>
<td>53.4</td>
<td>246</td>
<td>58.1</td>
<td>24.0</td>
</tr>
<tr>
<td>6 ( \times ) 6 Avg. ( K ), order 2</td>
<td>52.3</td>
<td>49.1</td>
<td>246</td>
<td>57.0</td>
<td>21.8</td>
</tr>
</tbody>
</table>

Table 3. Single step timing results, in seconds, for the quarter five-spot 2-D example.

<table>
<thead>
<tr>
<th>Case</th>
<th>Pressure</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>subgrid solution</td>
</tr>
<tr>
<td>30 ( \times ) 30 Fine</td>
<td>—</td>
</tr>
<tr>
<td>6 ( \times ) 6 Subgrid</td>
<td>0.14</td>
</tr>
<tr>
<td>3 ( \times ) 3 Subgrid</td>
<td>0.74</td>
</tr>
<tr>
<td>6 ( \times ) 6 Avg. ( K )</td>
<td>—</td>
</tr>
<tr>
<td>6 ( \times ) 6 Avg. ( K ), order 2</td>
<td>—</td>
</tr>
</tbody>
</table>
Fig. 12. Producing water cut vs. time for the 2-D quarter five-spot numerical example.

Fig. 13. Cumulative oil production vs. time for the 2-D quarter five-spot numerical example.

Fig. 14. Fine 30 × 30 saturation contours for the 2-D quarter five-spot numerical example.

Fig. 15. Subgrid 6 × 6 saturation contours for the 2-D quarter five-spot numerical example.

Fig. 16. Subgrid 3 × 3 saturation contours for the 2-D quarter five-spot numerical example.

Fig. 17. Average permeability coarse 6 × 6 saturation contours for the 2-D quarter five-spot numerical example.