Upscaling transmissivity in the near-well region for numerical simulation: A comparison on uncertainty propagation

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Abstract

Upscaling transmissivity near the wellbore is expected to be useful for well performance prediction. This article ascertains the need of upscaling by comparing several numerical schemes and presents an approach to upscale transmissivity in the near-well region. This approach extends the Laplacian method with skin, which was successfully applied to the parallelepiped flow case, to radial flow case in the vicinity of wellbore through a nonuniform gridding technique. Several synthetic fields with different stochastic models are chosen to check the efficiency of this method. Both flow and transport simulations are carried out in finite heterogeneous confined aquifers to evaluate the results. It is demonstrated that the proposed method improves the ability of predicting well discharge or recharge and solute transport on the coarse scale in comparison with other schemes by examining the uncertainty propagation due to upscaling.

Keywords: Geostatistics, Well capture zone, Radial flow, Stochastic model, Reservoir simulation, Gridding
1 Introduction

Stochastic modeling of reservoir parameters with the aid of geostatistical techniques can effectively provide high-resolution models of reservoir at the measurement scale (Fu and Gómez-Hernández, 2008, 2009a). Limitation in computer resources forces these models to be upscaled to a modeling (coarse) scale such that a numerical simulator can afford in practical engineering applications of computational fluid mechanics, e.g., petroleum engineering, hydrogeology, environmental engineering, CO$_2$ sequestration, etc. (see Li et al., 2007; Fu, 2008; Jenny et al., 2003). A large number of upscaling approaches have been developed to coarsen detailed aquifer or reservoir models into those at an appropriate scale for numerical simulations (Wen and Gómez-Hernández, 1996; Renard and de Marsily, 1997). Many of them are proven quite efficient for upscaling under the uniform flow condition, where the local piezometric head or pressure values vary normally slowly, or say “linearly” (Durlofsky et al., 2000).

For the immediate vicinity of a well, however, these existing upscaling approaches may not completely apply due to the fact that the flow pattern is no longer uniform but convergent around a pumping well or divergent near an injection well. The pressure gradient typically increases close to the well and becomes highly sensitive to the spatial variation of hydraulic conductivity (Desbarats, 1992; Fiori et al., 1998) and especially to the difference between the global mean conductivity and the value at the wellbore (Axness and Carrera, 1999). Moreover, the distribution of concentration and the breakthrough curve of conservative tracers are typically different from those of the uniform flow cases. An effective upscaling scheme should be able to capture this character of pressure gradient distribution around the well and honor the statistical structure of conductivity in order to provide accurate coarse models. Basically, there are two problems needed to be addressed for upscaling: (1) Can the coarse grid account for the flow geometry in the near-well region? and (2) Can block transmissivity adequately honor heterogeneities of the aquifer or reservoir?

As for the first problem, several authors have already presented some approaches to address it. Ding (1995) proposed an upscaling procedure which consists of upscaling transmissivity and numerical productivity index. An obvious improvement over traditional methods is observed. Durlofsky et al. (2000) further extended Ding’s technique to the 3D case. Muggeridge (2002) assessed Ding’s method in a variety of case studies with partially penetrating wells and non-vertical wells of both two- and three-dimensional problems. Wolfsteiner and Durlofsky (2002) developed an upscaling approach for a near-well radial grid on the basis of the so-called multiblock-grid simulation technology. Such grid is globally unstructured but maintains locally structured. However, a drawback of them is that they use a regular coarse grid, either rectangular or almost rectangular, although a non-uniform grid is applied among them, e.g., Durlofsky et al. (1997), which is considered to be efficient for dealing with the case of connected region with high conductivity values. The influence of regular gridding lies in that, once the simulation grid becomes very coarse or the upscaling ratio is quite high, a significant or even intolerable loss of information will arise. That is because the
regular coarse grid has a rather limited flexibility to capture the feature of pressure gradient variation near the wellbore. An improvement in grid design and/or refinement is expected to enhance the accuracy of upscaling under the condition of preserving the coarsening ratio, or upscaled factor, so as to produce a coarsened model at a reasonable scale as the input to the flow simulator.

The second problem is concerned about the computation of the equivalent transmissivity for the coarse-scale model based on the fine-scale model. It is known that using the upscaled transmissivity for the coarse-scale flow simulation is much more accurate than the upscaled conductivity (e.g., Jenny et al., 2003); it is also well known that the block transmissivity is not only an intrinsic property of the porous media but also depends on the flow geometry and boundary conditions (e.g., Gómez-Hernández and Journel, 1994). Typical numerical approaches to computing the equivalent or block transmissivity call for solving flow problems over the local fine-scale block, which includes all the cells that are embedded in the corresponding coarse grid. One of the most striking considerations is the configuration of boundary conditions for the coarse grid (e.g., Jenny et al., 2003; Fu et al., 2010). This is of paramount importance because different boundary specifications will produce quite distinct equivalent block values. White and Horne (1987) computed the block conductivity from the solutions of flow for several alternative boundary conditions. Gómez-Hernández and Journel (1994) applied a skin surrounding the coarse grid as the boundary configuration, which accounts for the influences from the neighbor cells but without resorting to solve flow problems over the entire field. Durlofsky (1991) developed an approach to yield a set of symmetric, positive-definite block conductivity tensors by applying periodic boundary conditions. All these methods assume that the block transmissivity is a local or extendedly local property of porous media, i.e., the effects of neighbors can be ignored. Global approaches, on the other hand, consider such influences by solving flow problems on the global fine scale, e.g., Holden and Nielsen (2000). However, this method may be extremely computationally expensive. More recently, to overcome this shortcoming, Chen et al. (2003) developed a technique that couples local and global approaches with the aid of iterative solutions to flow and/or transport problems on the global coarse scale and the local fine scale.

In addition, an extra technical detail is how to choose a neighbor size for solving local flow problems with specified boundary conditions. That is, should the neighbor cells be included when solving local fine-scale flow problems? Mascarenhas and Durlofsky (2000) proposed a near-well upscaling method extending the local fine regular grid to include neighbor regions when the transmissivity tensors are computed for a coarse regular grid, while a traditional upscaling procedure does not do so. Their numerical results from single- and multi-phase flow experiments display a significant improvement compared to the conventional methods by analyzing inflow profile and water cut parameters based on the fine-scale simulation against those based on the coarse-scale simulation. Actually, a similar enhancement was observed when upscaling transmissivity for the uniform flow, where the Laplacian method with skin (Gómez-Hernández and Journel, 1994) or border region (Wen et al., 2003; Chen et al., 2003) is named. In the present study, we include this idea into our upscaling approach with a slight
modification to the coarse radial grid.

This paper proceeds as follows: the next section gives details of the proposed upscaling method. Then, we outline the assessment criteria for subsequent comparisons and the numerical methods for flow and transport simulations. In the fourth section, the numerical results are compared with several existing upscaling techniques. It follows by some discussion on the upscaling techniques. Finally, the paper ends up with a summary of the proposed method for upscaling transmissivity in the near-well region.

2 Upscaling

An upscaling procedure in numerical simulation typically consists of three steps: first, using geostatistical techniques, a series of fine, detailed model parameters (i.e., hydraulic conductivity) are generated, each representative of the geology and hydrology of the area. Then a coarse grid is designed to capture main characteristics of flow and transport accordingly at the coarse scale. Finally, an equivalent value, either scalar or vectorial, calculated from the fine model of scalar parameters is assigned to the coarse model.

2.1 Generation of conductivity fields at the fine scale

The currently existing geostatistical techniques allow for generating property fields of reservoir at a point scale or for a uniform grid. The former can generate parameter fields of reservoir at any location for an arbitrary grid. This is a simple but fast scheme since only a small quantity of data are necessary for the near-well region. Moreover, it circumvents the scaling problem. The deficiency, however, lies in that it can not capture the detailed spatial variation of reservoir parameters. The latter, on the contrary, generates data at the specified position in a regular grid frame. This requires a very fine grid for the entire field such that the spatial fluctuations in the near-well region are adequately represented. However, it inevitably brings up a scaling problem in order to produce a proper coarse model as input to the expensive flow simulator. For the purpose of checking the necessity of upscaling in the near-well region, we compare these two types of techniques subject to an identical flow and transport scenario.

The sequential simulation algorithm is a powerful stochastic simulation technique and has been applied in many studies (Gómez-Hernández and Journel, 1993). It can be used to generate conditional or unconditional realizations from either multi-Gaussian or non-Gaussian random functions. Following the two-point geostatistics, the multi-Gaussian hydraulic conductivity field, \( \ln K(x) \), is modeled through a normally distributed random space function, \( Y(x) \equiv \ln K(x) \), with an exponential semivariogram specified by,

\[
\gamma_Y(r) = \sigma_Y^2 \left\{ 1 - \exp \left( -\frac{r}{\lambda_Y} \right) \right\},
\]

(1)
where $r$ is the two-point separation distance, $\sigma_Y^2$ is the variance, and $\lambda_Y$ is the correlation length. For the non-Gaussian model, the indicator semivariogram for a continuous variable is defined as,

$$\text{ind}_i = \begin{cases} 1 & \text{if } Y_i \leq \text{cut}_k; \\ 0 & \text{otherwise,} \end{cases}$$

where the subscript $i$ refers to a specific location, and the cutoff $\text{cut}_k$ is a threshold that is specified for the $k$th class of the continuous variable $Y$ to create the indicator transform.

Once the semivariogram model is specified, the corresponding covariance is easily obtained through $C(r) = C(0) - \Gamma(r)$ if the lag effect is ignored and the stochastic samples may be efficiently generated by the sequential simulation approach. The public domain codes GCOSIM3D (Gómez-Hernández and Journel, 1993) and ISIM3D (Gómez-Hernández and Srivastava, 1990) are used to generate hydraulic conductivity fields at the fine scale. Figure 1 gives three typical realizations of log-conductivity field with square cells 801×801: Figure 1(A) a multi-Gaussian field with statistically isotropic structure, Figure 1(B) a multi-Gaussian field with statistically anisotropic structure, and Figure 1(C) a non-Gaussian field with statistically anisotropic structure. Each cell has a dimension of 0.25×0.25. The wellbore is assumed to locate at the center of computational domain, i.e., with the planar coordinate index (401, 401). We assume that the property value at the wellbore center is known, so the generation of log-conductivity field belongs to a conditional simulation.

Due to the limitation of GCOSIM3D, however, the existing code has no ability of generating data with arbitrarily irregular grids although the sequential simulation algorithm maybe allow one to do so in theory. But we can address this problem by resampling from the fine-scale field. That is, we first generate a fine-scale field by assuming a statistical structure, and then resample the property values from this field and assign them to proper locations in the irregular grid. The resampled field shares the same statistical structures as that of the fine scale, even though it ignores some details of the spatial variations. One of the advantages of this way is that its results can be used to compare with those of responding field directly. We denote it as a non-upscaled field.

### 2.2 Design of the coarse grid

The Thiem solution to a 2D steady-state head field in a homogeneous medium with prescribed heads at the well radius and at an exterior circular boundary can be written as,

$$h(r) = h_w + \frac{h_e - h_w}{\ln (r_e/r_w)} \ln (r),$$

where $r$ represents the normalized radius $r = r'/r_w$; $r_w$ is the wellbore radius; $r'$ is the radius away from the well axis; $h_e$ and $h_w$ are the heads at the outer and inner radii, respectively. Although this solution is only applicable for homogeneous media, it may be shown that gridding with respect to $\ln (r)$ other than $r$ minimizes the error in the resulting hydraulic
Figure 1: Several typical realizations of $\ln K(x)$ field: (A) a multi-Gaussian field with isotropic structure, (B) a multi-Gaussian field with anisotropic structure, and (C) a non-Gaussian field with anisotropic structure.
head (Axness et al., 2004). The coarse grid design in terms of log-scale, therefore, will have more advantages than that of the natural scale. The former is expected be able to capture main features of gradient variations better than the latter even for highly heterogeneous media.

In this study, we first normalize the well radius to one unit, i.e., $r_w = 1$, while the exterior radius is equal to 100 units, i.e., $r_e = 100$. The whole circular field is divided into ten annuli excluding the well block. The ratio of radius increment of every annulus compared to that of the previous inner annulus is 1.584893, i.e., $\frac{\Delta r_{i+1}}{\Delta r_i} = 1.584893$. Those rings are further divided into twelve segments along the ray direction. This ensures the grid very fine close to the wellbore but quite coarse far away from it (see Figure 2(A)). The original $801 \times 801$ fine square grid is upcaled into a $12 \times 10$ coarse radial, circular grid. In order to validate the efficiency of this grid design, we have compared its results with those of equal radius increment (Figure 2(B)).

![Figure 2: Geometry of the coarse grid: (A) non-equal radius increment and (B) equal radius increment](image)

2.3 Computation of equivalent transmissivity

We extend the concept of skin (Gómez-Hernández and Journel, 1994) to the computation of the equivalent transmissivity for the coarse grid blocks. Two sets of boundary conditions are considered for each coarse block: one is in the $x$ direction and the other in the $y$ direction (Figure 3). By doing so, we can obtain $T_{xx}$ and $T_{yy}$ correspondingly through solving (extendedly) local flow problems at the fine scale. In addition, one of the crucial problems is the configuration of boundary conditions for each coarse rectangular block. We approximate the head values for each side by assigning a Thiem solution to the middle point of each edge such that the computationally demanding global problem for boundary condition configuration is avoided but a relatively accurate result is still achieved. The method developed here closely follows the seminal idea of the work that was presented in a conference paper (Fu et al.,...
Figure 3: Configuration of boundary conditions for the coarse grid: (A) $T_{V,xx}$, (B) $T_{V,yy}$.

The procedure to calculating the equivalent transmissivity is as follows (Figure 3):

- define a rectangular block $R$ that includes the non-rectangular target block $B$;
- solve the flow problem with a specified boundary condition system which consists of non-flow boundary and prescribed head. The prescribed heads are approximated by the analytical solution for the homogeneous media as described as above;
- evaluate the average flow rate $Q_V$ and the average head gradient $\Delta h_V$ over the non-rectangular target block $B$ both in $x$ and $y$ directions;
- compute the equivalent transmissivity by,

$$T_{V,xx} = -\frac{Q_{V,xx}}{\Delta h_{V,xx}}, \quad (4)$$
$$T_{V,yy} = -\frac{Q_{V,yy}}{\Delta h_{V,yy}}. \quad (5)$$

3 Numerical Simulation

We choose two types of metrics for assessing upscaled results: one is water injections (or, similarly, yields for a pumping well) into the wellbore, and the other is travel time of conservative tracers. The former are achieved by calculating the flow rate in the wellbore after solving flow equations. The latter can be accomplished by solving transport equations with the aid of the random walk particle tracking method.
3.1 Flow and transport simulation

The flow problem is solved by tailoring the block-centered finite-difference simulator to the radial flow case (Fu, 2008). To avoid the discretization error, all the simulations are performed at the fine scale. The transmissivity at the interface between cells are computed using harmonic averages of the adjacent blocks. We model the radial flow to a well by specifying fixed heads at the wellbore and at the exterior circular boundary, i.e., \( h_w = 10 \) and \( h_e = 0 \), respectively. It models the case of an injection well or source configuration since the pressure at wellbore is higher than that of the surroundings.

We prefer an injection well to a pumping well on the basis of the consideration that the latter will have a rather lower particle capture rate when we employ the particle tracking scheme to solve transport problems, which will undermine the reliability of solutions. Our experience shows that characteristics of flow and transport around an injection well have no obvious difference than those surrounding a pumping well. Second, we intentionally impose a prescribed head boundary at the exterior radius since we reason that its influence is not significant when the outer radius is placed a few correlation lengths, e.g., ten as in this study, away from the well (Arness and Carrera, 1999). Finally, we choose a constant head at the wellbore as an evaluation criterion not only because it is widely used in the engineering practice but also because it is the easiest way to modeling the steady flow.

The transport equation solver adopts the random walk particle tracking scheme developed by Fu (2008). Two thousand particles released from the wellbore are followed until they arrive at the control circle, which is set 100 units away from the wellbore. Because the pressure near the wellbore is higher than that far away, it ensures that all particles move from the wellbore to the exterior boundary. It evidently overcomes the difficulty of rather lower particle capture rate, which is common in employing the particle tracking scheme to solve the transport equations. By doing so, a stable solution can be obtained.

3.2 Computation of well discharge/recharge

In this study, the well injection is calculated by numerically integrating Darcyian velocity along the surface surrounding the wellbore,

\[
Q_w = \int_0^{2\pi} q(\theta) d\theta,
\]

where \( q(\theta) \) is the component of Darcyian velocity around the wellbore. To ease the complexity of problems, we assume that property values at the wellbore are the same as those at the well block.

3.3 Computation of travel time

Well discharge and recharge can reasonably reflect an average effect of heterogeneities in porous media, but they can not sufficiently display the detailed variation of heterogeneity.
Travel time of passive tracers computed by the particle tracking scheme has a more powerful ability to do so. This is because the evolution of a solute plume is more sensitive to fluctuations of the hydraulic conductivity and head fields. There exist two ways to evaluate the upscaling approaches in terms of travel time: (1) compare travel time of individual realizations to check their accuracy; and (2) compare an ensemble of realizations to check their robustness. We use both of them in this work for comparison. Due to the difficulty of comparing the whole breakthrough curve for all realizations, we only sample several typical points from the breakthrough curve (BTC), e.g., $t_{5\%}$, $t_{25\%}$, $t_{50\%}$ and $t_{95\%}$, which account for the early, middle and late arrival time.

It is worth emphasizing that the selection of the most appropriate part of BTC is of importance in engineering environmental operations, e.g., to monitor the extent and degree of groundwater contamination from a known source (Fu and Gómez-Hernández, 2009b). $t_{5\%}$ better reproduces the earliest part of the reference BTC and represents the fastest particles arriving to the control plane as needed for the design of radioactive underground repository. The earliest arrivals in the BTC follow the fastest pathways between the release source and the control plane, which are dominated by preferential flow and reactive transport paths. Failing to account for such case will yield a too conservative conclusion in risk analysis in that the real arrival time may be too much faster than that estimated (Gómez-Hernández and Wen, 1998). $t_{25\%}$ and $t_{75\%}$ reproduce the middle part of the reference BTC and reflect the portion of particles arriving to the control plane with high frequency. Public officials assessing health risks associated with contaminant exposure in a drinking water supply system may be most concerned with this parameter. $t_{95\%}$ reproduces the tail part of the reference BTC and denotes the slowest particles arriving to the control plane as needed for mass removal calculations in remediation problems. The late travel times reflect a more integral behavior, or even flow and reactive transport barriers. An aquifer remediation design without considering such feature may fail because the resident contaminants will be removed more slowly than expected (Wagner and Gorelick, 1989).

4 Results

Four types of coarsening techniques are assessed in this section: (a) the proposed scheme as previously stated (named as proposal), (b) traditional geometric average from the fine scale in the framework of non-uniform grid (called as GM), (c) non-upscaled method (noted as NP), and (d) the geometric average from the fine scale in the framework of uniform grid (shorted as UG). The second one, GM, assigns the coarse conductivity to be the geometric mean of support cells contained in the coarse element. This is the simplest and traditional method for upscaling uniform flow. We present the results here with two aims: one is to check its efficiency under the radial flow conditions, and the other is to compare its results with the proposed approach. The third one, NP, simply assigns the conductivity in the coarse element to be the point value at the coarse element centroid or to be the conductivity of the support scale element closest to the centroid when the field is generated at a fine support
scale. The last technique, UG, is done the same way as the second but only distinctly with uniform grid intervals.

Flow and transport results from four types of techniques are presented in three scenarios: the first is multi-Gaussian conductivity fields with isotropic structures; the second is multi-Gaussian conductivity fields with anisotropic structures; and the last one is non-Gaussian conductivity fields with anisotropic structures.

4.1 Multi-Gaussian fields with isotropic structures

One hundred \(\ln K(x)\) fields are generated by GCOSIM3D, each of them with a correlation length equal to ten in both \(x\) and \(y\) directions, i.e., \(\lambda_x = \lambda_y = 10\), and the expected variances more than one, i.e., \(\sigma^2_{\ln K(x)} = 2\). The \(\ln K(x)\) fields are scalar, that is, the conductivity value in the \(x\) direction are the same as those in the \(y\) direction, i.e., \(\ln K_{xx} = \ln K_{yy}\). The mean of \(\ln K(x)\) is set to zero, i.e., \(E[\ln K(x)] = 0\). The flow rates of the wellbore are computed on the fine scale named as reference values. Then, the fine-scale fields with 801\(\times\)801 grids are upscaled to those with 12\(\times\)10 coarse grids in the same way as described above. The fluxes over the wellbore are computed at the coarse scale for four different coarsening approaches, named as proposal, GM, NP and UG, respectively.

Figure 4 shows the relationship of wellbore fluxes between the reference fluxes and those using the four different approaches. Figure 4(A) displays one hundred wellbore fluxes by the proposed upscaling approach compared to those of reference fine scales. The average flux of one hundred realizations is 17.537 for reference fields and 17.012 for the proposal method with relative error only 3%. It shows that the upscaled values are a rather reasonable approximate. Note, moreover, that the \(x\) and \(y\) mean values are close to each other, meaning that this method is unbiased, that is, the upscaled \(Q\) tends to be close to the reference \(Q\) in the mean. Figure 4(B) shows the performance of the traditional geometric mean method. The correlation and rank correlation coefficient have a slight decrease but still more than 99%. It seems that the geometric mean method is quite efficient and robust in upscaling radial flow in the near-well region for a scalar log-conductivity field with isotropic structures. Figure 4(C) plots the result of the NP method. The correlation coefficient is only 73%. Obviously the reproduction ability is worse than the GM method. Figure 4(D) is the result of the UG method, i.e., that with a uniform interval grid. The wellbore fluxes calculated from the coarse grid are severely deviated from those of the fine grid. Although the correlation coefficient arrives to 88%, the mean of wellbore fluxes has a rather high error up to 20%. Such apparent error is intolerable for the computation of well yields.

The similar average effect of upscaled heterogeneities can be observed from the mean breakthrough time after solving solute transport problems. Figure 5 plots the average breakthrough times of different coarse fields versus those of fine reference fields. Figure 5(A) shows the average breakthrough times of one hundred realizations calculated by the proposed upscaling approach. Their correlation coefficient with those of reference fields is up to 99% and the relative average error is only 1.3%. The GM method also provides rather acceptable
Figure 4: Wellbore fluxes cross relationship between the fine scale and the coarse scale of multi-Gaussian $\ln K(x)$ fields with isotropic structures
results as shown in Figure 5(B). The relative average error is 1.2% and the correlation coefficient is a little bit worse but still more than 98%. The NP method produces unpleasant results as shown in Figure 5(C). Although the average error is quite low only with 1% owing to the randomness of sampling, the solutions are too unstable: the correlation coefficient is only 68%. The UG method has the results with a better correlation coefficient, almost of 85%, but the average error up to 16%. In summary, the first two upscaling schemes produce quite satisfactory results.

Figure 5: Average breakthrough time cross relationship between the fine scale and the coarse scale of multi-Gaussian ln$K(x)$ fields with isotropic structures

Unlike the well discharge or recharge and the average breakthrough time, which are only comprehensive effects of field fluctuations, a whole breakthrough curve can provide a better review of upscaling results because it can adequately sample the spatial variation of heterogeneous fields. Figure 6 gives a comparison of five breakthrough curves from a typical realization. The results from the NP and UG methods obviously deviate from that of the reference fine scale. The proposed approach has a slightly better capability of reproduction than the GM method. For this single realization, the proposed method produces better results at the early and middle arrival time.
Figure 6: Typical breakthrough curves of the fine scale and the coarse scales of multi-Gaussian \( \ln K(x) \) field with isotropic structures
In order to further inspect the stability of the proposed approach in reproducing the breakthrough curve of reference fields, we sample four typical points, i.e., \( t_{5\%}, t_{25\%}, t_{75\%}, \) and \( t_{95\%} \), from the breakthrough curve and compare them with those of the GM method. Figure 7 illustrates the comparison of the breakthrough curve matching with reference fields between these two methods. The left column plots the matching of the proposed method from all one hundred realizations, and those of the GM method are listed in the right column. As for the average breakthrough error, the proposed approach gains some advantage over the GM method: almost all four sample points of the former have closer values to the reference ones than those of the latter. Moreover the stability of the latter has no diminishment in general; their correlation coefficients maintain about 90%.

4.2 Multi-Gaussian fields with anisotropic structures

One hundred \( \ln K(x) \) fields are generated by GCOSIM3D, each of them with a correlation length equal to ten in the \( x \) direction and five in the \( y \) direction, i.e., \( \lambda_x = 10 \) and \( \lambda_y = 5 \). Their expected variances are kept more than one, i.e., \( \sigma^2_{\ln K(x)} = 2 \). The \( \ln K(x) \) fields are constant vectorial, that is, the \( \ln K(x) \) values in the \( y \) direction are one half of those in the \( x \) direction, i.e., \( \ln K_{yy} = \frac{1}{2} \ln K_{xx} \). The mean of \( \ln K(x) \) is set to zero, i.e., \( E[\ln K(x)] = 0 \).

The fluxes over the wellbore are computed for the fine-scale field and four different coarse fields in the same manner as in the first scenario.

Figure 8 shows the cross relationship of the wellbore fluxes between the reference and four different upscaling approaches. Figure 8(A) exhibits one hundred wellbore fluxes via the proposed upscaling approach with comparison to those of the reference fine scales. The average flux of one hundred realizations is 12.469 for reference fields and 12.077 for the proposed method with a relative error only three percent. It shows that the upscaled values are quite close to the actual case. Moreover, the correlation and rank correlation coefficient between two types of fluxes are more than 99%, which demonstrates the robustness of this approach. Figure 8(B) indicates the performance of the traditional geometric mean method. The correlation and rank correlation coefficient have a little decrease but still close to 99%. Figure 8(C) plots the result of the NP method. The correlation coefficient is only 71%. Figure 8(D) is the results of the UG method, which obviously underestimates the actual values. The mean of wellbore fluxes has a rather high error up to 19%, although the correlation coefficient reaches 88%.

Figure 9 plots the average breakthrough times of different upscaled fields versus those of reference fields with anisotropic structures. Figure 9(A) shows the average breakthrough times of one hundred realizations from the proposed upscaling approach. The correlation coefficient with those of the reference fields is up to 99% and the relative average error is only 1.3%. The GM method also provides acceptable results as shown in Figure 9(B). The relative average error is 1.2% and the correlation coefficient is a little bit worse but still more than 98%. The NP method produces unpleasant results as in Figure 9(C). Although the average error is only 1% owing to the randomness of sampling, the stability is too low: the
Figure 7: Comparison of breakthrough curve reproduction of multi-Gaussian $\ln K(x)$ fields with isotropic structures: (A) the proposed method (left column), (B) the geometrical mean (right column).
Figure 8: Wellbore fluxes cross relationship between the fine scale and the coarse scale of multi-Gaussian $\ln K(x)$ fields with anisotropic structures.
correlation coefficient is only 68%. The UG method has results with correlation coefficient better, almost of 85%, but with the average error up to 16%. The last two methods are obviously unsuccessful in reproducing the average breakthrough time.

Figure 9: Average breakthrough time cross relationship between the fine scale and the coarse scale of multi-Gaussian \( \ln K(\mathbf{x}) \) fields with anisotropic structures

Figure 10 gives a comparison of the all five breakthrough curves from a typical realization. The NP method fails to reproduce the result of the reference field. The proposed method has a better result than the GM method, especially at the late arrival time for this typical realization.

Figure 11 shows the comparison of the breakthrough curve matching with reference fields between the proposed method and the GM method. As for the average breakthrough error, the proposed approach has a quite noticeable gain over the GM method. All four sample points of the former have closer values to the reference ones than those of the latter. The error reduction is 16.6% for the first point \( (t_{5\%}) \), 39.9% for the second \( (t_{25\%}) \), 25.6% for the third \( (t_{75\%}) \), and 51.4% for the fourth \( (t_{95\%}) \). This enhancement is not observed so clearly as in the case of isotropic structure. Moreover the stability of the proposal has no decrease. The correlation coefficients are over 90% for all sample points. This is true especially for
Figure 10: Typical breakthrough curves of the fine scale and the coarse scales of multi-Gaussian $\ln K(\mathbf{x})$ field with anisotropic structures
reproducing the early arrival particles.

4.3 Non-Gaussian fields with anisotropic structures

One hundred $\ln K(x)$ fields are generated by ISIM3D, each of them with a correlation length equal to ten in the $x$ direction and three in the $y$ direction, i.e., $\lambda_x = 10$ and $\lambda_y = 3$. The mean and variance of $\ln K(x)$ is set to zero and two, i.e., $E[\ln K(x)] = 0$ and $\sigma^2_{\ln K(x)} = 2$. The $\ln K(x)$ fields are constant vectorial, that is, the $\ln K(x)$ values in the $y$ direction are one half of those in the $x$ direction, i.e., $\ln K_{yy} = \frac{1}{2} \ln K_{xx}$. The fluxes over the wellbore are computed for the fine-scale field and four different coarse fields in the same manner as in the first scenario.

Figure 12 shows the cross relationship of wellbore fluxes between the reference and four different upscaling approaches. Figure 12(A) exhibits one hundred wellbore fluxes via the proposed upscaling approach with compared to those of reference fine scales. The average flux of one hundred realizations is 43.976 for reference fields and 35.292 for the proposed method with relative error 19.7%. The correlation and rank correlation coefficient between two types of fluxes are more than 99%, which demonstrates the robustness of this approach. Figure 12(B) indicates the performance of the traditional geometric mean method. The correlation and rank correlation coefficient have a little decrease but still close to 99%. Figure 12(C) plots the result of the NP method. The average flux of one hundred realizations is 43.976 for reference fields and 42.483 for the proposed method with relative error 3%. The correlation coefficient is still up to 90%. Figure 12(D) is the results of the UG method which distinctly underestimates the actual values.

Figure 13 plots the average breakthrough times of different upscaled fields versus those of reference fields with anisotropic structures. Figure 13(A) shows the average breakthrough times of one hundred realizations from the proposed upscaling approach. The correlation coefficient with those of the reference fields is up to 97% and the relative average error is only 1.4%. The GM method also provides acceptable results as shown in Figure 13(B). The relative average error is 9.3% and the correlation coefficient is a little bit worse but still more than 97%. The NP method produces unpleasant results as in Figure 13(C). The UG method has the results almost the same as the NP method. The last two methods are obviously unsuccessful in reproducing the average breakthrough time.

Figure 14 gives a comparison of the all five breakthrough curves from a typical realization. Due to the peculiarity of non-Gaussian field, there is no upscaling method capable of reproducing the reference field completely. The results of four coarse grids, as shown in this typical realization, obviously deviate from the reference fine grid. But generally, the proposed method produces a better result than the others.

Figure 15 shows the comparison of the breakthrough curve matching with reference fields between the proposed method and the GM method. As for the average breakthrough error, the proposed approach has a quite noticeable gain over the GM method. All four sample points of the former have closer values to the reference ones than those of the latter. The
Figure 11: Comparison of breakthrough curve reproduction of multi-Gaussian ln$K(x)$ fields with anisotropic structures: (A) the proposed method (left column), (B) the geometrical mean (right column).
Figure 12: Wellbore fluxes cross relationship between the fine scale and the coarse scale of non-Gaussian lnK(x) fields with anisotropic structures
Figure 13: Average breakthrough time cross relationship between the fine scale and the coarse scale of non-Gaussian $\ln K(x)$ fields with anisotropic structures
Figure 14: Typical breakthrough curves of the fine scale and the coarse scales of non-Gaussian ln$K(x)$ field with anisotropic structures
error reduction is 47% for the first point ($t_{5\%}$), 63.3% for the second ($t_{25\%}$), 37.3% for the third ($t_{75\%}$), and 83.4% for the fourth ($t_{95\%}$). This enhancement is more obvious than that of multi-Gaussian fields.

5 Discussion and Conclusions

This paper compares two types of geostatistical techniques for generating hydraulic conductivity fields at the near-well region by assuming spatial structures. One creates property fields directly on the coarse scale and the other reconstructs such fields via upscaling from the corresponding fine scale. The results from flow and transport simulations demonstrate that the directly generated field can not effectively capture the spatial variations of hydraulic conductivity while the upscaled field can do. Therefore, the requirement of upscaling in the near-well region is apparent.

Previous studies indicate that the flow to a well is strongly influenced by local conditions around the wellbore and is less influenced by fluctuations far away from the well, e.g., Sanchez-Vila et al. (1997) and Axness and Carrera (1999). At the radial flow zone near the wellbore, therefore, one should adopt a different upscaling scheme than that of the linear flow zone far away from the well in order to correctly answer this response. This study shows that the non-uniform radial coarse grid can effectively capture this character.

We further extend the upscaling approach of Gómez-Hernández and Journel (1994), which was proved efficient under the uniform flow condition, to the convergent or divergent flow case by designing proper radial grids that sufficiently account for the flow patterns in the near-well region. The proposed scheme gains some improvement over the simple geometric mean. Several stochastic models with a single well system are chosen to illustrate the efficiency and robustness of this method.

Numerical simulations from mass transport demonstrate that the simple geometric mean can reasonably reproduce the results of the reference fine scale for multi-Gaussian models, either statistically isotropic or weakly anisotropic. But it fails to do so for non-Gaussian models. The multi-Gaussian model implies the minimal spatial correlation of extreme values, which is critical for mass transport and may be in contradiction with some geological reality, e.g., channeling. Connectivity patterns of extreme conductivity values can not be represented by a multi-Gaussian model. Gómez-Hernández and Wen (1998) proved that the groundwater travel time predicted by the multi-Gaussian model could be ten times slower than that by non-Gaussian models. The reason is that, for a non-Gaussian model, the simple geometric mean weakens the heterogeneity of conductivity field while the proposed upscaling method effectively preserves such extreme values, with high connectivity either at the extremely high values or at the extremely low values, by solving local flow problems. On the other hand, for a multi-Gaussian model where the spatial variability is not so high as in the non-Gaussian model, the simple geometric mean can produce quite similar results as the proposed approach. We notice that the proposed method has not sufficiently reproduced the result of reference field. A promising improvement is to use full tensorial transmissivity fields, i.e.,
Figure 15: Comparison of breakthrough curve reproduction of non-Gaussian ln$K(x)$ fields with anisotropic structures: (A) the proposed method (left column), (B) the geometrical mean (right column).
with the introduction of $T_{V,xy}$ and $T_{V,yx}$.

Finally, several conclusions from this study are worth repeating as follows: (1) Upscaling transmissivity in the near-well region can efficiently preserve the main features of flow and transport in the heterogeneous media. (2) The proposed method improves the ability of predicting well discharge or recharge and solute transport in terms of the coarse grid. Several synthetical examples prove that the proposed upscaling approach is efficient and robust both for flow simulation and for transport simulation. (3) The geometric mean of log conductivity is an alternative approach in upscaling transmissivity in the near-well region. This is true especially when the log-conductivity is a multi-Gaussian field. (4) Uniform grid fails to capture the flow and transport features in the near-well region.

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