Density Functional Hydrodynamics in Multiscale Pore Systems: Chemical Potential Drive

Oleg Dinariev1, Nikolay Evseev1,*, and Denis Klemin2

1 Schlumberger, Schlumberger Moscow Research, 113, Pudovkina Street, Moscow 119284, Russia
2 Schlumberger, Schlumberger Reservoir Labs, 6350 West Sam Houston Pkwy N, Houston, TX 77041, Houston, USA

Abstract. We use the method of density functional hydrodynamics (DFH) to model compositional multiphase flows in natural cores at the pore-scale. In previous publications the authors demonstrated that DFH covers many diverse pore-scale phenomena, starting from those inherent in RCA and SCAL measurements, and extending to much more complex EOR processes. We perform the pore-scale modelling of multiphase flow scenarios by means of the direct hydrodynamic (DHD) simulator, which is a numerical implementation of the DFH. In the present work, we consider the problem of numerical modelling of fluid transport in pore systems with voids and channels when the range of pore sizes exceed several orders of magnitude. Such situations are well known for carbonate reservoirs, where narrow pore channels of micrometer range can coexist and interconnect with vugs of millimeter or centimeter range. In such multiscale systems one cannot use the standard DFH approach for pore-scale modeling, primarily because the needed increase in scanning resolution that is required to resolve small pores adequately, leads to a field of view reduction that compromises the representation of large pores. In order to address this challenge, we suggest a novel approach, in which transport in small-size pores is described by an upscaled effective model, while the transport in large pores is still described by the DFH. The upscaled effective model is derived from the exact DFH equations using asymptotic expansion in respect to small-size characterization parameter. This effective model retains the properties of DFH like chemical and multiphase transport, thus making it applicable to the same range of phenomena as DFH itself. The model is based on the concept that the transport is driven by gradients of chemical potentials of the components present in the mixture. This is a significant generalization of the Darcy transport model since the proposed new model incorporates diffusion transport in addition to the usual pressure-driven transport. In the present work we provide several multiphase transport numerical examples including: a) upscaling to chemical potential drive (CPD) model, b) combined modeling of large pores by DFH and small pores by CPD.

1 Introduction

In recent years we witnessed an increasing level of capability and acceptance of digital rock as a complementary tool in core analysis. This tendency is following a gradual increase in both resolution of rock imaging techniques and the power of high-performance computing that are the key to unlock the potential in the core analysis by digital rock. As an answer to the growth in the hardware possibilities various modeling methods applicable in digital rock also evolve; see reviews in [1-6]. One of such methods, which the authors are actively developing, is the density functional hydrodynamics (DFH).

DFH is essentially a pore scale multi-phase, multi-compositional approach. As it has been demonstrated in previous publications [7-10], various multiphase compositional problems can be described in the frame of the DFH. The DFH method has found many applications in direct pore-scale modeling of various hydrodynamic and petrophysical phenomena on digital rock models obtained by X-ray micro-CT and SEM [11-21]. The practical usefulness of a standard workflow is very much dependent on the possibility to resolve necessary pore structure adequately, as well as on the rock sample being representative for the considered formation. There are specific ways to meet both of these requirements, but in this work, we discuss the resolution problem only.

In many cases the natural rocks are characterized by a wide range of pore sizes making the choice of X-ray micro-CT resolution very difficult if not altogether impossible. Indeed, if pores of both micrometer and millimeter range coexist within one piece of rock then, at present, it is not always possible to develop 3D model of rock microstructure, which can represent all pores. In situations like this we propose a combined approach, when several 3D digital rock models having different resolutions are integrated into one synthetic model. Combined multiscale approach is also being developed in the frame of pore-network modelling [1], but we follow image-based approach as it is inherent in DFH. High-resolution sub-models with explicitly resolved
small pores are used for direct modeling of transport processes by DFH to obtain effective flow characteristics. Then these results are incorporated into low-resolution model with explicit imaging of large pores while small pores are not necessarily resolved. At this stage the hydrodynamics in large pores is still simulated directly by DFH, while the transport in matrix is simulated using the previously computed effective transport properties. All these operations are done by the DFH pore-scale simulator called DHD (Schlumberger).

In effect, the described procedure represents upscaling from pore-scale DFH equations to macroscopic porous medium equations. As concerned with the single-phase flow, our idea is similar to the one described in [22, 23]. However, in addition to being applicable in general multiphase compositional scenarios, the rigorous upscaling starting with the DFH equations leads to the transport equations, which constitute the significant extension of the Darcy model. The driving force for the fluid flow becomes gradients of chemical potentials of fluid chemical components that gives the model its name Chemical Potential Drive (CPD). Because change in pressure is directly related to change in chemical potential, this new model is consistent with pressure-drive Darcy approach as will be shown below. On the other hand, the CPD approach covers certain physical phenomena outside the scope of the Darcy model. Here are two examples in relation to the latter assertion. First, there are cases, when definition of pressure in confined fluid is problematic, e.g., when gas mean free path length is comparable to pore size (high Knudsen number), or liquid in pores with significant disjoining pressure; at the same time, the definition of chemical potential is still correct in both said instances. Second, there is an easy to derive exact solution for two-phase water-oil transport velocities in a hydrophilic circular capillary

\[
    u_w = -\frac{8\eta_w}{R} R^2 (1 - \frac{r}{R})^2 G
\]

\[
    u_o = -\frac{8\eta_o}{R} R^2 (2\frac{r}{R})^2 (1 - \frac{r}{R})^2 G
\]

\[
    (\frac{\eta_w}{\eta_o} + \frac{r}{R})^4 G
\]

where the water and oil velocities \( u_w, u_o \) are defined as phase fluxes per capillary cross-section, \( \eta_w, \eta_o \) are the shear viscosities of water and oil, respectively, \( R \) is the capillary radius, \( r \) is the radius of the central region occupied by oil, and \( G \) is the pressure gradient. An important observation following from Eqs. (1), (2) is that the transport of oil is influenced by viscosity of water, because oil, being the central phase, is carried by water. This cross-phase fraction is in direct contradiction with the routine concept of phase permeabilities indicating the possible role of cross-terms in phase permeability matrix [24]. At the same time the cross-component influence is inherent in CPD, as it will be demonstrated below.

Limitations of the Darcy law are discussed in the literature and deviations from it are observed in both high-accuracy 4D visualization of pore-scale fluid dynamics as well as in numerical simulations of pore-scale flows [25]. One may expect more deviations when dealing with complex fluids.

In Sec. II we give a brief reminder of the DFH equations, as well as a derivation of upscaling from DFH to CPD. Numerical demonstration of combined DFH and CPD models is presented in Sec. III. The overall summary and discussion of results is in Sec. IV.

The summation over repeated indices is implied everywhere. The indices \( a, b, c = 1, 2, 3 \) are related to Cartesian coordinates \( x^a \), the indices \( i, j, k = 1, ..., M \) are related to chemical components in the fluid mixture. We consider isothermal processes, so the temperature is assumed to be fixed, and dependence of certain variables on temperature is omitted. We use short symbols for partial derivatives: \( \partial_a = \partial / \partial x^a \) and \( \partial_t = \partial / \partial t \), where \( t \) is time.

### 2 Theoretical concepts and equations

#### 2.1 Density functional hydrodynamics

Here we provide only some of the basic definitions necessary to evolve the step from the DFH equations to the CPD ones. A detailed exposition of the DFH can be found in [7-10].

We consider continuum mechanics description of a mixture of \( M \) chemical components present inside a spatial region \( D \) having volume \( V_D \). The region contains \( N_{ib} \) of each type of molecules. To avoid large numbers, the quantities \( N_{ib} \) are measured in moles. In case of homogeneous and static mixture we define chemical component molar densities by \( n_i = N_{ib} / V_D \). If the mixture is inhomogeneous and possibly evolving, one can define \( n_i \) as a dynamic variable at time \( t \) and spatial point \( x^a \) by establishing a small volume limit, such as \( n_i(t, x^a) = \lim_{\delta x \to 0} (N_{ib} / V_D) \). Like elsewhere in continuum mechanics, the small volume limit is understood as the convergent procedure with \( V_D \) being small, but still larger than the molecular volume.

By counting the flow rate of molecules through a small area inside the mixture, one can define the component flux \( I_{ia} = I_{ia} (t, x^a) \). The component fluxes are used to calculate the mass flux \( I_a = m_i I_{ia} \), where \( m_i \) is the molar mass of the \( i \)-th component. By introducing mass density \( \rho = m_i n_i \) it is possible to define mass velocity \( v_a = \rho^{-1} I_a \). Component flux \( I_{ia} \) can be represented as a combination of transport term \( n_i v_a \) and diffusion flux \( Q_{ia} \):

\[
    I_{ia} = n_i v_a + Q_{ia}
\]
where by definition diffusion flux does not influence net mass transfer \( m_i Q_{ia} = 0 \). We assume the existence of the Helmholtz energy functional

\[
F_D = F_D [n_i] = \int_D \omega dV + \int_{\partial D} f_x dA, \tag{4}
\]

\[
\omega = \kappa_i n_i - p, \quad d\omega = \kappa_i d n_i, \quad d\rho = p d\kappa_i, \tag{6}
\]

where \( \kappa_i \) is the chemical potential of the \( i \)-th component, and \( p \) is the hydrostatic pressure.

The model in Eqs. (4) and (5) is adequate for description of many important phenomena involving multiphase multicomponent mixtures. Up to now, it was successfully used to simulate multiphase multiphase phenomena with or without phase transitions, surfactants, and mixtures with solid phases such as gas hydrates or solid particles [9-21, 26].

Now we list the DFH statements necessary to move forward to CPD.

The multiphase compositional transport in case of isothermal flow of fluids with Newtonian rheology is governed by the following equations written in a form of conservation laws for chemical components and momentum

\[
\begin{align*}
\partial_t n_i + \nabla \cdot (n_i v_i + Q_{ia}) &= 0, \tag{7} \\
\partial_t (\rho v_i) + \nabla \cdot (\rho v_i v_i - p_{ab}) &= 0, \tag{8}
\end{align*}
\]

together with the relations

\[
\begin{align*}
p_{ab} &= \sigma_{ab} + \tau_{ab}, \tag{9} \\
\sigma_{ab} &= (\omega - \Phi_i n_i) \delta_{ab} - v_{ij} \delta_{a} n_i \delta_{b} n_j, \tag{10} \\
\tau_{ab} &= \eta_v \delta_{ab} v_i v_i + \eta_s (\delta_{ab} v_i + \delta_{b} v_i - \frac{2}{3} \delta_{ab} \nabla^2 v_i), \tag{11} \\
Q_{ia} &= -D_{ij} \delta_{ia} \Phi_i, \tag{12} \\
\Phi_i &= \kappa_i + 2 \left( \frac{\partial v_{ab}}{\partial n_i} \right) \delta_i n_i \delta_{ab} n_j - \delta_{ab} (v_i \nabla^2 n_i), \tag{13}
\end{align*}
\]

and subject to the boundary conditions at \( \partial D \), when it is a contact surface with some immobile solid

\[
v_a = 0, \tag{14}
\]

\[
I^a Q_{ia} = 0, \quad \frac{\partial f_x}{\partial n_i} - I^a v_i \nabla^2 n_i = 0, \tag{15}
\]

where \( p_{ab} \) is the stress tensor, \( \sigma_{ab} \) is the static stress tensor, \( \tau_{ab} \) is the viscous stress tensor, \( \Phi_i \) is the generalized chemical potential of the \( i \)-th mixture component, \( \eta_v \) and \( \eta_s \) are nonnegative bulk and shear viscosity coefficients, respectively, \( D_{ij} \) is the nonnegative definite symmetric matrix subject to \( D_{ij} m_j = 0 \), \( I^a \) is the internal normal unit vector at the boundary surface \( \partial D \), and \( \delta_{ab} \) is the Kronecker symbol. We also note that there is the identity \( \nabla^2 \delta_{ab} = -n_j \delta_{a} \Phi_j \), which reduces to \( \nabla^2 \delta_{ab} = 0 \) in static equilibrium.

It is possible to derive the differential equation for the total energy (that includes both Helmholtz and kinetic energy) with energy flux \( J_a \) and dissipation rate function \( \chi \):

\[
\begin{align*}
\partial_t (\omega + 2^{-1} \rho v_i v_i) &= -\partial_a J_a + \chi, \tag{17} \\
J_a &= \frac{1}{2} \rho v_i v_i - v_i n_i \delta_{a} n_j - \tau_{ab} v_b + \Phi_i, \tag{18} \\
\chi &= -\delta_{ab} \delta_{a} v_i + Q_{ia} \delta_{ia} \Phi_i \tag{19}
\end{align*}
\]

In accordance with Eqs. (11), (12), and (22) the inequality \( \chi \leq 0 \) holds. Also, in accordance with Eqs. (14)-(16) the energy flux vanishes at the boundary \( \partial D \) :

\[
l^a Q_{ia} = 0. \tag{15}
\]

This signifies that in absence of external sources and forces the total energy decreases \( \frac{d}{dl} \int_D (\omega + 2^{-1} \rho v_i v_i) dV \leq 0 \) in consistency with the second law of thermodynamics.

2.2 Upscaling to macroscopic transport in porous medium: chemical potential drive

Now we consider transition from microscopic DFH to macroscopic CPD description of fluid transport in porous medium. We use variables related to microscopic pore-scale description in parallel to similar variables related to macroscopic description. If \( a \) is a microscopic variable, then similar macroscopic variable is denoted by symbol \( \overline{a} \). The latter variable is usually obtained from the former by using certain averaging procedure. For example, we define component molar density by

\[
\pi_i (t, x) = \lim_{r \to 0} \left( N_i / r^3 \right), \quad \text{where the pore volume} \ V_D \ \text{is now sufficiently large to contain representative piece of saturated porous medium. The respective spatial region} \ D \ \text{encompasses pores belonging to a porous medium region} \ \Delta \ \text{containing both solid skeleton and pores. This porous medium region} \ \Delta \ \text{is supposed to be representative of pore}
\]
structure in statistical terms but not necessarily identical to the neighbor regions of porous medium. This is different from homogenization approach with periodic geometry of pore system [27]. The volumes of two regions $D$ and $\Delta$ are related by $DV_{\phi}=\phi V_{\Delta}$, where $\phi=\phi(x^+)$ is porosity. The processes in a porous medium are assumed to be sufficiently slow in respect to the changing macroscopic component molar densities, so that the local distribution of chemical components and phases in pores are close to the equilibrium. Therefore, it is possible to define the macroscopic Helmholtz energy density $\mathcal{D}^V=\frac{\phi}{V}$ since the equilibrium state of the mixture is determined by the functional (4) together with the components conservation condition $\int_{D} n_i dV=0$.

Using $\mathcal{D}$ it is possible to define macroscopic chemical potentials $\mathcal{F}_i=\frac{\partial \mathcal{D}}{\partial \pi_i}$. Let us study transport through the spatial porous region $D$. Transport is possible only if there are free subregions of the boundary $\partial D$, which are not associated with solid surface. We designate $\partial D=D_\Gamma \cup D_\Gamma^s$, where $\Gamma$, $\Gamma^s$ are free and solid boundary, respectively. At the free boundary $\Gamma$, the so-called free boundary condition for velocity is used in the form of $l^b \partial_b v_a=0$ instead of the no-slip condition in Eq. (14) of the DFH. The diffusion transport is generated by gradients of generalized chemical potentials $\partial_a \Phi_j$ (see Eq. (12)). The convective transport is determined by mass velocity, which under current assumptions satisfies the reduced momentum equation

$$\partial_b \tau_{ab}=n_i \partial_a \Phi_j,$$  (20)

with mixed no-slip and free boundary conditions. In terms of differential operations Eq. (20) can be represented as follows

$$L_{ab} v_b=-\eta_a=-n_i \partial_a \Phi_j,$$  (21)

where the left side is positive elliptic self-adjoint operator in $L_2(D)$ acting on the velocity field. It admits the solution with a Green’s function, which can also be represented in operator form

$$v_a=-G^{ab} \eta_b=-G^{ab} (n_i \partial_a \Phi_j).$$  (22)

Here the integral operator in $G^{ab}$ is positive definite, and also $\partial_a \Phi_j=n_i \partial_a \Phi_j$.

Averaging Eq. (23) over different cross-sections of the region $\Delta$ produces the macroscopic flux of components generated by gradients of chemical potentials (Chemical Potential Drive or CPD)

$$\overline{\mathcal{F}}_{ia}=<I_{ia}>=-K_{ij}^{ab} \partial_a \Phi_j,$$  (24)

where the transport matrix $K_{ij}^{ab}=K_{ij}^{ba}(x^+)$ is positive definite and satisfies the symmetry conditions $K_{ij}^{ab}=K_{ij}^{ba}=K_{ji}^{ab}$. The component conservation equation together with the CPD transport law in Eq. (24) that constitutes the closed formulation of macroscopic fluid transport model is as follows

$$\partial_t (\phi \pi_i) + \partial_a \mathcal{F}_{ia}=0.$$  (25)

Direct calculations lead to the macroscopic equivalent of Eq. (17)

$$\partial_t (\phi \mathcal{F})=-\partial_a \mathcal{F}_a+\mathcal{X},$$  (26)

$$\mathcal{F}_a=\overline{\mathcal{F}}_{ia}+\mathcal{X}=\overline{\mathcal{F}}_{ia} \partial_a \Phi_i \leq 0.$$  (27)

2.3 Boundary conditions between DFH and CPD domains

In highly heterogeneous porous material there can be situations when there are adjacent spatial regions where one can use alternatively DFH or CPD. It is necessary to discuss what boundary conditions should be set at the common boundary $\partial D_1 \cap \partial D_2$ between the DFH region $D_1$ and the CPD region $D_2$. Eq. (16) is still valid at $\Gamma$ in $D_1$, while Eqs. (14) and (15) are not because there can be transport of components through the boundary.

One evident boundary condition follows from conservation of component fluxes

$$I^a_{ia}=I^a_{ia},$$  (29)

The next condition can be derived from energy conservation at the boundary

$$l^a J_a=l^a \mathcal{F}_a.$$  (30)

Using Eqs. (18), (29), and (30) one can satisfy Eq. (27) by imposing boundary condition for chemical potentials

$$\mathcal{K}_i=\Phi_i - m_i X,$$  (31)

where variable $X$ is determined by the additional boundary equation

$$X \rho \frac{l^a v_a}{2} + \frac{1}{2} \rho v_k v_k l^a v_a - l^a \tau_{ab} v_b = 0.$$  (32)
2.4 Examples of CPD models

For the first example we assume that the porous medium is homogeneous and isotropic, and there is single-phase single-component flow. In this case the CPD transport law is equivalent to the Darcy law (see the last equation in (6))

$$T_{wa} = -K \partial_a \kappa_1 = -K \pi_1^{-1} \partial_a \pi_1$$

with CPD transport coefficient $K$ being proportional to the absolute permeability $k$, $K = k \pi_1^{-1} \eta_1^{-1}$, where coefficient $\eta_1$ is the shear viscosity.

Second, we consider two-phase two-component mixture modeling water and oil. Here we have the following definitions

$$\tilde{n}_1 = \tilde{s}_w \tilde{n}_w, \quad \tilde{n}_2 = (1-\tilde{s}_w) \tilde{n}_o,$$  \hspace{1cm} (34)

$$\tilde{\bar{f}} = \bar{f}(\tilde{n}_w) = \tilde{s}_w \bar{f}_w (\tilde{n}_w) +$$

$$(1-\tilde{s}_w) \bar{f}_o (\tilde{n}_o) + \bar{f}_{\text{cap}} (\tilde{s}_w),$$  \hspace{1cm} (35)

$$\tilde{\bar{p}}_w - \tilde{\bar{p}}_o = \bar{f}_{\text{cap}} (\tilde{s}_w) = \bar{f}_{\text{cap}}' (\tilde{s}_w),$$  \hspace{1cm} (36)

where $\tilde{s}_w$ is the water saturation, $\tilde{\bar{f}}_w, \tilde{\bar{p}}_w$ are the Helmholtz energy density and pressure of water, $\tilde{\bar{f}}_o, \tilde{\bar{p}}_o$ are the Helmholtz energy density and pressure of oil, and $\bar{f}_{\text{cap}} (\tilde{s}_w), \tilde{\bar{p}}_{\text{cap}} (\tilde{s}_w) = \bar{f}_{\text{cap}}' (\tilde{s}_w)$ are the Helmholtz energy density and pressure for capillary forces. Eqs. (34)-(36) determine variables $\tilde{s}_w, \tilde{\bar{f}}_w,\tilde{\bar{p}}_w$ when variables $\tilde{n}_1, \tilde{n}_2$ are known. The transport model in Eq. (24) is consistent with the generalized Darcy law if there are the following additional conditions:

$$K_{12} = K_{21} = 0, \quad K_{11} = k_w \pi_1^2 \eta_1^{-1},$$

$$K_{22} = k_o \pi_2^2 \eta_2^{-1},$$  \hspace{1cm} (37)

where $k_w, k_o$ are water and oil phase permeabilities. If conditions in Eq. (37) are not satisfied, CPD model can demonstrate some phenomena outside traditional Darcy transport.

3 Numerical examples

To demonstrate application of a workflow based on the joint DFH modeling within resolved porosity and CPD modeling within matrix containing unresolved smaller pores (DFH+CPD workflow) we chose a heterogeneous, but relatively well characterized chalk sample. The sample is grainstone composed of pelloids, skeletal grains and ooids. The grains are cemented by equant calcite spar cement. Intergranular porosity is the dominant pore system, additionally some moldic and micropores are present in partially leached grains. Pore throats are ranging between approximately 1 and 10 microns. Pore throat distribution is skewed to smaller pore throat sizes. Measured gas porosity of the sample is 26.2%, and gas permeability is 19.4 mD.

Fig. 1. Micro-CT grayscale cross-section of the dry chalk (a) 8 mm mini plug at 2.46 um resolution, (b) a 2.46 mm x 2.46 mm fragment of a cross-section at 2.46 um resolution, and (c) a 0.82 mm x 0.82 mm fragment of a cross-section at 0.82 um resolution; the dashed line in (a) shows the position of the maximum size parallelepiped model extracted from the mini plug.
An 8 mm diameter mini plug was scanned with 2.46 μm per voxel resolution, and a portion of the mini plug was scanned with higher resolution of 0.82 μm per voxel (Fig. 1).

Two 3D digital rock models (DRM) with these two different resolutions were constructed (Fig. 2). Both models cover the same spatial region with the size 0.82 mm x 0.82 mm x 0.82 mm, but they have different number of cells (voxels). The high-resolution DRM has $1000^3$ (1000x1000x1000) cells and the low-resolution DRM has $333^3$ (333x333x333) cells so that the cell sizes are 0.82 μm and 2.46 μm, respectively. In either model some portion of pores fell below resolution. However, it is evident from Fig. 2 that significantly higher proportion of pores remains below resolution in the low-resolution DRM than it does in the high-resolution DRM. The data on the resolved and unresolved porosity is shown in Table 1.

![Fig. 2](image)

**Fig. 2.** 3D view of the pore space of the (a) high-resolution $1000^3$ and (b) low-resolution $333^3$ DRMs. The resolved pores are shown in green, the unresolved pores (subresolution matrix) are shown in red, and the skeleton is made transparent.

| Model       | Resolved porosity | Subresolution matrix |
|-------------|-------------------|----------------------|
| $1000^3$, 0.82 μm | 0.166              | 0.073                |
| $333^3$, 2.46 μm  | 0.091              | 0.157                |

Table 1. The portion of resolved porosity and unresolved porosity (subresolution matrix) within digital rock models of different spatial resolution.

In addition to X-ray micro-CT, we have done scanning electron microscopy (SEM) of parts of the sample to reveal the pores below micro-CT resolution down to 50 nm. Using both micro-CT high-resolution (0.82 μm) and SEM data we performed digital modeling of MICP experiment and reconstructed pore throat size distribution that is compared to the standard experimental MICP measurements in Fig. 3. The two curves in Fig. 3 match well meaning that most of the controlling pore throats have been adequately resolved on the corresponding scales.

In modeling on the high-resolution DRM, we assumed that the subresolution matrix portion of the model is effectively impermeable for oil at relevant oil-water capillary pressure levels so that only the DFH equations need to be solved within the resolved pore space. On the other hand, in modeling on the low-resolution DRM, we assumed that the subresolution matrix is permeable and therefore the full DFH+CPD simulation needs to be done. To set up a single-phase DFH+CPD simulation it is necessary to provide absolute permeability in the subresolution matrix cells so that the single-phase CPD transport law given in Eq. (33) could be resolved. In order to do this, we first simulated single-phase flow on the high-resolution DRM to obtain velocity distribution (Fig. 4a).

![Fig. 3](image)

**Fig. 3.** Reconstructed numerically from the micro-CT and SEM data (green curve) and obtained from the MICP measurements (blue curve) pore throat size distributions.

Then, by interpreting the simulation results in terms of Eq. (33) and linking them with the gray scale distribution from the high-resolution images such as those shown in Fig. 1c, we established a correlation between the gray scale and absolute permeability values. We then applied the same correlation to the low-resolution gray scale.
images to inform the low-resolution DRM about the subresolution matrix permeability. The velocity distribution obtained on this 333³ low-resolution DRM is presented in Fig. 4b. The absolute permeabilities calculated on the high- and low-resolution DRMs are 17.3 mD and 18.2 mD, correspondingly.

The next step was simulating two-phase flow and the steady-state relative permeability experiment. The two immiscible phases were assumed to be water and oil with the properties as follows: ρ_w = 1000 kg/m³, ρ_o = 800 kg/m³, η_w = 0.001 Pa·s, η_o = 0.004 Pa·s, γ_{ow} = 0.03 N/m, where ρ_w, ρ_o are the water and oil mass densities, η_w, η_o are the water and oil viscosities, and γ_{ow} is the oil-water interfacial tension. Since the sample is carbonate, it was assumed that it has mixed wettability. To model this, we first injected oil into initially water-wet 100% water saturated high-resolution DRM using our standard DFH workflow [11, 14-17, 19-21]. (In all cases here and below injection was arranged in direction from bottom to top as shown in the 3D views.) Then at those places where oil touched the pore walls we switched wettability to moderately oil-wet while leaving the untouched pore walls water-wet. In this way most of the smaller pores as well as the pores isolated by thin pore throats remained water-wet, while the larger pores attained wettability towards oil. The distribution of water and oil at the residual water saturation state is shown in Fig. 5a. Using the obtained mixed-wet wettability distribution we simulated the steady-state relative permeability experiment by injecting oil and water mixture in several steps different in water content in the influx; at the final step, with only water injected, the residual oil saturation was obtained (Fig. 5c).

Fig. 4. 3D view of the velocity absolute value distribution in (a) the high-resolution 1000³ DRM and (b) the low-resolution 333³ DRM of the same location within the core. The shown distribution was scaled to the [0; 1] range. The velocity corresponds to flow in vertical (z) direction.
Fig. 5. 3D view of the distribution of water and oil in the **high-resolution DRM** (left column) and the **low-resolution DRM** (right column) at the initial conditions corresponding to the residual water saturation (a, b), and the final state corresponding to the residual oil saturation (c, d). Water is shown in semitransparent blue, oil is shown in red, and skeleton is completely transparent.

To run DHD+CPD two-phase simulation, it is necessary to provide CPD transport matrix of Eq. (24) in subresolution matrix cells. Alternatively, when dealing with immiscible phases like oil and water that are each described by just one pseudo-component, it is possible to use simplified CPD transport model provided in Eq. (37) and based on the concepts of absolute and relative permeabilities and capillary pressure. We already discussed how absolute permeability was assigned to the subresolution matrix cells of the low-resolution DRM. The relative permeability was assigned in a similar way since all the velocity distribution, pressure, and water saturation for the subresolution matrix cells are known from the simulation on the high-resolution model. The capillary pressure function was constructed using the Young-Laplace equation and then scaled to the pore size distribution correlated by the gray scale values.

The distributions of oil and water obtained in simulation of the two-phase steady-state relative permeability experiment modeled using the DFH+CPD workflow on the low-resolution DRM are presented in Fig. 5b,d. The chart in Fig. 6 compares the relative permeability curves obtained on both high- and low-resolution DRMs.

The final example we wish to present here is the simulation of both absolute and steady-state relative permeabilities on another low-resolution DRM with the size of $4.92 \times 4.92 \times 2.46$ mm approximated with $500 \times 500 \times 250$ cells with the cell size of 9.84 um. This model covers maximum size parallelepiped that can be extracted from the micro-CT scans of the 8 mm mini plug as shown by the dashed line in Fig. 1a. All of the parameters necessary to set up the DFH+CPD model were taken from the previous calculations on the high-resolution DRM as described previously.

Simulated absolute permeability of this model is 19.8 mD. The distributions of water and oil corresponding to the residual water saturation and residual oil saturation as obtained during the steady-state relative permeability simulation are shown in Fig. 7. The simulated relative permeability curves are presented in Fig. 8.

In this work, the simulation results were interpreted in consistency with the conditions in Eq. (37) that is with the traditional Darcy transport. However, in complex fluid problems deviations from relative permeability concept are expected, which then must be treated in the frame of the wider CPD model.
Fig. 7. 3D view of the distribution of water and oil in the upscaled low-resolution DRM at (a) initial conditions corresponding to the residual water saturation, and (b) final state corresponding to the residual oil saturation. Water is shown in semitransparent blue, oil is shown in red, and skeleton is transparent.

Fig. 8. Simulated steady-state relative permeability curves obtained on the upscaled low-resolution model.

2 Conclusion

We developed a new digital rock workflow applicable for simulation of multiphase flow on multiscale models representing heterogeneous core samples. The workflow called DFH+CPD is based on solving the standard DFH equations within resolved porosity and solving the new CPD equations within unresolved porosity or subresolution matrix. The CPD equations are derived in mathematically rigorous way from the DFH equations averaged over statistically representative ensemble of pores. The new workflow amounts to an effective upscaling procedure allowing multiphase simulation on coarse resolution digital rock models that include information that can be obtained on high-resolution models. The DFH+CPD workflow was implemented within the Schlumberger DHD simulator.

To validate the new workflow, we compared absolute and steady-state relative permeabilities simulation results on the high-resolution model and the low-resolution model of the same region within the heterogeneous carbonate core. Then we demonstrated simulation of the steady-state relative permeabilities on the upscaled digital rock model that covers almost entire 8 mm mini plug.

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