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A new estimation of equivalent matrix block sizes in fractured media with two-phase flow applications in dual porosity models

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Abstract

Single and multiphase flows in fractured porous media at the scale of natural reservoirs are often handled by resorting to homogenized models that avoid the heavy computations associated with a complete discretization of both fractures and matrix blocks. For example, the two overlapping continua (fractures and matrix) of a dual porosity system are coupled by way of fluid flux exchanges that deeply condition flow at the large scale. This characteristic is a key to realistic flow simulations, especially for multiphase flow as capillary forces and contrasts of fluid mobility compete in the extraction of a fluid from a capacitive matrix then conveyed through the fractures. The exchange rate between fractures and matrix is conditioned by the so-called mean matrix block size which can be viewed as the size of a single matrix block neighboring a single fracture within a mesh of a dual porosity model.

We propose a new evaluation of this matrix block size based on the analysis of discrete fracture networks. The fundamentals rely upon establishing at the scale of a fractured block the equivalence between the actual fracture network and a Warren and Root network only made of three regularly spaced fracture families parallel to the facets of the fractured block. The resulting matrix block sizes are then compared via geometrical considerations and
two-phase flow simulations to the few other available methods. It is shown that the new
method is stable in the sense it provides accurate sizes irrespective of the type of fracture
network investigated. The method also results in two-phase flow simulations from dual
porosity models very close to that from references calculated in finely discretized networks.
Finally, calculations of matrix block sizes by this new technique reveal very rapid, which
opens the way to cumbersome applications such as preconditioning a dual porosity approach
applied to regional fractured reservoirs.

**Keywords**

Fractured porous media; Matrix block sizes; Dual porosity models; Multiphase flow.
1. Introduction

Conventional oil reservoirs are often housed in fractured rocks, especially in carbonates environments, and one can estimate that more than 30% of world oil reserves are concealed in densely fractured systems, oil being mainly trapped in the host rock matrix. Paradoxically, these geological structures may trigger contrasted effects on large-scale two-phase flow patterns by increasing oil recovery due to high local permeability values, or on the opposite, by decreasing oil extraction rates because of early water invasion, viscous fingering etc. The same type of behavior is also encountered in the context of water decontamination and can become even more complex if oil (and/or water) is swept by injections of miscible gas.

Modeling two-phase flow in fractured reservoirs is now often employed for the purpose of various applications, for instance to assess the relevance of different oil recovery strategies or to investigate on the feasibility of in-situ water decontamination processes (Bourbiaux, 2010). This fact makes that modeling single phase or multiphase flow in fractured media is still a fertile research domain even though pioneering works on the topic started in the early sixties (e.g., in Lemonnier et al., 2010a, b).

In this context, flow simulations relying upon finely gridded discrete fracture networks and their associated (discretized) matrix blocks are becoming increasingly popular because of the availability of high performance computers, the progress in algorithms for meshing complex geometries, and the availability of sophisticated numerical techniques for solving partial differential equations (Landereau et al., 2001; Noetinger et al., 2001; Adler et al., 2005; Matthäi and Nick, 2009; Fourno et al., 2013). This exhaustive approach is critical to bring us reference solutions and various benchmarks with which simpler approaches can be compared. Nevertheless, gridded discrete fracture networks may be poorly documented and include flawed information in the case of real-world applications. In addition, finely gridded
systems remain hardly usable for current practical applications to large-scale systems that
result in cumbersome model parameterizations and heavy computations. This downside is
emphasized in the domain of petroleum engineering usually dealing with both non-linear
multiphase flow and dense fracture networks requiring huge discretization efforts (Landereau
et al., 2001; Adler et al., 2005; Fournon et al., 2013). Applicability is also hindered by
duplicated calculations if the study encompasses tests of various model designs, various
model parameterization and various flow scenarios.

Fortunately, dense fracture networks are also good candidates to homogenization at
the scale of reasonable elementary mesh sizes (on the order of 5-100 m) by resorting for
example to the dual porosity approach to fractured media initially developed by Barenblatt et
al. (1960). The dual porosity formulation conceptualizes a fractured system as two
overlapping continua merging a fracture medium and a matrix medium with contrasts of
hydraulic properties between the two continua. Flow is then described by a set of equations in
each continuum (this set depends on the type of flow and the fluid phases present in the
system) associated with an exchange term ruling the fluid fluxes percolating between
continua.

This exchange term is all the more important that in general fractures are conveying
flow as the matrix stores fluid volumes. In transient problems as for example forced flow
between injecting and extracting wells, the way the relationship establishes between storage
capacity and conduction property conditions the overall response of the reservoir (e.g., Acuna
and Yortsos, 1995). In the specific cases of two-phase flow (water and oil), the absence or the
weak incidence of capillary forces in open fractures makes that flow is locally mainly of
single-phase type conveying either oil or water (with sharp saturation fronts in between) at
different locations in the fracture network. For its part, the way the matrix blocks are soaked
(water invades the matrix and pushes oil away) or drained (oil pushes water) strongly depends
on matrix block sizes and on the petro-physics properties of the matrix, making that extraction
to the matrix of a fluid by the other is mainly driven by capillary forces or by capillary
forces plus viscous forces (single-phase Darcian flow to make it simple).

When a discrete fracture network is connected enough and handled at an elementary
scale larger than a representative elementary volume, the exchange term in the dual porosity
models is proportional to an equivalent matrix block size. Intuitively, a REV for a fracture
network is a volume within which mean properties of the network such as fracture density,
fracture aperture, fracture hydraulic conductivity have some statistical meaning (Long et al.,
1982; Neuman, 1988). In a dual porosity model, the REV is also associated with the
capability to represent the actual fracture network as a synthetic network made (in three-
dimensional problems) of three regularly spaced fracture families, each family developing
fracture planes normal to one of the three main directions of flow. The so-called DFN
homogenized as a "sugar-cube" model (Warren and Root, 1963) is at the origin of the notion
of the equivalent matrix block size in relation with the dimensions of the elementary "sugar
piece" separating neighbor fractures in the homogenized DFN (Kazemi et al., 1976).

There exist two types of methods to evaluate the elementary matrix block size. The
first type relies upon exercises matching actual well test drawdown curves with analytical
solutions that inherit from rigorous mathematical homogenization or large-scale averaging
techniques (Arbogast 1990; Quintard and Whitaker, 1993; Noetinger et al., 2001; Unsal et al.,
2010; Noetinger and Jarrige, 2012). The downside of these techniques is that sometimes
actual well testing in fractured rock do not exist and when these tests exist, the damaged zone
in the close vicinity of a well may not fully reflect flow conditions in the natural fracture
network. The second type of methods is based on geometrical considerations regarding the
fracture network. These considerations led to three geometrical approaches that are the
geometrical imbibition method (GI, Bourbiaux et al., 1997), the enhanced general imbibition
method (EGI, Bourbiaux et al., 2006), and the mean spacing method (MS, Narr, 1996). It is obvious that these approaches can only be applied if a minimum prior knowledge about the fracture network geometry is available.

In this contribution we propose a new geometrical method that can to some extent overlook the actual geometry of the fracture network because the method relies upon the identification of a sugar-cube DFN equivalent to the actual network (see details hereafter). The method also allows us to calculate matrix block sizes along directions parallel to the main flow directions that are conditioned by the geometry of the fracture network (or its equivalent as a sugar-cube model). Section 2 (and Appendix A) is focused on the theoretical framework we rely upon to build the so-called oriented block size (OBS) method that we propose. For the sake of clarity, a few features about dual-porosity models are also reminded. The matrix block sizes stemming from the OBS technique are then compared to that from the other geometrical techniques (GI, EGI, and MS, see above). The comparison is performed by way of a suite of calculations applied to synthetic random fracture networks for which we explicitly control both the geometric and hydraulic properties of the networks and the mean size of matrix blocks between fractures. As told earlier, only dense and well-connected fracture networks are considered because sparse networks cannot be homogenized via a dual porosity model at the scale of a complete underground reservoir. Section 4 evaluates the OBS technique and also the other geometrical approaches within the framework of a dual-porosity model compared with exhaustive calculations discretizing the fracture network and the matrix blocks. The two-phase flow scenarios are either dominated by capillary forces or viscous forces in an exercise which consists in draining oil from matrix blocks by injecting water in fractures. These complex flow scenarios are performed over synthetic test cases in which we control the reference calculations (in a fully discretized system). This procedure enable us to clearly
emphasizes the main theoretical findings regarding matrix block size in dual porosity models before envisioning further concrete field-scale applications.

2. Theoretical background

In various approaches to fractured systems, the duality of fracture networks embedded in a host rock matrix is often represented as two overlapping continua merging a fracture medium and a matrix medium. In a so-called dual porosity – single permeability model, the fractures are usually highly conductive and poorly capacitive as the matrix is highly capacitive but with negligible flow triggered by fluid pressure gradients (weak permeability). As an example, single-phase Darcian flow in a dual continuum approach results in the resolution of two equations in the form

\[
\frac{\partial (\rho \phi^f)}{\partial t} + \nabla \left( -\rho \frac{k^f}{\mu} \nabla (P^f + \rho g z) \right) - E^{m\to f} = 0
\]

(1)

\[
\frac{\partial (\rho \phi^m)}{\partial t} + E^{m\to f} = 0 ; E^{m\to f} = \rho \sigma \frac{k^m}{\mu} (P^m - P^f)
\]

(2)

For the sake of simplicity, references to space (x) and time (t) for parameters and state variables have been dropped. The indexes f and m refer to fracture and matrix continua, respectively. \( \rho \) [ML\(^{-3}\)] is the mass density of the fluid, \( \mu \) [ML\(^{-1}\)T\(^{-1}\)] is the dynamic viscosity of the fluid, \( k^\lambda \) [L\(^2\)] is the permeability of the continuum \( \lambda \), \( \phi^\lambda \) [-] is the porosity of the continuum \( \lambda \), \( P^\lambda \) [ML\(^{-1}\)T\(^{-2}\)] is the fluid pressure in the continuum \( \lambda \), \( g \) [LT\(^{-2}\)] is the scalar value of the gravity acceleration \( g \), and \( z \) [L] is the local elevation taken from an arbitrary reference and counted positive upward. \( E^{m\to f}_p \) [ML\(^3\)T\(^{-1}\)] is the exchange rate (a mass fluid flux per unit volume of medium) between the fracture and the matrix continua.
In (2), the exchange rate is of pseudo steady-state type meaning that the relationship between matrix and fractures depends on pressure gaps between the continua and not on a convolution product of their derivatives with respect to time. In (2), the matrix permeability $k^m$ is assumed small enough to neglect Darcian fluxes in the matrix (compared to that in the fractures) but large enough to enable fluid flux percolation between the matrix and the fractures. Therefore, the matrix permeability is an entry of the exchange rate via the term $\sigma k^m, \sigma [L^{-2}]$ being a shape factor tensor that quantifies the mean size of the matrix blocks associated with the fractures included in an elementary volume (for example, the volume corresponding to the elementary meshing employed when solving numerically Eqs (1) and (2)). By construction, the pseudo steady-state assumption in (2) ignores the early transient flow regime between matrix and fractures which may result in erroneous evaluations of exchanged fluid fluxes, especially in the case of weakly permeable matrix media requiring long times for equilibrating their fluid pressure fields with that of fractures (e.g., as in shale gas and shale oil extraction problems). Transient exchange rates between fractures and matrix are the natural outcome of Multiple INteracting Continua (MINC approaches) initially developed in the late eighties (e.g., Pruess and Narasimhan, 1985; Pruess et al., 1990) and more recently reassessed and improved (e.g., Karimi-Fard et al., 2006; Tatomir et al., 2011, de Dreuzy et al., 2013). The MINC models are not incompatible with the notion of mean matrix block size in homogenized fractured systems as each matrix block is viewed as an entity of prescribed size enclosing a nested heterogeneity.

Various formulations of the shape factor have been proposed for many modeling applications (Kazemi et al., 1976; Thomas et al., 1983; Coats 1989; Ueda et al., 1989; Lim et al. 1995; Quintard and Whitaker, 1996; Noetinger and Estebenet., 2000) amongst which the formulation proposed by Kazemi et al. (1976) is the one used in this study. This choice is motivated by a quite simple formulation which allows for dealing with diagonal tensors, and
also introduces the mean matrix block size as a quantity weighting the influence of the matrix permeability tensor to control the fluid fluxes exchanges between matrix and fractures. For diagonal permeability and shape factor tensors, the product $\mathbf{\sigma k^m}$ is developed as

$$\mathbf{\sigma k^m} = \begin{pmatrix} k_x^m/s_x^2 & 0 & 0 \\ 0 & k_y^m/s_y^2 & 0 \\ 0 & 0 & k_z^m/s_z^2 \end{pmatrix}$$

(3)

with $s_i$ [L] ($i=x, y, z$) the mean matrix block size along the main flow direction $i$. As the exchange rate between the fractures and the matrix is a key feature to the behavior of a dual continuum and some other homogenized approaches (Lemonnier et al., 2010a, b), it makes sense to revisit the item especially regarding the mean matrix block size (which rules the fluxes, provided the fluid pressure fields are correctly calculated).

The Oriented Block Size (OBS) technique that we develop below infers the mean matrix blocks sizes $s_i$ ($i=x, y, z$) from a fractured system by assuming that a rock block enclosing an actual fracture network with various characteristics (e.g., Fig 1) can be turned into a simplified block with an equivalent fracture network composed of three families of planar fractures.

Fig. 1. A fractured rock block at the scale of a reservoir grid cell with references to main flow directions and facets of block normal to flow directions.
Each family is defined by a uniform spacing between fractures and a fracture plane normal to one direction of flow (or including the two other directions). This equivalent fracture network (e.g., Fig. 2) which draws the so-called "sugar-cube" configuration as proposed by Warren and Root (1963) and referred to as the WR model hereafter, is conceptually compatible with the notion of mean matrix block size. The three families of WR fractures delimit a parallelepiped elementary block separating neighbor fractures that should coincide with the shape factor as defined in Eq (3). If the whole WR block is wide enough, the three fracture families can be aggregated as a single fracture permeability tensor (or value) and a single fracture porosity for the whole block or its facets. These parameters depend on the size of the elementary matrix block separating the WR fractures. By comparing, or more exactly by identifying permeability and porosity properties of a WR block with that of an actual fractured block, one is able to define the equivalent mean matrix block size of the actual fractured block.

Fig. 2. A regular fracture network of three fracture families (a Warren and Root (WR) model) at the scale of a reservoir grid cell with reference to main flow directions, facets of block normal to flow directions, and spacing between fractures.
Let us take a parallelepiped block housing an actual fracture network as depicted in Fig. 1. The first way to identify a single macroscopic permeability tensor $\mathbf{k}^f$ for the block is to employ upscaling approaches, multiple continua theory (Karimi-Fard et al., 2006; Tatomir et al., 2011; Jourdain et al., 2014), analytical solutions (Oda, 1985) or simply to conjecture the entries of the tensor as could be done, for example, in parameterizing a dual porosity approach after having postulated that the approach was convenient for the problem under investigation.

A second way is to extract the (diagonal) tensor from the structural properties of the fracture network and its relationships with the homogenization volume (the block) concealing it.

The actual fractured block as depicted in Fig 1 is oriented with its main directions along the main directions of flow indexed by $i=1,2,3$ (here completely equivalent to $i = x, y, z$ for locations in space denoted $\mathbf{x} = (x, y, z)$ but easier to manipulate when incrementing the index). The block size in direction $i$ is denoted $\Delta_i$ and the sides delimiting the block are also indexed by $i$ but for limits normal to the main direction $i$. In addition, block sides are labelled $i-$ or $i+$ according to their respective location upstream or downstream along direction $i$.

Assuming that the fractured block is well connected, the mean permeability of the block along a direction $i$ can be calculated as the average over the sides $i-$ and $i+$ of the local permeability of fractures intercepting the sides. This yields

$$k_{i}^{FN-S} = \frac{1}{2(\Delta_{i+1}\Delta_{i+2})} \left( \sum_{n=1}^{N_{f_i-}} k_n l_n e_n + \sum_{n=1}^{N_{f_i+}} k_n l_n e_n \right)$$

(4)

In the above equation, $i$ is a cycling index such that, e.g., $i+1 = 3$ when $i = 2$ and $i+1$ returns to 1 when $i=3$. $k_{i}^{FN-S}$ [L$^2$] is the macroscopic fracture permeability of the fractured block along direction $i$, $\Delta_{i+1}\Delta_{i+2}$ [L$^2$] is the total surface area of sides $i-$ and $i+$ these being intercepted by a number of fractures $N_{f_i-}$ and $N_{f_i+}$. $k$ [L$^2$] is the local permeability of a
fracture seen as intercepting the side of the block over an apparent length $l$ and with apparent fracture aperture $e$ [L].

By re-using the same notations for directions and sides in a rock block modeled as a WR network (Fig. 2), one can also calculate the entries $k_{i}^{WR}$ of the diagonal fracture permeability tensor of the WR block along directions $i$. The three fracture families of the WR network are also indexed by $i$ with the same notation as for the block sides, i.e., a fracture family $i$ corresponds to fracture planes normal to direction $i$. A family $i$ is of uniform spacing $s_i$ [L] ($s_i$ is measured along direction $i$, see Fig. 2), counts $Nf_i$ fractures with a uniform scalar permeability $k_i$ and a uniform fracture aperture $e_i$. With these settings and the assumption that flow only occurs in the fractures, the total flow rate entering or exiting the WR fractured block through a side $i$ normal to the direction $i$ can be expressed as

$$Q_i = -\nabla_i P \left( \sum_{j \neq i} Nf_j e_j \frac{k_j}{\mu} \right) = -\nabla_i P \frac{k_{WR}^{i}}{\mu} \Delta_{i+1} \Delta_{i+2}$$ \hspace{1cm} (5)

For the sake of simplicity, the gravity components of flow have not been accounted for in (5). $Nf_j e_j \Delta_{k,j,s,i,k,k,j}$ represents the total surface of flow developed by the family fracture $j$ through the side $i$ of the block, $\Delta_{i+1} \Delta_{i+2}$ is the total surface area of the side $i$, and $k_{i}^{WR}$ is the macroscopic fracture permeability of the WR block along direction $i$. The equality in (5) comes down to a direct identification of the three terms of the macroscopic permeability $k_{WR}^{i}$ as

$$\begin{bmatrix} k_{1}^{WR} \\ k_{2}^{WR} \\ k_{3}^{WR} \end{bmatrix} = \begin{bmatrix} 0 & 1/\Delta_2 & 1/\Delta_3 \\ 1/\Delta_1 & 0 & 1/\Delta_3 \\ 1/\Delta_1 & 1/\Delta_2 & 0 \end{bmatrix} \begin{bmatrix} Nf_1 e_1 k_1 \\ Nf_2 e_2 k_2 \\ Nf_3 e_3 k_3 \end{bmatrix}$$ \hspace{1cm} (6)

In the context of scaling the values $k_{i}^{WR}$ so they become equivalent to calculated values in a rock block encapsulating an actual fracture network, the number $Nf_i$ of WR fractures, their
aperture $e_i$ and their permeability $k_i$ become the unknowns of the problem. Therefore, it makes sense to invert the linear system of equation (6) which yields

$$N_{ij}e_ik_i = \frac{\Delta_i}{2} \sum_{j=1}^{3} (-1)^{\delta_{i,j}} k_j^{WR}$$

(7)

with $\delta_{i,j}$ the Kronecker delta function, $\delta_{i,j} = 1$ if $i = j$ and $\delta_{i,j} = 0$ if $i \neq j$.

By considering the structure of a WR fracture network, one can write

$$N_{ij}(e_i + s_i) = \Delta_i \Rightarrow \frac{N_{ij}e_i}{\Delta_i} = \frac{e_i}{e_i + s_i}$$

(8)

Note in the above expression that counting $N_{ij}$ fractures assumes the presence of $N_{ij} - 1$ fractures inside the block and that the two sides $i$ of the block are each bounded by half a fracture of family $i$ with half the aperture $e_i$ counted in the block. Introducing (8) in (7) results in

$$k_i = \frac{1}{2} \left( 1 + \frac{s_i}{e_i} \right) \sum_{j=1}^{3} (-1)^{\delta_{i,j}} k_j^{WR}$$

(9)

The expression (9) will be used later for the purpose of identification between an actual fractured block and a WR block.

Let us look at the porosity properties of the WR block. The fracture porosity $\phi^{WR}$ [-] of the whole WR block and the fracture density $\phi^{WR-S}$ [-] at a side $i$ defined as the porosity of fracture network at a side of the block (the ratio of the surface area of open fractures at a side to the total surface of the side) can also be derived as

$$\phi^{WR} \approx \sum_{i=1}^{3} \frac{N_{ij}e_i\Delta_{i\Delta_2}\Delta_{i\Delta_3}}{\Delta_1\Delta_2\Delta_3} = \frac{3}{2} \sum_{i=1}^{3} \frac{N_{ij}e_i}{\Delta_i}$$

(10)

$$\phi^{WR-S}_i \approx \sum_{i=1}^{3} \frac{N_{ij}e_i}{\Delta_i}$$

(11)
Notably, the expressions in (10) and (11) are rather simple but are approximations since the intersections of fractures are counted twice in the porosity values. This was found of negligible influence for classical block sizes and fracture apertures. Subtracting (11) from (10) returns the term \( N_f c_i / \Delta_i \) which also appears in Eq (8). Therefore, another way to express the relationship between the local WR fracture permeability \( k_i \) and the macroscopic permeability \( k^{WR} \) in (9) is

\[
k_i = \frac{1}{2} \left( \phi^{WR} - \phi^{WR-S}_i \right) \sum_{j=1}^{3} (-1)^{\delta_{ij}} k^{WR}_j
\]  

(12)

Both expressions (9) and (12) are employed to define the matrix block size \( s_i \) (in 9).

If the WR network is equivalent regarding its hydraulic properties to the actual fracture network, it is expected that \( \phi^{WR}, \phi^{WR-S}_i, \) and \( k^{WR}_j \) are similar to the equivalent properties in the actual block of fracture network, respectively denoted as \( \phi^{FN}, \phi^{FN-S}_i, \) and \( k^{FN-S}_j \) (see (4) for the latter term). It is also expected that the WR network, while being still equivalent to the actual fractured block, can inherit some properties (parameters) of a homogenized model such as the mean matrix block sizes of the medium and the permeability tensor at the macroscopic scale of a fractured block. By imposing these properties in (9) and (12), and after a few algebraic manipulations (see Appendix A for details), an expression of the mean matrix block sizes in a homogenized fractured block can be written as

\[
s_i \approx \frac{\sum_{j=1}^{3} (-1)^{\delta_{ij}} k^{FN-S}_j}{\left( \phi^{FN} - \phi^{FN-S} \right) \sum_{j=1}^{3} (-1)^{\delta_{ij}} k^f_j}
\]  

(13)

\( k^{FN-S}_j (i = 1, 2, 3) \) are the permeability values at the sides \( i \) of the actual fractured block, \( k^f_j (i = 1, 2, 3) \) are the entries of the diagonal permeability tensor of an homogenized medium equivalent to the fractured block (e.g., that of a dual porosity model), and \( \phi^{FN+}, \)
\( \phi_{ij}^{FN-S^*} (i = 1, 2, 3) \) are rescaled block and side porosities of the actual fractured block. These rescaled porosities of dimension \([L^{-1}]\) (a porosity per unit fracture aperture) are calculated over the skeleton of the actual fracture network to which each fracture is assigned a unit fracture aperture.

In addition to postulating the equivalence between a WR block and the actual fractured block, the assumptions allowing us to derive (13) are twofold. First, the actual fracture network is a good candidate for homogenization with the meaning that there exist macroscopic properties as mean matrix block size and diagonal permeability tensor characterizing the hydraulic behavior of the network at the large scale (at least, the scale of a mesh of a homogenized model). Second, a WR network exists (as that investigated by way of equations 5 to 12) but with uniform fracture aperture \( e_f \) over its three fracture families and still equivalent to the actual fracture network (see Appendix A for details). There is no clear criterion (except dealing with a dense and well-connected network) allowing us to state beforehand whether or not a given fracture network would follow the above assumptions. Eventual criteria would also depend on the flow processes and mechanisms targeted for further applications at the large scale.

It is worth to note that Eq. (13) depends on both the facet permeability values of the actual fractured block \( k_j^{FN-S} \), and the structural properties of the actual fracture network skeleton in the form of porosities \( \phi^{FN} \) and \( \phi_i^{FN-S^*} \). These features make that applicability of (13) is conditioned by a good knowledge of the actual fracture network geometry and, as a downside, renders the method hardly applicable to poorly-known natural systems. In the end, Eq. (13) should be mainly used in problems dealing with homogenization of systems with well-known geometry and discretization of synthetic fracture networks (as done for instance in reservoir engineering when passing from a geological model to a tractable flow model).
This notwithstanding, the OBS technique can also deliver another form of the mean matrix block size. By manipulating the expression of the side block permeability of the actual fractured block in (4), scaling the subsequent expression with the side block porosities $\phi_{i}^{FN-S}$ and making use of (13) (details are provided in Appendix A), another form of the mean matrix block size comes up as

$$s_i \approx \frac{2e_j \bar{k}}{\sum_{j=1}^{i} (-1)^{i-j} k_j}$$

This form introduces the existence of a mean single fracture aperture $e_j$ [L] (which is also the uniform aperture mentioned above for the WR network) and a mean single-fracture permeability $\bar{k}$ [L$^2$] at the scale of the whole actual fractured block. These two quantities are additional assumptions to that discussed regarding (13) for the applicability of (14).

Even though these assumptions may appear very restrictive, they give the possibility to infer mean matrix block sizes from poorly known and hardly accessible fracture networks as often encountered in field case applications. The entries $k_j$ of the permeability tensor of the whole fractured block can be evaluated by way of hydraulic tests; preferably interference testing between distant wells that avoid bias stemming from an environment close to the tested well that would not be representative of the fracture network at a larger scale. Values of uniform single-fracture aperture $e_j$ and uniform single-fracture permeability $\bar{k}$ are harder to infer because data obtained for instance from optic imaging of boreholes (for $e_j$) and flow or production logs (for $\bar{k}$) may reveal not representative of the whole network. It remains that the matrix block size calculation in (14) is feasible without resorting to any knowledge on the structure of the actual fracture network. It is obvious that the subsequent inferred value of mean matrix block size should be taken as an order of magnitude (then refined for instance by model inversion) instead of a robust pinpoint value.
In the following comparing: 1- the OBS technique with other geometrical techniques, and 2- the dual porosity approach (handling matrix block sizes $s_j$) with finely discretized networks, we address the relevance of the simplified expression in (14) under the assumption that the skeleton of the fracture network is known (as is the case with other geometrical methods). We prescribe to each fracture a uniform aperture and a uniform fracture permeability. The skeleton is then discretized and the entries $k_{ij}$ of the permeability tensor are calculated by performing numerical "permeameter" experiences (i.e., calculating fluid fluxes between opposite facets of the fractured block under prescribed Dirichlet boundary conditions while the other facets of the block are of no-flow type).

3. Comparison with structural-geometrical approaches

As shown from a theoretical standpoint, the OBS technique defines a mean matrix block size as a measure drawn from geometrical and structural properties of a discrete fracture network and its equivalent representation via a regular "sugar cube" network. In theory, no reference to any calculation of flow at the large scale is evoked in obtaining the OBS matrix block size, which renders the technique comparable in its spirit to other previous approaches also based on geometrical-structural characteristics of the discrete fracture network.

In the following, the OBS evaluation of matrix block sizes is compared with three other types of geometrical calculations, namely: the geometrical imbibition (GI) technique (Bourbiaux, 1997), the extended geometrical imbibition (EGI) technique (Bourbiaux et al., 2006), and the mean spacing (MS) technique (Narr, 1993). The main backgrounds of GI, EGI, and MS are summarized (sometimes slightly enhanced, as for EGI) and presented with notations consistent with that of the present work in Appendix B. GI and EGI techniques are only applicable (in their original version) to two-dimensional fracture networks and model the distance between any location in the matrix and the nearest fracture of the DFN. MS is
available for two- and three-dimensional systems and infers the mean lag distance between two neighbor fractures along the main directions of flow in a fractured block. All the geometrical methods need the detailed geometry of the DFN, although OBS could be used without it (See Section 2). But for a fair comparison we assume for all methods that the skeleton of the fracture network is known.

The comparison of OBS, GI, EGI, and MS is conducted for the two horizontal directions of a three-dimensional fractured block (100 m on a side) consisting of two families of near-vertical fracture planes. In the first test, a fracture family denoted A, is oriented with an azimuth of 100° counted positive anticlockwise from the main direction $x$ of the fractured block. The second fracture family denoted B is oriented 10°. The spacing between fractures of family $B$ is kept constant at 7 m, as the spacing of family $A$ is varied between 2 and 8 m for different realizations of the DFN (two examples reported in Fig. 3).

![Fig. 3. Examples of random discrete fracture networks (DFN) with two near-vertical fracture families at the scale of a reservoir grid cell. The azimuths of family $A$ and $B$ are 100° and 10°, respectively. DFN a: family $A$ (resp. $B$) with mean spacing of 2 m (resp. 7 m); DFN b: families $A$ and $B$ with mean spacing of 7 m.](image-url)
If we denote as $s_x$ and $s_y$ the mean matrix block sizes along the $x$ and $y$ horizontal directions of the fractured block, in view of the orientations of fracture families $A$ and $B$, $s_x$ should be close to the mean spacing of $A$ (i.e., 2 – 8 m), and $s_y$ close to the spacing of $B$ (i.e., 7 m). Fig. 4 reports on sought values of $s_x$ and $s_y$ for different methods of calculation with specifically the EGI technique rendering two sets of measures (see Appendix B) - small-EGI, large-EGI – as the technique assumes the existence of two types of matrix block interacting with the fracture network during flow.

![Diagram](image.png)

**Fig. 4.** Mean matrix block sizes $s_x$ and $s_y$ as functions of the spacing of fracture family $A$ (fracture networks in Fig. 4) for different methods of calculation. OBS = oriented block size method, GI = general imbibition method, EGI = enhanced general imbibition method (with "small" and "large" sizes of matrix blocks), and MS = mean spacing method.

In general, the OBS calculations retrieve the expected values of $s_x = 2 - 7$ m as a function of the spacing of fracture family $A$ (Fig. 4a). The size $s_y$, which should be constant at...
7 m, actually evolves with the spacing of family A and is overestimated of 10% to 50% (Fig. 4b). This overestimation cannot be the consequence of an actual fracture network that would be far from a regular WR representation since the actual network is simple and made of two perpendicular fracture families with directions almost parallel to the x and y directions of the fractured block. Nevertheless, we noted that increasing the spacing of the fracture family A also diminished the connectivity of the DFN with a few subdomains almost free of any fracture and poorly connected to the facets of the fractured block. It is noteworthy that estimates of effective properties of the DFN, especially porosities (or their influence on macroscopic permeabilities in Eq. (15)), both at the facets and inside the block are key features to the OBS calculations (see Section 2). Since less connected networks return weaker porosity values, the equivalent WR network assigned with those porosities will contain less fractures and result in increased matrix block sizes extracted from the equivalence between the WR network and the DFN.

Compared with the expected values, matrix block sizes $s_x$ and $s_y$ extracted from the GI technique tend to be overestimated. This result is foreseeable because GI usually experiences some difficulties when dealing with DFN encompassing both small and large matrix blocks. These difficulties are the consequence of the oversimplified fitting with a second degree polynomial of the so-called invasion area curve calculated by the method as the surface area in the matrix domain located at a given distance from the closest fracture of the system (see Appendix B). Regarding EGI, the "small block" estimates $s_x$ are in the correct range 2-8 m when the size $s_y$ is always overestimated. For their part, the "large block" estimates in EGI are always more than twice the expected values. Finally, the MS method infers correct values of $s_x$ and $s_y$ whichever the investigated DFN and the spacing of fracture families A and B. Notably, the MS method is weakly influenced by the fracture network connectivity which might become a drawback when dealing with sparse and poorly connected
fracture networks. In that case MS will still measure the mean lag distance separating two
eighbor fractures, as a poorly connected network tends to conceal a few cluster of large
matrix blocks in the system. In that case mean matrix block sizes from MS would be
underestimated.

In the OBS technique, whose specificity is seeking the equivalence between the actual
DFN and a regular WR network, this equivalence seems intuitively easier to achieve for
DFNs with fracture families whose principal orientations are close to the main directions of
the whole block. Therefore, it makes sense to address the capabilities of the method under less
favorable conditions where actual fractures do not line up with the main block directions. We
re-handled the comparison of matrix block sizes drawn from fracture networks still made of
two almost vertical fracture families, but this time with a constant spacing of 3 m for family
A, 5 m for family B, and varying the orientation of the families with respect to the main
directions x and y of the block. The fracture family A is still oriented 100° (counted positive
anticlockwise) with respect to the x direction and the orientation of family B is varied between
0 and 70° with respect to x (Fig. 5). In view of the geometrical settings of the DFNs, the
matrix block size $s_x$ should be close to 3 m and $s_y$ close to 5 m when the fracture family B is
almost orthogonal to family A (azimuth of $B = 0-10^\circ$). Block sizes $s_x$ should then slightly
decrease as $s_y$ should increase when the direction of fracture family B departs from
orthogonality with A.

The GI method systematically overestimates both $s_x$ and $s_y$ in each configuration of
the fracture network. The EGI technique still tends to overestimate $s_x$ and $s_y$ with its "large
block" measure while correct or slightly underestimated values are found with the "small
block" measure. In any case, both GI and EGI are weakly sensitive to the fracture family
orientations with almost constant values $s_x$ and $s_y$ irrespective of the azimuth prescribed to
fracture family $B$ in the DFN (Fig. 6). This result is consistent with the fact that both techniques model the surface occupied by matrix domains in the fractured block as a function of the distance to the nearest fracture (Appendix B). This measure reveals far less sensitive to fracture orientations than to fracture spacing.

Fig. 5. Examples of random discrete fracture networks (DFN) with two near-vertical fracture families at the scale of a reservoir grid cell. The mean spacing of fracture families $A$ and $B$ are prescribed at 3 m and 5 m, respectively, while the azimuth of family $A$ is kept at 100° and the azimuth of family $B$ is varied between 0° (DFN a) and 70° (DFN b).

Fig. 6. Mean matrix block sizes $s_x$ and $s_y$ as functions of the azimuth of fracture family $B$ (fracture networks in Fig. 5) for different methods of calculation. OBS = oriented block size method, GI = general imbibition method, EGI = enhanced general imbibition method (with "small" and "large" sizes of matrix blocks), and MS = mean spacing method.
The MS and OBS techniques infer very similar matrix block size values, these being sometimes slightly underestimated by MS and slightly overestimated by OBS. For azimuths of the fracture family $B$ between 0 and $45^\circ$, the estimated $s_x$ with both MS and OBS are close to the expected value of 3 m and stay almost constant whichever the orientation of family $B$. Concerning $s_y$, the expected value of 5 m is retrieved by OBS and underestimated at 3-4 m by MS. For azimuths of the fracture family $B$ between $45$ and $70^\circ$, both methods return, as expected, $s_x$ values that slightly decrease, as $s_y$ values increase from approximately 5 m up to 8 m. OBS mainly captures the projection of the fracture planes onto the facets delimiting the fractured block (see Section 2 and Appendix A), which is obviously sensitive to fracture orientations. In the same vein, MS evaluates the mean distance between fractures along the main directions of the fractured block with the obvious consequence of increasing the apparent distance when fracture planes are not normal to the direction of measure. Nevertheless, both methods provide valuable results for dense fracture networks or fractured blocks wide enough to enclose a large number of fractures allowing for significant statistical measures of fracture spacing (MS) or block-side and inner-block hydraulic properties (OBS).

Notwithstanding other considerations such as computation times (see hereafter), OBS and MS techniques seem to outperform GI and EGI in extracting mean matrix block sizes from fractured system. We noted however that OBS is sensitive to the loss of connectivity in a fracture network with the consequence of increasing the inferred matrix block size. This artificial increase might result in biased evaluations of fluid flux exchanges between fracture and matrix media. Numerical exercises comparing discrete fracture network outputs and their dual porosity representation with OBS-sized matrix blocks are conducted to answer this question. The other geometrical techniques GI, EGI, and MS are also tested. We remind that these three numerical techniques are in essence only applicable when a prior knowledge of the fracture network geometry is available, while the OBS technique might by applied either on
known or unknown geometries (see Section 2). For a fair comparison of all techniques hereafter, we consider that the fracture network geometry is known.

4. Two-dimensional numerical test cases

As already mentioned, dual continua representations of discrete fracture networks are conducive to drastic reductions in computation costs but require carefully designed settings to adequately represent both conductive and capacitive properties of a fractured porous medium subject to Darcian flow. We address here two phase flow in both DFN and dual porosity models. The setup of calculations is dimensioned to represent large laboratory analogs of flow in fractured media as conducted for instance in "Hele-Shaw" cells (e.g., Park and Homsy, 1984; Folch et al., 1999). We remind that we are interested in the assessment of mean matrix block size from different geometrical-structural techniques that always manipulate relative quantities as the spacing of fractures compared to the block size, or fracture traces intercepted by block facets. Therefore, our findings from numerical experiments at the scale of a lab device should not be hampered by loss of generality. In addition, we perform calculations, especially in the context of DFN discretization, over synthetic fracture networks with regular fracture orientations. This choice reduces discretization efforts but is mainly employed herein because it ensures accuracy of reference calculations in a DFN compared with that from a dual porosity model. Even though sophisticated meshing techniques and advanced numerical methods exist, it was found that thin fracture elements in unstructured meshing tend to smear the calculation of their state variables over the large matrix blocks. This feature is not suited to compare (local) DFN and (large scale) dual porosity calculations of diffusive flow.

Numerical simulations are performed over two-dimensional horizontal fractured systems (of unit thickness) that only neglect gravity-driven flow. Notably, the various
techniques employed in this study to calculate mean matrix block sizes are not sensitive to
gavity-driven flow and only manipulate geometrical considerations on the fracture network
or equivalences in permeability-porosity between an actual fractured block and a sugar-cube
model. Two-phase flow in a DFN is performed over a fractured system of 3 m length and 1.5
m width finely discretized by 11590 square elements for an accurate representation of both
the fracture network and matrix. The system is also roughly discretized by only 920 square
elements of a dual-porosity, single-permeability model with matrix block sizes extracted from
the DFN via the EI, EGI, MS and OBS techniques (see Section 3). Two-phase flow is
numerically solved by means of a finite volume technique and uses an implicit-in-time
scheme for time integration of the pressure equation while an explicit-in-time scheme is used
for time integration of either the water or oil mass balance. To avoid unfair comparisons
between GI, EGI, MS, and OBS, a simple fracture network is delineated with fractures only
parallel to the main flow directions $x$ and $y$ of the system. Dead ends of the fracture network
are also removed since in essence they are always accounted for in the fracture-matrix
relationship by GI and EGI methods when MS and OBS might not see these dead-ends
because they are not counted in MS or do not participate to side-block properties in OBS.

The first fractured system investigated (Fig. 7) is initially saturated with oil and
percolated by water injected from the western boundary taken as a Neumann condition
prescribing a constant-in-time water flux. The eastern boundary of the system is of Dirichlet
type while North and South boundaries are of no-flow type. Table 1 indicates the local
hydraulic properties of each medium (fractures, matrix) in the DFN, Table 2 reports on inner-
block and block-side properties used by the OBS method to calculate matrix block sizes, and
Table 3 gathers the various matrix block sizes $s_x$ and $s_y$ obtained from the GI, EGI, MS and
OBS methods.
Fig. 7. Two-dimensional fracture network serving as a system finely discretized or handled as a dual-porosity model for the purpose of flow dynamics comparison. The size of matrix blocks in a dual porosity approach are reported as colored frames, from left to right: Red = oriented block size method, Blue = general imbibition method, Orange = mean spacing imbibition method, Green and Purple = small and large sizes from enhanced general imbibition method.

<table>
<thead>
<tr>
<th></th>
<th>Matrix medium</th>
<th>Fractures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Porosity $\phi$ [-]</td>
<td>0.1</td>
<td>1</td>
</tr>
<tr>
<td>Permeability $k$ [$10^{-15}$ m$^2$]</td>
<td>1 and 10</td>
<td>10000</td>
</tr>
<tr>
<td>Relative permeability $k_r$ [-]</td>
<td>Brooks Corey ($\lambda=2$)</td>
<td>&quot;Cross&quot; $k_r$</td>
</tr>
<tr>
<td>Capillary pressure $P_c$ [bar = $10^4$ kgms$^{-2}$]</td>
<td>Brooks Corey ($\lambda=2$)</td>
<td>Null capillary pressure</td>
</tr>
</tbody>
</table>

Table 1. Set up of main flow parameters for calculations of two-phase flow in fractured systems depicted in Fig. 7 and 10. The relative permeability and capillary pressure as functions of water saturation in the matrix obey the Brooks and Corey model (1964) with $\lambda$ (=2) the so-called pore-size distribution index.
Table 2. Main macroscopic parameters of the fractured block in Fig. 7 to infer via the oriented block size technique the mean matrix block size of a dual porosity model. $k_f$, $\phi_f$ respectively are the permeability and porosity of the whole block, $k^{FN-S}$, $\phi^{FN-S}$ respectively are the permeability and porosity of the fracture network at the sides (normal to $x$ and $y$ directions) of the block.

<table>
<thead>
<tr>
<th>Block sizes</th>
<th>OBS</th>
<th>GI</th>
<th>MS</th>
<th>EGI-large</th>
<th>EGI-small</th>
</tr>
</thead>
<tbody>
<tr>
<td>$s_x$ [m]</td>
<td>0.360</td>
<td>0.6</td>
<td>0.35</td>
<td>0.696</td>
<td>0.257</td>
</tr>
<tr>
<td>$s_y$ [m]</td>
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<td>0.3</td>
<td>0.345</td>
<td>0.494</td>
<td>0.257</td>
</tr>
</tbody>
</table>

Table 3. Mean matrix block sizes of a dual porosity model as a surrogate to the discrete fracture network in Fig. 7.

Two different types of flow are simulated, the first one with low water injection rate of 0.1 m/day in the fractures and low matrix permeability of $10^{-15}$ m$^2$, the second one with higher injection rate of 1 m/day and higher matrix permeability of $10^{-14}$ m$^2$. On the one hand, the first scenario with small water fluxes in the fractures and weakly permeable matrix enhances capillary effects as the origin of pressure gradients between fracture and matrix and subsequent exchange rates between both media. Due to capillary effects in the matrix and absence of these in the fractures, the oil pressure in the matrix is higher than that in the fractures and oil is ejected from the matrix (or water invades the matrix). On the other hand, the second scenario with high injection velocities favors "piston" flow in the fractures and enhances fracture-matrix exchanges as the consequence of the excess of water pressure in fractures compared with oil pressure in the matrix. Water invades the matrix and the process...
is enhanced by the contrast of mobility (the ratio $kr/\mu$) between oil and water phases which triggers rapid water invasion along the fractures and early leaching of matrix blocks.

To reinforce these assertions about flow scenarios with contrast between capillary and viscous forces to extract oil from matrix blocks, we also calculated a dimensionless capillary number based on the evaluation of water fluxes invading matrix blocks versus expulsion of oil from the matrix to fractures by capillary pressure contrasts. With steady-state flow sweeping oil from the system by forced water injection at one side of the fractured block, the mean water pressure gradient in the system is evaluated as

$$\left| \nabla P_w \right| = \frac{V_{\text{inj}} \mu_w}{k^f}$$  \hspace{1cm} (15)

$P_w$ [ML$^{-1}$T$^{-1}$] is the water pressure in both the fractures and the matrix, $V_{\text{inj}}$ [LT$^{-1}$] is the injection velocity of water at the upstream side of the fractured block, $\mu_w$ [ML$^{-1}$T$^{-1}$] is the dynamic viscosity of water, and $k^f$ [L$^2$] is the equivalent fracture permeability of the whole block along the direction $i$ of water injection. Regarding the capillary pressure gradient, we assume a null capillary pressure in the fractures (open medium of unit porosity) and we take in the matrix the maximal capillary pressure $P^c_{\text{max}}$ given by relationships capillary pressure – saturation (see, e.g., Table 1). The capillary pressure gradient between matrix and fractures is then approximated as

$$\left| \nabla P^c \right| = \frac{P^c_{\text{max}}}{s_{\text{min}}/2}$$  \hspace{1cm} (16)

with $s_{\text{min}}$ [L] the smallest dimension (in either directions $x$, or $y$ or $z$) of the mean matrix block size. A dimensionless capillary number balancing capillary gradient with water pressure gradient can be expressed as

$$n_c = \frac{\left| \nabla P^c \right|}{\left| \nabla P_w \right|} \approx \frac{2P^c_{\text{max}}k^f}{s_{\text{min}}\mu_w V_{\text{inj}}}$$  \hspace{1cm} (17)
This capillary number is larger than one for flow conditions dominated by capillary forces as it becomes close to one or less than one when viscous forces condition flow in the fractured block.

Fig. 8. Maps of water saturation in a water-flooding two-phase flow scenario. Calculations are performed over a fine grid discretizing both the fracture network and matrix (system in Fig. 7). The system is initially saturated in oil and water is injected in the fractures at a constant flow rate at the western boundary of the system. Oil recovery is monitored at the eastern boundary (see Fig. 9). The fluid exchange between fractures and matrix is dominated by capillary forces in map a as both capillary and viscous forces are active in map b.

In the DFN approach where matrix-fracture exchanges are dominated by capillary effects \( n_c = 4.1 \) with the settings of the simulations, water does not deeply invade the matrix (Fig 8.a,) while for the same injected water pore volume, high injection velocity and piston flow \( n_c = 0.41 \) maintains higher water pressure gradients that help to a deeper water invasion of the matrix (Fig. 8.b). Calculations in the DFN serve as reference to the comparison of flow scenarios between dual porosity models assigned with matrix block size from the GI, EGI, MS and OBS methods (sizes of blocks are pictured in Fig. 7). The comparison is here performed by way of a single indicator defined as the evolution in time (precisely, the evolution with the water pore volume injected in the system) of the oil recovery ratio at the outlet of the fractured system. This oil recovery corresponds to the ratio of the cumulative volume of oil exiting the system to the total initial volume of oil in the system. This indicator is obviously macroscopic, with the meaning that it monitors the
behavior of the system at the large scale (at least the homogenization scale of the fracture network). It would not make sense to compare a local feature of the fracture network (e.g., the pressure transients in a single fracture) with averaged behaviors obtained for the large blocks (cells) of a dual porosity approach.

Fig. 9. Oil recovery ratio versus water injected pore volumes at the eastern boundary of a fractured network (in Fig. 7). The so-called reference is calculated by means of a finely discretized network as the other curves are drawn from a dual porosity model with various mean matrix block sizes. Results from the mean spacing technique for matrix block size evaluation are not reported because they are merged with those from the oriented block size technique. Capillary forces dominate the exchange rate between fractures and matrix in plot a, as both capillary and viscous forces are active in plot b.

Fig. 9 presents two plots of the oil recovery ratio as a function of the injected pore volume and stemming from flow scenarios with low and high injection velocities. The same oil recovery ratio of approximately 60% is reached for both flow scenarios, but with only 5 pore volumes in the case of high injection velocity compared with the 50 pore volumes required by the case of low injection velocity. No dual porosity model with their different matrix block size renders results that completely depart from the reference calculations in the DFN. Since matrix block sizes calculated with OBS and MS techniques are quite similar (see...
Fig. 7 and Table 3), the results from dual porosity model simulations do not differ significantly and only outputs from the OBS technique are reported in Fig. 9.

The OBS technique tends to slightly underestimate the matrix block size which triggers a quicker oil extraction from the matrix and produces recovery curves slightly shifted toward short injection times. The simulations handling the GI matrix blocks are also in very good agreement with references, especially in the case of fracture-matrix exchanges enhanced by high water injection rate. The matrix block sizes of EGI are still underestimated by the "small block" measure and overestimated by the "large block" measure giving rise to respectively faster and slower evolutions of the oil recovery ratio with respect to time. As such, the EGI technique is not the most accurate to calculate matrix block sizes and should be employed as a convenient way to provide minimal and maximal bounds to these sizes.

Notably, the fractured system discussed above does not significantly distinguish between GI and OBS in terms of accuracy whichever the mechanism prevailing in fluid flux exchanges between fractures and matrix. Nevertheless, we are reminded that the reference fracture network was built to mitigate GI downsides. Fracture dead-ends were removed from the network and the two fractures families were set parallel to the $x$ and $y$ directions of the fractured block, thus allowing the GI method to infer a precise "invasion curve" ($A(X)$ in Appendix B). This is why GI shows good performances in the present test cases as it exhibited more discrepancies in the geometrical test cases discussed in Section 3.

At this stage, it must be raised that the OBS technique partly relies upon evaluations of block-side properties such as fracture porosity and permeability, the latter being eventually not representative of inner-block quantities when the portion of fractures intercepting the block sides are not representative of the network geometry inside the block. To address the eventual influence of this downside, we recalculated the two flow scenarios discussed above for another fractured system (Fig. 10) which comprises a few long fractures located very close
to the sides of the system. These fractures delimit a few very elongated matrix blocks close to
the boundaries of the system (those encircled in Fig 10) as the majority of matrix blocks
inside the system are rectangular with a ratio length to width barely exceeding a factor 3. As
expected, the inner-block and block-side properties used by the OBS method (Table 4) differ
from that of the fractured "regular" system previously discussed.

Fig. 10. Two-dimensional fracture network serving as a system finely discretized or handled as a dual-porosity
model. Fractures close to the boundaries delimit very narrow matrix blocks (encircled) that depart from the shape
of blocks within the fracture network. The identified sizes of matrix blocks in a dual porosity approach are
reported as colored frames, from left to right: Red = oriented block size method, Blue = general imbibition
method, Orange = mean spacing imbibition method, Green and Purple = small and large sizes from enhanced
general imbibition method.

<table>
<thead>
<tr>
<th></th>
<th>x direction</th>
<th>y direction</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k_f$ [10^{-15} m^2]</td>
<td>107.72</td>
<td>240</td>
</tr>
<tr>
<td>$k^{FN-S}$ [10^{-15} m^2]</td>
<td>106.66</td>
<td>240</td>
</tr>
<tr>
<td>$\phi^{FN-S}$ [-]</td>
<td>0.0106</td>
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</tr>
<tr>
<td>$\phi_f$ [-]</td>
<td>0.024</td>
<td></td>
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</tbody>
</table>

Table 4. Main macroscopic parameters of the fractured block in Fig. 10 to infer via the oriented block size
technique the mean matrix block size of a dual porosity model. $k_f$, $\phi_f$ respectively are the permeability and
porosity of the whole block, $k^{FN-S}$, $\phi^{FN-S}$ respectively are the permeability and porosity of the fracture
network at the sides (normal to x and y directions) of the block.
Comparing reference calculations performed over the DFN (maps of water saturation in the system reported in Fig. 11, capillary number $n_c$ of 19 for Fig. 11a and of 1.9 for Fig 11b) and calculations in the dual porosity models reveals that the oil recovery ratio is still of approximately 60% after 4-5 injected pore volumes for high injection velocity and 40 pore volumes under low injection velocity conditions (Fig. 12).

Fig. 11. Maps of water saturation in a water-flooding two-phase flow scenario. Calculations are performed over a fine grid discretizing both the fracture network and matrix (system in Fig. 10). The system is initially saturated in oil and water is injected in the fractures at a constant flow rate at the western boundary of the system. Oil recovery is monitored at the eastern boundary (see Fig. 12). The fluid exchange between fractures and matrix is dominated by capillary forces in map a as both capillary and viscous forces are active in map b.

Fig. 12. Oil recovery ratio versus water injected pore volumes at the eastern boundary of a fractured network (in Fig. 10). The reference curve is calculated by means of a finely discretized network as the other curves are drawn from a dual porosity model with various mean matrix block sizes. Results from the mean spacing technique are not reported because they are merged with those from the oriented block size technique. Capillary forces dominate the exchange rate between fractures and matrix in plot a, as both capillary and viscous forces are active in plot b.
This result confirms that the macroscopic behaviors of both the DFN and its representation as a dual porosity system are changed much by the few fractures that do not obey the general geometric and structural settings of the whole fractured block. This feature is also evidenced by the comparison between the maps of water saturation in Fig. 8 and Fig. 11 that only differ by the locations of fractures underlined by high water saturations. However, discrepancies between the reference (taken as the DFN) and the dual porosity approximations increase. As for the preceding example, OBS and MS techniques provide very similar matrix block sizes (these sizes are pictured in Fig. 10 and reported in Table 5) and similar dual-porosity behaviors making that MS results are not discussed in the following.

<table>
<thead>
<tr>
<th>Block sizes</th>
<th>OBS</th>
<th>GI</th>
<th>MS</th>
<th>EGI-large</th>
<th>EGI-small</th>
</tr>
</thead>
<tbody>
<tr>
<td>$s_x$ [m]</td>
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<tr>
<td>$s_y$ [m]</td>
<td>0.149</td>
<td>0.21</td>
<td>0.19</td>
<td>0.325</td>
<td>0.138</td>
</tr>
</tbody>
</table>

Table 5. Mean matrix block sizes of a dual porosity model as a surrogate to the discrete fracture network in Fig. 7.

In the case of matrix-fracture exchanges dominated by capillary forces (Figs 11a, 12a) the OBS technique overestimates the leaching of matrix block (and oil production at the outlet of the system) because the smallest matrix block size (here along the $y$ direction) is underestimated. Whichever the algebraic form chosen in OBS to infer the matrix block size (See Section 2), the method is in essence sensitive to fracture densities close to the boundaries of the whole fractured block, either in regard of porosities at the sides of the block or of permeability values in a "permeameter" type system. If the actual matrix block sizes close to the boundaries of the block are smaller than inside the block, as is the case with the present example, the smallest matrix block size (here along $y$, see Table 5) is underestimated which
favors rapid imbibition under capillary forces (see above the capillary number $n_c$). Notably, the GI technique is not sensitive to the few small matrix blocks of the DFN because it treats the shell and inner parts of the block exactly the same way. For its part, the "small" EGI technique underestimates the mean matrix block size as the "Large" EGI overestimates it ("small" EGI overestimates matrix imbibition and "Large" EGI underestimates imbibition, see oil recovery in Fig. 12a compared with reference).

When matrix-fracture exchanges occur as a conjunction of viscous and capillary forces (see the capillary number in (17) and subsequently evaluated for DFN simulations), the OBS technique renders results the closest to reference. The key is that rapid water invasion of the fractured block through permeable fractures (see Fig. 11b) and subsequent viscous effects between matrix and fractures are dominated by percolation through the large fractures and their (large) neighbor matrix blocks. As the OBS technique identifies the correct largest matrix block size (here along the $x$ direction, see Table 5), flow simulations with a dual porosity model are convincing. This time, the GI technique underestimates oil recovery, as "Large" EGI does too, because the overestimated matrix block size (especially along the $x$ direction, see Table 5) is favorable to capillary imbibition but hampers water invasion along fractures and matrix block leaching at early injection times.

Finally, regarding performances in terms of computation costs, the different geometric methods were applied to a large DFN represented as a synthetic dual porosity reservoir of 1.05 million grid cells. For OBS and MS methods, matrix block size calculations were performed for each elementary cell and duplicated over all cells of the reservoir with total CPU times coming up as: 230 s for OBS and 1120 s for MS. Notably, the time counted for GI and EGI is that of calculations over a limited number of cells "strategically" sampled in the whole grid of the dual porosity reservoir, yielding a fair representation of the system after 1800 s of calculation. With approximately 4 s of calculation per cell and $10^6$ cells, identifying
a matrix block size for each cell with GI and EGI methods would render impracticable evaluations exceeding 45 days. When applied to known DFNs, both OBS and MS require a pre-evaluation of the diagonal permeability tensor of the fractured block; by construction for OBS (see Section 2) and to identify main flow directions in MS for which random lines counting the spacing of fractures (see Appendix B) are parallel to these directions. Differences of computations times between methods are in the straightforward (and fast) application of an analytical solution for OBS opposed to the need for many random draws in MS.

5. Conclusions

The Oriented Block Size (OBS) technique has been developed as a new way to infer the mean matrix block sizes in porous fractured media with application to dual porosity models of flow at the large scale. Matrix block sizes are calculated by seeking the equivalence in terms of fracture permeability and fracture porosity between a fractured block and a Warren and Root discrete fracture network made of three fracture families with regular spacing and fracture planes normal to the main flow directions.

Two expressions of the OBS are available according to which type of fractured block the method is applied. The first expression is well suited to infer matrix block sizes over synthetic discrete fracture networks or well-known actual networks since it requires identifying fracture porosity of the network, fracture porosity at the sides of the fractured block, and the diagonal permeability tensor of the whole block (which can be calculated analytically or numerically). This first expression is based on a rigorous algebraic development which reveals precise and renders matrix block sizes close to expectations drawn from various synthetic discrete fracture networks. The second expression is derived from the first one via assumptions on the fracture porosities of the block. It has the advantage of being
applicable to hardly accessible fracture network as encountered in the field. This second expression is compatible with an inference from field measurements such as hydraulic tests and observations in wells but should only render orders of magnitude instead of pinpoint values. Further works should address how matrix block sizes are influenced by uncertainty on available field data.

The OBS technique revealed much faster in terms of computation times compared with other available geometrical techniques developed to infer matrix block sizes. This feature is a promising avenue for tentative applications of the method in up-scaling the representation of huge fractured reservoirs as done for instance in the oil industry when optimization of oil recovery from various exploitation scenarios is planned. In this context, OBS and its precise evaluation of matrix block sizes is useful to the parameterization of dual porosity models for two phase flow either dominated by capillary forces or viscous forces. However, as the other methods, the OBS technique may fail in retrieving matrix block sizes within poorly connected fracture networks. It is worth to note however that poorly connected networks are not valuable candidates to homogenization into a dual porosity model.

Finally, it must also be raised that OBS is associated with the identification of large scale permeability tensors that are mostly sensitive to the backbone of a fractured network and do not see fracture dead-ends. In the case of applications relying upon data from hydraulic well tests, the type of occurring flow should be carefully considered. Two phase flow, mostly witnessed by the propagation of an oil/water saturation front, will mainly record the effects of the backbone, as single phase flow, mainly monitored by the transient evolution of water pressure heads, would also be sensitive to dead-ends. It deserves some additional synthetic test cases or confrontation to actual field data to see whether or not the OBS technique reveals suited in these instances.
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Appendix A. Matrix block sizes extracted from the equivalence between an actual fractured block and a Warren and Root (WR) block.

We remind that an actual and well connected fractured block oriented with its main directions along the main directions of flow indexed by $i=1,2,3$, can be characterized by mean permeabilities $k_{i}^{FN-S}$ along the sides of the block as

$$k_{i}^{FN-S} = \frac{1}{2(\Delta_{i+1}\Delta_{i+2})} \left( \sum_{n=1}^{N_{i-}} k_{n} l_{n} e_{n} + \sum_{n=1}^{N_{i+}} k_{n} l_{n} e_{n} \right)$$  \hspace{1cm} (A1)

The block size in direction $i$ is denoted $\Delta_{i}$ and the sides delimiting the block are also indexed by $i$ but for limits normal to the main direction $i$. In addition, block sides are labelled $i-$ or $i+$ according to their respective location upstream or downstream along direction $i$. In (A1), $i$ is a cycling index such that, e.g., $i+1 = 3$ when $i = 2$ and $i+1$ returns to 1 when $i=3$. The sides $i-$ and $i+$ of the block are intercepted by a number of fractures $N_{i-}$ and $N_{i+}$, and $k$ is the local permeability of a fracture intercepting the side of the block over an apparent length $l$ and with apparent fracture aperture $e$.

We also remind that a Warren and Root (WR) block concealing a regular fracture network of three fracture families can be characterized by two expressions associating: 1- the diagonal tensor of permeability of the whole block $k^{WR}_{i} (i = 1,2,3)$, 2- the mean porosity of the block $\phi^{WR}$, and 3- the porosity of the block sides $\phi^{WR-S}_{i} (i = 1,2,3)$, with the spacing $s_{i} (i = 1,2,3)$, the aperture $e_{i} (i = 1,2,3)$, and the local permeability $k_{i} (i = 1,2,3)$ of the three fracture families composing the WR block (for details, see Section 2). These expressions are

$$k_{i} = \frac{1}{2} \left( 1 + \frac{s_{i}}{e_{i}} \right) \sum_{j=1}^{3} (-1)^{\delta_{ij}} k^{WR}_{j}$$  \hspace{1cm} (A2)

$$k_{i} = \frac{1}{2(\phi^{WR} - \phi^{WR-S})} \sum_{j=1}^{3} (-1)^{\delta_{ij}} k^{WR}_{j}$$  \hspace{1cm} (A3)
where \( \delta_{i,j} \) is the Kronecker symbol, \( \delta_{i,j} = 1, i = j; \delta_{i,j} = 0, i \neq j \).

Following the idea that one can establish the equivalence between a WR network and an actual fractured block regarding their hydraulic properties, it is assumed that \( \Phi^\text{WR}, \Phi^\text{WR-S} \), and \( k^\text{WR} \) are similar to the equivalent properties in the actual fractured block, respectively denoted as \( \Phi^\text{FN}, \Phi^\text{FN-S} \), and \( k^\text{FN-S} \) (see A1, for the latter). In the same vein, if a WR network serves as reference for fixing model parameters of homogenized approaches to fractured media, the characteristics of a WR network can be substituted by parameters of the homogenized model. For example, the characteristics \( s_i, e_i, \) and \( k^\text{WR}_j \) in (A2) are respectively substituted by a mean matrix block size (also denoted \( s_i \) as defined in (3)), a mean fracture aperture \( e_f \), and the entries of a diagonal tensor \( k^f_j \) of the homogenized model. With these transformations, equating (A2) and (A3) results in

\[
\left(1 + \frac{s_i}{e_f}\right) \sum_{j=1}^{3} (-1)^{\delta_{i,j}} k^f_j = \frac{1}{(\Phi^\text{FN} - \Phi^\text{FN-S})} \sum_{j=1}^{3} (-1)^{\delta_{i,j}} k^\text{FN-S}_j ; \text{i.e.,} \ 
\]

\[
s_i = e_f \left( \frac{\sum_{j=1}^{3} (-1)^{\delta_{i,j}} k^\text{FN-S}_j}{(\Phi^\text{FN} - \Phi^\text{FN-S}) \sum_{j=1}^{3} (-1)^{\delta_{i,j}} k^f_j} \right) - 1 \quad (A4)
\]

For the sake of simplification (see hereafter), the term \(-1\) in the expression of \( s_i \) can be dropped by considering that the term in \((\Phi^\text{FN})^{-1}\) is much larger than one for usual fracture porosity of a rock block barely exceeding a few percent. Stated differently, one might also consider in (A4) that the matrix block size \( s_i \) is much larger than the fracture aperture \( e_f \) and results in

\[
s_i \approx e_f \frac{\sum_{j=1}^{3} (-1)^{\delta_{i,j}} k^\text{FN-S}_j}{(\Phi^\text{FN} - \Phi^\text{FN-S}) \sum_{j=1}^{3} (-1)^{\delta_{i,j}} k^f_j} \quad (A5)
\]
The mean matrix block size $s_i$ in (A5) depends on both the mean fracture aperture $e_f$ and the fracture permeability of a homogenized model $k_f^j$. It is noteworthy that $e_f$ is usually not a parameter of a homogenized approach, and it makes sense to render (A5) (partly) independent of any conjecture on the value of $e_f$. To this end, it is reasonably assumed that a WR network has its matrix block sizes separating neighbor fractures independent of the apertures $e_i$ of the fractures. Stated differently, it is assumed that a WR network with a uniform aperture $e_f$ for its three fracture families can be found as equivalent to a WR with its three fracture families with apertures $e_i$. With a uniform aperture $e_f$, a WR network would render a value $\phi^{WR} - \phi_i^{WR-S} = e_f N_f / \Delta_i$ with $N_f$ the number of fractures in the family $i$, and $\Delta_i$ the size of the whole fractured block along direction $i$. If the values $\phi^{FN} - \phi_i^{FN-S}$ were not replacing their equivalent $\phi^{WR} - \phi_i^{WR-S}$ in Eq (A5), the latter would no longer depend on $e_f$.

Hence, our proposal is to calculate porosities of the actual fracture network by assigning the whole skeleton of the network with a constant single-fracture aperture $e_f$. The fracture network porosities for a constant aperture $e_f$ would write as

$$\left(\phi^{FN} - \phi_i^{FN-S}\right) = e_f \left(\phi^{FN*} - \phi_i^{FN-S*}\right)$$

(A6)

The terms $\phi^{FN*}$, $\phi_i^{FN-S*}$ [$L^{-1}$] denote porosities of the actual fracture skeleton per unit fracture aperture (that can be calculated by assigning a uniform fracture aperture of 1 to the whole fracture network). Substituting (A6) in (A5) simplifies the formulation of the matrix block size into

$$s_i \approx \frac{\sum_{j=1}^{3} (-1)^{3-j} k_f^{FN-S}}{\left(\phi^{FN*} - \phi_i^{FN-S*}\right) \sum_{j=1}^{3} (-1)^{3-j} k_f^{j}}$$

(A7)
The main characteristic of (A7) is that the mean matrix block size depends: 1- on a mean permeability tensor \( k_{ij} \) of fractures at the scale of a (mesh of a) homogenized model of the system (e.g., a conjecture of the fracture permeability in a dual porosity model), 2- on the facet permeability values of the actual fracture network \( k_{ij}^{FN-S} \), and 3- on structural properties of the actual network resulting in fracture porosity values of the whole fractured block and its sides \( \phi_{FN}^* \) and \( \phi_{FN-S}^* \), respectively. These features make that the form in (A7) is hardly applicable to poorly-known natural systems and should be mainly used in problems dealing with homogenization of systems with well-known geometry and discretization of synthetic fracture networks and matrix blocks.

Nevertheless, another form of the mean matrix block size can be proposed. By manipulating (A1), the permeability of the actual fracture network at the facets of the whole fractured block can be rewritten as

\[
k_{ij}^{FN-S} = \frac{\left( \sum_{n=1}^{N_f} l_n e_n + \sum_{n=1}^{N_i} l_n e_n \right)}{2 (\Delta_{j+1} \Delta_{j+2})} \times \frac{\left( \sum_{n=1}^{N_f} k_n l_n e_n + \sum_{n=1}^{N_i} k_n l_n e_n \right)}{\left( \sum_{n=1}^{N_f} l_n e_n + \sum_{n=1}^{N_i} l_n e_n \right)} ; \text{ i.e., (A8)}
\]

\[
k_{ij}^{FN-S} = \phi_{ij}^{FN-S} \bar{k}_i \quad \text{with} \quad \bar{k}_i = \frac{\left( \sum_{n=1}^{N_f} k_n l_n e_n + \sum_{n=1}^{N_i} k_n l_n e_n \right)}{\left( \sum_{n=1}^{N_f} l_n e_n + \sum_{n=1}^{N_i} l_n e_n \right)}
\]

The tensor components \( \bar{k}_i (i=1,2,3) \) in (A8) are an arithmetic mean of single-fracture permeability values weighted by open fracture surface areas at the sides of the whole fractured block. If we assume that these mean values are equal, irrespective of the facet of the fractured block (which also can go with fractured systems candidates to homogenization), it also means that the eventual anisotropy of permeability in the fracture network is just the consequence of fractures densities normal to the flow directions, i.e., \( k_{ij}^{FN-S} = \phi_{ij}^{FN-S} \bar{k} \).

Notably, this strong assumption stating that one can define a constant single-fracture
permeability value $\kappa$ also goes with the existence of an equivalent uniform single-fracture
aperture $e_f$ for the whole fracture network. Reintroducing in (A5) the expression (A8) with a
constant value $\kappa$ and making use of rescaled porosities defined in (A6) as $\phi_i^{\text{FN-S}} = e_f \phi_i^{\text{FN-S^*}}$
comes down to

$$s_i \approx \frac{e_f \kappa \sum_{j=1}^{3} (-1)^{\delta_{i,j}} \phi_j^{\text{FN-S^*}}}{(\phi_i^{\text{FN-S^*}} - \phi_j^{\text{FN-S^*}}) \sum_{j=1}^{3} (-1)^{\delta_{i,j}} k_f^j} \tag{A9}$$

It can also be shown that a WR network with constant aperture $e_f$ for its three fracture
families has block and side porosities following the relation $\sum_{j} \phi_j^{\text{WR-S^*}} = 2 \phi_i^{\text{WR^*}}$. If the rescaled
DFN is equivalent to the WR network, then one can state that $\sum_{j} \phi_i^{\text{FN-S^*}} = 2 \phi_i^{\text{FN^*}}$. Noting that

$$\sum_{j=1}^{3} (-1)^{\delta_{i,j}} \phi_j^{\text{FN-S^*}} \text{ can also be rewritten as } \sum_{j=1}^{3} \phi_j^{\text{FN-S^*}} - 2 \phi_i^{\text{FN-S^*}} \text{ and reintroducing the preceding}$$

relationship between block and side porosities in (A9) results in

$$s_i \approx \frac{2 e_f \kappa}{\sum_{j=1}^{3} (-1)^{\delta_{i,j}} k_f^j} \tag{A10}$$

In the case of field applications with poorly known and hardly accessible fracture
networks, (A10) returns the mean matrix block sizes in a fracture network based on the field
evaluations of the permeability tensor $k^f$ of a whole fractured block, the average uniform
aperture $e_f$ and permeability $\kappa$ of a single fracture. Because the entries of (A10) are not
straightforward to obtain and may also be associated with important measurement errors, it is
expected that (A10) will only render orders of magnitude of mean matrix block sizes.
Appendix B. Structural-geometrical evaluations of a mean matrix block size in a dual continuum flow model

The geometrical imbibition (GI) method

The method has been developed for two-dimensional flow models only. Three-dimensional approaches are therefore handled as multilayer systems. For a two-dimensional image of an actual or synthetic fracture network, the first task to handle consists in mapping the image on a regular grid of square pixels. Each pixel is then assigned a value $d_f$ that represents the distance between the center of the pixel and the closest fracture of the network. One sums up the area of pixels whose distance $d_f$ is less than a prescribed value $X$, and the area is then normalized by the total surface area of the image to form the quantity $A(X)$. The resulting measure $A(X)$ (Fig. B1) is modeled as

$$ A(X) = \frac{2X}{a} + \frac{2X}{b} - \frac{4X^2}{ab} $$

with $a$ and $b$ the resulting mean size of the matrix block of a two-dimensional dual porosity model. $a$ and $b$ are obtained by minimizing the sum of squared errors between the model in (B1) and the actual measures of $A(X)$.

![Fig. B1. General imbibition technique to mean matrix block size identification. Normalized invaded matrix area $A(X)$ as a function of the distance $X$ between a location in the matrix and the closest fracture.](image)
**The extended geometrical imbibition (EGI) method**

This method improves the two-dimensional GI technique by assuming that two mean matrix block sizes characterize the relationships between fractures and matrix. For locations in the matrix close to fractures, two types of matrix block interact with fractures, whereas locations far from fractures are influenced by a single size of matrix block. This feature makes that the quantity $A(X)$ drawn from the mapping of the actual fracture network (see above the GI technique) is modeled by a discontinuous curve in the form

\[ A(X) = \alpha_1 \left( \frac{2X}{a_1} + \frac{2X}{b_1} - \frac{4X^2}{a_1b_1} \right) + \alpha_2 \left( \frac{2X}{a_2} + \frac{2X}{b_2} - \frac{4X^2}{a_2b_2} \right) ; \quad X \leq \frac{a_1}{2} \]

\[ A(X) = \alpha_2 \left( \frac{2X}{a_2} + \frac{2X}{b_2} - \frac{4X^2}{a_2b_2} \right) ; \quad X > \frac{a_1}{2} \]

with $(a_1, b_1), (a_2, b_2)$ the size of the small and large matrix blocks respectively. $\alpha_1, \alpha_2$ are the proportions of small (type 1) blocks and large (type 2) blocks with $\alpha_2 = 1 - \alpha_1$. The distance $X = a_1/2$ is the threshold beyond which a single type of large matrix block is sufficient to model interactions between fractures and matrix.

The inference of a single set of parameters $(\alpha_1, a_1, b_1, \alpha_2, a_2, b_2)$ by minimizing errors between the model in (B2) and actual measures of $A(X)$ is not straightforward because the subsets of parameters $(\alpha_1, a_1, b_1)$ and $(\alpha_2, a_2, b_2)$ are partly interchangeable to shape the same function $A(X)$. It is better suited to analyze the derivative $A'(X)$

\[ A'(X) = \frac{dA(X)}{dX} = \alpha_1 \left( \frac{2}{a_1} + \frac{2}{b_1} - \frac{8X}{a_1b_1} \right) + \alpha_2 \left( \frac{2}{a_2} + \frac{2}{b_2} - \frac{8X}{a_2b_2} \right) ; \quad X \leq \frac{a_1}{2} \]

\[ A'(X) = \frac{dA(X)}{dX} = \alpha_2 \left( \frac{2}{a_2} + \frac{2}{b_2} - \frac{8X}{a_2b_2} \right) ; \quad X > \frac{a_1}{2} \]

This derivative appears as a decreasing piece-wise linear function of $X$ which can be fitted by hand or numerically on the plot of actual values $A'(X)$ (see Fig B2). The parameter $a_1$ is set
so that the break point of the model $A'(X)$ located in $a_i/2$ matches with the change of slope of actual data. The parameter $a_2$ is defined as the length (distance) for which $A'(a_2/2) = 0$ (see Fig. B2).

Fig. B2. Enhanced general imbibition technique to mean matrix block size evaluation. First-order derivative of the normalized invaded matrix area $A(X)$ as a function of the distance $X$ between a location in the matrix and the closest fracture. The derivative with respect to $X$ is modelled as a piecewise linear function allowing to infer a small and a large matrix block size.

The threshold $a_i/2$ separates the linear function $A'(X)$ in two portions with slopes

$$A''_1 = \frac{dA'(X)}{dX} = -\frac{8\alpha_1}{a_1b_1} - \frac{8\alpha_2}{a_2b_2}; \quad X \leq \frac{a_i}{2}$$

$$A''_2 = \frac{dA'(X)}{dX} = -\frac{8\alpha_2}{a_2b_2}; \quad X > \frac{a_i}{2}$$

(B4)

The difference of slopes on a plot of $A'(X)$ can be identified with the expression of

$$A''_1 - A''_2 = -8\alpha_i/a_ib_i$$

which in turn fixes the ratio $\alpha_i/b_i$ since $a_i$ has been previously prescribed.

The height of the step between the two linear portions of $A'(X)$ can be calculated as

$$\delta A' = A'(\frac{a_i}{2}) - A'(\frac{a_i}{2})^+ = \frac{2}{\alpha_i} \left( 1 - \frac{1}{b_i} \right)$$

(B5)
Identifying (B5) with the value of the plot and associating the result with the identified value $A''_1(X) - A''_2(X)$ renders two equations allowing for the calculation of both $\alpha_1$ and $b_1$ values.

Finally, the expression of $A'(X)$ in $X = 0$ which writes as

$$A'(0) = \alpha_1 \left( \frac{2}{a_1} + \frac{2}{b_1} \right) + (1 - \alpha_1) \left( \frac{2}{a_2} + \frac{2}{b_2} \right)$$  \hspace{1cm} (B6)

is identified via the equivalent value observed on the plot of actual data (Fig. B2) and returns the value of $b_2$.

**The mean spacing (MS) technique**

The principle of MS is sketched in Fig. B3.

For each main direction $i$ of a fractured block with length $l_i$, random lines parallel to direction $i$ and crossing the whole block are drawn. For each line, one counts as $n_i$ the number of intersections between the line and any fracture plane (or trace in a two-dimensional problem) of the fracture network. For each random line in the direction $i$, the mean distance
between two successive intersections is \( l_i / (n_i + 1) \). The mean size of the matrix block in the direction \( i \) is defined as

\[
s_i = l_i \left( \frac{1}{n_i + 1} \right)
\]  

(B7)

where averaging \( \langle \ \rangle \) is conducted over the whole set of random lines in the direction \( i \).


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