An effective criterion to prevent injection test numerical simulation from spurious oscillations

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An Effective Criterion to Prevent Injection Test Numerical Simulation from Spurious Oscillations

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Résumé — Un critère efficace pour prévenir les oscillations parasites dans la simulation numérique du test d’injection — Les tests d’injection/fall-off sont une des alternatives les plus prometteuses à la conventionnelle séquence de production/build-up, car ils éliminent les émissions de surface et peuvent réduire considérablement les coûts des essais. Ce type de test est caractérisé par la présence de deux phases mobiles, le fluide initialement en place (hydrocarbure) et le fluide injecté (Diesel, eau ou azote). L’approche classique d’analyse utilisée pour décrire le comportement transitoire de pression n’est plus appropriée en raison des variations de la saturation en fluide pendant le test. Bien qu’applicable en théorie, l’approche analytique implique souvent des simplifications excessives du comportement du système réel comme la description du déplacement des fluides en forme de piston. Ainsi, seules les simulations numériques peuvent complétement décrire les phénomènes qui se produisent lors de l’injection. Cependant, la réponse calculée numériquement de la pression et de sa dérivée montre souvent des oscillations physiques au cours de la phase d’écoulement radial, lorsque la dérivée de la pression devrait être horizontale. Il a été constaté que ces oscillations parasites se posent en problèmes dominés par la convection et sont associées à des fronts de saturation tranchants. Dans cet article, une méthodologie efficace pour le calcul d’une time-step adaptative est présentée, avec l’objectif d’éviter les oscillations de pression. La sélection de time-step que l’on propose est en même temps efficace et appropriée pour capturer la physique du système.

Abstract — An Effective Criterion to Prevent Injection Test Numerical Simulation from Spurious Oscillations — Injection/fall-off tests are one of the most promising alternatives to the conventional production/build-up sequence because they eliminate surface emissions and can significantly reduce testing costs. This kind of test is characterized by the presence of two mobile phases, the fluid originally in place (hydrocarbon) and the injected fluid (Diesel, brine or nitrogen). The conventional analytical approach used to describe the transient pressure behavior is no longer suitable due to the variations in fluid saturations during the test. Although applicable in theory, the analytical approach often implies excessive simplifications of the real system behavior, such as piston-like displacement. Thus only numerical simulations can thoroughly describe the phenomena occurring during the injection process. However, the pressure and pressure derivative response calculated numerically often shows non-physical oscillations during the radial flow phase, when the pressure derivative is expected to be horizontal. It was found that these spurious oscillations arise in convection-dominated problems and are associated with sharp saturation fronts. In this paper, an effective methodology, based on an adaptive time-step calculation, is presented so as to avoid pressure oscillations. The proposed time-step selection is both computationally efficient and suitable to capture the physics of the system.
LIST OF SYMBOLS

\( a_r \)  
Grid progression (-)

\( a_t \)  
Time progression (-)

\( \gamma \)  
Gradient = \( \rho g \) (Pa/m)

\( \phi \)  
Porosity

\( \mu \)  
Viscosity (Pa s)

\( \rho \)  
Density (kg/m³)

\( A \)  
Interface area orthogonal to the flux direction (m²)

\( B \)  
Formation volume factor (m³/Sm³)

\( c \)  
Compressibility (Pa⁻¹)

\( h \)  
Pay thickness (m)

\( h_w \)  
Perforated interval (m)

\( k \)  
Absolute permeability (m²)

\( k_h \)  
Horizontal absolute permeability (m²)

\( k_r \)  
Relative permeability (-)

\( k_z \)  
Vertical absolute permeability (m²)

\( \Delta L \)  
Distance connecting two nodes (m)

\( M \)  
Mobility ratio (-)

\( n_o \)  
Oil relative permeability exponent (-)

\( n_r \)  
Number of cells in the radial direction (-)

\( n_w \)  
Water relative permeability exponent (-)

\( p \)  
Pressure (Pa)

\( q \)  
Rate for unit volume (1/s)

\( Q \)  
Water injection rate (Sm³/day)

\( r \)  
Nodal coordinate in the radial direction (m)

\( r_1 \)  
Radial position of first cell node (m)

\( r_e \)  
External radius (m)

\( r_w \)  
Wellbore radius (m)

\( S \)  
Saturation (-)

\( \Delta t \)  
Time-step length (s)

\( \Delta t^0 \)  
Initial time-step length (s)

\( V \)  
Volume (m³)

\( z \)  
Vertical coordinate (m)

INTRODUCTION

Well testing is a process widely used in the oil industry for evaluating the well productivity and the formation damage, estimating the reservoir characteristics such as initial pressure, fluid type and effective permeability, and identifying the reservoir heterogeneities, which are key information for field development and facilities design (Coelho et al., 2005). Well tests, usually performed during the exploration and appraisal phases of a reservoir, consist in introducing abrupt changes in the surface production rates and recording the associated changes in bottomhole pressure. The pressure disturbance induced by production travels into the formation and is affected by the rock features and fluid properties in various ways. Therefore, a record of the pressure response over time produces a curve whose shape is defined by the reservoir’s unique characteristics (Schlumberger, 1998).

The selection of the test type to assess the fluid nature and the reservoir potential must be balanced against operational risk, environmental constraints and value derived from affecting early decisions on project appraisal or development. Typically, conventional testing methods involve surface production of fluids. However, in exploration and often in appraisal scenarios, surface facilities to store the reservoir fluid are not available and hence the fluid is discharged or flared. Burning hydrocarbons produces significant amounts of emissions, which in turn produce acid rain, smog, ozone at ground levels and greenhouse gases in the upper atmosphere. The demands to reduce emissions during well testing put enormous pressure to avoid these tests altogether. Alternative testing procedures have thus been investigated and attempted for reservoir appraisal so as to have sufficient information to evaluate the investment risk and make a decision whether to sanction a project or to develop the field. A valid contribution to the review and discussion of technologies such as wireline formation tests, closed chamber tests, production/reinjection tests and injection tests as viable alternatives to conventional well testing can be found in the technical literature (Coelho et al., 2005; Woie et al., 2000; El-Khazindar et al., 2002; Hollaender et al., 2002; Banerjee et al., 1998; Beretta et al., 2007; Levitan, 2002; Verga and Rocca, 2010).

Among these unconventional methodologies injection testing is considered very interesting. An injection test consists substantially in injecting a fluid, typically brine, in a potential pay zone and monitor the pressure response during the injection period and the subsequent fall-off period, in which the well is closed and the pressure tends to return to the equilibrium value. Although an injection/fall-off test is similar to a conventional drawdown/build-up test, a distinction between the two is necessary when the properties of the injected and reservoir fluids are different (Gunawan et al., 2002). In fact, the physics of injection tests is characterized by the presence of two phases in the reservoir, the hydrocarbons originally in place and the injected fluid; therefore, fluid saturations change dynamically during injection in both space and time and the permeability of the reservoir rock to each fluid will be dependent on saturations.

If convection dominates the immiscible displacement of hydrocarbons by the injected fluid (i.e. capillary pressures are negligible), the saturation profile in the porous medium is characterized by a steep transition
between the flushed and the unflushed zone (Buckley and Leverett, 1941). When the process is simulated numerically, such sharp transition can induce non-physical oscillations in the pressure response (Forester, 1977). Such oscillations, called spurious in the technical literature (Le Veque, 1990), are due to the truncation error which arises from the numerical discretization of the partial derivatives (Lantz, 1971). Besides being responsible of the spurious oscillations in the pressure response, this truncation error acts as an artificial dispersion term, often denoted as numerical diffusion (Lantz, 1971), which tends to artificially flatten the sharp saturation profile. If the magnitude of the artificial dispersion term is comparable with the convection one, the description of the transition zone can be inaccurate, compromising the correct computation of the fluid mobilities at the interface and, in turn, the pressure response computed during the simulated injection test.

Several methods to reduce the truncation error were presented in the literature. Some methods act on the differentiation scheme, such as the transfer overshoot (Peaceman and Rachford, 1962), the two-points upstream approximations (Todd et al., 1972), the filtering techniques (Van Leer, 1977) the higher order variational approximations (Settari et al., 1977) or the addition of terms for truncation error cancellation (Laumbach, 1975). Others focus on a local grid refinement, in some cases with moving grids which follow the frontal advance (Han et al., 1987). Eventually, others adjust the time-step size in order to restrict the variation of each variable over the time-step (Jensen, 1980) or to cancel the first order term of the truncation error (Aziz and Settari, 2002).

This paper presents an effective and computationally efficient criterion to select the proper time-step size in order to avoid spurious oscillations of the pressure derivative in injection testing simulation. A comprehensive numerical 2D axial-symmetric near wellbore model was implemented to reproduce the pressure response registered by a gauge positioned in the wellbore at the depth of the tested zone (Verga et al., 2011). The model simulates the flow of two immiscible phases as an implicit finite difference problem. The simulated pressure transient curves are subsequently post-processed in order to extract the contained reservoir information, i.e. mimicking a test interpretation.

1 NUMERICAL SIMULATION OF INJECTION TESTS

Simulating an injection test consists in reproducing the pressure response registered by a gauge positioned in the wellbore just above the tested zone. The simulated pressure signal is subsequently processed in order to extract reservoir information such as the formation permeability-thickness product (kh) and the well damage, expressed in terms of mechanical skin (S_m), which are key parameters for the well productivity estimate. The pressure increments (Δp) and the pressure derivative (Δp') are displayed on a log-log diagnostic plot. Details on the calculation of the pressure derivative are given in Appendix A.

Although the model has the extension of the typical drainage area of a well, the attention was focused on the near wellbore zone, thus the bi-phase flow model for simulation of injection tests was developed under the assumption of radial flow geometry. This hypothesis limits the application of the model to the case of vertical wells intercepting a hydrocarbon-bearing formation of constant thickness. However, exploration and appraisal wells, for which injection tests are of great interest, generally have a vertical or a slightly slanted trajectory, thus this assumption is not very restrictive. Fluid injection or production can occur along the entire thickness of the reservoir or through a limited interval in order to simulate partial penetration or perforation. The injection fluid and the reservoir dominant fluid are immiscible.

The model, described in detail in Appendix B, is made up of two convection-diffusion equations, one for each fluid phase. The equations arise from the conservation equations, assuming Darcy’s flow in the porous medium. Several non-linearities occur, due to the dependency of the model coefficients on pressure and saturation.

The 2D Cartesian axial symmetric model was formulated in finite differences, where a multiplication for the block volume was introduced in order to conserve flow rates (Aziz and Settari, 2002). The simulator follows a “SSimp” scheme (Aziz and Settari, 2002). In fact, the algorithm is fully implicit in pressure and saturation and the flow equations are solved iteratively with the Simultaneous Solution (SS) method. The detailed numerical formulation is shown in Appendix C. The use of an implicit formulation with simultaneous solution in pressure and saturation avoids stability restrictions on the time-step size, i.e. numerical errors are not amplified from time-step to time-step, independently from the chosen time-step length (Aziz and Settari, 2002). Furthermore, an upstream weighting scheme was adopted for the evaluation of the saturation dependent variables, such as the relative permeabilities, at the cell interfaces. Conversely, a centered approximation was applied to the pressure dependent variables, such as the PVT properties. The midpoint weighting of the relative permeability, although most appropriate from the numerical analysis standpoint because inducing a second order truncation error, was discarded because it could give erroneous solutions. When the capillary
pressure values are small or negligible, the SSimp equations result in an almost hyperbolic problem, with the true solution very close to the Buckley-Leverett solution; conversely, the numerical solution using midpoint weighting converges to a different solution, which is mathematically possible but physically incorrect (Aziz and Settari, 2002). Due to the non-linear nature of the scheme, a Newton-Raphson iterative solution was adopted to solve the set of coupled equations. Finally, an automatic time-stepping selection was adopted so that shorter intervals are used when rate changes occur and, then, the time-steps increase with a geometric progression during the flow period. This option provides a reliable calculation of the pressure derivative in a reasonable computational time. Criteria for the selection of the minimum time-step length ($\Delta t^0$) are discussed in Appendix C.

The chosen discretization scheme leads to a first order truncation error on a uniform grid. Since the solution of the studied PDE (Partial Differential Equation) is calculated in correspondence of a non-uniform grid, with the cell dimension increasing with a fixed geometric progression, the analytical evaluation of the corresponding truncation error is not trivial. The complexity of the evaluation is increased by the presence of diffusion and advection coefficients which vary with the independent variables (pressure and saturation). Thus an effective theoretical support was impossible to achieve.

2 SPURIOUS OSCILLATION ANALYSIS

The equations describing immiscible displacement in a porous medium (i.e. convection-diffusion equations) convey some numerical issues when convection predominates. In fact, when the equations are solved with differentiation schemes, non-physical oscillations can be observed near steep gradient regions (Forester, 1977).

A numerical simulation of an injection test can be affected by these spurious oscillations when capillary pressures are negligible. In fact, the saturation distribution during injection can be described by the presence of three zones (Sosa et al., 1981), as shown in Figure 1:

- water zone near the wellbore, where water has completely displaced the oil and the saturation reaches the constant value $1 - S_{hr}$;
- transition zone, where the water saturation progressively decreases from the maximum value, $1 - S_{hr}$, to the irreducible value, $S_{wi}$;
- undisturbed zone, where the water saturation is equal to the irreducible value, $S_{wi}$.

When convection is dominant (i.e. negligible capillary pressure), the saturation profile of an immiscible displacement in a porous medium is characterized by a steep transition between the flushed and the unflushed zone (Buckley and Leverett, 1941). This transition becomes steeper and steeper if the mobility ratio ($M$) between fluids is lower than 1 or if the relative permeability curves are described by polynomials of high order ($n > 2$). A moving front is observed during injection, while only little changes in the saturation values are registered during the pressure fall-off subsequent to the injection period (Levitan, 2002).

Azarkish et al. (2006) compared Levitan’s analytical solution for injection test (Levitan, 2002) with a numerical model, discretized with an areal 2D, unstructured Voronoy grid and solved with finite elements, where the time-step duration had an almost constant progression. The authors observed some oscillations in the late time period of the pressure derivative of the injection period. These oscillations had not been observed on the pressure derivative calculated with Levitan’s model, thus their nature is clearly numerical. The authors verified that spurious oscillations tend to disappear when the mobility ratio between the injected and reservoir fluids increases.

Spurious oscillations are due to the local truncation error (Lantz, 1971), which arises from the numerical discretization of the partial derivatives appearing in the mass conservation laws describing the physical system. When a uniform time and space discretization is assumed in a numerical differentiation scheme, it is possible to analytically calculate the time-step length that improves the solution accuracy, i.e. reduces the discrepancy between the theoretical and the numerically approximated solutions (Higham, 1996). This is obtained by calculating the local discretization error.
via the Taylor series and by imposing a time-step which makes the term of lowest order to cancel (Aziz and Settari, 2002). Unfortunately, the physics of the displacement process taking place during an injection test requires irregular time and space discretization. In fact, a centimetric gridding around the well is needed so as to obtain a detailed description of the solution in the near wellbore zone, while the domain affected by the pressure disturbance is almost kilometric. On such a domain, the use of a centimetric uniform grid is strongly inefficient from the computational point of view, while a grid with local refinement defined by a logarithmic progression is more appropriate (Ertekin et al., 2001). Analogous problems are encountered for time discretization. Very small time-steps (few seconds) are necessary at the beginning of the flow period in order to obtain a refined pressure derivative plot on the log-log scale, but the overall test can last hours or even a day. As a consequence, a constant time-step duration would be impractical. Thus, since the analytical approach for truncation error calculation can be applied only under simplified assumptions (i.e. uniform space and time discretization), the optimal time-step length able to reduce the local truncation error was obtained as a result of sensitivity analyses to both physical and numerical parameters. In order to do this the 2D axial symmetric Cartesian finite difference discretization introduced in Section 2 was exploited. Details of the discretization are given in Appendices B and C.

An undersaturated oil-bearing homogeneous reservoir, the properties of which are summarized in Table 1, was selected as the base case for sensitivity analysis. Three reservoir fluids were considered (light, medium and heavy oil, respectively) with the PVT properties summarized in Table 2; the injected water properties are summarized in Table 3. Relative permeability curves were assumed Corey-shaped with the parameters listed in Table 4. The rate history consisted of 8 h of injection followed by 24 h of fall-off; rates, summarized in Table 5, were chosen so as to induce a pressure drop of about 30 bar in each case. It should be noted that these oscillations do not jeopardize the interpretability of the simulated pressure response since they merely affect the injection period (Fig. 2).

Furthermore, it is possible to reduce, and sometimes eliminate, the oscillation amplitude on the log-log plot making use of conventional smoothing algorithms in the pressure derivative calculation. However, an insight of these phenomena was necessary in order to verify whether the numerical errors affect the reliability of the interpretation results.

Firstly, sensitivities to numerical parameters such as grid refinement and time-step length were performed on a base case \((M = 1.19, n_w = n_\phi = 2)\) discretized into 40 cells \((x_r = 1.259)\) with a time-step length starting from 3 seconds and growing by a factor \(x_t = 1.01\). Results confirmed that spurious oscillations are related to the local truncation error, which is decreased by a finer grid and increased by finer a time-step (Todd and Longstaff, 1972). In fact, once the time progression is fixed, a finer grid discretization \((n_r)\) reduces the oscillation amplitude and increases the oscillation frequency (Fig. 3); in addition the frequency is not constant in time but decreases logarithmically (Fig. 3c), thus looking constant on a log-log scale (Fig. 3b). At the same time, once the grid is fixed, reduced amplitudes correspond to faster time-step progression \((x_t)\) (Fig. 4). No significant influence of the minimum time-step length \((\Delta t^0)\) was observed, except at the very beginning \((t \leq 0.01 \text{ h})\) of the test (Fig. 5).

### Table 1
Reservoir properties

| Property                     | Value |
|------------------------------|-------|
| Permeability (mD)            | 100   |
| Porosity                     | 0.2   |
| Rock compressibility (bar\(^{-1}\)) | \(4 \times 10^{-5}\) |
| Initial pressure (bar)       | 350   |
| Temperature (°C)             | 100   |
| Pay thickness (m)            | 10    |
| Wellbore radius (m)          | 0.1   |
| External radius (m)          | 1000  |

### Table 2
Oil properties at 350 bar and 100°C

| Property                      | Heavy | Medium | Light |
|-------------------------------|-------|--------|-------|
| Density (°API)                | 17    | 28     | 37    |
| Viscosity (cP)                | 8     | 0.95   | 0.2   |
| Compressibility (bar\(^{-1}\)) | \(9 \times 10^{-5}\) | \(1.3 \times 10^{-4}\) | \(2.9 \times 10^{-4}\) |
| Formation volume factor \((\text{m}^3/\text{m}^3\text{ST})\) | 1.2   | 1.3    | 1.7   |
| \(R_w @ \text{BP}\) \((\text{m}^3/\text{m}^3\text{ST})\) | 50    | 120    | 250   |
| Gas gravity                   | 0.7   | 0.7    | 0.7   |
| Mobility ratio                | 10    | 1.19   | 0.25  |
Secondly, sensitivities on reservoir and fluid properties were performed in different scenarios, starting from a reference case \( M = 0.25, \) \( n_w = n_o = 2, \) \( n_r = 40, \) \( D_{t_0} = 3 \) s, \( a_t = 1.01 \). Sensitivities on reservoir properties such as permeability, thickness and porosity, along with the injection rate, showed a negligible impact:

- the oscillation amplitude and frequency were not affected by permeability (Fig. 6a) nor pay (Fig. 6b), neither was a time shift observed at late time;
- amplitude and frequency were not influenced by porosity (Fig. 6c) nor injection rate (Fig. 6d); only a time shift was observed.

On the contrary, from sensitivities on fluid mobilities a relevant impact on the oscillation behavior was noticed. It was observed that spurious oscillations due to numerical errors can appear at each mobility ratio (Fig. 7b). Oscillations are generally not visible on the pressure response, except for the case of high Corey exponents, \( i.e. n_w = n_o > 2 \) (Fig. 7a, 8a). Conversely, oscillations could be clearly detected from the derivative of the pressure response in all cases, except for linear relative permeability curves (Fig. 7b, 8b). Sensitivity analyses, along with the complete absence of oscillations in the fall-off derivative, showed that there is a link between spurious oscillations and water front velocity. In fact, oscillations appear later if the water velocity is reduced, either due to a low permeability or to mobility ratio \( M \) lower than 1. Furthermore, larger oscillations were observed in the case of slow oil velocity either due to reduced mobility in the transition zone \( i.e. \) Corey exponent \( n_o > 2 \) or to mobility ratio \( M > 1 \). The following results were found:

- fixing the injection pressure drop, the lower the mobility ratio the higher the oscillation amplitude (Fig. 7);
- oscillations appeared later in the case of higher mobility ratio (Fig. 7); however, the oscillation frequency did not change;
- the higher the Corey exponents of the relative permeability curves, the larger the oscillation amplitude (Fig. 8b,c). A time delay was observed for low exponents (Fig. 8b), but the oscillation frequency remained exactly the same (Fig. 8c).

Since the grid has a geometric progression, the transition zone moves in time from very fine cells to larger ones. As a consequence, at fixed time-step length, the truncation error is expected to affect the

---

**TABLE 3**

| Water properties                        |       |
|-----------------------------------------|-------|
| Density (kg/m³)                        | 1.031 |
| Viscosity (cP)                          | 0.4   |
| Compressibility (bar⁻¹)                 | \(3.1 \times 10^{-5}\) |
| Formation volume factor (m³_R/m³_ST)    | 1     |
| Salinity (kg/m³)                        | 80    |

**TABLE 4**

| Corey parameters for relative permeability curves |
|--------------------------------------------------|
| \(k_{w,\text{max}}\)                            | 0.4   |
| \(k_{o,\text{max}}\)                            | 0.8   |
| \(S_{wi}\)                                       | 0.2   |
| \(S_o\)                                          | 0.3   |
| \(n_w, n_o\)                                     | 2     |

**TABLE 5**

| Rate history                             |
|------------------------------------------|
| Water injection rate (Sm³/day)           |
| \(M = 10\)                               | 45   |
| \(M = 1.19\)                             | 190  |
| \(M = 0.25\)                             | 335  |
| \(M = 1.19, k = 10 \text{ mD}\)         | 19   |

---
solution accuracy more severely when the transition zone reaches coarse grid cells. Thus, for the fixed time-step cases, oscillations are expected to appear at different times, depending on the velocity of the saturation front advance. In turn, the front velocity depends on the fluid mobility and thus on absolute permeability, mobility ratio and relative permeability. Figures 6a, 7b and 8b seem to confirm such considerations. Furthermore, the mobility ratio and the curvature of the relative permeability functions influence the shape of the displacing front, which is characterized by smaller saturation gradients for mobility ratio $M > 1$ and low Corey exponents ($n < 2$) and by higher saturation gradients (almost piston like displacement) for mobility ratio $M < 1$ and high Corey exponents ($n > 2$) as shown in Figure 7c and 8d. For a specified time-step size, the magnitude of the truncation error increases for increasing front velocity and saturation gradient at the front (Todd and Longstaff, 1972), thus a higher truncation error is expected for
$M < 1$ and $n > 2$. This is confirmed by the increasing oscillation amplitude observed in such cases (Fig. 7, 8).
As it will be empirically demonstrated further on in this section, the only assumptions are that the well trajectory is vertical and that the numerical grid follows a geometrical progression, as expressed by equations (C4)-(C7). The values assigned to fluid properties, i.e. mobility and compressibility, and to reservoir properties, such as permeability, porosity, heterogeneity, anisotropy, do not jeopardize the validity of the relation. Furthermore, the well completion and test procedure, such as partial penetration/perforation of the well or the imposed rate, are not subject to any limitation as well.

It was verified that the time-step selection according to Equation (1) successfully prevented spurious oscillations to arise in all the considered scenarios. By way of example, the effectiveness of the proposed time-step selection ($z_i = z_f$) is shown for the case most affected by oscillations, corresponding to a mobility ratio $M = 0.25$ and high Corey exponents ($n_w = n_o = 3$) (Fig. 9). It was noticed that the obtained time-step length corresponded to the oscillation semi-period (Fig. 9d).

The proposed time-step selection scheme is both computationally efficient and physically reliable. In fact, on the one hand the introduced numerical
formulation ensures accuracy and stability without requiring an excessive number of time-steps or an extremely fine mesh resolution. In fact, local grid refinement and adaptive time-step length are imposed via logarithmic progressions, thus efficiently reducing the overall computational cost with respect to a uniform time discretization with the same degree of detail. Furthermore, since the length of each time-step is univocally determined before the calculations over the time-step are performed, no additional computations are required; on the opposite, in methodologies based on constraints over variable variations, the time-steps need to be recalculate if the trial time-step length proves to be too long to comply with the imposed criterion. Besides preventing oscillations to occur, the proposed discretization does not significantly flattens the physically sharp saturation front, as it can be observed comparing the saturation profiles at the end of the injection period with the corresponding analytical Buckley-Leverett curves (Fig. 9b). Therefore, a reliable description of the solution is obtained even in the transition zone, with
a consequent correct description of the changing fluid mobilities and, in turn, of the pressure response.

It was also verified that the progression in time (1) does not involve any stability problem because the scheme is fully implicit. Furthermore, the required time resolution of the pressure response reduces logarithmically in time. In fact, as the diagnostic plot used for interpretation is log-log and the flow periods typically last a few hours, very short time-steps are needed at the beginning of each flow period (i.e. in the order of magnitude of seconds), while long time-steps are adequate at the end of the flow period (i.e. in the order of magnitude of hours). Therefore, a geometric progression in time would provide a good log-log representation.

Figure 8
Sensitivity analyses to the Corey exponents \( (n_w = n_o = 1, 2, 3) \) for the mobility ratio \( M = 0.25 \): a) simulated pressure response and b) pressure derivative for the injection period; c) zoom of the oscillations during the horizontal stabilization of the pressure derivative; d) water saturation profile at the end of the injection period. Oscillations maintain exactly the same frequency, but grow in amplitude with the Corey exponents; moreover, a shift in time can be observed. Spurious oscillations on the pressure profile are also visible for high Corey exponents \( (n_w = n_o = 3) \).
Relation (1) is still valuable in heterogeneous cases. Exemplifying that, two heterogeneous cases were considered: a radial composite reservoir (Fig. 10a) and a layered reservoir (Fig. 11a). For a more general validation, the fluid properties that induced the most evident oscillations were considered, i.e., mobility ratio $M = 0.25$ and high exponent of relative permeability curves ($n = 3$). The pressure derivatives of the injection period are shown in Figure 10b and 11b, respectively. In both cases oscillations were successfully removed.

The introduced relation (1) is generally suitable to eliminate spurious oscillations also when a limited entry well is considered, even though the uniform grid discretization adopted in the vertical direction is coarser than that in the radial direction in the near wellbore zone (i.e. decametric vs centimetric). In fact, as injection begins, the vertical velocity is small compared to the radial velocity and an almost radial flow occurs in front of the perforated interval. In such cases diffusion in the vertical direction can be negligible (Lantz, 1971). After a short time (a few minutes in the presented case), when spherical flow begins, vertical diffusion is no longer negligible, but a time-step progression greater than or equal to the grid factor alpha is generally sufficient. Firstly, an oil-bearing homogeneous reservoir ($M = 0.25$) produced by a well with limited entry (perforations corresponded to 30% of the net pay) was considered (Fig. 12a). The effectiveness of the time discretization was also verified in potentially critical cases (even though
they have limited practical interest), when the vertical permeability is higher than the horizontal one. Three cases were addressed:

– anisotropy ratio 0.1 and high order relative permeability curves ($n = 3$),
– anisotropy ratio 10 and high order relative permeability curves ($n = 3$),
– anisotropy ratio 10 and low order relative permeability curves ($n = 2$).

For high order Corey functions ($n = 3$) oscillations were successfully removed for an anisotropy ratio of 0.1 (Fig. 12b) and significantly reduced for anisotropy ratio of 10 (Fig. 12c). Difficulties related to anisotropy ratio were not encountered for Corey exponent $n = 2$, as shown in Figure 12d.

Afterwards, an isotropic heterogeneous reservoir with different permeability layers (Fig. 13a) was investigated ($M = 0.25, n = 3$), where only the fraction of pay corresponding to the higher permeability layer was perforated. Again the oscillations were successfully removed (Fig. 13b).
Figure 12
a) Single layer reservoir with limited entry well; b, c) removal of spurious oscillations from the pressure derivative of the injection period for the case $M = 0.25$ and different combinations of anisotropy ratio ($k_z/k_h = 0.1$ and 10) and Corey exponents $n_w = n_o = 2$ and 3.

Figure 13
a) Layered reservoir with limited entry well; b) removal of spurious oscillations from the pressure derivative of the injection period for the case $M = 0.25$ and Corey exponents $n_w = n_o = 3$. 
CONCLUSIONS

A numerical solution is required for an adequate simulation of the injection/fall-off tests. Although applicable in theory, the analytical approach often implies excessive simplifications of the real system behavior, thus it fails to properly describe the reservoir characteristics and the phenomena occurring during the injection process. However, it was observed that numerical solutions can be affected by spurious oscillations, particularly evident on the pressure derivative calculated for the injection period at late time. This non-physical response arises in convection-dominated problems, which are characterized by the presence of a sharp saturation front. Sensitivity analyses proved that spurious oscillations depend on the parameters affecting the injected fluid velocity.

Oscillations can be mitigated, or even eliminated, by a reduction of the local truncation error, typically achieved by highly refined grids at the cost of remarkably increasing the computational effort. In this paper, a valuable alternative, based on an adaptive time-step calculation, was presented. It was observed that a time progression factor equal to the nodal grid progression keeps the truncation error efficiently under control. In fact, spurious oscillations were successfully avoided in all the considered scenarios, given that the only required assumptions are that the well is vertical and that the spatial discretization follows a geometric progression. The proposed time-step selection scheme is both physically reliable and computationally efficient, avoiding time consuming sensitivities for the identification of a suitable time-stepping. In fact, the presented time-step length which increases logarithmically provides a nice log-log representation of the pressure derivative, free from spurious oscillations also in convective-dominated problems. Furthermore, the computation of the saturation profile is not significantly affected by numerical errors, such as an artificial flattening of the front corresponding to an overestimation of the transition zone extension. Finally, accuracy and stability of the solution are ensured without requiring an excessive time or mesh resolution. Adaptive time-step length and local grid refinement are imposed via logarithmic progressions, significantly reducing the overall computational cost with respect to a uniform discretization with the same degree of detail. Opposite to methodologies based on fixed maximum variable variations, no additional computation is required because the length of each time-step is univocally determined before the calculations over the time-step are performed.

To some extent the proposed solution might appear pragmatic because it was derived empirically and it has a very simple form: it avoids oscillation by calculating the model solution with a time-step length equal to the oscillation semi-period. Nevertheless, it was shown that its validity is quite general in the context of injection test application. In fact, it was successfully applied to a variety of scenarios, from light oils to heavy oils, from fully perforated to limited entry wells, from homogeneous to heterogeneous cases (either radial composite or layered reservoirs). In addition, since injection rates and formation properties (such as permeability, thickness and porosity) do not significantly influence the oscillation phenomena, the applicability of the methodology is not limited to a range of parameters. For these reasons, the simplicity of the proposed methodology should be valued and not regarded as a limiting aspect.

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APPENDIX A: Injection test interpretation

An injection test interpretation consists in analyzing the pressure transient recorded during the injection of a fluid into the reservoir and subsequent shut-in phase (termed fall-off period). A log-log graphical representation of the pressure increments ($\Delta p$) and pressure derivative ($\Delta p'$) is used. The pressure increments express the pressure change from the beginning of the flow period ($t^0$):

$$\Delta p = p(t) - p(t^0)$$  \hspace{1cm} (A1)

while the pressure derivative is essentially the rate of pressure change with respect to time (Bourdet, 2002):

$$\Delta p' = \frac{\partial \Delta p}{\partial \log (t - t^0)}$$  \hspace{1cm} (A2)

A horizontal straight line on the pressure derivative diagnostic plot corresponds to a radial flow regime; the ordinate value, $y$, of these stabilization corresponds to (Bourdet, 2002):

$$y = \frac{1}{2} \frac{QB}{2\pi h \lambda}$$  \hspace{1cm} (A3)

where $Q$ is the rate at standard conditions, $B$ is the volume factor, $h$ is the pay thickness and $\lambda$ is the fluid total mobility, defined as:

$$\lambda(S_w) = k \left[ \frac{k_{rh}(1 - S_w)}{\mu_h} + \frac{k_{rw}(S_w)}{\mu_w} \right]$$  \hspace{1cm} (A4)

where $S_w$ is the water saturation, $k$ is the absolute permeability, $k_{rw}$ and $k_{rh}$ are water and hydrocarbon relative permeability, respectively, and $\mu_w$ and $\mu_h$ are water and hydrocarbon viscosity, respectively.

During the injection phase the pressure derivative at early time reflects the mobility in the oil zone ahead of the transition zone while at late time it reflects the mobility in the water region behind the transition zone. The opposite is observed during the fall-off phase (Levitan, 2002). The $kh$ product is estimated from (A3). Furthermore, the distance between the derivative stabilization and the pressure increment curve is related to the mechanical skin, which is a fundamental well testing target, essential to estimate the well productivity. More in detail, in the absence of fractures the total skin obtainable from the test interpretation can be expressed as a linear composition of three components (Verga et al., 2012): the mechanical skin ($S_m$) due to permeability damage in the near wellbore zone, the geometrical skin ($S_c$), due to well partial penetration/perforation and the bi-phase skin ($S^*$) due to the presence of two fluid phases:

$$S_t = \frac{h}{h_w} \left( \frac{S_m}{M} + S^* \right) + S_c$$  \hspace{1cm} (A5)

where $M$ is the ratio between the mobility of the injected phase and the mobility of the resident phase (evaluated at the critical saturation) and $h_w/h$ is the perforated fraction of the net pay.

APPENDIX B: Model equations

The coupled model consists of two flow equations, one for each phase: water (B1) and hydrocarbon (B2) (Aziz and Settari, 2002):

$$\nabla \cdot \left[ \frac{1}{B_w \mu_w} (\nabla p_w - \gamma_w \nabla z) \right] = \frac{\partial}{\partial t} \left( \frac{S_w \phi}{B_w} \right) + q_w$$  \hspace{1cm} (B1)

$$\nabla \cdot \left[ \frac{1}{B_h \mu_h} (\nabla p_h - \gamma_h \nabla z) \right] = \frac{\partial}{\partial t} \left( \frac{S_h \phi}{B_h} \right) + q_h$$  \hspace{1cm} (B2)
where water pressure \((p_w)\) and water saturation \((S_w)\) were chosen as the unknowns. If the capillary pressures are zero, the hydrocarbon pressure \((p_h)\) coincides with the water pressure. The other parameters are porosity \((\phi)\), hydrocarbon \((B_h)\) and water \((B_w)\) formation volume factor, absolute permeability \((k)\), hydrocarbon \((k_{rh})\) and water \((k_{rw})\) relative permeability, hydrocarbon \((\gamma_h)\) and water \((\gamma_w)\) gradient, hydrocarbon \((\mu_h)\) and water \((\mu_w)\) viscosity, hydrocarbon \((q_h)\) and water \((q_w)\) rate per unit volume.

The initial conditions are obtained imposing the static equilibrium of the system at the initial time-step. The boundary conditions adopted in the model are: constant rate for each flow period at the inner radius of the system (well radius); either constant pressure or no-flow boundary at the external radius of the system.

The model is characterized by several non-linearities due to the dependency of the model coefficients on pressure and saturation. These dependences were accounted for by adopting correlations available in the technical literature. In particular, the permeability curves were assumed Corey-shaped (Chen et al., 2006):

\[
\begin{align*}
  k_{rw}(S_w) &= k_{rw\text{max}} \left( \frac{S_w - S_{wir}}{1 - S_{hr} - S_{wir}} \right)^{n_w} \\
  k_{rh}(S_w) &= k_{rh\text{max}} \left( \frac{1 - S_w - S_{wir}}{1 - S_{hr} - S_{wir}} \right)^{n_s}
\end{align*}
\]

where \(S_{wir}\) is the irreducible water saturation, \(S_{hr}\) is the hydrocarbon residual saturation and \(k_{rw\text{max}}\) and \(k_{rh\text{max}}\) are the relative permeability end points of water and hydrocarbon, respectively.

According to the hypothesis that fluids are slightly compressible, porosity and oil formation volume factor were assumed linearly dependent on pressure (Aziz and Settari, 2002):

\[
\begin{align*}
  \phi &= \phi^0 \left( 1 + c_f (p_w - p_w^0) \right) \\
  \frac{1}{B_w} &= \frac{1}{B_w^0} \left( 1 + c_w (p_w - p_w^0) \right) \\
  \frac{1}{B_h} &= \frac{1}{B_h^0} \left( 1 + c_h (p_h - p_h^0) \right)
\end{align*}
\]

where constant water \((c_w)\), oil \((c_h)\) and formation \((c_f)\) compressibility were assumed.

**APPENDIX C: Numerical formulation**

According to the fully implicit simultaneous solution method (Aziz and Settari, 2002), each quantity appearing in the flux terms is expressed at the current time-step. The discretization scheme for the generic phase \(\ell\) (water or hydrocarbon) at the generic cell \((i,j)\) reads:

\[
\begin{align*}
  \left[ v \nabla \cdot \frac{k_{w} c_{w}}{B_{\ell} \mu_{\ell}} (\nabla p_{\ell} - \gamma_{\ell} \nabla z) \right]_{i,j} &\approx T_{\ell}^{n+1}_{i+1/2,j} \left( p_{\ell}^{n+1}_{i+1,j} - p_{\ell}^{n+1}_{i,j} \right) - T_{\ell}^{n+1}_{i-1/2,j} \left( p_{\ell}^{n+1}_{i,j} - p_{\ell}^{n+1}_{i-1,j} \right) \\
  &\quad + T_{\ell}^{n+1}_{i,j+1/2} \left( p_{\ell}^{n+1}_{i,j+1} - p_{\ell}^{n+1}_{i,j} - \gamma_{\ell}^{n+1} \frac{\Delta z_{j+1}}{2} \left( z_{j+1} - z_j \right) \right) \\
  &\quad - T_{\ell}^{n+1}_{i,j-1/2} \left( p_{\ell}^{n+1}_{i,j-1} - p_{\ell}^{n+1}_{i,j} - \gamma_{\ell}^{n+1} \frac{\Delta z_{j-1}}{2} \left( z_j - z_{j-1} \right) \right)
\end{align*}
\]

where the generic transmissibility term \((T)\) is define by:

\[
T_{\ell} = \frac{k_{w} c_{w}}{B_{\ell} \mu_{\ell} \Delta L}
\]

where \(\Delta L\) indicates the distance connecting the nodes opposite to the interface.

As explained in Section 2, the transmissibility terms \((T)\) is computed at the cell interfaces following the one point upstream weighting for relative permeability value and the midpoint weighting for PVT properties.
The cumulative terms were discretized following the conservative expansion (Ertekin et al., 2001), thus obtaining:

\[
\left[ V \frac{\partial}{\partial t} \left( \frac{S_i \phi_i}{B_i} \right) \right]_{i,j} \approx V \frac{\partial}{\partial t} \left( \phi_i B_i \left( \frac{P_{i,j}^{n+1} - P_{i,j}^n}{\Delta t} \right) \right) + \left( \frac{\phi_i}{B_i} \right)^{n+1} \left( S_{i,j}^{n+1} - S_{i,j}^n \right)
\] (C3)

In order to truly reconstruct the pressure log response, a very fine grid is necessary near the wellbore. Since the pressure distribution around the wellbore is logarithmic, a logarithmically-spaced mesh keeps the pressure drops across all grid blocks approximately the same (Ertekin et al., 2001):

\[
r_{i+1} = \alpha r_i
\] (C4)

where:

\[
\alpha = \left( \frac{r_e}{r_w} \right)^{\frac{1}{n r}}
\] (C5)

and \(r_i\) the position of the first cell center.

In order to ensure that the flux across the grid boundaries is identical for the continuous and discrete form of Darcy’s law, block interfaces were defined by (Ertekin et al., 2001):

\[
r_{i+1/2} = \frac{(r_{i+1} - r_i)}{\ln \left( \frac{r_{i+1}}{r_i} \right)}
\] (C6)

By imposing that \(r_{1/2} = r_w\), from Equation (C6) it follows that:

\[
r_1 = \frac{(r_w \ln n_r)}{1 - \left( \frac{1}{n_r} \right)}
\] (C7)

It is pointed out that grid cells must be big enough not to violate the continuum hypothesis. In fact, the macroscopic approach to porous medium states that volume average quantities are associate with each point in space, where a REV (Representative Elementary Volume) was chosen as the averaging volume (Bear and Bachmat, 1990). Below REV, the parameters are not defined and the rock cannot be treated as a continuum. In the case of reservoir simulation, the sand grain size can range from 1/16 mm to 1-2 mm. An acceptable REV should be one order of magnitude larger than the grain size. The grid cell volume, in turn, has to be larger than the REV, otherwise meaningless results are obtained. Thus 1 cm can be assumed as the lower bound for the cell radial dimension. On the other hand, the dimension of the cell next to the wellbore wall must be small enough to truly describe the pressure behavior at the well, since this is indeed the final goal of the simulation. Extended simulations showed that the correct order of magnitude for the radial dimension of that cell is centimetric. Therefore, a very nice correspondence exists between the different requirements. According to the current pressure gauge specifications, a lower bound of 0.1 second was considered for the duration of the first time-step (\(\Delta^0_t\)), which is the shorter one over the simulation. Furthermore, an upper bound was imposed based on the injected volumes:

\[
\Delta^0_t \leq \min_{1 \leq j \leq n_z} \frac{Q_j}{\phi_{1,j} V_{1,j} (1 - S_{hr})}
\] (C8)

where \(Q_j\) is the inflowing rate of layer \(j\), \(n_z\) is the number of the vertical grid subdivisions and \((\phi_{1,j} V_{1,j})\) is the pore volume of the cell which borders the well at layer \(j\).

Condition (C8) was introduced because the saturation at initial conditions is often assumed to be irreducible in the formation. Therefore, the initial water transmissibility is zero everywhere except at the well-reservoir interface. Hence, since water cannot flow, if the volume injected at the first iteration of the first time-step overcomes the available porous volume of the near wellbore cell, the cell saturation exceeds the \(1 - S_{hr}\) value and the convergence of Newton method is compromised.