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## Article

# Gridding Effects on CO<sub>2</sub> Trapping in Deep Saline Aquifers

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**Abstract:** Three-dimensional numerical models of potential underground storage and compositional simulation are a way to study the feasibility of storing carbon dioxide in the existing geological formations. However, the results of the simulations are affected by many numerical parameters, and we proved that the refinement of the model grid is one of them. In this study, the impact of grid discretization on CO<sub>2</sub> trapping when the CO<sub>2</sub> is injected into a deep saline aquifer was investigated. Initially, the well bottom-hole pressure profiles during the CO<sub>2</sub> injection were simulated using four different grids. As expected, the results confirmed that the overpressure reached during injection is strongly affected by gridding, with coarse grids leading to non-representative values unless a suitable ramp-up CO<sub>2</sub> injection strategy is adopted. Then, the same grids were used to simulate the storage behavior after CO<sub>2</sub> injection so as to assess whether space discretization would also affect the simulation of the quantity of CO<sub>2</sub> trapped by the different mechanisms. A comparison of the obtained results showed that there is also a significant impact of the model gridding on the simulated amount of CO<sub>2</sub> permanently trapped in the aquifer by residual and solubility trapping, especially during the few hundred years following injection. Conversely, stratigraphic/hydrodynamic trapping, initially confining the CO<sub>2</sub> underground due to an impermeable caprock, does not depend on gridding, whereas significant mineral trapping would typically occur over a geological timescale. The conclusions are that a fine discretization, which is acknowledged to be needed for a reliable description of the pressure evolution during injection, is also highly recommended to obtain representative results when simulating CO<sub>2</sub> trapping in the subsurface. However, the expedients on CO<sub>2</sub> injection allow one to perform reliable simulations even when coarse grids are adopted. Permanently trapped CO<sub>2</sub> would not be correctly quantified with coarse grids, but a reliable assessment can be performed on a small, fine-grid model, with the results then extended to the large, coarse-grid model. The issue is particularly relevant because storage safety is strictly connected to CO<sub>2</sub> permanent trapping over time.



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**Keywords:** CO<sub>2</sub> storage; numerical simulation; grid discretization; trapping mechanisms; deep saline aquifer

## 1. Introduction

Geological carbon dioxide sequestration offers a promising solution to reduce greenhouse gases in the atmosphere. The experience gained from CO<sub>2</sub> injection in many existing enhanced oil recovery (EOR) projects [1–4], as well as from operating CO<sub>2</sub> storage sites [5,6], indicates that it is feasible to store CO<sub>2</sub> in geological formations as a mitigation option to climate change [7–11].

Geological storage sites should fulfill the following conditions: the capacity to store an adequate volume of CO<sub>2</sub> under the overlying impermeable layers; the ability to accept the injected CO<sub>2</sub> at the rate that is supplied; and the seismic stability and geomechanical integrity of the storage site [12–16]. The carbon capture and storage (CCS) technology in deep geological formations involves the injection of CO<sub>2</sub> in the supercritical phase in order to achieve high fluid density and therefore maximize the storage capacity. Both

depleted reservoirs and saline aquifers can provide adequate storage, with the depleted reservoirs typically having smaller capacity but confirmed confinement ability and the aquifers potentially offering much larger storage capacity but requiring system assessment to ensure the existence of suitable conditions for CO<sub>2</sub> injection and containment. At the end of 2021, more than 50% of the existing CCS projects involved deep saline aquifers [17–21]; out of the 27 CCS network facilities operating at the end of 2021 or planned for operation before 2030, some 80% are saline aquifers and only 20% are depleted reservoirs [22].

Fluid dynamics and geochemical phenomena at the thermodynamic conditions of the storage site, especially when a massive injection of CO<sub>2</sub> occurs into the deep saline aquifers, are responsible for the development of different trapping mechanisms [23–26], namely structural/stratigraphic and hydrodynamic trapping, residual or capillary trapping, solubility trapping, ionic trapping, mineral trapping.

A way to evaluate the feasibility and safety of a CO<sub>2</sub> storage project, including the effects of each trapping mechanism under the conditions of interest, is the use of numerical models [27].

In ref. [28,29], the key parameters affecting solubility trapping and the effect of convective flow were investigated. Structural or hydrodynamic trapping was investigated by [30–33], among others. Capillary trapping, which has a fundamental contribution to sedimentary formations, was studied by [34,35], while dissolution and mineral trapping, which can be observed in both basaltic and sedimentary rocks, were studied by [36–38].

In ref. [39], the authors focused their study on the distribution of CO<sub>2</sub> between the aqueous and the gaseous phases for several aquifer scenarios. In ref. [40], the authors demonstrated the importance of considering hysteresis effects in saline aquifers since they contribute to reducing CO<sub>2</sub> migration and accumulation along seal layers. In ref. [41–43], the authors investigated both mineral and solubility trapping during CO<sub>2</sub> disposal in aquifers, analyzing porosity and permeability variations. In ref. [44], the authors carried out sensitivities analyses to study the contribution of solubility and mineral trapping under different pressure and temperature conditions. In ref. [4,45–47], the authors conducted simulation studies to investigate the CO<sub>2</sub> plume behavior and trapping under different injection strategies.

The results of these studies confirmed that the contribution of each trapping mechanism largely depends not only on the fluid-rock and mineral properties of the aquifer under consideration but also on the CO<sub>2</sub> injection strategy. Furthermore, trapping mechanisms are dependent on one another [48].

However, one aspect that—to the best of the authors' knowledge—was not thoroughly investigated by previous works concerns how spatial discretization of the volume of interest affects the simulation of CO<sub>2</sub> storage. While discretization is recognized as strongly impacting simulated pressure and saturations when fluids are produced from, or injected into, a reservoir, the impact on the simulated CO<sub>2</sub> migration and trapping mechanisms is uncertain. In this study, the effects of grid discretization on CO<sub>2</sub> solubility and residual trapping were investigated. Initially, the sensitivities were performed on the well bottom-hole pressure during CO<sub>2</sub> injection to identify which grids could correctly reproduce the pressure increase. Then, the same grids were used to simulate the CO<sub>2</sub> trapping by solubility and residual trapping during the 100, 200, and 500 years after injection. The target was to assess if and to what extent block dimensions could affect the accuracy of the simulated quantity of the CO<sub>2</sub> permanently captured underground. Given that permanently captured CO<sub>2</sub> is no longer of concern because it cannot leak to the surface, the implication is that adequate model gridding can contribute to correctly framing the safety issues potentially related to geological storage.

## 2. Model Description

A simplified 3D numerical model representing a portion of a deep saline aquifer was set up using compositional commercial software, namely GEM by CMG, which is able to

simulate coupled geochemical reactions and fluid flow. The GEM software has been used for simulations of CO<sub>2</sub> storage by various authors, such as ref. [49,50], among others.

The model dimensions were selected in order to represent all the phenomena of interest and are  $1350 \times 1350 \times 31 \text{ m}^3$ . The extension of the model size to that of a full aquifer, with a consequent significant increase in the simulation run time, was unnecessary because the target was the investigation of the solubility and the residual trapping mechanisms, which are independent of the system dimensions.

The aquifer thickness is 20 m, which is a typical value for a sedimentary formation, and this ensures that the CO<sub>2</sub> upward migration due to gravitational segregation can be observed. The overlying caprock thickness is 10 m and is sufficient to provide the aquifer sealing under the simulated operational conditions. However, a monitoring layer above the caprock was also defined to monitor any possible leak of the injected CO<sub>2</sub> through the caprock. The monitoring layer is 1 m thick.

A single injector well was defined; it was located in the middle of the model to avoid any boundary effects on the CO<sub>2</sub> plume migrating through the aquifer.

The depth of the aquifer top was set at 1500 m. The initial conditions applied to the aquifer were a pressure equal to 15 MPa and a temperature equal to 50 °C, both set at the reference depth of 1500 m. Therefore, the CO<sub>2</sub> in the aquifer can only exist as a supercritical fluid, as is recommended to grant maximum storage capacity.

Four different grid discretization options were considered. The simulations were performed with the aim of assessing the impact of discretization on the CO<sub>2</sub> fluid-flow behavior and the trapped quantities according to the different mechanisms. Initially, the bottom-hole pressure during the CO<sub>2</sub> injection was monitored. Under the assumption that multiphase fluid flow prevails during the CO<sub>2</sub> early injection, the commercial black-oil simulator Eclipse 100 (Schlumberger) was also used to benchmark the obtained results, while the commercial black-oil simulator Rubis (Kappa) was used to assess more gridding options. Then, the impact of the model discretization on the CO<sub>2</sub> trapped by the different mechanisms was assessed by simulating CO<sub>2</sub> injection and a subsequent monitoring period using GEM.

The petrophysical characteristics and the rock-fluid interaction properties were the same for all the models. Conversely, as described in the following, the boundary conditions and the PVT properties had to be set differently in the used software due to their different capabilities.

### 2.1. Rock and Fluid Properties

The rock and fluid properties used for characterizing the synthetic model are typical values for a 1500 m-deep sandstone aquifer [51,52] and are summarized in Table 1.

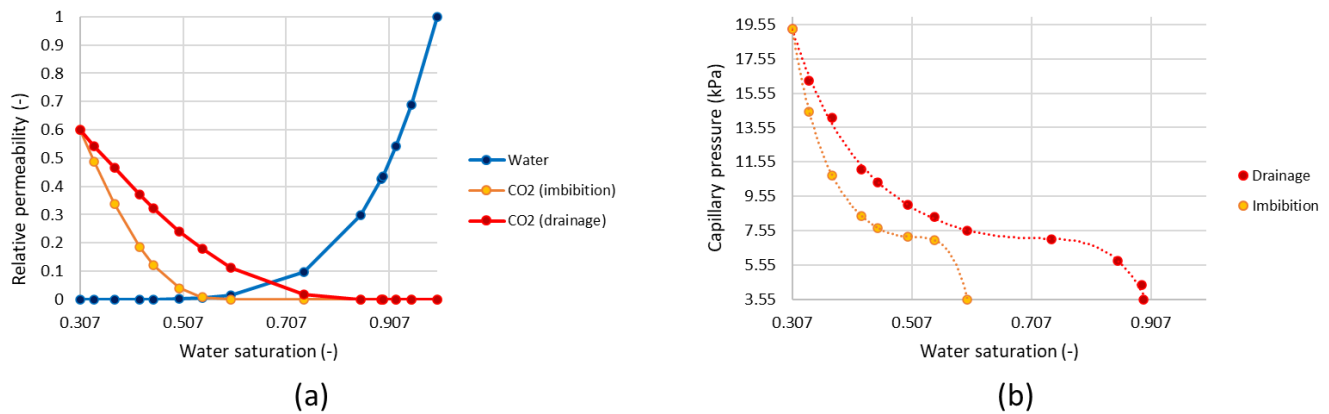
**Table 1.** Rock and fluid properties.

Aquifer porosity	20%
Aquifer permeability	200 mD
Anisotropy ratio	1.0
Rock compressibility	$4.5 \cdot 10^{-4} \text{ MPa}^{-1}$
Water compressibility	$4.35 \cdot 10^{-4} \text{ MPa}^{-1}$
Water density (at standard conditions)	1020 kg/m <sup>3</sup>
Water salinity (NaCl)	10,000 ppm

The porosity and the permeability of the caprock were set to be equal to 20% and  $10^{-7}$  mD, respectively [53]. The porosity and permeability of the monitoring layer were set to be equal to those of the aquifer (Table 1).

Water salinity is compatible with a deep aquifer; yet, it was assumed to be low in order to maximize the amount of CO<sub>2</sub> that could be solubilized [54,55].

The relative permeability curves were calculated according to the Corey–Brooks method [56], using the following exponents and endpoints from the literature:  $N_W = 5$  [57],  $N_{CO_2} = 2$  [58],  $S_{wi} = 0.307$ ,  $S_{gc} = 0.106$ , and  $S_{ghys} = 0.4$  [59]. The capillary pressure curves were also taken from the experimental data reported in the literature [60]. The relative permeability and capillary pressure curves are shown in Figure 1.



**Figure 1.** (a) Relative permeability curves; (b) capillary pressure curves.

The rock mineralogy was selected among the default options available in GEM as the rock composition was irrelevant to the scope of this study. In fact, mineral trapping was not taken into consideration as it typically occurs over thousands of years after injection.

## 2.2. PVT Properties

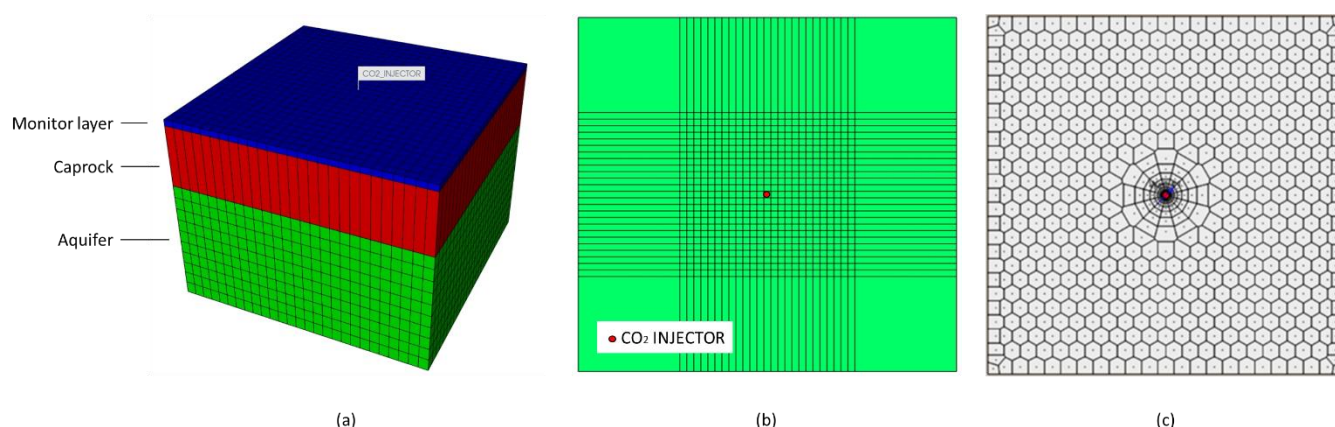
In GEM, the CO<sub>2</sub> PVT properties were calculated by the software based on the CO<sub>2</sub> critical properties and acentric factor [61]; the Peng–Robinson [62] equation of state was used. The same rock–fluid properties, as well as the PVT model used in GEM, were also adopted in Eclipse 100. In Rubis, the CO<sub>2</sub> PVT properties were taken from the software’s internal database for pure CO<sub>2</sub>.

## 2.3. Gridding Options

The effect of grid discretization on the simulated bottom-hole pressure and trapping mechanisms was evaluated using GEM by considering four grids with different discretizations (G1, G2, G3, and G4). The vertical layering of the aquifer was kept constant, with each layer having a thickness of 2 m. In grids G1, G2, and G3, the block areal dimensions were constant in the area close to the well (Figure 2a). Conversely, G4 was built using blocks whose dimensions increased according to their distance from the well: in the area surrounding the well, the blocks had a  $2 \times 2$  m<sup>2</sup> areal dimension, which progressively increased as they moved towards the model boundaries, up to  $145 \times 145$  m<sup>2</sup> (Figure 2b). The grid parameters for the four investigated cases are reported in Table 2.

The bottom of the aquifer was set as a no-flow boundary. An analytical infinite Carter–Tracy aquifer was adopted as a lateral boundary condition allowing water to exit and enter the model domain. Because the water is the only component allowed to flow through the boundary, a sufficient distance between the CO<sub>2</sub> plume and the interface with the analytical aquifer was guaranteed so as to prevent the accumulation of CO<sub>2</sub> in the external blocks of the model.

The same grid options used in GEM (Table 2), as well as the same aquifer initial conditions, were also used in Eclipse 100. The grids were named E1, E2, E3, and E4.



**Figure 2.** Gridding: (a) 3D synthetic model used in GEM showing the three zones: aquifer, caprock and monitoring layer, and the uniform discretization; (b) grid refinement for grid G2 (top view); (c) unstructured Voronoi grid R4 used in Rubis (top view).

**Table 2.** Grid parameters.

Grid	G1	G2	G3	G4
Top grid depth			1500 m	
Aquifer dimensions			$1350 \times 1350 \times 20 \text{ m}^3$	
N° of blocks			7290 ( $27 \times 27 \times 10$ )	
Block aerial dimensions	$50 \times 50 \text{ m}^2$	$25 \times 25 \text{ m}^2$	$15 \times 15 \text{ m}^2$	$2 \times 2 \div 145 \times 145 \text{ m}^2$
Block thickness			2 m	

A different approach was used for Rubis as the software does not allow the definition of an analytical aquifer. In order to reproduce an infinite aquifer, the model dimensions were increased to  $300 \times 300 \text{ km}^2$ . The software uses an unstructured Voronoi grid with a refinement in the near-wellbore area (Figure 2c). The grid refinement area developing around the well is controlled by the progressive ratio (PR): by decreasing the PR value the refinement increases. This grid refinement approach is particularly effective in simulating the pressure history during  $\text{CO}_2$  injection. Four different grids (named R1, R2, R3, and R4), were generated adopting four different progressive ratios (Table 3). The depth of the aquifer top as well as the initial conditions of the aquifer used in Rubis are the same as those used in GEM.

**Table 3.** Grids parameters in Rubis.

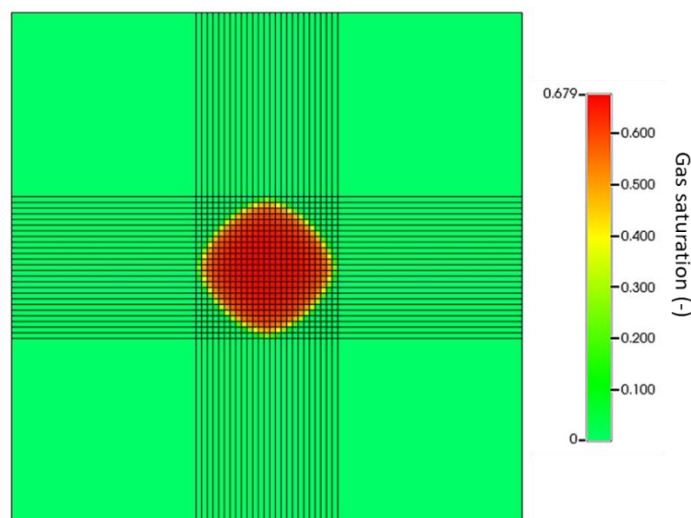
GRID	PR	Number of Angular Divisions	Number of Horizontal Layers	Total Number of Blocks
R1	4.0	12	10	6510
R2	2.5	12	10	6930
R3	1.7	12	10	7890
R4	1.4	12	10	9090

#### 2.4. Simulation Strategy

In every simulation, the perforations of the injector well, which was placed at the center of the model, were open in the 1510–1518 m depth interval. The well radius was equal to 0.0762 m, while the rate of injected  $\text{CO}_2$  was constant and equal to  $200,000 \text{ sm}^3/\text{day}$ , which is equivalent to a mass rate of 0.136 Mt/y. The duration of the injection was 120 days



so that the CO<sub>2</sub> plume did not reach the model boundary over the entire simulation period (Figure 3). The same simulation strategy was adopted in Eclipse 100 and in Rubis.



**Figure 3.** CO<sub>2</sub> saturation at the end of injection in grid G3 (top layer of the aquifer).

### 3. Methodology

Numerical simulations were aimed at assessing how spatial discretization of the volume of interest affects permanent CO<sub>2</sub> trapping by solubility and capillarity, which are the mechanisms ensuring safe CO<sub>2</sub> storage in the short term. In fact, the following considerations can be provided on the main trapping mechanisms occurring when CO<sub>2</sub> is injected into deep saline aquifers (the trapping mechanisms and a brief explanation of their effects are summarized in Table 4). In the short term, stratigraphic–structural trapping is the dominant mechanism and relies on the geological characteristics of the system. Then, when brine replaces the volume previously occupied by the injected CO<sub>2</sub>, which is thus partially trapped by capillary forces, residual trapping occurs. Solubility trapping is triggered by CO<sub>2</sub> dissolution into the brine. Even though the kinetics of this reaction is extremely fast, the dissolution of the injected CO<sub>2</sub> is a rather slow process: only a fraction of the injected volume is sequestered due to aqueous solubility over hundreds of years [48,63]. Mineral trapping occurs when the carbonate minerals precipitate due to the reaction of the dissolved CO<sub>2</sub> and the aquifer minerals or dissolved solids. Dissolution of supercritical CO<sub>2</sub> into brine controls the rate of dissolution and the precipitation of the minerals constituting the porous rocks. Volume changes in the solid phase can modify the pore structure, affecting both porosity and permeability and, therefore, altering the pathways of the injected CO<sub>2</sub>. However, mineral trapping is typically not considered in storage capacity estimation due to the complexity of the process and because it usually becomes significant only over thousands of years [64,65]. Therefore, while the extension of the CO<sub>2</sub> plume mainly depends on the aquifer geometry and the petrophysical properties and on the injection strategy, only solubility and residual trapping concur to permanently block CO<sub>2</sub> underground in the short term.

To estimate the impact of the model discretization on the simulation results when CO<sub>2</sub> is injected into the aquifer, numerical simulations were carried out using the different grids described in Section 2.1. The results were analyzed in terms of pressure response at the well and in the aquifer, the amount of CO<sub>2</sub> confined by stratigraphic–structural trapping, and the amount of CO<sub>2</sub> sequestered by solubility and residual trapping.

**Table 4.** CO<sub>2</sub> trapping mechanisms.

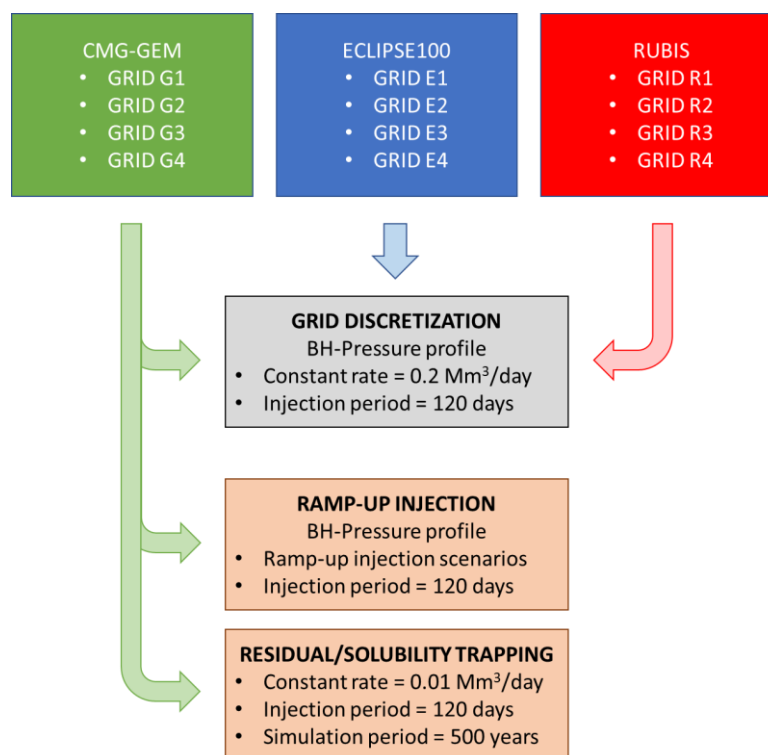
Structural/stratigraphic and hydrodynamic trapping	The buoyant CO <sub>2</sub> remains as a mobile fluid but is prevented from flowing back to the surface by an impermeable cap rock (structural/stratigraphic trapping). For pressure and temperature conditions above the critical point ( $p = 7.3$ MPa and $T = 31$ °C), which correspond to a depth of approximately 1000 m, CO <sub>2</sub> is a supercritical fluid. While the viscosity values remain gas-like, the CO <sub>2</sub> density is liquid-like. However, CO <sub>2</sub> the density is still significantly smaller than the density of the saline formation water. As a consequence, CO <sub>2</sub> rises towards the caprock due to buoyancy forces. When it reaches the top of the formation, it continues to slowly flow laterally (hydrodynamic trapping).
Residual or Capillary trapping	This is the disconnection of the CO <sub>2</sub> phase into an immobile (trapped) fraction. In sedimentary rocks that do not contain organic material, including sandstones and carbonates, carbon dioxide is the non-wetting fluid relative to brine. As a result, during the injection stage CO <sub>2</sub> displaces the formation water (drainage process). However, after injection, the buoyant CO <sub>2</sub> migrates laterally and upward, and water displaces CO <sub>2</sub> in an imbibition-like process. This leads to the disconnection of the once-continuous plume and to the formation of residual gas (trapped CO <sub>2</sub> ). Larger grain sizes correspond to increased hydraulic conductivity but negatively affect residual trapping efficiency. Relative permeability hysteresis is a key factor in the assessment of CO <sub>2</sub> residual trapping.
Solubility trapping	This refers to the dissolution and hydration of CO <sub>2</sub> in the brine to form carbonic acid ( $\text{CO}_2 + \text{H}_2\text{O} \leftrightarrow \text{H}_2\text{CO}_3$ ), possibly enhanced by gravity instabilities due to the larger density of the brine–CO <sub>2</sub> liquid mixture. In fact, the brine with dissolved CO <sub>2</sub> is slightly denser (by almost 1%) than the brine without dissolved CO <sub>2</sub> . CO <sub>2</sub> solubility in water increases with increasing pressure and decreases with increasing temperature and water salinity. When CO <sub>2</sub> dissolves in brine it forms a weak acid. The time scale of the process is hundreds of years.
Ionic trapping	This refers to the reactions due to the ion exchange between the carbonic acid and the formation liquid ( $\text{H}_2\text{CO}_3 \leftrightarrow \text{H}^+ + \text{HCO}_3^-$ and then $\text{HCO}_3^- \leftrightarrow \text{H}^+ + \text{CO}_3^{2-}$ ). It occurs for pH values higher than 6. It is a preliminary step to mineralization, which will take place if Ca, Fe, and Mg cations are available.
Mineral trapping	This involves the reactions between the dissolved CO <sub>2</sub> and the formation rock minerals with the geochemical binding to the rock due to mineral precipitation. The CO <sub>2</sub> –brine–rock interaction enables both mineral dissolution and the generation of secondary minerals from precipitation. The duration of this process depends on various factors such as the mineral kinetic properties and reactive surface areas [66], and it can last up to thousands of years.

The first step of the study focused on the impact of the discretization on the well bottom-hole pressure profiles. To this end, numerical simulations were performed using GEM and using Eclipse 100 as a benchmark, based on the assumption that multiphase fluid flow dominates during the very first stage of injection; thus, the compositional capabilities would have a limited impact on the wellbore pressure trend. Rubis was also used due to the possibility of defining very refined grids in the near-wellbore area. In the second step, three different injection history scenarios were assigned to the model. The analysis of the simulation results concentrated on the induced well bottom-hole pressure profiles because it is well-known that coarse grids can generate unrealistically high pressure values, invalidating the representativeness of the whole simulation [67,68].

Then, the impact of discretization on residual and solubility trapping was investigated by simulating 100, 200, and 500 years after CO<sub>2</sub> injection, respectively, and monitoring the CO<sub>2</sub> distribution and dissolution. In these cases, the rate of CO<sub>2</sub> injection was reduced to 10,000 sm<sup>3</sup>/day (the injection period was kept equal to 120 days) in order to limit the extension of the plume of CO<sub>2</sub>, both in free phase and in solution, to the near-wellbore area. This is where the grid refinements were applied with respect to the base case scenario (having uniform grid blocks) and thus where the potential impact of gridding on the simulation results was expected to be detectable.

Mineral trapping was not accounted for because of its negligible contribution to the total amount of CO<sub>2</sub> trapped in the considered time frame [48]. The different scenarios are summarized in Figure 4.





**Figure 4.** Rate injection scenarios.

## 4. Results

### 4.1. Grid Discretization

The results of the simulations performed with GEM show that when injection starts, the well bottom-hole pressure reaches a peak; then, the pressure progressively decreases until stabilizing (Figure 5). The value of this peak and the time to reach it are proportional to the dimension of the blocks (Table 5). The pressure peak ranges from 113% (G4) to 125% (G1) of the initial aquifer pressure; at the end of the injection, the bottom-hole pressure reaches values ranging between 106.2% (G4) and 107.5% (G1).

The same pressure trends observed with GEM were confirmed by the Eclipse100 simulation results, even though the values of the overpressure peaks were different (Table 5 and Figure 6). Furthermore, the reservoir pressure reached at the end of injection, ranging from 104.6% (E4) to 108.7% (E1), also showed significant dependency on the gridding.

When Rubis was used for the simulation of the same scenarios, the bottom-hole pressure peaks were much lower with respect to the ones observed in the GEM and Eclipse 100 simulations, as well as to the time needed to reach the peak values (Figure 7). By increasing the near-wellbore grid refinement through the PR parameter, the bottom-hole pressure peak decreases from 110.4% (grid R1, PR = 4.0) to 108.5% (grid R4, PR = 1.4) of the initial aquifer pressure. No significant effects on the bottom-hole pressure at the end of the CO<sub>2</sub> injection were detected for the four grids (Figure 7).

In summary, all the sensitivities showed that the grid block dimensions always have a significant impact on the simulated overpressure during injection, and in most cases, the maximum bottom-hole pressure largely exceeded the final overpressure value. To further confirm these results, it can be observed that the simulations performed with Rubis, which allowed the adoption of a very refined grid around the wellbore, were those providing the lower values of the local pressure increase at the wellbore.

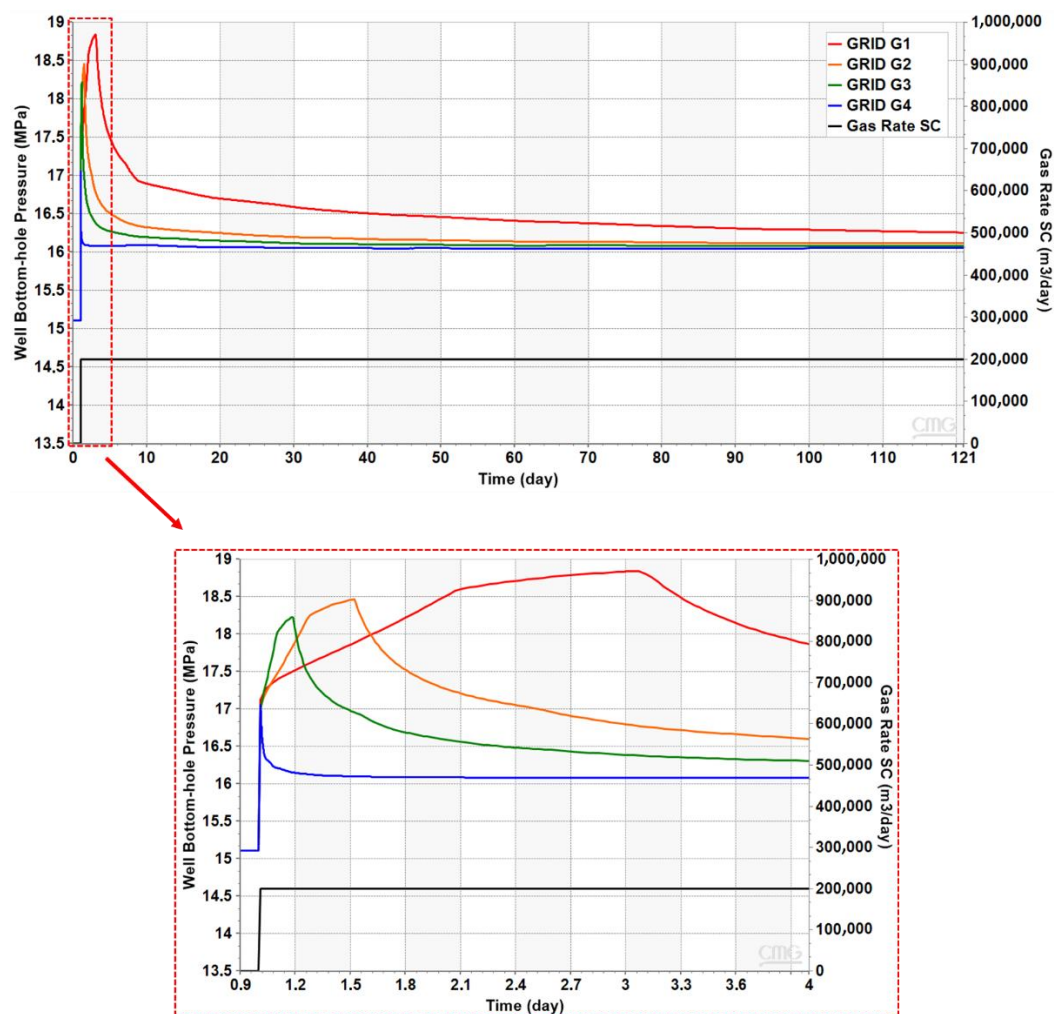
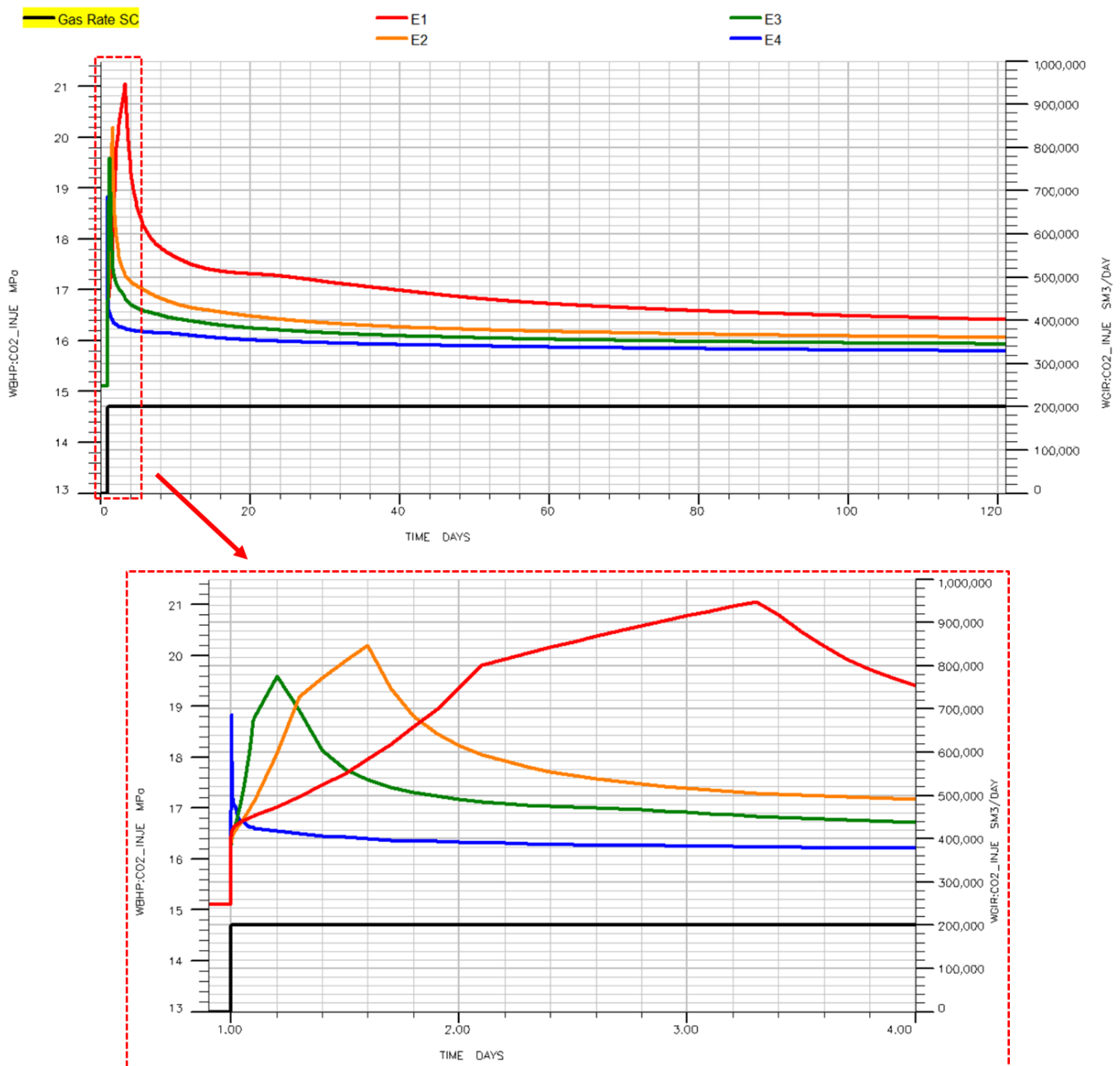


Figure 5. Well bottom-hole pressure profiles for grids G1, G2, G3, and G4.

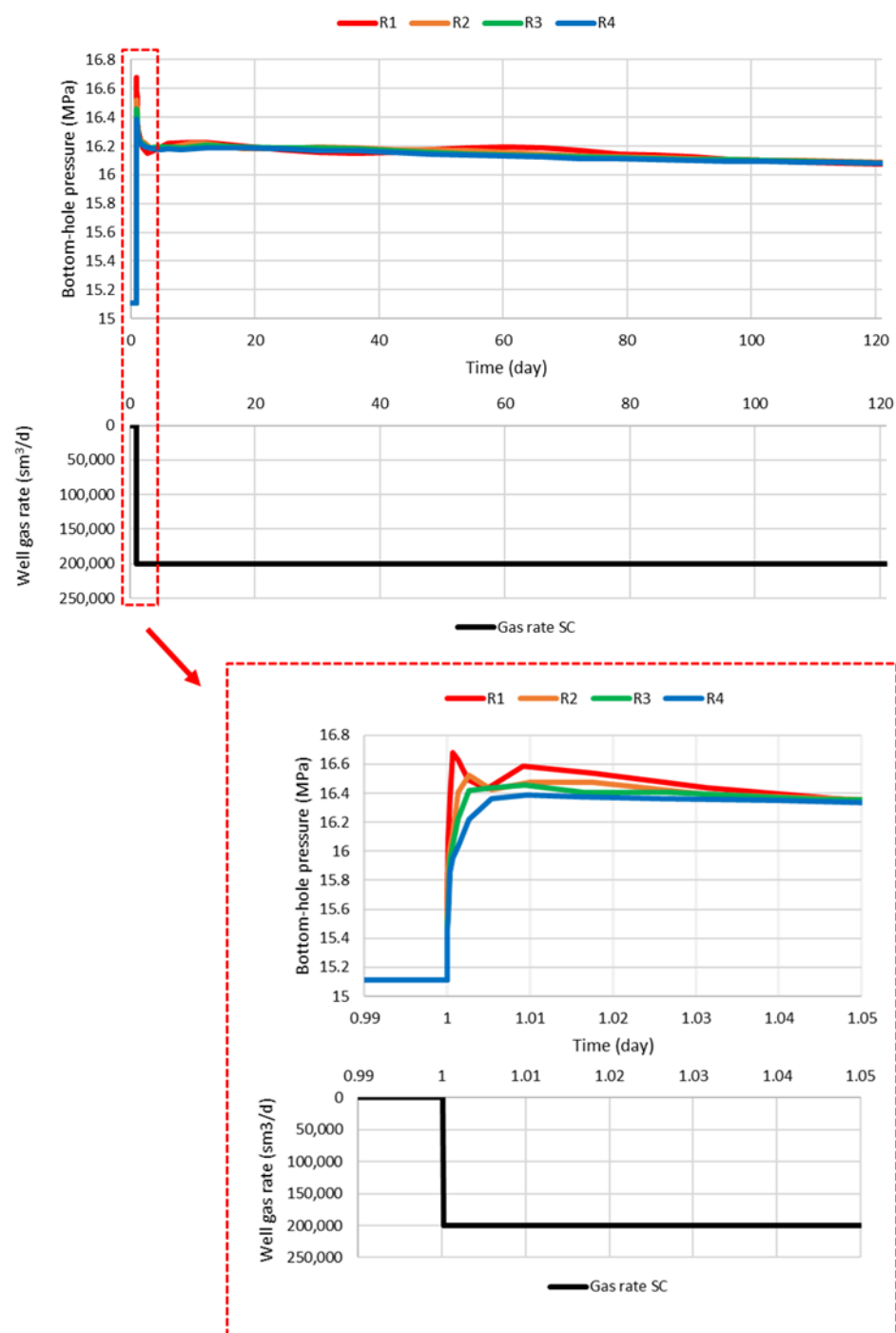
Table 5. Grid discretization effects on well bottom-hole pressure profiles.

GRID	Bottom-Hole Pressure Peak (MPa)	Peak Overpressure (Mpa)	Time Needed to Reach the Pressure Peak (Days)	Final Bottom-Hole Pressure (MPa)
G1	18.84	3.73	2.08	16.25
G2	18.46	3.35	0.52	16.11
G3	18.23	3.12	0.19	16.08
G4	17.06	1.95	<0.01	16.05
E1	21.06	5.95	2.3	16.43
E2	20.17	5.06	0.6	16.08
E3	19.58	4.47	0.2	15.93
E4	18.85	3.74	<0.01	15.80
R1	16.68	1.57	<0.01	16.07
R2	16.52	1.41	<0.01	16.09
R3	16.45	1.34	<0.01	16.08
R4	16.39	1.28	<0.01	16.08



**Figure 6.** Well bottom-hole pressure profiles for grids E1, E2, E3, and E4.

Even though no CO<sub>2</sub> above the caprock was detected due to leakage induced by overpressure conditions, it should be noted that the effects of the caprock threshold pressure and most of the geomechanics of the system were not simulated. Should a more comprehensive assessment be performed, the simulated peak pressure values would likely not be acceptable. However, these pressure peaks at the wellbore are not due to physical phenomena but are only due to modeling issues; thus, they are not representative of the response of the real system. Furthermore, while the results on the pressure trend induced during injection were expected, the question is the extent to which the model discretization can affect the quantity of CO<sub>2</sub> trapped in the aquifer by the different trapping mechanisms.



**Figure 7.** Well bottom-hole pressure profiles for the grids R1, R2, R3, and R4.

#### 4.2. Ramp-Up Injection Scenarios

Ideally, a highly refined grid should be used to correctly model the pressure trend during  $\text{CO}_2$  injection. However, considering that a number of injector wells would be typically simulated in a full reservoir study, the need for grid refinement might significantly increase the number of grid blocks and thus also dramatically increase the simulation run time and/or the computing requirements.

A good strategy to compromise between grid refinement and the fictitious local pressure increase at the wellbore could be to inject  $\text{CO}_2$  according to a rate ramp-up. To this end, three scenarios were simulated. Each scenario consisted in ramping the rate up to the original value of  $200,000 \text{ sm}^3/\text{day}$ , with a different duration and increase in the

subsequent rate steps. The three ramp-up scenarios are reported in Table 6 along with the corresponding values of the maximum well bottom-hole pressure and of the well bottom-hole pressure after injection. The results show that a ramp-up injection rate can effectively mitigate the non-physical well pressure increase as well as a progressive convergence between the maximum well bottom-hole pressure and the final aquifer overpressure after injection. The results are also shown in Figure 8.

**Table 6.** Ramp-up injection scenarios.

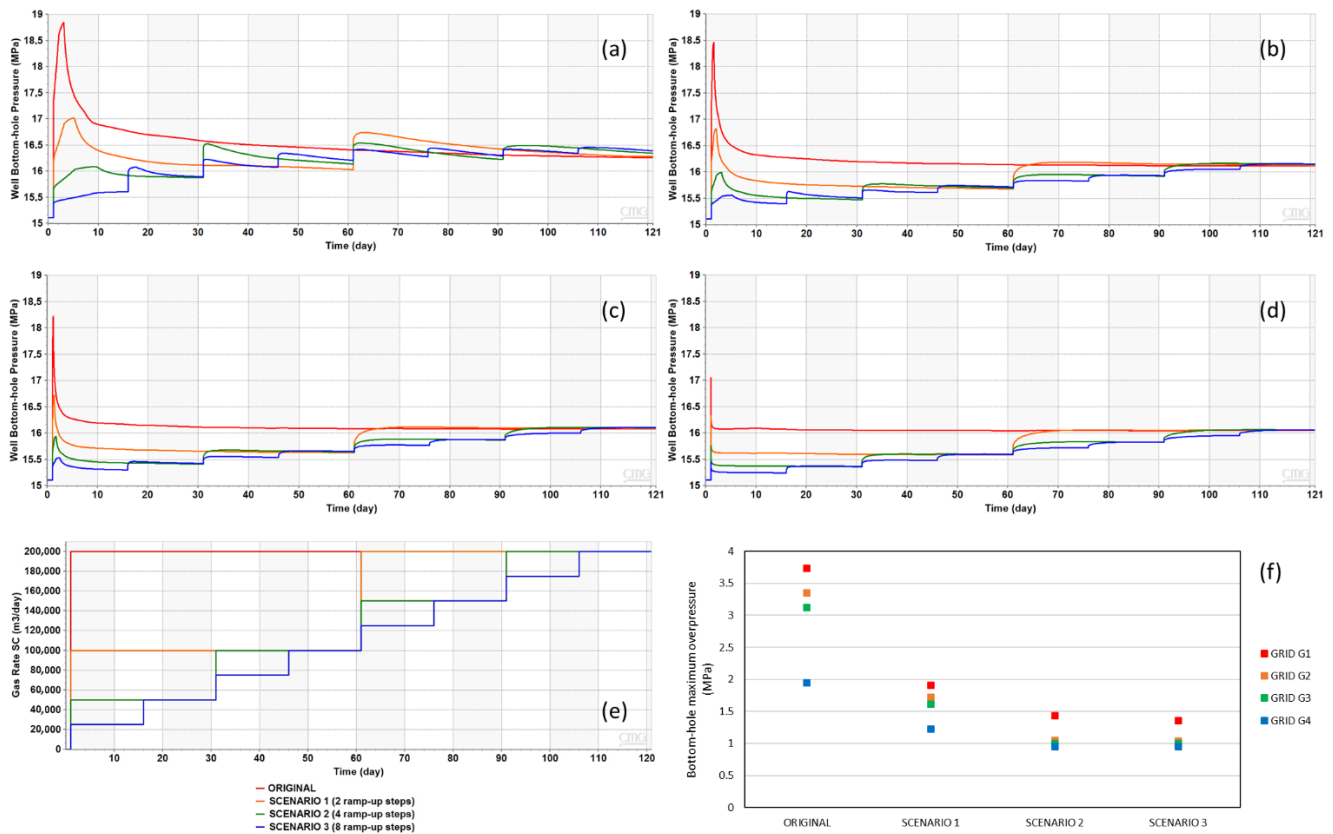
Scenario	ORIGINAL	1	2	3
Steps	1	2	4	8
Time step (days)	120	60	30	15
Rate step (sm <sup>3</sup> /day)	200,000	100,000	50,000	25,000
GRID G1				
Maximum well bottom-hole pressure (MPa)	18.84	17.02	16.54	16.45
Maximum well bottom-hole overpressure (MPa)	3.73	1.91	1.43	1.35
Well bottom-hole pressure after injection (MPa)	16.25	16.28	16.34	16.39
GRID G2				
Maximum well bottom-hole pressure (MPa)	18.46	16.83	16.16	16.15
Maximum well bottom-hole overpressure (MPa)	3.35	1.72	1.05	1.04
Well bottom-hole pressure after injection (MPa)	16.11	16.13	16.14	16.15
GRID G3				
Maximum well bottom-hole pressure (MPa)	18.23	16.72	16.11	16.11
Maximum well bottom-hole overpressure (MPa)	3.12	1.61	1.00	1.00
Well bottom-hole pressure after injection (MPa)	16.08	16.09	16.10	16.11
GRID G4				
Maximum well bottom-hole pressure (MPa)	17.06	16.34	16.06	16.06
Maximum well bottom-hole overpressure (MPa)	1.95	1.23	0.95	0.95
Well bottom-hole pressure after injection (MPa)	16.05	16.06	16.06	16.06

#### 4.3. Gridding Effects on CO<sub>2</sub> Solubility and Residual Trapping

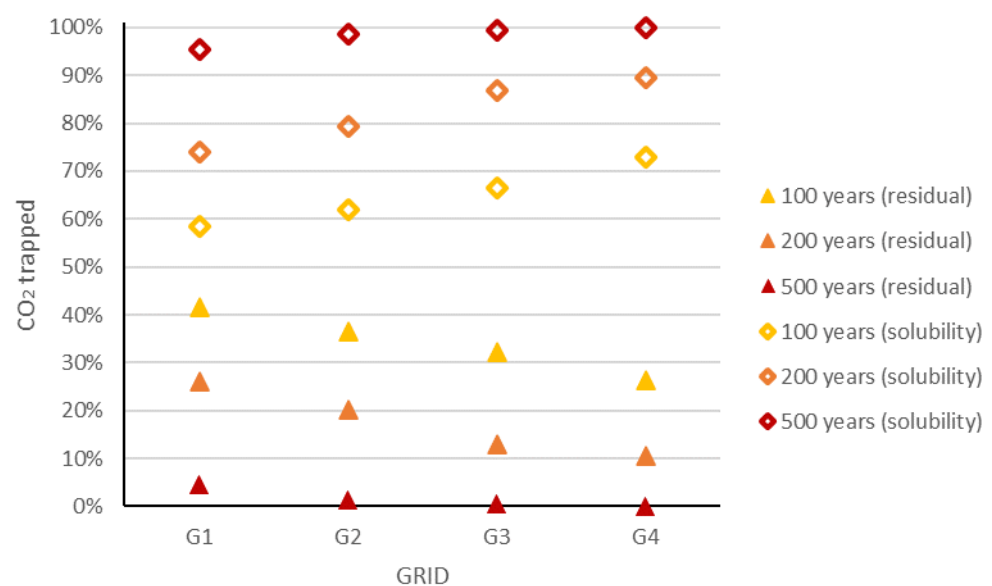
The grid discretization also proved to strongly affect CO<sub>2</sub> trapping. The simulations were performed for monitoring periods of 100, 200, and 500 years, respectively, after injection. The three monitoring periods were defined so as to validate the obtained results.

As expected, stratigraphic/hydrodynamic trapping does not depend on gridding, whereas mineral trapping does not concur to constrain the CO<sub>2</sub> underground in the short term. However, given that only a small volume of CO<sub>2</sub> was injected into the aquifer, in our simulations all the CO<sub>2</sub> was either dissolved into the water or trapped by capillary forces some 100 years after injection. The results showed that the model discretization strongly affects the solubility trapping and, consequently, the residual trapping of the CO<sub>2</sub>. A different trend is observed between these trapping mechanisms (Figure 9). The CO<sub>2</sub> immobilized by residual trapping decreases from the coarse grid to the fine grid (i.e., from G1 to G4), while an opposite trend is observed for solubility trapping. This translates into an average of a 15% discrepancy between the trapped CO<sub>2</sub> by both residual and solubility trapping for the monitoring periods of 100 and 200 years. After 500 years of simulation, the CO<sub>2</sub> was almost completely dissolved into the brine for all the simulated grids, except for G1 in which 5% of the CO<sub>2</sub> remained trapped by capillary forces. This is due to the supercritical CO<sub>2</sub>-formation water interface surface, which is larger in fine grids compared to coarse grids; as supercritical CO<sub>2</sub> dissolves into the aqueous phase only at the CO<sub>2</sub>-water

interface, the solubility trapping of CO<sub>2</sub> is strongly dependent on the extension of this interface [48]. The higher the amount of CO<sub>2</sub> dissolved into the formation water, the lower the CO<sub>2</sub> which remains trapped as a supercritical phase by residual trapping.



**Figure 8.** Well bottom-hole pressure profiles for the grids (a) G1, (b) G2, (c) G3, and (d) G4 for ramp-up injection scenarios 1, 2, and 3; (e) well gas rate SC for ramp-up injection scenarios 1, 2, 3; (f) well bottom-hole maximum overpressure for the grids G1, G2, G3, and G4 for ramp-up injection scenarios 1, 2, and 3.



**Figure 9.** Percentage of CO<sub>2</sub> trapped by residual and solubility trapping mechanisms for grids G1, G2, G3, and G4 after 100, 200, and 500 years, respectively.



## 5. Discussion and Conclusion

In the oil field, reservoir simulation has been used for decades; thus, extensive knowledge has been gained. Model discretization is recognized to strongly affect the simulation results, with fine grids leading to more reliable results than coarse grids but requiring significant—and sometimes prohibitive—simulation run time and/or computational resources. For this reason, when we simulated CO<sub>2</sub> injection into a deep saline aquifer, the impact of the model discretization on the simulated pressure values was expected. Conversely, the potential impact of model discretization on the amount of permanently trapped CO<sub>2</sub> could not be anticipated because dedicated studies were not found in the technical literature.

Our simulation results showed that a coarse grid in the near-wellbore area generates an unrealistically high bottom-hole pressure profile at the beginning of injection, as well as a higher average aquifer pressure over the years with respect to a fine grid. The reason for this is that, initially, there is no CO<sub>2</sub> in the aquifer (i.e., the CO<sub>2</sub> saturation is zero); thus, the effective permeability is also null. For CO<sub>2</sub> to become mobile, the saturation must reach a critical value; however, the larger the grid blocks, the larger the volume of CO<sub>2</sub> needed to reach this critical saturation and thus the higher the grid block pressure. This is why fine grids exhibit a much smaller pressure increase than coarse grids. Even if to a slightly different extent, this phenomenon is well represented by all the software used in this work for the simulations. The main issue related to a fictitious overpressure in the near-well grid blocks is that the pressure might exceed the limiting value, which is usually imposed to avoid damaging the well completion and fracturing the formation. If the assigned pressure constraint is exceeded, the CO<sub>2</sub> injection rate is automatically reduced so that the induced pressure decreases below the constraint. As a consequence, the rate at which CO<sub>2</sub> can be injected is underestimated. Furthermore, an overestimation of the average aquifer pressure might be misleading when the integrity of the aquifer and caprock is investigated under operational conditions through geomechanical analyses. In this study, we demonstrated that the maximum pressure values at the well and the aquifer average pressure can be much more realistic if an injection strategy with a rate ramp-up is implemented. By increasing the number of rate steps and decreasing the rate increments, the fictitious increase in the well bottom-hole pressure, especially during early CO<sub>2</sub> injection, can be avoided even if adopting a coarse discretization. Thus, a well-calibrated ramp-up injection strategy can be very efficient in obtaining consistent simulation results without excessive grid refinement in the near-wellbore area. Avoiding fine grids or grid refinements, and thus reducing the number of grid blocks, is particularly important in view of large models and multiple injectors.

Discretization also proved to impact the amount of CO<sub>2</sub> permanently trapped in the aquifer. In fact, our results showed that the CO<sub>2</sub> trapped by residual and solubility trapping is highly affected by the block dimensions. Smaller blocks lead to a higher quantity of CO<sub>2</sub> dissolved into the formation water and to a lower amount of CO<sub>2</sub> trapped by residual trapping. Based on the simulation results obtained for the pressure increase induced by the CO<sub>2</sub> injection, we deem it reasonable to confirm that fine grids provide more reliable results. Therefore, it can be affirmed that coarse grids lead to an underestimation of the CO<sub>2</sub> trapped by solubility and to an overestimation of the CO<sub>2</sub> trapped by residual trapping. Even though mineral trapping was not simulated, because it typically occurs over a very long timescale and it was beyond the scope of this study, it is evident that an underestimation of the CO<sub>2</sub> trapped by solubility will also lead to an underestimation of the quantity of mineralized CO<sub>2</sub> in the long term. This underestimation of the amount of CO<sub>2</sub> trapped by solubility and mineral trapping compromises the reliability of the entire CCS simulation and directly translates into an incorrect evaluation of the storage safety.

In conclusion, high-resolution models would be needed to reliably simulate the pressure evolution during CO<sub>2</sub> injection and accurately describe the migration paths of the injected CO<sub>2</sub> in the aquifer and its trapping by the different mechanisms. However, as already discussed, discretization also plays a key role in the simulation run time and/or

computational requirements. Thus, a compromise is needed between these two conflicting targets. The use of a CO<sub>2</sub> injection strategy with a rate ramp-up and sensitivities on a small model such as the one we presented in this paper can help to improve the confidence in the simulation results from large full-aquifer models when a coarse discretization is adopted.

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