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Title: Quantification of CO2 generation in sedimentary basins through Carbonate/Clays Reactions with uncertain thermodynamic parameters

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Abstract: We develop a methodological framework and mathematical formulation which yields estimates of the uncertainty associated with the amounts of CO2 generated by carbonate-clays reactions (CCR) in largescale subsurface systems to assist characterization of the main features of this geochemical process. Our approach couples a one-dimensional compaction model, providing the dynamics of the evolution of porosity, temperature and pressure along the vertical direction, with a chemical model able to quantify the partial pressure of CO2 resulting from minerals and pore water interaction. The modeling framework we propose allows (i) estimating the depth at which the source of gases is located and (ii) quantifying the amount of CO2 generated, based on the mineralogy of the sediments involved in the basin formation process. A distinctive objective of the study is the quantification of the way the uncertainty affecting chemical equilibrium constants propagates to model outputs, i.e., the flux of CO2. These parameters are considered as key sources of uncertainty in our modeling approach because temperature and pressure distributions associated with deep burial depths typically fall outside the range of validity of commonly employed geochemical databases and typically used geochemical software. We also analyze the impact of the relative abundancy of primary phases in the sediments on the activation of CCR processes. As a test bed, we consider a computational study where pressure and temperature conditions are representative of those observed in real sedimentary formation. Our results are conducive to the probabilistic assessment of (i) the characteristic pressure and temperature at which CCR leads to generation of CO2 in sedimentary systems, (ii) the order of magnitude of the CO2 generation rate that can be associated with CCR processes.

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1	Quantification of CO2 generation in sedimentary basins through Carbonate/Clays Reactions
2	with uncertain thermodynamic parameters
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Abstract

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40

41 Introduction

42 Natural accumulations of CO_2 are commonly observed in sedimentary basins. The carbon 43 dioxide occurs as a gaseous phase with proportions ranging from 5% to 100% of the total gas phase 44 volume. These CO_2 accumulations are exploited in several sectors, including, for example, the food 45 industry (e.g., Broadhead et al., 2009) or within the context of Enhanced Oil Recovery (EOR; e.g., Allis et al., 2001 and references therein) operations. They are also investigated as natural analogs for 46 47 improving our understanding and design of subsurface CO₂ storage protocols/technologies and for 48 the assessment of the ensuing environmental risks associated with diverse migration pathways 49 connecting sources with receptors (e.g., Metz et al., 2005). Accumulation of CO_2 in sedimentary 50 basins can lead to dilution of valuable hydrocarbon gas mixtures (e.g., methane and propane), thus 51 reducing energy storage in a reservoir and resulting in an increased production cost (Imbus et al., 52 1998).

53 Various authors indicate diverse organic and/or inorganic processes as possible causes of 54 natural CO2 accumulation (e.g., Higgs et al., 2013; Hutcheon and Abercrombie, 1990; Clayton et al., 55 1990; van Berk et al., 2013; Smith and Ehrenberg, 1989; Chiodini et al., 2007; Ballentine et al., 2001; 56 Cooper et al. 1997; Dubacq et al., 2012; Fischer et al. 2006; Houtcheon et al., 1980; Imbus et al, 1998; 57 Cathles and Schoell, 2007; Kotarba and Nagao, 2008; Li et al, 2008; Mayo and Muller. 1997; 58 Wycherley et al. 1999; Cai et al. 2001; Farmer, 1965; Goldsmith 1980 Arnórsson, 1986, Chiodini et 59 al., 2000; Fischer et al., 2006). Among these sets of processes, in this study we focus on CO_2 60 generation in sedimentary formations through the Carbonate/Clay Reaction (CCR) mechanism. The 61 role of CCR as a possible relevant CO_2 generating mechanism in sedimentary systems is originally 62 suggested by Hutcheon and Abercrombie (1990), Hutcheon et al. (1980), Hutcheon et al. (1990), 63 Hutcheon et al. (1993). The feasibility of CCR occurrence in a sedimentary environment is supported 64 by a significant amount of studies (e.g., Coudrain-Ribstein and Gouze, 1993, Coudrain-Ribstein et 65 al., 1998, Cathles and Schoell, 2007; Giggenbach, 1980; Smith and Ehrenberg, 1989; Chiodini et al.,

2007; Xu and Pruess, 2001; van Berk et al., 2009). These works document a series of field data about 66 CO_2 partial pressure and/or pore-water chemical compositions sampled in real sedimentary basins 67 68 and/or computed through geochemical speciation models which are compatible with the CCR 69 mechanism. Cathles and Schoell (2007) propose a clear and schematic conceptual model of CO_2 70 generation through CCR and provide a mathematical formulation relying on a chemical equilibrium 71 model for the identification of the environmental conditions (temperature and pressure) at which CO_2 72 may be generated as a separate gas phase. These authors illustrate the use of their model through an 73 exemplary setting assuming a time-invariant linear relationship between temperature and pressure, 74 along the lines of Smith and Ehrenberg (1989). The results of this study suggest that CCR may 75 become a relevant process for gaseous CO₂ generation at a temperature of about 330 °C. Even as the 76 results of the illustrative example of Cathles and Schoell (2007) are not directly transferable to a real 77 sedimentary basin setting (where temperature and pressure vary with time according to a higher 78 complexity pattern), they clearly suggests that the CO_2 gas generation associated with CCR is 79 expected to occur at very high temperatures and pressures.

Uncertainties associated with thermodynamic parameters characterizing CCR are virtually ubiquitous. This is the consequence of a variety of factors (including, e.g., intrinsic natural variability of mineral compositions, non-ideal behavior of multiphase solutions, paucity and/or inaccuracy of available experimental data) and constitutes a critical challenge for the robust characterization of geochemical processes taking place at high temperature and pressure which are typically observed in deep sedimentary formations.

In this context, the major objective of our study is to propose a general framework within which we develop a modeling approach which incorporates the uncertainty associated with the thermodynamic parameters characterizing the CCR mechanism to yield a quantitative estimation of the amount of CO_2 released from CCR in sedimentary formations. Our approach is grounded on two coupled components: (*i*) a compaction model, simulating the burial history of a sedimentary basin; and (*ii*) a geochemical model which quantifies the amount of generated CO_2 (as a dissolved or

92 separate gaseous/supercritical phase) on the basis of thermodynamic equilibrium concepts. For the 93 purpose of demonstrating our approach, we consider the one-dimensional compaction model 94 presented by Formaggia et al. (2013), Porta et al. (2014) and Colombo et al. (2016), other numerical 95 models (eventually characterized by an increased degree of complexity) being fully compatible with 96 our methodological framework. Quantification of CO_2 in aqueous and gaseous phase in surface 97 environments or shallow subsurface systems is generally tackled through a hydro-geochemical 98 speciation software (e.g., Phreeqc, Parkhurst and Appelo, 2013). Available databases supporting 99 these software are typically considered as reliable within a range of temperatures lower than 300 °C. 100 Settings of the kind investigated in this study are characterized by temperatures larger than 300 °C 101 and pressure values significantly larger than those typically found in shallow aquifer systems. Hence, 102 we employ here an *ad-hoc* geochemical model which is consistent with the formulations proposed by 103 Giggenbach (1981), Coudrain-Ribstein et al. (1998) and Cathles and Schoell (2007) and can be 104 applied in the presence of temperature/pressure conditions taking place in deep sedimentary 105 formations.

106 We highlight that a major element of novelty of our work is the analysis of the way 107 uncertainties associated with the thermodynamic parameters employed to characterize the CCR 108 mechanism, i.e., the mineral solubility and phase equilibrium constants, propagate to the final model 109 outputs. These parameters are viewed as random model inputs characterized by a given probability 110 density function (*pdf*). As a consequence, all outputs are considered in a probabilistic framework. A 111 variety of additional sources of model and parametric uncertainty (Neuman, 2003) may affect the 112 outputs of the proposed modeling approach. These include, e.g., the salinity of the brine and the 113 feedback with other geochemical processes which may take place in sedimentary systems. In this 114 work we focus on the characterization of parametric uncertainty related to thermodynamic 115 equilibrium constants, because these parameters are not firmly constrained at the pressure and 116 temperature conditions of interest. To the best of our knowledge, an assessment of this kind is still lacking in the context of basin scale modeling of CCR processes. 117

118 Key target quantities that we consider as model outputs are the amount of CO_2 produced in 119 the system and its temporal dynamics resulting from the compaction processes of the sediment 120 evolving along geologic time scales. Results stemming from our approach include an explicit quantification of the depth at which the source of gaseous CO_2 is located and of the impact of the 121 122 relative abundance of primary phases affecting the generation of CO_2 . As a first test bed to illustrate 123 our methodology, we implement the conceptual and numerical model proposed on a realistic 124 sedimentary basin setting in terms of temperature-pressure-porosity, upon considering multiple 125 scenarios in terms of relative abundance of CCR primary phases in the mineralogical assemblage. 126 We base this study on a streamlined conceptual and numerical model of the system to allow (i) 127 focusing on the stochastic analysis of selected uncertain quantities and (ii) comparing our results 128 against available literature data. As such, we consider uncertainty to be embedded in the effects of 129 the temperature on the thermodynamic constants regulating the equilibrium between CO_2 - water -130 mineral phases, all of the remaining model features being treated as deterministic (see also Section 3 for a detailed discussion). The methodological framework we propose is then portable to scenarios 131 132 characterized by an increased level of complexity and in the presence of a variety of sources of 133 uncertainty.

134 The work is structured as follows: in Section 1 we provide a brief overview of the CCR 135 process; Section 2 illustrates the theoretical framework and modeling workflow as well as the coupled 136 formulation of the geochemical and basin models we employ to quantify the CO_2 generated during the basin evolution; in Section 3 we illustrate the main sources of uncertainty which can arise in our 137 138 modeling procedure and classify these into modeling and parametric uncertainties; in Section 4 we 139 present the main results obtained by the implementation of the modeling workflow for a basin-scale 140 case study; Section 5 is devoted to a detailed discussion and analysis of the results. We provide 141 conclusions and an overview on future perspectives in Section 6.

142 **1** Overview of CCR processes

143 Previous works (e.g. Giggenbach, 1980) have shown that the presence of carbonate phases 144 along with clavs and/or alumino-silicates in high-temperature geothermal or sedimentary systems acts 145 as a buffer system for the pore-water and might then control the partial pressure of CO_2 . Assuming 146 that the rock-fluid system attains an equilibrium, we can model the interaction between carbonates, alumino-silicates, clays and CO_2 as a single equilibrium reaction. The latter is typically termed 147 148 Carbonate/Clays Reaction (CCR), following the nomenclature introduced by Hutcheon et al. (1980). Several authors (Giggenbach, 1978; Giggenbach, 1981; Giggenbach, 1984, Coudrain-149 150 Ribstein et al., 1998; Cathles and Schoell, 2007; Hutcheon and Abercrombie, 1990; Hutcheon et al., 151 1980; Hutcheon at al., 1989; Hutcheon et al., 1993; Zhang et al., 2000; Huang and Longo, 1994; Ueda 152 et al., 2005) suggest a variety of chemical equilibrium relationships to depict the stoichiometry of 153 CCR. Table 1 lists a set of CCRs following the study of Coudrain-Ribstein et al. (1998). These can 154 be generalized through a chemical equilibrium relationship of the kind

155

$$\alpha_1 M_1 + \dots + \alpha_m M_m = \alpha_{m+1} M_{m+1} + \dots + \alpha_{n+m} M_{n+m} + \alpha_0 CO_{2(g)}$$
(1)

156 where M_k (k = 1, ..., n+m) represents the k^{th} mineral phase involved in the process and α_k are 157 stoichiometric coefficients.

158 Since the generation of CO_2 is the result of the consumption of the mineral phases of the left 159 hand side of (1), we indicate M_k with $1 \le k \le m$ or $m + 1 \le k \le m + n$ as primary or secondary phases 160 of the system, respectively. Primary phases include a carbonate mineral (which represents the source of CO_2) together with other cations (e.g., Mg^{+2} and Ca^{+2}), and additional clay/alumino-silicate phases. 161 The latter act as source of other ions (e.g., OH^2 , Al^{3+} and K^+) when dissolved in water. All reactions 162 listed in Table 1 include dolomite as carbonate mineral. The secondary phases include CO_2 , clay 163 164 minerals (e.g., chlorite, phoglopite, illite; Bergaya and Lagaly, 2013) and other species (e.g., calcite) which act as sinks for the ions released by the primary phases and represent a more stable 165 mineralogical assemblage at large temperature (e.g., T > 300 °C), as compared to primary phases 166

167 (Giggenbach, 1981; Hutcheon and Abercrombie, 1990; Smith and Ehrenberg, 1989). Partitioning 168 among primary and secondary phases in the system can be described through the equilibrium constant 169 (K_R). All phases appearing in (1) are in pure liquid or solid phases, CO_2 being the only gaseous 170 phase. The logarithmic transform of K_R is

171
$$\log K_R = \alpha_0 \log \eta_{CO2(g)} + \sum_{k=m+1}^{n+m} \alpha_k \log a_{Mk} - \sum_{k=1}^m \alpha_k \log a_{Mk}$$
(2)

172 a_{Mk} and η_{CO2} respectively representing the activity of species M_k and the CO_2 fugacity. Assuming 173 that the fugacity coefficient of CO_2 is equal to one (Hutcheon, 1990, Chiodini et al., 2007; Cathles 174 and Schoell, 2007; Coudrain-Ribstein et al., 1998) yields

$$\log P_{CO2} = \log \eta_{CO2(g)} = \frac{\log K_R}{\alpha_0}$$
(3)

176 P_{CO2} being the partial pressure of CO_2 either in the gaseous or supercritical phase. Note that, according 177 to our assumption, the numerical values of CO_2 fugacity and partial pressure coincide (Anderson, 178 2009). We provide additional discussion about the assumption of $\eta_{CO2} = P_{CO2}$ in Electronic Annex II. 179 The value of log K_R (and therefore P_{CO2}) is a function of the local conditions of pressure and 180 temperature, as discussed in Sections 2.2.

181 The CCR process can be summarized by the phenomenological scheme illustrated in Fig. 1182 and described in the following.

1831. Given a sedimentary rock containing carbonates and clays/alumino-silicates, the amount of184dissolved CO_2 in the pore water is regulated by the chemical equilibrium among all phases185(Fig. 1a). Even as a separate gas phase is not formed, the concept of partial pressure associated186with gaseous species can be still preserved if referred to a fictive gas phase hypothetically at187equilibrium with the pore water (Coudrain-Ribstein et al., 1998).

Pressure and temperature typically increase throughout the burial process. Under these
 conditions, the sum of the partial pressures associated with gaseous species (*CO*₂ and possibly

190 other species including, e.g., $H_2O_{(g)}$, $CH_{4(g)}$) might exceed the fluid environmental pressure. 191 When this happens, a separate gas phase is generated (Fig.1b). In this work we consider only 192 CO_2 and H_2O as possible gaseous species.

When CO₂ (possibly mixed with other gases) is released as a gas phase, the difference
between gas and fluid phase densities promotes upward migration of CO₂. As a consequence
of this migration, the equilibrium reaction (1) is shifted towards its right side (Fig. 1c) and the
reactions listed in Table 1 can be considered as a quantitative transformation of the reactants
(primary phases) into the products (secondary phases), as seen, e.g., in Cathles and Schoell
(2007).

Note that supercritical CO_2 is likely to be expected at the pressure and temperature conditions characterizing sedimentary formations. We assume that the conceptual model proposed by Cathles and Schoell (2007) still holds when CO_2 is in supercritical conditions. Supercritical CO_2 is always characterized by lower density when compared to water and buoyancy effects always force the CO_2 rich separate gas phase to migrate upwards as soon as it is generated (Battistelli et al., 2016; Span and Wagner, 1996; Johnson et al., 1992). For simplicity, we refer in the following to the separate CO_2 rich phase as gaseous CO_2 .

When the conditions for the generation of a separate gas phase are not attained, CCR leads only to the formation of aqueous CO_2 which remains dissolved in the pore-fluid. Dissolved CO_2 can constitute a significant fraction of the overall CO_2 amount released by CCR and its occurrence can be a relevant aspect to consider when the characterization of flow processes in sedimentary formations is of concern (Coudrain-Ribstein and Gouze, 1993; Chiodini et al., 2000; Farmer, 1965).

211 2 Modeling workflow

We illustrate here a procedure to compute the time, depths and temperature at which the process described in Fig. 1, i.e. the activation of the gaseous CO_2 source, takes place. The two main constituents of the numerical modeling procedure we propose are: a basin compaction model, providing the temporal dynamics of porosity, temperature,
 pressure and basin stratigraphy along the vertical direction in the presence of mechanical
 compaction;

2. a geochemical model which allows computing the partial pressure of CO_2 (P_{CO2} [Pa]) and the 2. concentration of dissolved CO_2 (C_{CO2} [mol/L]) as a function of temperature and pressure.

220 Our modeling strategy focuses on the uncertainty associated with the identification of CO_2 221 sources and with the quantification of the resulting CO_2 fluxes. Characterization of migration of CO_2 222 after its generation is beyond the scope of our study. Fig. 2 illustrates the key steps of the workflow, 223 which is subdivided in three blocks: *i*) implementation of the burial model (Block 1), described in 224 Section 2.1; *ii*) computation of the CO_2 pore-water concentration and CO_2 partial pressure (Block 2), 225 illustrated in Section 2.2; and *iii*) estimation of CO₂ generation rate and source location (Block 3), 226 detailed in Section 2.3. All details on the computational steps of the model are reported in the 227 Electronic Annex I where a step-by-step illustration of the procedure is included to assist reproducibility of the model implementation. 228

229 **2.1 Basin Model**

230 The quantification of the amount of CO_2 generated in sedimentary systems requires the quantification 231 of (i) porosity (ϕ), temperature (T), pressure (P) distributions and burial velocity of sediments (V_{SED} , 232 i.e., rate at which the sediments are displaced along the vertical direction) as a function of depth and 233 time; and of (*ii*) the temporal evolution of the stratigraphy. In this study, we obtain these quantities 234 through the one-dimensional compaction model proposed and tested by Formaggia et al. (2013), Porta 235 et al. (2014), and Colombo et al. (2016). Further details related to the burial model implemented in 236 this work can be found in Electronic Annex I. We highlight here that any type of compaction/diagenesis model (e.g., a three-dimensional model) is compatible with the proposed 237 238 procedure, provided it renders a characterization of the dynamics of temperature, pressure, porosity 239 and sediment burial velocity in the system. In this study, we consider the outputs of the basin model

(e.g., temperature and pressure distributions) as deterministic quantities, consistent with our focus on
the quantification of the parametric uncertainty related to the geochemical model. Possible sources
of uncertainty affecting the burial model are explicitly discussed in Section 3.

243 **2.1.1** Basin scale case study and compaction setting

We illustrate the applicability of our methodological framework for the quantification of the uncertainty associated with estimates of CO_2 generation at basin scale by focusing on an exemplary system inspired to a realistic compaction setting.

247 We consider a basin deposition over a period of 135 Ma (Millions of years before present), 248 from time t = 135 Ma, to present day (i.e., t = 0 Ma). According to our simplified compaction model, we assume the basin to be described as a one-dimensional system along the vertical direction. The 249 250 paleo-bathimetry is constant and set equal to an elevation of 106 m (the Z-axis is considered to point 251 downwards and the sea level to correspond to Z = 0). Carbonate sediments are deposited within the interval ranging from time t = 135 Ma and t = 23 Ma, leading to the formation of carbonate rock 252 253 layers. Shale and sandy shale sediments are deposited within the period ranging from t = 23 Ma and 254 t = 0 Ma, leading to mudrock after compaction. Sediment deposition rate (V_D) at the basin top is imposed as boundary condition. We assume that it varies in time and can be described by a piecewise 255 256 constant function of time across six time intervals as indicated in Table 2. A given temperature gradient of 32 °C /km is prescribed at the basement. Each sediment type is characterized by a given 257 258 thermal conductivity of the solid matrix (K_T) , initial porosity (i.e., porosity at sediment deposition time, ϕ_0), and vertical compressibility coefficients (β). We set the parameters K_T , ϕ_0 and β to the 259 260 values listed in Table 3.

We analyze the two possible scenarios of mineral composition associated with carbonate rock
listed in Table 4. We highlight that: (*i*) Scenario S_{dol} considers dolomite as the only carbonate mineral

present in the rock; (ii) Scenario Scal is characterized by the presence of magnesiac limestone where 263 calcite is the prevailing carbonate (73% in weight) and the fraction of dolomite is lower than 10. 264

265

Geochemical modeling under uncertainty 2.2

The main physical quantities which allow quantifying the CO_2 generated by CCR are the 266 partial pressure of the gas phases and the concentration of the CO_2 dissolved in the fluid phase. We 267 268 obtain these outputs starting from (1)-(3) and relying on the assumption that the activity of the pure 269 solid mineral and liquid phases are set equal to unity (Giggenbach, 1980; Giggenbach, 1984).

270 The dependence of P_{CO2} on temperature in (3) is assessed by relying on the thermochemical 271 parameters collected in a thermodynamic database. Among the databases available in the literature 272 (e.g., LLNL, Delany and Lundeen, 1990; Vminteq, Peterson, 1987; SOLMINEQ, Kharaka et al., 273 1988), we select the Thermoddem database (Blanc et al., 2012) due to its completeness, traceability 274 of data, and proven internal thermodynamic consistence, especially for the aluminum silicate phases 275 (Blanc et al., 2015).

276 In the remainder of the work, an uncertain (i.e., random) quantity ζ is identified with the notation $\tilde{\zeta}$. Our operational procedure relies on the following steps: 277

278 1. A set of basis species is selected coherently to the chosen thermodynamic database (e.g.,

279
$$H_2O_{(l)}, H^+_{(aq)}, HCO_{3(aq)}, Ca^{2+}_{(aq)}, Mg^{2+}_{(aq)}, Al^{3+}_{(aq)}, K^+_{(aq)}, H_4SiO_{4(aq)}).$$

2. A set of stoichiometric coefficients are defined to honor mass and charge balances, i.e., 280

281
$$\begin{cases} \gamma_k = \alpha_k & \text{if } 1 \le k \le m \\ \gamma_k = -\alpha_k & \text{if } m+1 \le k \le m+n \end{cases}$$
(4)

282 3. A speciation reaction S_k is defined for each M_k phase (k=1, ..., n+m) involved in reaction (1) 283 to describe speciation in the formation fluid (which we consider as water) of M_k through the basis species selected in step 1. Uncertain chemical equilibrium constants \tilde{K}_{sk} are quantified 284 285 to characterize the speciation reaction S_k of phase M_k at temperature T. We do so by employing the following equation derived from the Maier-Kelley heat capacity definition (Maier and
Kelley, 1932; Parkhurst and Appelo, 2013; van Berk et al., 2009)

$$\log \tilde{K}_{Sk} = \tilde{A}_k + \tilde{B}_k T + \frac{\tilde{C}_k}{T} + \tilde{D}_k \log T + \frac{\tilde{E}_k}{T^2}$$
(5)

where $(\tilde{A}_k, \tilde{B}_k, \tilde{C}_k, \tilde{D}_k, \tilde{E}_k)$ is a vector of uncertain quantities which are treated as independent random variables/parameters with an assigned probability density function (*pdf*). The characterization of the uncertainty of these parameters is discussed in Section 2.2.1. Note that the (5) allows evaluating the equilibrium constant \tilde{K}_{sk} as a function of temperature while keeping the pressure of the system at a constant reference value of 1 bar.

4. The equilibrium constant $\log \tilde{K}_{R,T,1}$ associated with reaction (1) is computed as (Coudrain-Ribstein et al., 1998)

296
$$\log \tilde{K}_{R,T,1} = \sum_{k=1}^{n} \gamma_k \log \tilde{K}_{Sk} - \alpha_0 \log \tilde{K}_{S,CO2(g)}$$
(6)

where $\tilde{K}_{S,CO2(g)}$ is the equilibrium constant associated with the reaction defining the CO_2 (either gaseous or supercritic) in terms of its basis species. Note that $\tilde{K}_{S,CO2(g)}$ is considered to be characterized by a relationship having the same format as (5) and is also considered as random. Subscript 1 appearing in $\log \tilde{K}_{R,T,1}$ indicates that the value of the equilibrium constant evaluated through (6) is associated with the reference pressure of 1 bar.

302 5. We compute $\tilde{K}_{R,T,P}$ as a modification of $\tilde{K}_{R,T,1}$ to account for the high pressure at which the 303 CCR process occurs according to the procedure proposed by Millero (1982). Further details 304 can be found in Electronic Annex I.

305 6. CO_2 partial pressure \tilde{P}_{CO2} is evaluated upon replacing $\log K_R$ with $\log \tilde{K}_{R,T,P}$ in (3).

306 The activity $\tilde{a}_{CO2(aq)}$ of carbon dioxide dissolved in the liquid phase can be estimate by considering

307 the equilibrium as an effective model

$$CO_{2(g)} = CO_{2(aq)}$$
 (7).

308

309

Assuming a unit coefficient activity associated with $[CO_{2(aq)}]$ (see Electronic Annex II for additional details about this assumption), we can directly derive the molar concentration of aqueous CO_2 ($\tilde{C}_{CO2(aq)}$ [mol/1]) from the CO_2 activity. A description of the detailed steps leading to the quantification of aqueous CO_2 through our computational procedure are included in Electronic Annex I.

Values of the uncertain quantities $\tilde{C}_{CO2(aq)}$ and \tilde{P}_{CO2} may be constrained by the effect of 314 limiting reactants, as a consequence of relative abundancy of diverse primary phases. Given a mineral 315 316 composition, the generation of CO_2 takes place according to the equilibrium relationship (3) until one of the involved primary mineral phases vanishes. We then verify that the computed $\tilde{C}_{CO2(aq)}$ is 317 318 compatible with the maximum CO_2 concentration (C_{max}) associated with the complete depletion of 319 the limiting reactant across all primary phases. We set $C_{CO2(aq)} = C_{max}$ at locations where $C_{CO2(aq)}$ is 320 larger than C_{max} , and accordingly correct the associated value of P_{CO2} . In the following we denote as $\tilde{C}_{CO2(aq)}|C_{max}$ and $\tilde{P}_{CO2}|C_{max}$ the values of dissolved CO_2 and CO_2 partial pressure conditional to the 321 effect of limiting reactant. Additional computational details related to C_{max} , $\tilde{C}_{CO2(aq)}|C_{max}$, and 322 $\tilde{P}_{_{CO2}}|C_{_{max}}$ are included in Electronic Annex I. 323

324 **2.2.1** Characterization of uncertain model inputs

325 Here we illustrate the stochastic characterization of the chemical equilibrium constants. We 326 assess the consistency of the results stemming from the proposed procedure with available data of 327 CO_2 partial pressure in Sections 4.2 and 5.1.

328 For the purpose of illustration of our uncertainty quantification procedure, hereinafter we329 focus on the following CCR (see Table 1)

330
$$5CaMg(CO_3)_2 + Al_2Si_2O_5(OH)_2 + SiO_2 + 2H_2O = 5CO_2 + 5CaCO_3 + Mg_5Al_2Si_3O_{10}(OH)_8 (8)$$

dolomite kaolinite silica clicite clinochlore

We select this equilibrium reaction among those listed in Table 1 because all of the involved mineral phases are commonly found in sedimentary system (e.g., Hutcheon, 1990; Coudrain-Ribstein and Gouze, 1993) and it is in agreement with the mineralogical assemblage alteration observed in the Kootenay Formation studied by Hutcheon et al. (1980). We follow the procedure outlined in Section 2.2 and start by selecting the following basis species

336
$$H_2O_{(l)}; H^+_{(aq)}; HCO^-_{3(aq)}; Ca^{2+}_{(aq)}; Mg^{2+}_{(aq)}; Al^{3+}_{(aq)}; K^+_{(aq)}; H_4SiO_{4(aq)}$$
(9)

337 We then write the following system governing speciation of all liquid and solid phases involved and 338 of the gaseous CO_2

$$\begin{cases} CaMg(CO_{3})_{2}+2H^{+}=2HCO_{3}^{-}+Ca^{2+}+Mg^{2+} \\ Al_{2}Si_{2}O_{5}(OH)_{4(s)}+6H^{+}_{(s)}=2Al^{3+}_{(s)}+2H_{4}SiO_{4(aq)}+H_{2}O_{(l)} \\ SiO_{2(s)}+2H_{2}O_{(l)}=H_{4}SiO_{4(aq)} \\ CaCO_{3(s)}+H^{+}_{(aq)}=HCO_{3}^{-}_{(aq)}+Ca^{2+}_{(aq)} \\ Mg_{5}Al(AlSi_{3})O_{10}(OH)_{8(s)}+16H^{+}_{(aq)}=2Al^{3+}_{(aq)}+5Mg^{2+}_{(aq)}+3H_{4}SiO_{4(aq)}+6H_{2}O_{(l)} \\ H_{2}O_{(l)}=H_{2}O_{(l)} \\ CO_{2(g)}+H_{2}O_{(l)}=H^{+}_{(aq)}+HCO_{3}^{-}_{(aq)} \end{cases}$$
(10)

339

We characterize the equilibrium constants associated with (10) by relying on Thermoddem as a reference database (Blanc et al., 2012). We employ (5) to account for the influence of temperature, where parameters $(\tilde{A}_k, \tilde{B}_k, \tilde{C}_k, \tilde{D}_k, \tilde{E}_k)$ are assumed to be uncertain.

To streamline the uncertainty quantification procedure, we perform a preliminary sensitivity analysis by means of a numerical Monte Carlo procedure. This enables us to single out the contributions of the five parameters appearing in (5) to the variability of \tilde{K}_{sk} characterizing the reactions S_k presented in (10). In this framework, the five parameters $(\tilde{A}_k, \tilde{B}_k, \tilde{C}_k, \tilde{D}_k, \tilde{E}_k)$ associated with the quantification of each log \tilde{K}_{sk} are randomly sampled from uniform distributions centered on the reference value reported in Thermoddem and of width equal to ± 20 % of such reference value. We compute the first order sensitivity index (SI_h , h = A, B, C, D, E) for each parameter, according to

the variance-based method described, e.g., in Sobol (2001), Saltelli et al. (2008), and Razavi and 350 Gupta (2015). These sensitivity indices represent the relative contribution of each uncertain parameter 351 in (5) to the variance of $\log \tilde{K}_{sk}$. These metrics are widely employed in diverse fields (e.g., Saltelli et 352 353 al., 2008; Formaggia et al., 2013; Porta et al., 2014; Riva et al., 2015) to perform global sensitivity 354 analysis of the output of a model as driven by a set of uncertain (random) model parameters. The results of this analysis reveal that the variability of $\log \tilde{K}_{sk}$ computed through (8) is mainly due to the 355 random variability of parameters \tilde{A}_k and \tilde{D}_k (details not shown). These parameters are always 356 associated with first order sensitivity indices (SI_A and SI_D) larger than 0.43 for 0 °C \leq T \leq 400 °C, SI_B, 357 SI_C and SI_E always being lower than 0.02. On these bases, we set parameters B, C and E to the values 358 listed in Thermoddem and consider \tilde{A}_k and \tilde{D}_k as random input parameters in (5). 359

We rely on laboratory scale mineral solubility experiments to characterize the uncertainty 360 associated with \tilde{A}_k and \tilde{D}_k . As observed by Blanc et al. (2013), only a few experimental studies 361 reporting values of \tilde{K}_{sk} as a function of temperature are available, particularly with reference to clay 362 minerals. The estimation of the coefficients embedded in Thermoddem is mostly based on 363 thermodynamic calculations, which are associated with an uncertainty level which is difficult to 364 quantify. Here, we employ the mineral solubility data for calcite and kaolinite reported by Plummer 365 and Busenberg (1982) and Blanc et al. (2013). Plummer and Busenberg (1982) provide observations 366 of the calcite speciation constant for a series of temperatures ranging from 0.1 °C to 89.7 °C. Blanc 367 368 et al. (2013) collect a set of solubility experimental data related to kaolinite previously presented by various authors and associated with temperature values ranging between 25 °C and 300 °C. 369

- 370 We employ the following procedure to quantify uncertainties associated with \tilde{A}_k and \tilde{D}_k . 371 (with *i* = kaolinite, calcite) using the solubility experimental data indicated above:
- 372 1. We calibrate model (5) against available experimental observations K_{Sk}^* upon estimating the
- 373 parameters \tilde{A}_k and \tilde{D}_k (with k = calcite, kaolinite) through a standard least square criterion.

As indicated above, parameters B_k , C_k , E_k are set to the corresponding values reported in Thermoddem. This procedure yields best estimates (\hat{A}_k, \hat{D}_k) of parameters $(\tilde{A}_k, \tilde{D}_k)$ and the related uncertainty expressed in terms of a 2×2 symmetric covariance matrix $\hat{\Psi}_k$. The results of these calculations are listed in the first two rows of Table 5.

- 378 2. We consider that the entries of the uncertain parameter vectors $\tilde{\boldsymbol{u}}_k = (\tilde{A}_k, \tilde{D}_k)$ can be described
- 379 through a bivariate Gaussian distribution with mean $\mu(\tilde{\boldsymbol{u}}) = (\hat{A}_k, \hat{D}_k)$ and covariance matrix

380
$$\boldsymbol{\Psi}_{k} = \boldsymbol{\Psi}_{k}$$
 (with $k = \text{calcite, kaolinite}$).

No direct references are reported in the Thermoddem database to characterize the uncertainty associated with the equilibrium constants related to the remaining phases included in (8) (dolomite, clinochlore, quartz, $CO_{2(g)}$, $CO_{2(aq)}$). In our illustrative example we resort to the following set of assumptions to characterize uncertainties associated with \tilde{A}_k and \tilde{D}_k (with k = dolomite, clinochlore, quartz, $CO_{2(g)}$, $CO_{2(aq)}$):

- 1. The vector of parameters $\tilde{\boldsymbol{u}}_k = (\tilde{A}_k, \tilde{D}_k)$ (with k = dolomite, clinochlore, quartz, $CO_{2(g)}$, $CO_{2(aq)}$) is associated with a bivariate Gaussian distribution. Here, we assume that the entries of the vector of mean values $\mu(\tilde{\boldsymbol{u}}_k)$ coincide with the values included in Thermoddem for each phase k (see Table 5).
- 390 2. Affine minerals are characterized by the same parametric uncertainty, i.e., we set 391 $\Psi_{dolomite} = \hat{\Psi}_{calcite}$ (as dolomite and calcite are both carbonates minerals) and 392 $\Psi_{clinochlore} = \hat{\Psi}_{kaolinite}$ (as clinochlore and kaolinite are both clay minerals, Bergaya and 393 Lagaly, 2013).
- 394
 3. The parameters describing the solubility of quartz and the water transition phase
 approximate equilibrium are characterized by negligible uncertainty when compared against the
 approximate uncertainty level of the equilibrium constants discussed above.

397 4. We set $\Psi_{CO2(g)} = \Psi_{CO2(aq)} = \hat{\Psi}_{kaolinite}$, as $\hat{\Psi}_{kaolinite}$ renders the highest level of uncertainty 398 following estimation of the coefficients of (5) through the experimental data employed 399 (i.e., solubility data of calcite and kaolinite).

Mean values of the parameter distributions are listed in Table 5 together with the associated covariance matrix entries and the set of assumptions illustrated above. Note that these assumptions are not strictly required for the applicability of the proposed methodology and are here considered solely for illustrative purposes. In this sense, measurements on mineral solubility or equilibrium constants can readily be integrated in the proposed workflow when available.

405 **2.3** Quantitative assessment of *CO*₂ generation and CCR mechanism activation at

406 **basin scale**

407 The basin compaction and geochemical models illustrated in Sections 2.1 and 2.2 allow 408 assessing the desired dynamics of the CCR mechanism and quantifying the amount of CO_2 generated 409 (as gaseous or dissolved species) during the diagenetic process.

410 Generation of a separate gas phase at a location *Z* and time *t* takes place when

411
$$\tilde{R}(Z,t) = \frac{P_{gas}(Z,t)}{P(Z,t)} = \frac{\tilde{P}_{CO2}(Z,t) + P_{H2O}(Z,t)}{P(Z,t)} \ge 1$$
(11)

412 where P(Z,t) and $\tilde{P}_{gas}(Z,t)$ respectively are the fluid pressure and the partial pressure of the gas 413 phase. Note that P(Z,t) is rendered by the basin compaction model of choice (see Section 2.1) while 414 \tilde{P}_{CO2} is evaluated through the procedure illustrated in Section 2.2. The computation of partial pressure 415 of water vapor, $P_{H2O}(Z,t)$, is detailed in Electronic Annex I. According to criterion (11), the space-416 time locations at which the generation of gaseous CO_2 may take place can be identified through the 417 local values of the ratio \tilde{R} . For a given time level *t*, the activation of the mechanism is assigned to 418 the location

419
$$\tilde{Z}_{act}(t) = \begin{cases} \varnothing & \text{if } \tilde{R} < 1 \text{ for all } Z \in \Omega_Z \\ \min\left\{ Z \in \Omega_Z \left| \tilde{R}(Z, t) \ge 1, m_{CO2} > 0 \right\} & \text{if } \exists Z \in \Omega_Z \left| \tilde{R}(Z, t) \ge 1 \right. \end{cases}$$
(12)

i.e., the location of the CO_2 source at time t, $\tilde{Z}_{act}(t)$, is assumed to correspond to the shallowest depth 420 at which $\tilde{R} \ge 1$, given that the mineral composition is compatible with CCR. Note that $\tilde{Z}_{act}(t)$ is a 421 422 function of time because of the temporal variability of vertical profiles of temperature and pressure. 423 Definition (12) is consistent with the assumption that CO_2 migrates instantaneously upwards when a 424 gas phase is formed (see Fig. 1). Under such conditions, the primary phases of the equilibrium 425 reaction (1) are progressively consumed because one of the secondary phases (CO_2) is continuously 426 driven away. This behavior is observed until the limiting reactant in (1) vanishes. We assume that the 427 complete consumption of at least one primary phase takes place on a time scale that is considerably 428 smaller than the one associated with the basin evolution. Therefore, the burial velocity of the sediment, $V_{SED}(\tilde{Z}_{act}, t)$, is a limiting factor for the generation of CO_2 as a gas phase through a CCR 429 430 mechanism. Under this assumption, we can then evaluate the rate of CO_2 generation as

431
$$\tilde{F}_{CO2}(t) = m_{CO2} \cdot V_{SED}\left(\tilde{Z}_{act}, t\right) \cdot \left[1 - \phi\left(\tilde{Z}_{act}, t\right)\right] \cdot L \cdot \rho$$
(13)

Here, \tilde{F}_{co2} [kg/Ma] is the CO_2 mass generation rate; L [m²] is the planar cross sectional area of the basin/reservoir; and m_{co2} [-] is maximum amount of mass of CO_2 released by unity mass of sediment (see Electronic Annex I for further detail about the computation of m_{CO2}). Note that, following (8), the limiting reactant is dolomite in the two mineralogical scenarios investigated in this work (Table 3). When the gas generation mechanism is activated, the reaction evolves over time until at least one primary phase is exhausted (see Section 1). We note that $\tilde{F}_{co2} \equiv 0$ when the mechanism is not activated (i.e., when $\tilde{Z}_{act}(t) = \emptyset$).

According to the conceptual model described above, at least one of the mineral phases involved in the CCR mechanism is expected to be exhausted at locations below the activation depth (i.e, for $Z > \tilde{Z}_{act}$) and the mineral phases equilibrium (1) leading to dissolved CO_2 is no longer

possible. We therefore assume that the dissolved amount of CO_2 is zero at all locations $Z > \tilde{Z}_{act}$ (see 442 443 Electronic Annex I for additional details about the computational procedure).

444 3 Analysis of sources of uncertainty

445 Any model which aims at quantifying CO_2 generation in sedimentary basin is subject to 446 considerable uncertainties. These are due to our incomplete knowledge of the processes involved and 447 of the initial/boundary conditions together with the lack of information resulting from the large space-448 time scales, which are characteristic of the evolution of sedimentary systems. Upon following 449 Neuman (2003), we distinguish in the following sections between modeling and parametric 450 uncertainties. This work is keyed to the development and implementation of a methodology for the 451 quantification of the uncertainty stemming from our incomplete knowledge of equilibrium reaction 452 constants. In this section we frame this choice within the context of uncertainty quantification and 453 discuss a variety of possible sources of uncertainty which may be relevant to our setting.

454

3.1 Model uncertainties

455 Investigation of complex settings in earth and environmental sciences typically relies on the formulation of a conceptual-mathematical model which is consistent with available information on 456 457 the system investigated. Multiple and competing conceptual models can be formulated, according to 458 diverse interpretations of the processes underlying the target scenario.

- 459 We list here key model uncertainties and the related assumptions associated with our setting.
- 460 While we focus on the occurrence of reaction (8), other geochemical processes may • 461 take place simultaneously during basin compaction. Different competing models could 462 therefore be formulated according to which reaction (8) occurs jointly with a set of diagenetic processes (e.g., dolomitization, albitization, illitization, cracking of 463 464 biological matter and many others). All these processes can jointly contribute to CO_2

465 partial pressure and to increase/decrease or to the amount of CO_2 which can be found 466 in the system. The selection of the geochemical processes which should be considered 467 and the formulation of a related model is not a trivial task and constitutes a remarkable 468 source of model uncertainty.

- 469 The selection of the primary phases considered in the mineralogical assemblage is a 470 key input to our methodology. This information is typically uncertain and various 471 admissible hypotheses may be formulated, consistent with geological and 472 sedimentological conceptual models and interpretations. Companion considerations hold on the assumed initial interstitial fluids composition (i.e., gas phase and brine). 473 474 Our approach rests on the assumption that i) gaseous phase are CO_2 and H_2O_2 , and ii) 475 the initial pore-brine is pure water and the primary phases appearing in (8) are all available in the mineral composition. This is a simplification of the conditions 476 477 encountered in real cases, but does not disable the proposed methodology.
- The spatial arrangement of the mineral composition may be affected by heterogeneity
 at all scales. In our conceptual model we assume a uniform spatial distribution of
 primary mineral phases throughout the carbonate-rich sedimentary layers. The
 spatial/temporal distribution of minerals could alternatively be described as a
 stochastic process, whose main features should possibly be characterized through real
 mineralogical samples of a specific sedimentary basin case study.

484 Quantification of the modeling uncertainties listed above may be performed through dedicated 485 techniques (see, e.g., Neuman, 2003). While this task lies beyond the scope of the present work, we 486 remark that these types of uncertainties should be carefully considered prior to applying the procedure 487 outlined in Section 2 to the interpretation of observations from a real field site.

488 **3.2** Parametric uncertainties

489 An admissible conceptual/mathematical model of a process commonly includes a number of 490 parameters. These are in turn associated with a given level of uncertainty due to lack of information. 491 This incomplete knowledge about parameter values can be quantified through, e.g. statistical 492 characterization of available experimental data via parameter estimation techniques. In this work, we 493 a) present a rigorous methodology to account for parametric uncertainty associated with mineral 494 solubility equilibrium constants and b) propagate such uncertainty throughout our geochemical model 495 of choice, which is aimed at representing CCR. While the need to account for these parameters is 496 ubiquitous in geochemical models of environmental systems, a rigorous quantification of their 497 uncertainty and its ensuing effects is often neglected. To sharply delineate the effect of this specific 498 source of parametric uncertainty, we do not consider here other sources of parametric uncertainties 499 such as: a) properties of the sedimentary rocks, i.e., density, permeability, thermal diffusivity and 500 mechanical compressibility; b) boundary conditions of the compaction problem, i.e., heat flux at basin 501 basement, and temporal dynamics of sea level evolution and sediment deposition rate; and c) other 502 parameters of the geochemical model, including relative abundance of each mineral phase in the 503 primary assemblage (for the given the qualitative composition of the mineralogy, which falls into the 504 category of modeling uncertainties, as discussed in Section 3.1), molar volume change during the 505 reaction (8), and activity and fugacity coefficients.

506 The influence of each set of parameters may be assessed through local and/or global sensitivity 507 analysis techniques (e.g., Razavi and Gupta, 2015 and references therein), which we envision to 508 explore in future works.

509 4 Results

510 This Section is devoted to a synthetic illustration of the results stemming from the 511 implementation of the methodology proposed in Section 2.

512 **4.1 Basin evolution**

The results depicted in Fig. 3 are obtained through the numerical solution of the basin compaction model illustrated in Section 2.1. Fig. 3a depicts the space-time evolution of the vertical stratigraphic sequence of the basin (i.e., the system geo-history). The total basin thickness at present day is also shown. Fig. 3b-d respectively depict the space-time evolution of porosity, temperature and pressure with reference to the stratigraphy displayed in Fig. 3a. The black vertical lines identify the times when the sediment deposition rate (V_D) changes its value according to stepwise function described in Table 2 imposed at basin top.

520 4.2 Consistency of geochemical modeling results with field data

Here, we compare the results stemming from the application of our geochemical modeling approach against a set of field observations of CO_2 partial pressures reported by Coudrain-Ribstein et al. (1998). This comparison aims at assessing the robustness of our procedure and of the assumptions underlying the uncertainty quantification steps proposed in Section 2.2.1. We focus on the variation of $\log \tilde{K}_{R,T,P}$ and $\log \tilde{P}_{CO2}$ as a function of temperature and pressure.

To this end, we perform a Monte Carlo sampling of the parameter space to obtain Nrealizations (here, we consider $N = 10^5$) of $\log \tilde{K}_{R,T,P}$ (6) as function of temperature and pressure. In the context of our comparison between field data and geochemical model outputs, we assume the following relationship between temperature and pressure (Smith and Ehrenberg, 1989; Cathles and Schoell, 2007)

531

$$P[bar] = 6(T[K] - 298) \tag{14}$$

Fig. 4a depicts the dependence on temperature of the mean, median, and 1st- and 99th-percentiles of the sample distribution of $\log \tilde{K}_{R,T,P}$. Here and in the following we denote a percentile (or quantile) of the distribution of a random variable $\tilde{\zeta}$ as $p_W(\tilde{\zeta})$. The latter is defined as the value below which 535 a percentage equal to W of observations of $\tilde{\zeta}$ falls. Note that the mean and the median coincide in 536 Fig. 4a, $\log \tilde{K}_{R,T,P}$ being characterized by a symmetric sample distribution.

The Monte Carlo sample of $\log \tilde{P}_{CO2}$ values can be obtained from $\log \tilde{K}_{R,T,P}$ through (3). Fig. 4b depicts percentiles $p_1(\log \tilde{P}_{CO2})$, $p_{50}(\log \tilde{P}_{CO2})$, and $p_{99}(\log \tilde{P}_{CO2})$ as a function of temperature. These Monte Carlo - based results are juxtaposed in Fig. 4b to a set of available measurements of P_{CO2} reported by Coudrain-Ribstein et al. (1998) for sedimentary formations. The consistency of the results provided by our geochemical model and the field data depicted in Fig. 4b is discussed in Section 5.1.

543 **4.3** Quantitative assessment of CO₂ generation and CCR mechanism activation

We present here results associated with the way parametric uncertainty propagates to the outputs of the model described in Section 2.3, i.e., to the rate of generation of gaseous CO_2 and to the total dissolved CO_2 . Characterization of parameter uncertainty relies on the procedure described in Section 2.2.1. The results are related to the mineral compositions S_{dol} and S_{cal} (see Table 4) and are discussed in Section 5.2. All of the results presented are obtained upon relying on a sample of N = 10^5 Monte Carlo realizations.

Fig. 5 shows the vertical profiles of the percentiles of the partial pressure of CO_2 , $p_W \left(\log \tilde{P}_{CO2} | C_{max} \right)$, and of the ratio \tilde{R} as defined in (11), $p_W \left(\tilde{R} \right) (W = 1, 25, 50, 75, 99)$, at two selected time levels (t = 48, 0 Ma) and for scenarios S_{dol} and S_{cal}. To complement this result, Fig. 6 provides a comparison of the sample *cdf* (cumulative distribution function) of \tilde{R} (11) at Z = 8 km for scenarios S_{dol} and S_{cal}. Note that at t = 0 the top layer (0 < Z < 1.4 km) of the basin is formed by mudrocks (see Fig. 3a). Therefore, we set $P_{CO2} = 0$ at these locations (see Fig. 5b and d), as we assume CCR happens exclusively in carbonates layers.

557 The probability of activation $G_A(t)$ can then be as the sample probability (relative frequency) 558 of observing at least one point in the domain for which $\tilde{R} \ge 1$, i.e., the generation of CO_2 as a separate

559 phase through CCR is activated at time t. The procedure to compute GA is exemplified in Fig. 6, where the value R = 1 is identified by a vertical red line, which represents the conditions at which the 560 CO_2 generation as a separate gas phase is activated (see Section 2.3). The *cdfs* associated with the 561 two diverse mineral compositions intercept the threshold line corresponding to R = 1 (i.e., the 562 563 conditions at which the CO_2 generation as a separate gas phase is activated) at different points, i.e., 564 for R = 1 the *cdf* attains a value equal to 0.55 and 0.85, respectively for S_{dol} and S_{cal}, indicating a different probability of activation in the two scenarios. Fig. 7 depicts the temporal evolution of $G_A(t)$ 565 566 for S_{dol} and S_{cal} across the overall basin history.

Our procedure allows identifying not only the probability of activation at given time but also to estimate the location of the CO_2 sources through (12). Fig. 8 depicts the sample probability (relative frequency) f_{Zacbt} that the activation of gaseous CO_2 generation takes place at location \tilde{Z}_{act} at time *t*. In particular Fig. 8a displays f_{Zacbt} , i.e., f_{Zacbt} , for t = 10 Ma, where the domain is comprised between the sea bottom (at 106 m) and 7.6 Km. We note that this relative frequency is computed upon considering the complete set of Monte Carlo realizations, including those for which $\tilde{Z}_{act} = \emptyset$ according to (12). Thus, the function f_{Zacbt} integrates to the corresponding value of G_A at time *t* i.e.

574

$$\int_{\mathbf{\Omega}_{Z}(t)} f_{Zact,t} dZ = G_{A}(t)$$
(15)

For example, the integral (15) evaluated at t = 10 Ma for scenario S_{dol} is equal to 0.30, which corresponds to the value of $G_A(t=10 \text{ Ma})$ for the corresponding scenario reported in Fig. 7. Fig. 8b-c depict the temporal dynamics of the relative frequency $f_{Zact,t}$ for scenarios S_{dol} and S_{cal}. As anticipated by the temporal variation of $G_A(t)$ in Fig. 7 the nonzero values are obtained for t < 50 Ma in both scenarios. The generation of gaseous CO_2 takes place at $\tilde{Z}_{act} > 4.8$ km in the considered example for both mineral composition scenarios. 581 Figure 9-10 provide the probabilistic quantification of the generated CO₂ in terms of (i) flux of gaseous CO_2 generated as a result of the CCR process. \tilde{F}_{CO_2} , as defined in (13), and (ii) 582 concentration of dissolved $CO_2 \ \tilde{C}_{CO2(aq)} \left(C_{max}, \tilde{Z}_{act} \right)$. Fig. 9a depicts the relative frequency $f_{F,t}$ 583 associated with $\log \tilde{F}_{co2}$ at time t for the overall basin history of scenario S_{dol}. Corresponding results 584 for S_{cal} are depicted in Fig. 9b. We set here $L = 1 \text{ m}^2$ in (13) so that the reported values of \tilde{F}_{CO2} are 585 586 per unit (planar) area of the sedimentary basin. For completeness, Fig. 9a-b include the information (black solid curve) corresponding to the frequency of activation $G_A(t)$. Note that $G_A(t) \equiv 0$ for 587 $t \in]$ 45 Ma, 135 Ma], thus implying that $\tilde{F}_{CO2} = 0$ across all Monte Carlo realizations for these 588 simulation times. Indeed, $f_{F,t} = 0$ for all non-zero values of \tilde{F}_{CO2} for $t \in]$ 45 Ma, 135 Ma]. Fig. 9c 589 depicts the sample *cdf*s of \tilde{F}_{CO2} associated with the two time levels identified by the red dashed 590 vertical lines in Fig. 9a, i.e., t = 20, and 0 Ma. 591

Figure 10 reports the distribution along the basin depth of the relative frequency associated with the log-concentration $\log \tilde{C}_{CO2(aq)} | (C_{max}, \tilde{Z}_{act})$ (denoted as $f_{C,Z}$ in Fig. 10a-b). Introducing here $f_{C,Z}(0)$ to denote the relative frequency associated with $\tilde{C}_{CO2(aq)} | (C_{max}, \tilde{Z}_{act}) = 0$, Fig. 10c, d) display the variation of $f_{C,Z}(0)$ with Z for the two mineral compositions S_{dol} and S_{cal}, respectively.

596 **5 Discussion**

597 This Section is devoted to the discussion and interpretation of the results illustrated in Section 598 4. We focus in particular on the key results obtained in terms of the probabilistic assessment of CO_2 599 generation through CCR.

600 **5.1 Geochemical modeling results**

With reference to Fig. 4a, we observe that all percentiles associated with $\log \tilde{K}_{R,T,P}$ tend to increase with temperature and pressure. Fig. 4a shows that a negligible probability is associated with positive values of $\log \tilde{K}_{R,T,P}$ when T < 50 °C, i.e. the equilibrium (8) favors primary phases over secondary phases. Otherwise, our results indicate that a probability very close to 1 is associated with values of $\log \tilde{K}_{R,T,P} > 0$ for T > 100 °C. This finding is consistent with the results of Smith and Ehrenberg (1989) who suggest that CO_2 formation is typically favored above 100-120°C as a consequence of carbonate phase consumption.

Fig. 4b, shows that the partial pressure of CO_2 tends to increase with temperature as a direct 608 consequence of the trend of $\log \tilde{K}_{R,T,P}$ in Fig. 4a. The median value of $\log \tilde{P}_{CO2}$, $p_{50}(\log \tilde{P}_{CO2})$ is 609 consistent with field observations (e.g., Texas, Norway and Thailand basins in Fig. 4b) for 610 611 temperature values higher than 100 °C. Almost all of the field data reported by Coudrain-Ribstein et al. (1998) for this temperature range fall between $p_1(\log \tilde{P}_{CO2})$ and $p_{99}(\log \tilde{P}_{CO2})$, with the 612 613 exception of a very limited number of points. Otherwise, the majority of the field data (mainly 614 associated with Alberta, Paris, Arkansans and Medison basins in Fig. 4b) falls outside the range identified by $p_1(\log \tilde{P}_{CO2})$ and $p_{99}(\log \tilde{P}_{CO2})$ for $T < 100 \,^{\circ}\text{C}$. The median value of $\log \tilde{P}_{CO2}$ resulting 615 from our simulations tends to overestimate the field data in this temperature range. Giggenbach 616 (1981) suggests that dilution of aqueous CO_2 in the system at shallow depth (corresponding to low 617 618 temperature) can happen due to mixing of fresh and cold water (i.e., from meteoric precipitations) 619 with groundwater. Moreover, Coudrain-Ribstein et al. (1998) observe that complex minerals such as 620 illite or competing geochemical processes can play a relevant role at low temperature levels. The discussion of the consistency of the data with possible alternatives of physical and conceptual models 621 622 as the ones suggested above is beyond the scope of the present work as previously explained in 623 Section 3.1. Here, we can highlight that our procedure leads to results which are consistent with the degree of variability of P_{CO2} values observed in real systems at temperatures T > 100 °C. 624

625

5.2 CO₂ generation and CCR mechanism activation

We start our discussion by considering the characterization of \tilde{P}_{CO2} as function of depth. All 626 values of $p_{W} \left(\log \tilde{P}_{CO2} | C_{max} \right)$ display a monotonic increase with depth (Fig. 5a-b) at the considered 627 628 times and for both mineralogical composition scenarios. This behavior is consistent with the 629 observation that (i) temperature and pressure increase with depth at all times (see Fig. 3c and d); and (*ii*) the equilibrium constant log $\tilde{K}_{R,T,P}$ increases with temperature and pressure (see Fig. 4a), i.e., 630 formation of CO_2 is favored by the increase of temperature and pressure. Partial pressure of CO_2 is 631 632 computed only in those layers within which there is a mineral composition compatible with the CCR process, labeled as carbonate layers in Figure 3. The total basin thickness at t = 48 Ma is 633 approximately equal to 5.5 km, the basin being completely constituted by carbonates rocks (see Fig. 634 4a). Thus, we find $\tilde{P}_{CO2} | C_{max} > 0$ across the whole computational domain (Fig. 5a). The impact of the 635 limiting reactant associated with the two mineral composition scenarios is negligible at this time level 636 and no significant differences are detected between values of $p_W \left(\log \tilde{P}_{CO2} | C_{max} \right)$ computed for 637 scenarios S_{cal} and S_{dol}. We can then conclude that the dissolved CO₂ concentration values rendered 638 by the geochemical model at this time do not exceed the value of the maximum admissible 639 640 concentration associated with either S_{dol} or S_{cal} . Otherwise, the mineral composition at t = 0 Ma influences the statistical characterization of $\log \tilde{P}_{CO2} | C_{max}$ at large depths (Z > 6 km). We observe that 641 $p_{W}\left(\log \tilde{P}_{CO2} | C_{max}\right)$ displays a different trend for depths larger than 6 km, according to the 642 mineralogical composition considered. Fig. 6b suggests that the effect of limiting reactant affects all 643 probability levels, i.e., $p_{W}\left(\log \tilde{P}_{CO2} \left| C_{max}\left(S_{dol} \right) \right) > p_{W}\left(\log \tilde{P}_{CO2} \left| C_{max}\left(S_{cal} \right) \right)$ for all considered values 644

645 of W even as the value of C_{max} (quantifying the effect of limiting reactant) is a deterministically 646 imposed upper boundary (see the Electronic Annex III for additional details).

We then discuss the results obtained in terms of the activation of the generation of gaseous 647 CO₂. Values of \tilde{R} associated with all of the considered percentiles $p_{I}(\tilde{R})$ increase with depth for 648 649 both time levels considered (see Fig. 5c-d). This result indicates that the sum of gas partial pressures $(\tilde{P}_{CO2} \text{ and } P_{H2O})$ tends to increase with depth at a faster rate than does the fluid pressure P. The 650 difference $p_{99}(\tilde{R}) - p_1(\tilde{R})$ markedly increases with depth, suggesting that the level of uncertainty 651 associated with \tilde{P}_{CO2} tends to increase with temperature and pressure. Consistent with Fig. 5b, the 652 mineral composition scenario influences these results only for Z > 6 km. Fig. 6 presents the 653 comparison of the sample *cdfs* (cumulative distribution functions) of \tilde{R} (11) at Z = 8 km for scenarios 654 S_{dol} and S_{cal}. We observe that the relative proportions among the different minerals constituting the 655 sediments influences the statistical distribution of \tilde{R} and, consequently, the probability of generation 656 657 of gaseous CO₂.

The value of the sample probability of generation of gaseous $CO_2 G_A(t)$ increases with time (Fig. 7) and attains its highest value for the final simulation time (t = 0 Ma). It is possible to distinguish three stages according to the time evolution of $G_A(t)$: (*i*) for $t \in]$ 45 Ma, 135 Ma], where G = 0; *ii*) for $t \in]$ 20 Ma, 45 Ma], where 0 < G < 0.2, with comparable values for S_{dol} and S_{cal}; and (*iii*) for $t \in]$ 0 Ma, 20 Ma], where G_A continuously increases, with a trend which varies according to the mineralogical scenario. It can be noted that the probability of activation grows slower in time for scenario S_{cal} than for S_{dol}.

665 Our results suggest that the temperature range associated with locations where the activation 666 of the process is possible, i.e., at which $f_{Zact,\tau} > 0$, is comprised between 200 and 300 °C (compare 667 Fig. 8b-c with Fig. 4c). This information can be highly valuable, e.g., to assess the prior probability 668 of CCR being a key source of CO_2 in natural systems (e.g., Jarvie and Jarvie, 2007). Cathles and Schoell (2007) predict an activation temperature of 330° C through a deterministic approach similar to the one presented in Section 2 and a simple time-independent *P*-*T* relationship. Our results suggest that the generation of gaseous *CO*₂ by CCR might take place also at lower temperatures when the parametric uncertainty related to the geochemical model are considered.

With reference to the results depicted in Fig. 9 and related to the probabilistic analysis of \tilde{F}_{CO2} 673 , we note that nonzero (positive) values of the latter can be found only if the CCR mechanism is active 674 at a given time, i.e., if $\tilde{Z}_{act}(t) \neq \emptyset$, \tilde{F}_{CO2} being equal to zero otherwise. The contour lines describing 675 $f_{F,t}$ in Fig. 9a-b are qualitatively very similar. However, we observe a remarkable quantitative 676 difference between the two scenarios analyzed: non-zero values of \tilde{F}_{CO2} range between 12 and 31 677 ton/Ma in scenario S_{dol}, while these are comprised between 1.0 and 2.5 ton/Ma for S_{cal}. This result 678 679 can be ascribed to the effect of the diverse fractions of dolomite characterizing S_{dol} and S_{cal} and acting as the limiting reactant. The *cdfs* reported in Fig. 9c indicate that the nonzero values of \tilde{F}_{CO2} display 680 681 a modest variability for a given time level. This suggests that, even as the location of the source is 682 characterized by remarkable variability across the Monte Carlo sample (see Fig. 8), porosity and sediment velocity which contribute to \tilde{F}_{CO2} according to (13) display modest variability along the 683 region of vertical domain where $f_{Zact,t} > 0$. Our results also show that the nonzero values of \tilde{F}_{CO2} 684 685 observed at t = 20 Ma in the Monte Carlo sample are fewer than those obtained at t = 0 Ma. Note that the non-zero values of \tilde{F}_{CO2} detected at t = 20 Ma are larger than their non-zero counterparts arising 686 687 at t=0 Ma. For those realizations within which the generation of gaseous CO_2 is activated, we obtain 688 a CO_2 generation rate of about 27 and 18 ton/Ma, respectively at t = 20, and 0 Ma. This difference is a consequence of the diverse values of the sediment burial velocity $(V_{SED}(\tilde{Z}_{act},t))$ in (13)) at the 689 690 location where gaseous CO_2 is generated. We exclude that porosity can play a relevant role in the different \tilde{F}_{CO2} values obtained at t = 0 and 20 Ma as it is almost constant (approximately equal to 0.1) 691 692 for Z > 4 km, where the CO_2 source is located (see Fig. 3c).

We conclude our discussion by considering the distribution of $\log \tilde{C}_{CO2(aq)} \left| \left(C_{max}, \tilde{Z}_{act} \right) \right|$ 693 (Eq.I.23 in the Electronic Annex I) depicted in Fig. 10. We recall that quantity $f_{C,Z}(0)$ denotes the 694 relative frequency associated with $\tilde{C}_{CO2(aq)} | (C_{max}, \tilde{Z}_{act}) = 0$. Fig. 10c, d) respectively depict the 695 dependence of $f_{C,Z}(0)$ on Z for S_{dol} and S_{cal}. Note that, according to our conceptual model, 696 $\tilde{C}_{CO2(aq)} | (C_{max}, \tilde{Z}_{act}) = 0$ at all locations where mudstone layers are found and below the depth Z_{act} . 697 As such, we find $f_{C,Z}(0) = 1$ at 0 < Z < 1.4 km. We observe that $f_{C,Z}(0) = 0$ at 1.4 < Z < 4.5 km, 698 699 suggesting that the concentration of dissolved CO₂ attains non-zero values across the complete Monte Carlo set. Finally, the relative frequency $f_{C,Z}(0)$ attains values higher than zero and lower than one 700 701 and increases with depth for Z > 4.5 km. This finding is consistent with results of Fig. 8, showing that (i) Z = 4.5 km is the shallowest location at which the activation of the CCR mechanism is possible at 702 703 t = 0 Ma and (*ii*) the probability to observe vanishing CO_2 concentrations at a given location increases with the relative frequency that the depth of such a location is larger than that corresponding to Z_{act} . 704 Calculated values for concentration of dissolved CO_2 display negligible dependence on 705 mineral composition scenario, in contrast with \tilde{F}_{CO2} (Fig. 9). The only impact of the mineral 706 composition scenario on $f_{C,Z}$ is due to the upper bound C_{max} imposed by the availability of reactants 707 which leads to an increase of the relative frequency $f_{C,Z}$ of values $\tilde{C}_{CO2(aq)} \left| \left(C_{max}, \tilde{Z}_{act} \right) = C_{max} \right|$ at large 708 709 depths within S_{cal} (see Fig. 10b). This behavior follows from the observation that the extent of the region where the reaction can occur is limited by the available dolomite volume fraction in S_{cal} (see 710 711 Fig. 10d).

712 6 Conclusions

713 We present a methodology conducive to a probabilistic assessment of the amount of CO_2 714 generated in sedimentary basins as consequence of the interaction between carbonate and clay minerals in the presence of pore-water. Our modeling strategy rests on the quantification of the uncertainty of chemical equilibrium parameters related to mineral solubility and the way it propagates to key model outputs. Application of the proposed workflow leads to a probabilistic assessment of: (*i*) the evolution of CO_2 partial pressure and dissolved CO_2 as a function of depth and time along the basin burial history; (*ii*) the location of the source where gaseous CO_2 is released from the sediments; (*iii*) the amount of gaseous CO_2 released per unit time.

721 We illustrate our approach upon relying on a realistic basin compaction history meaning that 722 temperature-pressure-porosity combinations are compatible with realistic fields. Our work provides 723 a first attempt to quantify CO₂ generation by CCR at geological scales with the explicit inclusion of a probabilistic assessment of the uncertainty stemming from the incomplete knowledge of mineral 724 725 solubility and phase equilibrium constants at high temperatures. Due to its flexibility, we envision 726 that the framework proposed here can be readily extended to include the uncertainty related to the 727 basin pressure and temperature dynamics. We envision that the proposed model may be extended in 728 future works to include other sources of model uncertainty, such as those associated with pore-water 729 chemistry (e.g., salinity).

Our uncertainty quantification is based on data of mineral solubility and phase equilibrium constants available at laboratory scales. We verify that the procedure we employ to characterize parametric uncertainty of the geochemical model leads to results which are consistent with field observations of CO_2 partial pressure in sedimentary formations reported in the literature.

Our study shows that the partial pressure of CO_2 displays a monotonic and increasing trend with depth. This suggests that the increase of temperature taking place during a basin burial history favors the progressive generation of CO_2 at the expense of carbonate mineral phases. CO_2 is generated as a separate phase only under specific conditions which depend on temperature and pressure distributions. The probability that these conditions are encountered tends to increase with time and attains its largest value (around 0.45 in the setting we analyze) at the end of the simulation period, which represent the present day. In our example we find that generation of CO_2 through CCR can

741 become effective at temperatures comprised between 200 and 300 °C. These specific results are 742 conditional to the given compaction history of the basin and of the geochemical model structure 743 selected in this study are therefore not amenable to direct transferability to diverse geological settings. 744 Mineral compositions associated with sediments largely affect the flux of generated CO_2 . In 745 the case we examine, the key driver is the amount of dolomite associated with the sediments and 746 representing the source of CO_2 . In our illustrative example, the impact of model parameter uncertainty 747 is stronger on the activation depth than on the CO_2 generation rate. As a consequence, our findings 748 suggest that reliable estimates of CO₂ migration scenarios should rely on accurate characterization of 749 mineral composition as well as geochemical model parameters.

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752 **References**

- Allis R., Chidsey T., Gwynn W., Morgan C., White S., Adams M. & Moore J. (2001). Natural CO2
- reservoirs on the colorado plateau and southern rocky mountains: Candidates for CO2
- sequestration. *Proceedings of the First National Conference on Carbon Sequestration*, 14-17.
- 756 Anderson G. M. (2009). *Thermodynamics of natural systems* Cambridge University Press.
- Arnórsson S. (1986). Chemistry of gases associated with geothermal activity and volcanism in
 iceland: A review. J. Geophys. Res. B: Solid Earth, 91(B12), 12261-12268.
- 759 Ballentine C. J., Schoell M., Coleman D. & Cai B. A. (2001). 300-myr-old magmatic CO2 in
- natural gas reservoirs of the west texas permian basin. *Nature*, **409(6818)**, 327-331.

761	Battistelli A., Berry P., Bonduà S., Bortolotti V., Consonni A., Cormio C. Geloni C. & Vasini E. M.
762	(2016) Thermodynamics-related processes during the migration of acid gases and methane in
763	deep sedimentary formations. GREENH GASES.

- 764 Bergaya F. & Lagaly G. (2013). General Introduction: Clays, Clay Minerals, and Clay Science.
- 765 *Handbook of Clay Science*, 1, 1.
- Bianchi Jannetti E., Dror I., Riva M., Guadagnini A. & Berkowiz B. (2012). Estimation of Single-Metal and
 Competitive Sorption Isotherms through Maximum Likelihood and Model Quality Criteria. *Soil Sci. Soc. Am. J.* 76, 1229–1245.
- 769 Blanc P., Lassin A., Piantone P., Azaroual M., Jacquemet N., Fabbri A., & Gaucher E. C. (2012).
- Thermoddem: A geochemical database focused on low temperature water/rock interactions and
 waste materials. *Appl. Geochem.*, 27(10), 2107-2116.
- Blanc P., Vieillardb P., Gailhanoua H., & Gaboreaua S. (2013). Thermodynamics of clay minerals. *Handbook of Clay Science*, 6, 173.
- Blanc P., Vieillard P., Gailhanou H., Gaboreau S., Gaucher É., Fialips C. I., Madé B. & Giffaut E.
 (2015). A generalized model for predicting the thermodynamic properties of clay
- 776 minerals. Am. J. S., **315(8)**, 734-780.
- 777 Broadhead R. F., Mansell M. & Jones G. (2009). Carbon Dioxide in New Mexico: Geologic
- Distribution of Natural Occurrences. New Mexico Bureau of Geology and Mineral Resources
 Open-file Report 514.
- Cai C., Hu W., & Worden R. H. (2001). Thermochemical sulphate reduction in Cambro–Ordovician
 carbonates in central Tarim. *Mar. Pet. Geol.*, 18(6), 729-741.

782 Cathles L. & Schoell M. (2007). Modeling CO2 generation, migration, and titration in sedimentary 783 basins. Geofluids, 7(4), 441-450.

784	Chiodini G., Baldini A., Barberi F., Carapezza M., Cardellini C., Frondini F., Granieri D. & Ranaldi
785	M. (2007). Carbon dioxide degassing at Latera caldera (Italy): Evidence of geothermal
786	reservoir and evaluation of its potential energy. J. Geophys. Res. B: Solid Earth, 112(B12).
787	Chiodini G., Frondini F., Cardellini C., Parello F. & Peruzzi L. (2000). Rate of diffuse carbon
788	dioxide earth degassing estimated from carbon balance of regional aquifers: The case of central
789	apennine, Italy. J. Geophys. Res. B: Solid Earth, 105(B4), 8423-8434.
790	Clayton J., Spencer C., Koncz I. & Szalay A. (1990). Origin and migration of hydrocarbon gases
791	and carbon dioxide, Bekes basin, southeastern Hungary. Org. Geochem., 15(3), 233-247.
792	Colombo I., Porta G. M., Ruffo P., Guadagnini A. (2016). Assessment of overpressure buildup
793	through inverse modeling of compaction processes in sedimentary basins. Hydrol. J.
794	(accepted).
795	Cooper B., Raven M. & Samuel L. (1997). Origin and geological controls on subsurface CO2
796	distribution with examples from western Indonesia. Proceedings of an International
797	Conference on Petroleum Systems of SE Asia and Australasia, 877-892
798	Coudrain-Ribstein A. & Gouze P. (1993). Quantitative study of geochemical processes in the
799	dogger aquifer, paris basin, france. App. Geochem., 8(5), 495-506.
800	Coudrain-Ribstein A., Gouze P. & de Marsily G. (1998). Temperature-carbon dioxide partial
801	pressure trends in confined aquifers. Chem. Geol., 145(1), 73-89.
802	Delany J. & Lundeen S. (1990). The LLNL thermochemical database. Lawrence Livermore National
803	Laboratory Report UCRL-21658.

804	Dubacq B., Bickle M. J., Wigley M., Kampman N., Ballentine C. J. & Lollar B. S. (2012). Noble
805	gas and carbon isotopic evidence for CO 2-driven silicate dissolution in a recent natural CO 2
806	field. Earth Planet. Sci. Lett., 341, 10-19.
807	Farmer R. (1965). Genesis of subsurface carbon dioxide. Fluids in Subsurface Environments, A
808	Symposium.

- 809 Fischer M., Botz R., Schmidt M., Rockenbauch K., Garbe-Schönberg D., Glodny J., Gerling R.,
- Littke R. (2006). Origins of CO2 in permian carbonate reservoir rocks (zechstein, Ca2) of the
 NW-german basin (lower saxony).*Chem. Geol.*, 227(3), 184-213.
- 812 Formaggia L., Guadagnini A., Imperiali I., Lever V., Porta G., Riva M., Scotti A., Tamellini L.
- 813 (2013). Global sensitivity analysis through polynomial chaos expansion of a basin-scale

geochemical compaction model. *Comput. Geosci.*, **17(1)**, 25-42.

- 815 Giggenbach W. F. (1978). The isotopic composition of waters from the el tatio geothermal field,
 816 northern chile. *Geochim. Cosmochim. Acta*, 42(7), 979-988.
- 817 Giggenbach W. F. (1980). Geothermal gas equilibria. *Geochim. Cosmochim. Acta*, 44(12), 2021818 2032.
- 819 Giggenbach W. F. (1981). Geothermal mineral equilibria. *Geochim. Cosmochim. Acta*, 45(3), 393820 410.
- Giggenbach W. F. (1984). Mass transfer in hydrothermal alteration systems—a conceptual
 approach. *Geochim. Cosmochim. Acta*, 48(12), 2693-2711.
- 823 Goldsmith J. R. (1980). Thermal stability of dolomite at high temperatures and pressures. *J.*
- 824 *Geophys. Res. B: Solid Earth*, **85(B12)**, 6949-6954.

- Higgs K. E., Funnell R. H. & Reyes A. G. (2013). Changes in reservoir heterogeneity and quality as
 a response to high partial pressures of CO2 in a gas reservoir, New Zealand. *Mar. Pet. Geol.*,
 48, 293-322.
- Huang W. & Longo J. (1994). Experimental studies of silicate-carbonate reactions—I. applications
 to diagenesis. *App. Geochem.*, 9(5), 501-522.
- Hutcheon I. (1990). Clay carbonate reactions in the venture area, Scotian shelf, Nova Scotia,
 Canada. *Geo. Soc. S. P.*, 2, 199-212.
- Hutcheon I. & Abercrombie H. (1990). Carbon dioxide in clastic rocks and silicate hydrolysis. *Geology*, **18(6)**, 541-544.
- Hutcheon I., Abercrombie H. J. & Krouse H. (1990). Inorganic origin of carbon dioxide during low
 temperature thermal recovery of bitumen: Chemical and isotopic evidence. *Geochim. Cosmochim. Acta*, 54(1), 165-171.
- Hutcheon I., Abercrombie H. J., Putnam P., Gardner R. & Krouse H. R. (1989). Diagenesis and
 sedimentology of the clearwater formation at tucker lake. *B. Can. Petrol. Geol.*, 37(1), 83-97.
- Hutcheon I., Oldershaw A. & Ghent E. D. (1980). Diagenesis of cretaceous sandstones of the
- 840 kootenay formation at elk valley (southeastern british columbia) and mt allan (southwestern
- 841 alberta). *Geochim. Cosmochim. Acta*, **44(10)**, 1425-1435.
- 842 Hutcheon I., Shevalier M., & Abercrombie H. J. (1993). pH buffering by metastable mineral-fluid
- 843 equilibria and evolution of carbon dioxide fugacity during burial diagenesis. *Geochim*.
- 844 *Cosmochim. Acta*, **57(5)**, 1017-1027.
- 845 Imbus S. W., Katz B. J. & Urwongse T. (1998). Predicting CO2 occurrence on a regional scale:
- 846 Southeast Asia example. Org. Geochem., **29(1)**, 325-345.

847	Jarvie D. M. & Jarvie B. M. (2007). Thermal decomposition of various carbonates: kinetic results
848	and geological temperatures of conversion. 23th international meeting on Organic
849	Geochemistry (IMOG), Torquay, UK, 9-14 September 2007.

- Johnson J. W., Oelkers E. H. & Helgeson H. C. (1992). SUPCRT92: A software package for
- 851 calculating the standard molal thermodynamic properties of minerals, gases, aqueous species,
- and reactions from 1 to 5000 bar and 0 to 1000 C. *Comput. Geosci.*, **18(7)**, 899-947.
- 853 Kharaka Y. K., Gunter W. D., Aggarwal P. K., Perkins E. H. & De Braal J. D. (1988). SOLMINEQ.
- 854 88: A computer program for geochemical modeling of water-rock interactions. US Geological
- 855 Survey Water-Resources Investigation Report, 88, 4227.
- 856 Kotarba M. J. & Nagao K. (2008). Composition and origin of natural gases accumulated in the
- polish and ukrainian parts of the carpathian region: Gaseous hydrocarbons, noble gases, carbon
 dioxide and nitrogen. *Chem. Geol.*, 255(3), 426-438.
- Li M., Wang T., Liu J., Lu H., Wu W. & Gao L. (2008). Occurrence and origin of carbon dioxide in
 the fushan depression, beibuwan basin, south china sea. *Mar. Pet. Geol.*, 25(6), 500-513.
- Maier C. G. & Kelley K. (1932). An equation for the representation of high-temperature heat
 content data1. *J. Am. Chem. Soc.*, 54(8), 3243-3246.
- Mayo A. L. & Muller A. B. (1997). Low temperature diagenetic–metamorphic and magmatic
 contributions of external CO2 gas to a shallow ground water system. *J. Hydrol.*, **194(1)**, 286304.
- 866 Metz B., Davidson O., De Coninck H., Loos M., & Meyer L. (2005). IPCC, 2005: IPCC special
- 867 report on carbon dioxide capture and storage. prepared by working group III of the

- 868 intergovernmental panel on climate change. *Cambridge, United Kingdom and New York, NY,*869 USA, 442.
- Millero F. J. (1982). The effect of pressure on the solubility of minerals in water and seawater. *Geochim. Cosmochim. Acta*, 46(1), 11-22.
- Neuman S. P. (2003). Maximum likelihood Bayesian averaging of uncertain model predictions. *Stoch. Environ. Res. Risk Ass.*, 17, 291-305.
- 874 Parkhurst D. L. & Appelo C. (2013). Description of input and examples for PHREEQC version 3—
- 875 a computer program for speciation, batch-reaction, one-dimensional transport, and inverse
- geochemical calculations. US Geological Survey Techniques and Methods, Book, 6, 497.
- 877 Peterson S. R., Hostetler C. J., Deutsch W. J. & Cowan C. E. (1987). *MINTEQ user's manual* (No.
- 878 NUREG/CR-4808; PNL-6106). Pacific Northwest Lab., Richland, WA (USA); Nuclear
- 879 Regulatory Commission, Washington, DC (USA). Div. of Waste Management.
- 880 Plummer L. N. & Busenberg E. (1982). The solubilities of calcite, aragonite and vaterite in CO 2-H
- 2 O solutions between 0 and 90 C, and an evaluation of the aqueous model for the system
- 882 CaCO 3-CO 2-H 2 O. *Geochim. Cosmochim. Acta*, **46(6)**, 1011-1040.
- 883 Porta G., Tamellini L., Lever V. & Riva M. (2014). Inverse modeling of geochemical and
- mechanical compaction in sedimentary basins through polynomial chaos expansion. *Water Resour. Res.*, **50(12)**, 9414-9431.
- 886 Razavi S. & Gupta H. V. (2015). What do we mean by sensitivity analysis? the need for
- 887 comprehensive characterization of "global" sensitivity in earth and environmental systems
- 888 models. *Water Resour. Res.*, **51(5)**, 3070-3092.

- Riva M., Guadagnini A. & Dell'Oca A. (2015). Probabilistic assessment of seawater intrusion under
 multiple sources of uncertainty. *Adv. Water Resour.*, **75**, 93-104.
- Saltelli A., Ratto M., Andres T., Campolongo F., Cariboni J., Gatelli D., Saisana M., Tarantola S.
 (2008). *Global sensitivity analysis: The primer* John Wiley & Sons.
- Smith J. & Ehrenberg S. (1989). Correlation of carbon dioxide abundance wit1h temperature in
 clastic hydrocarbon reservoirs: Relationship to inorganic chemical equilibrium. *Mar. Pet. Geol.*, 6(2), 129-135.
- Sobol I. M. (2001). Global sensitivity indices for nonlinear mathematical models and their Monte
 Carlo estimates. *Mat. Comput. Simul.*, 55(1), 271-280.
- Span R. & Wagner W. (1996). A new equation of state for carbon dioxide covering the fluid region
 from the triple-point temperature to 1100 K at pressures up to 800 MPa. *J. Phys. Chem. Ref. Data*, 25(6), 1509-1596.
- 901 Stumm W. & Morgan J. (1996). Aquatic chemistry: Chemical equilibria and rates in natural waters.
 902 *John Wiley & Sons: New York, 1040.*
- Spycher N., Pruess K., & Ennis-King J. (2003). CO2-H2O mixtures in the geological sequestration
 of CO2. *Geochim. Cosmochim. Acta* 67,3015.
- 905 Ueda A., Kato K., Ohsumi T., Yajima T., Ito H., Kaieda H., Metcalfe R. & Takase H. (2005).
- 906 Experimental studies of CO2-rock interaction at elevated temperatures under hydrothermal
- 907 conditions. *Geochem. J.*, **39(5)**, 417-425.
- 908 van Berk W., Schulz H. & Fu Y. (2009). Hydrogeochemical modelling of CO2 equilibria and mass
- 909 transfer induced by organic–inorganic interactions in siliciclastic petroleum reservoirs.
- 910 *Geofluids*, **9(4)**, 253-262.

911	van Berk W., Schulz H. & Fu Y. (2013). Controls on CO2 fate and behavior in the gullfaks oil field
912	(norway): How hydrogeochemical modeling can help decipher organic-inorganic interactions.
913	AAPG Bull., 97(12), 2233-2255.
914	Wycherley H., Fleet A. & Shaw H. (1999). Some observations on the origins of large volumes of
915	carbon dioxide accumulations in sedimentary basins. Mar. Pet. Geol., 16(6), 489-494.
916	Xu T. & Pruess K. (2001). On fluid flow and mineral alteration in fractured caprock of magmatic
917	hydrothermal systems. J. Geophys. Res. B: Solid Earth, 106(B2), 2121-2138.
918	Yaalon D. H. (1962). Mineral composition of the average shale. Clay Miner., 5(27), 31-36.
919	Zhang S., FitzGerald J. D. & Cox S. F. (2000). Reaction-enhanced permeability during
920	decarbonation of calcite quartz \rightarrow wollastonite carbon dioxide. <i>Geology</i> , 28(10) , 911-914.
921	

	Reaction
R1	$ \begin{array}{c c} 5CaMg(CO_3)_2 + Al_2Si_2O_5(OH)_2 + SiO_2 + 2H_2O = 5CO_2 + 5CaCO_3 + Mg_5Al_2Si_3O_{10}(OH)_8 \\ \hline \text{dolomite} & \text{kaolinite} & \text{silica} & \text{calcite} & \text{chlorite} \\ \end{array} $
R2	$\begin{array}{c c} 3CaMg(CO_3)_2 + KAl_3Si_3O_{10}(OH)_2 + 2SiO_2 + 2H_2O = 3CO_2 + 3CaCO_3 + KMg_3(AlSi_3)O_{10}(OH)_2 + Al_2Si_2O_5(OH)_2\\ \hline \text{dolomite} \qquad \text{muscovite} \qquad \text{silica} \qquad \text{calcite} \qquad \text{phlogopite} \qquad \text{kaolinite} \end{array}$
R3	$\begin{array}{ c c c c c c c } \hline & 3CaMg(CO_3)_2 + KAlSi_3O_8 + H_2O = & 3CO_2 + 3CaCO_3 + KMg_3(AlSi_3)O_{10}(OH)_2 \\ \hline & \text{dolomite} & \textbf{K-feldspar} & \textbf{calcite} & \textbf{phlogopite} \\ \hline \end{array}$
R4	$\begin{array}{ c c c c c c c c c c c c c c c c c c c$
R5	$\begin{array}{c c c c c c c c c c c c c c c c c c c $
R6	$5CaMg(CO_3)_2 + 5Al_2Si_2O_5(OH)_2 + 24SiO_2 + 12KAl_3Si_3O_{10}(OH)_2 = 5CO_2 + 5CaCO_3 + 20K_{0.6}Mg_{0.25}Al_{1.8}Al_{0.5}Si_{3.5}O_{10}(OH)_2 + 2H_2O_{1.5}O_{1.6}O_$

Table 1: Examples of CCRs proposed in Coudrain-Ribstein et al. (1998)

Time interval	$V_D [\mathrm{m}\mathrm{Ma}^{-1}]$
$t \in] 96 \text{ Ma}, 135 \text{ Ma}]$	23.6
$t \in \left] 48 \text{ Ma}, 96 \text{ Ma} \right]$	140
$t \in]$ 34 Ma, 48 Ma]	89.8
$t \in]$ 23 Ma, 34 Ma]	110
$t \in]$ 5 Ma, 23 Ma]	132
$t \in] 0 \text{ Ma}, 5 \text{ Ma}]$	142

Table 2: Values of sediment depositional velocity (V_D) as a function of time. The depositional velocity is assumed constant within each time interval.

Type of sediment	$K_T [J m^{-1} s^{-1} K^{-1}]$	<i>ф</i> ₀ [-]	β [Pa]
Carbonate	2.93	0.42	4.6 x 10 ⁻⁸
Shale	1.90	0.78	25 x 10 ⁻⁸
Sandy Shale	2.26	0.78	26 x 10 ⁻⁸

Table 3: Main characteristics (thermal matrix conductivity K_T , porosity at deposition ϕ_0 and compressibility coefficients β) of the sediment types considered.

Scenario	Calcite	Dolomite	Kaolinite	Quartz	Siderite	Pyrite	Illite	Smectite	Chlorite
	[%wt]	[%wt]	[%wt]	[%wt]	[%wt]	[%wt]	[%wt]	[%wt]	[%wt]
S _{dol}	-	74	21	5	-	-	-	-	-
S _{cal}	74	6	3	4	2	2	1	8	1

Table 4: Mineral composition for the two scenarios considered.

Phase	$\mu(\tilde{A})$	$\mu(ilde{D})$	Ψ (1,1)	Ψ (1,1)	$\Psi(1,2) = \Psi(2,1)$	Criterion
Calcite	-851.1	310.1	0.0104	0.0017	-0.0042	Estimated by solubility data
Kaolinite	-984.2	353.8	13.19	1.92	-5.03	Estimated by solubility data
Dolomite	-1781.4	647.1	0.0104	0.0017	-0.0042	Belongs to the carbonate mineral class as calcite
Quartz	-19.9	-84503.0	0	0	0	No uncertainty
Clinochlore	-2858.9	1029.1	13.19	1.92	-5.03	Belongs to the clay mineral class as kaolinite
$CO_{2(g)}$	-593.1	212.8	13.19	1.92	-5.03	Largest uncertainty estimated
$H_2O_{(g)}$	-23.21	5.925	0	0	0	No uncertainty
$CO_{2(aq)}$	682.1	-246.5	13.19	1.92	-5.03	Largest uncertainty estimated

Table 5: Characterization of the uncertain (random) model inputs (see the text for notation).

Figure 5: Vertical distribution of the percentiles p_1 , p_{25} , p_{50} , p_{75} , p_{99} of (a-b) $\log \tilde{P}_{CO2}$ and (c-d) \tilde{R} at t = (a, c) 0, (b, d) 48 Ma for mineralogy S_{dol} (solid curves) and S_{cal} (dashed curves). Stratigraphic columns corresponding to t = 48 Ma (a-c) and t = 0 Ma (b-d) are also depicted for ease of reference.

Figure 8: (a) Sample probability (relative frequency) of \tilde{Z}_{act} at t = 10 Ma; color maps of sample probability $f_{Zact,t}$ as a function of time for scenarios (b) S_{dol} and (c) S_{cal}10

Figure 10: Color map of sample probabilities (relative frequencies) associated with vertical distributions of (a-b) $\log \tilde{C}_{CO2(aq)} | (C_{max}, \tilde{Z}_{act}) | = 0$ Ma, for mineral composition scenarios (a) S_{dol}, and (b) S_{cal}. The red dashed curve in (b) corresponds to the threshold value C_{max} . Panels (d) and (e) depict the vertical distribution of $f_{C,Z}(0)$, i.e., relative frequencies of the occurrence of $\tilde{C}_{CO2(aq)} | (C_{max}, \tilde{Z}_{act}) = 0$ computed at t = 0 Ma. The stratigraphic column at t = 0 Ma is also depicted for ease of reference.



Figure 1: Outline of the carbonate/clay reactions mechanism (Cathles and Schoell, 2009): (a) the chemical system is at equilibrium and a separate gaseous CO₂ phase is absent; (b) the system reaches a new chemical equilibrium where gas partial pressure is higher than environmental pressure; (c) a separate CO₂ phase is formed and starts to migrate upwards due to buoyancy. Symbols P and P_{gas} respectively denote the pressure of the interstitial water and the sum of the partial pressure of all gaseous species formed in the pore-water along with CO₂.



Figure 2: Outline of the key steps of the modeling workflow. The overall procedure is subdivided in three blocks. In Electronic Annex 1 the computational steps of the three blocks are detailed item by item.



Figure 3: Computed space-time evolution of (a) stratigraphic sequence, (b) porosity; (c) temperature; (d) pressure of the basin scale test considered. The stratigraphic column at t = 0 Ma (ages of sediment layers are explicitly indicated) is depicted for completeness (Figure 3a). Solid vertical lines indicate the times associated with variations in the depositional velocity imposed at the basin top.



Figure 4: (a) Sample mean (red curve) and percentiles p_1 , p_{50} , p_{99} (blue curves) of $\log \tilde{K}_{R,T,P}$ and (b) percentiles p_1 , p_{50} , p_{99} of $\log \tilde{P}_{CO2}$ as a function of temperature. Symbols in (b) indicate field data reported in Coudrain-Ribstein et al. (1998).



Figure 5: Vertical distribution of the percentiles p_1 , p_{25} , p_{50} , p_{75} , p_{99} of (a-b) log \tilde{P}_{CO2} and (c-d) \tilde{R} at t = (a, c) 0, (b, d) 48 Ma for mineralogy S_{dol} (solid curves) and S_{cal} (dashed curves). Stratigraphic columns corresponding to t = 48 Ma (a-c) and t = 0 Ma (b-d) are also depicted for ease of reference.



Figure 6: Sample *cdf* of \tilde{R} (14) at t = 0 Ma and Z = 8 Km for mineral composition scenarios S_{dol} and S_{cal}. The red vertical line identifies the threshold value R = 1 which indicates the formation of a CO_2 -rich separate phase.



Figure 7: Sample probability (relative frequency) of occurrence of CO_2 generation as separate phase (G_A) as a function of time for mineral composition scenarios S_{dol} (solid curve) and S_{cal} (dashed curve).



Figure 8: (a) Sample probability (relative frequency) of \tilde{Z}_{act} at t = 10 Ma; color maps of sample probability $f_{Zact,t}$ as a function of time for scenarios (b) S_{dol} and (c) S_{cal} .



Figure 9: Color map of relative frequencies of logarithmically transformed values of the CO_2 generation rate, \tilde{F}_{CO2} , as a function of time for mineral composition scenarios (a) S_{dol} and (b) S_{cal} ; (c) sample *cdfs* describing the statistical distribution of \tilde{F}_{CO2} at t = 20 and 0 Ma (these observation times are highlighted by red dashed lines in Figure 9 (a)) associated with S_{dol} . Solid black curves in (a)-(b) represent the activation frequency (G_A) of the CO_2 generating process depicted in Figure 7.



Figure 10: Color map of sample probabilities (relative frequencies) associated with vertical distributions of (a-b) $\log \tilde{C}_{CO2(aq)} | (C_{max}, \tilde{Z}_{act}) |$ at t = 0 Ma, for mineral composition scenarios (a) S_{dol}, and (b) S_{cal}. The red dashed curve in (b) corresponds to the threshold value C_{max} . Panels (c) and (d) depict the vertical distribution of $f_{C,Z}(0)$, i.e., relative frequencies of the occurrence of

 $\tilde{C}_{CO2(aq)} | (C_{max}, \tilde{Z}_{act}) = 0$ computed at t = 0 Ma. The stratigraphic column at t = 0 Ma is also depicted for ease of reference.

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