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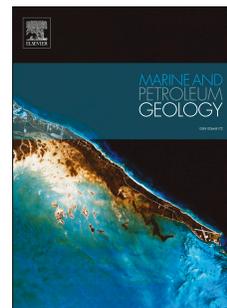
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Sample CRediT author statement

Amelie Cavelan : Investigation - Original draft preparation- Redaction - Writing - Review & Editing. **Mohammed Boussafir** : Conceptualization - Redaction -Supervision- Review & Editing - Funding acquisition. **Claude Le Milbeau** : Methodology – Validation – Reviewing. **Sandrine Delpeux** : Methodology – Validation – Reviewing. **Fatima Laggoun-Défarage** : Supervision – Project administration – Conceptualization .

Journal Pre-proof

Influence of experimental temperature and duration of laboratory confined thermal maturation experiments on the evolution of the porosity of organic-rich source rocks.

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Abstract: To investigate the influence of the duration-temperature pairs chosen for laboratory thermal maturations on the evolution of the organic matter (OM) and porosity of the clay-rich source rocks with increasing maturity, long duration (104 days) thermal maturations, bulk, molecular geochemical OM characterization (Rock Eval, GC-TCD, GC/MS) and nitrogen adsorption porosimetry were applied on six type II mudstones from the Kimmeridge Clay formation. These results were compared with shorter duration experiments (72 hours) previously carried out. The results show that the increase in the duration of thermal maturation experiments enhanced the degradation of heavy polar OM components into saturated and aromatic hydrocarbons, leading to the production of similar amounts of extractible OM (EOM) enriched in saturated and aromatic hydrocarbons. Then, in preventing the early breaking of low energy bonds, the lower temperature used for the longer duration experiments led to a less pronounced early cracking of OM resulting in a different timing of gas generation. This process did not influence the general evolution of the pore volume, which increased during gas generation. Whatever the duration-temperature pair chosen for maturation experiments highly oil-prone samples formed lower pore volumes during gas generation. Increasing the duration of thermal maturation seemed, nevertheless, to influence the amplitude and the timing of pore alteration events that occur during gas generation, leading to slightly different pore size distributions and pore volumes between

short and long duration experiments. However, these differences remain limited, suggesting that results from different time-duration experiments can be easily compared.

Key words: thermal maturation; porosity; source rocks; nitrogen adsorption; GC-MS

Introduction

The growing interest for unconventional resources has led to a considerable shift in the approach to characterize these reservoirs. The use of a combination of various cutting-edge technologies to quantify and image the complex structure of these fine-grained rocks has allowed a more precise understanding of the evolution of their porosity. The interparticle porosity associated with the mineral matrix generally decreases during burial, filled by entrapped EOM or reduced by compaction and OM ductile deformations (DiStefano et al., 2016; Furmann et al., 2016; Guo et al., 2018; Han et al., 2017; Katz and Arango, 2018; Loucks et al., 2009; Modica and Lapierre, 2012; X. Wang et al., 2020; Wang et al., 2019). Conversely, the progressive conversion of OM, the carbon loss and the significant variations of volume, which occur during oil and gas generation result in the growth of a secondary OM-hosted porosity (Bernard et al., 2012a; Chalmers and Bustin, 2008; Chen and Xiao, 2014; Curtis et al., 2012; Han et al., 2017; Jarvie et al., 2007; Ko et al., 2016, 2018; Loucks et al., 2012; X. Wang et al., 2020; Wu et al., 2019). In the vast majority of thermally mature rocks, the OM is the main contributor to the porosity (Bernard et al., 2012a; Cavelan et al., 2019a; Chalmers et al., 2012; Han et al., 2019; Juliao et al., 2015; Katz and Arango, 2018; Ko et al., 2018; Kuila et al., 2014; Loucks et al., 2009, 2012; Milliken et al., 2013; Pan et al., 2015). Thermal maturity and the initial OM composition, which controlled the amount of oil and gas generated during maturation, are thus widely considered as the key factors controlling the formation and the development of pores in organic-rich source rocks (Bernard et al., 2012a, 2012b; Cavelan et al., 2019a, 2020; Chalmers and Bustin, 2008; Chen and Xiao, 2014; Curtis et al., 2012; Ko et al., 2018; Loucks et al., 2009).

To investigate the relation between thermal maturity, OM composition and pore evolution, the use of laboratory thermal maturation has become more common these last years (Cavelan et al., 2019a, 2020; Chen and Xiao, 2014; Guo et al., 2017; Hu et al., 2015; Ko et al., 2016, 2018; X. Wang et al.,

2020; Y. Wang et al., 2020; Zhang et al., 2013). As demonstrated by these previous works, the use of laboratory thermal maturations could be a key method to obtain rocks of various maturities from the same initial material. However, various duration-temperature pairs can be selected for these experiments. While the effects of laboratory thermal maturation conditions on OM thermal degradation processes have been well studied (Behar et al., 2003; Landais et al., 1994; Michels et al., 1995, 1992), the effects on the evolution and the development of porosity have never been assessed. As OM thermal degradation processes are governed by the first order Arrhenius law, a plethora of many duration-temperature pairs should be able to be used to reach the same stage of maturity (Landais et al., 1994; Wood, 1988). However, previous studies show that the experimental temperature and duration of these laboratory simulations are not really exchangeable parameters (Landais et al., 1994), as increasing the thermal maturation duration and lowering the experimental temperatures could significantly influence the yield, the composition and the timing at which liquid and gaseous hydrocarbons are generated. In view of the strong relation existing between organic-rich source rocks porosity, OM thermal degradation processes, and oil and gas generation, these differences may influence the formation and the development of pores with increasing maturity. It seemed interesting therefore to test how the differences observed between short- and long-duration experiments can influence the formation and the development of organic-rich source rocks porosity. This may help to assess whether artificial maturation can accurately simulate the porosity evolution of natural source rocks.

To this end, long duration (104 days) gold-tube anhydrous confined thermal maturation experiments were carried out on six immature organic-rich source rocks from the Kimmeridge Clay formation. The concomitant evolutions of the bulk, molecular organic geochemistry and porosity during maturation were determined using Rock Eval[®] pyrolysis, GC-TCD, GC/MS and nitrogen adsorption measurements. The results were then compared with the shorter duration experiments (72 hours) previously carried out (Cavelan et al., 2019a, 2020). The present study aimed particularly to better understand the relation between the duration-temperature pair of anhydrous confined thermal maturation and the ability of rocks to develop and preserve pores with increasing maturity.

1. Samples and methods

1.1. Geological background

The late Jurassic Kimmeridge clay formation (KCF) consists of organic-rich marine shales deposited from the Kimmeridgian to the Tithonian (Herbin et al., 1995; Rawson and Riley, 1982). Onshore, this formation outcrops across England from the Dorset to the north Yorkshire; offshore the KCF is largely distributed throughout the Northern, Central and Southern North Sea (Fig. 1, Rawson and Riley, 1982).

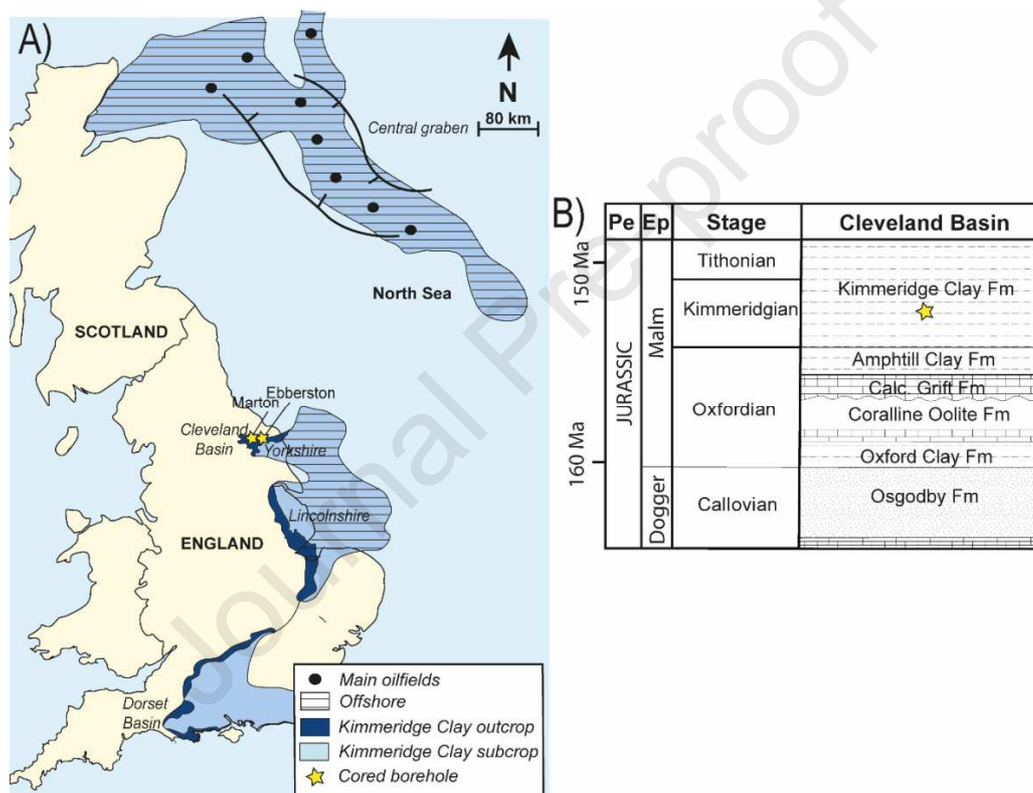


Fig. 1. A) Kimmeridge clay formation across England and North Sea; location of the studies boreholes (modified from Gallois, 2004). B) Simplified stratigraphic column of the Cleveland Basin (modified from Powell, 2010).

For this study, six immature KCF mudstones (Vitrinite reflectance ca. 0.42%, Cavelan et al., 2020) were selected from Marton and Eberston boreholes between 69.98 and 128.78 m deep (Yorkshire, UK, Fig.1). The organic geochemistry, the mineralogy and the porosity of this geological section were well-documented in the past twenty years (Boussafir et al., 1995a, 1995b; Boussafir and Lallier-Vergès, 1997; Cavelan et al., 2020; Desprairies et al., 1995; Ramanampisoa and Disnar, 1994;

Ramanampisoa and Radke, 1995). The studied rocks consist in finely laminated marine argillaceous mudstones with total organic carbon ranges between 3 to 15.4 wt.% (Rock Eval[®] 6 TOC) and a high OM oil generation potential varying between 418 to 617 mg of HC/g of TOC (Rock Eval[®] 6 hydrogen index, Cavelan et al., 2019b). The OM is mainly composed of amorphous organic matter (AOM) including in the order of decreasing abundance: highly oil-prone phytoplanktonic orange AOM, algal brown AOM and black AOM. This OM is accompanied by a minor contribution of structured components derived from zooplankton (pores, pollens,...) and plant tissues (Boussafir et al., 1995b; Boussafir and Lallier-Vergès, 1997).

1.2. Bulk and molecular OM composition

The devices used for anhydrous confined thermal maturation experiments were described in Cavelan et al. (2019a). About 1g of each sample was crushed into powder (particle size: 250 μm to 180 μm), loaded into gold cells (dia: 0.50 mm, len: 6.5 cm) and introduced into high-pressure stainless-steel autoclaves pressurized with water. A constant confining external pressure was maintained at 600 bar during the experiments. Samples were isothermally heated at 283, 338 and 390°C for 104 days in order to obtain the three ranks of maturity previously reached by the short-duration experiments of Cavelan et al., (2019a): the early oil generation, the condensate wet gas zone and the dry gas zone (Fig.2). The increasing rate of the temperature used for the experiments may impact thermal maturation results. To avoid any effect other than the experimental temperature and duration of the experiments on the results, the same temperature increasing rate was used for the long and short-duration experiments as well as for each stage of maturity: 5.5°C.min⁻¹. At the end of the experiments, the oven was turned off and the autoclaves were slowly depressurized. After maturation, the gold cells were sealed in separate 25 cm³- glass bottles and pierced under vacuum. Approximate 2 cm³ of gases were sampled using a gas tight syringe and injected into a gas chromatography coupled with a thermal conductivity detector (GC-TCD). Gas composition was determined using a Perkin Elmer[®] Clarus 580 gas chromatograph (Supelco[®] Carboxen 1010 Plot[®] Capillary column: 30 m \times 0.53 mm i.d., 0.15 μm film thickness) using the same operating conditions described in Cavelan et al. (2019a). Individual gas components were identified by comparison with the relative retention times of

standard. Each gas content was quantified using its respective peak surface area and calibration curves by assuming ideal gas behavior.

Between 50 and 60 mg of dried and powdered samples were analyzed using a Rock-Eval® 6 pyrolyzer (Vinci Technologies, Rueil Malmaison). For the pyrolysis, samples were first isothermally heated for 2 min at 200°C under helium atmosphere. The oven temperature was then raised with a ramp of 30°C.min⁻¹ to 650°C and held for 3 min. For the oxidation, samples were isothermally heated under purified air at 400°C. The temperature was then raised to 850°C at 30°C.min⁻¹ and held 5 min. The parameters used in this study are the TOC (wt. %), the hydrogen index (HI, mgHC/g TOC) and the oxygen index (OI, mgCO₂/g TOC). The significance of these parameters was described by Espitalie et al. (1985a, b) and Lafargue et al. (1998).

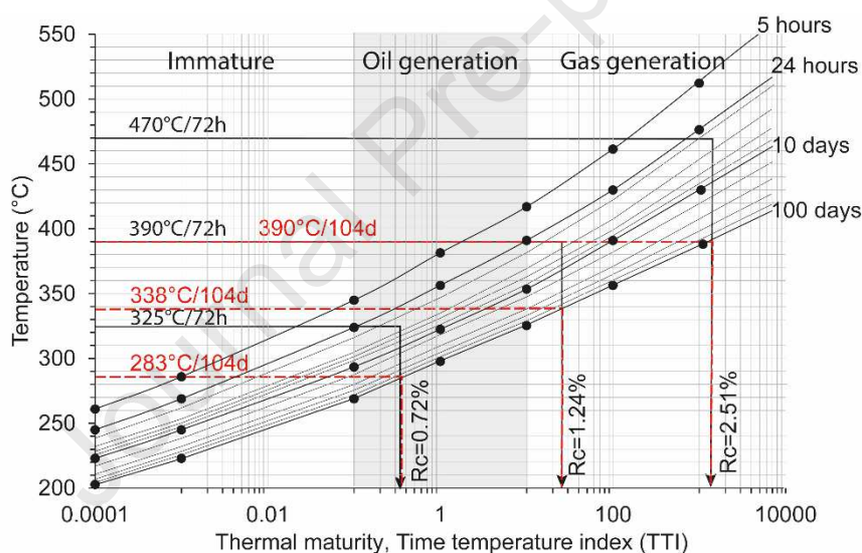


Fig. 2. Relation between maturity, temperature and duration of thermal maturations. In red: long-duration experiments (this study). In black: short-duration experiments and calculated vitrinite reflectance from Cavelan et al., (2019a). This chart was drawn from the data of Landais et al. (1994).

About 1g of powdered samples was then extracted with a solution of dichloromethane/methanol (1/1) using ultrasonication (10 min). The total extractible OM (EOM) was recovered by centrifugation, dried under nitrogen, weighed and desulfurized with copper shavings. The EOM was separated using deactivated silica gel (5% with water) chromatography column. The saturated hydrocarbons (SAT) were recovered by elution with heptane. The aromatic hydrocarbons

(ARO) were recovered by the subsequent elution of a mixture of heptane/toluene (3/1) and (2/2). The polar hydrocarbons were recovered by methanol elution. Each dried fraction was weighed. Asphaltenes remain trapped in the chromatography columns. The mass of asphaltenes was thus determined by difference in mass between the mass of the total EOM extract and the mass of the recovered fractions (SAT+ARO and polar). Some polar compounds (resins) can be trapped with asphaltenes in the columns. The mass of asphaltenes was thus counted with the mass of polar hydrocarbons to obtain the total mass of the NSO fraction. SAT and ARO were analyzed by gas chromatography-mass spectrometry (GC-MS) using a Trace-GC Ultra gas chromatograph equipped with a Thermo Trace-Gold 5 MS capillary column (60 m \times 0.25 mm i.d, 0.25 film thickness) and coupled to a TSQ Quantum XLS mass spectrometer (MS). The apparatus was fitted with an AS 3000 auto sampler (Thermo Scientific). The operating conditions were described in Cavelan et al., (2019a). The identification of each molecule presents in the SAT and ARO fractions was done by comparing retention times and mass spectra with previously published data. Their quantification was based on the integration of their peak area. Squalane was used as internal standard.

1.3. Pore volume, specific surface area and pore size distribution

The pore volume, the specific surface area and the pore size distribution (PSD) were determined by low-pressure nitrogen adsorption on a Micromeritics[®] ASAP -2420 apparatus. Before analysis, the samples (1g) were firstly crushed into powder (grain size: 180-250 μ m; 80-60 mesh size), oven-dried 48 hours at 105°C and outgassed at 65°C for 4 hours under vacuum (1 μ m Hg) in the apparatus. Adsorption and desorption isotherms were obtained at 77K. The relative pressure (p/p_0) ranges from 0.001 to 0.993 and an equilibration time of 10 s was applied during the analysis. Total pore volumes (V_{tot}) were determined after the liquid molar volume adsorbed at $p/p_0 = 0.993$. To compare the results of two common models, the PSD was determined using both the Barrett, Johner and Halenda (BJH) and the density functional theory (DFT) methods. The isotherms were treated with the DFT model using a slit-pores model based on a 'generic surface', which is an average of non-carbon materials listed for this type of geometry. The associated potential is an averaged potential of different non-carbon materials. This model was chosen by comparing the correspondence between the

Cylindrical model (using an Oxide surface or the Tarazona model) and the slit-pores model. The second model was selected due to a better match with the isotherms and the lowest RMS error. The regularization rate (Smooth or Lambda) was determined individually by the intersection of the 2 curves of "RMS error of fit" and "Roughness of distribution" for each sample but was of the order of 0.3 to 1 for each. An adjustment rate R of 0.01 was applied in order to eliminate the data out-of-model (zero point of the distribution). To avoid the tensile strength effect phenomenon that occurs during desorption, the PSD_{BJH} were determined from the adsorption isotherm (Barrett et al., 1951) between $p/p_0 = 0.042$ and 0.993. The Specific surface areas were calculated using the Brunauer, Emmett and Teller method (BET) for adsorption (Brunauer et al., 1938) between $p/p_0 = 0.011$ and 0.20. The external and micropore specific surface areas and pore volumes were determined using the Dubinin-Radushkevich model (DR) between p/p_0 ranging from 0.0001 and 0.01.

2. Results

2.1. Evolution of OM geochemical composition

After 104 days of thermal maturation at 282 to 390°C, the HI and the OI values follow the same evolution trend as those observed after 72 hours of maturation at 325°C to 470°C (Fig. 3). After maturation at 283°C, the mean calculated vitrinite reflectance (R_c) reached 0.69% ± 0.07 , indicating that the early oil generation stage had been reached (Tab. 1). All R_c values were confirmed by punctual vitrinite reflectance measurements taken at different stages of maturity. Hence, after thermal maturation, the measured vitrinite reflectance (R_o) is 0.71 % ± 0.04 . At this stage of maturity SAT+ARO hydrocarbon contents are relatively high (> 80 mg/g of TOC, > 45% of EOM) with only low concentrations of C_1 - C_5 and CO_2 (Tab. 1, Figs. 4, 5). The amount of SAT and ARO hydrocarbons increased from M2 to E40, as a function of their hydrogen index (HI, Tab.1, Fig. 5).

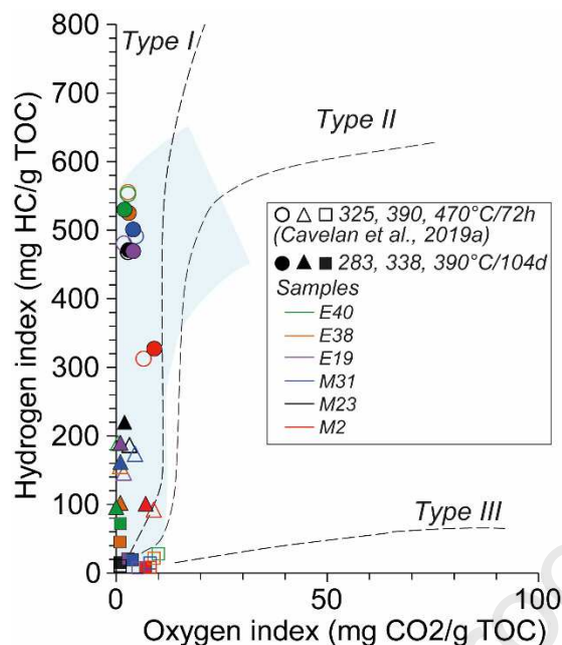


Fig. 3. Pseudo-Van Krevelen diagram showing the evolution of the hydrogen index and the oxygen index after 104 days (this study, filled symbols) and 72 hours (open symbols) of thermal maturation.

After heating at 338 and 390°C for 104 days the increase of the R_c to $1.22\% \pm 0.10$ and $2.17\% \pm 0.12$ (corresponding measured vitrinite reflectance is $1.30\% \pm 0.08$ and $2.14\% \pm 0.05$, respectively) indicates that the gas generation stages were reached (Tab. 1). These two stages are accompanied by a progressive and a strong increase in the concentration of CO_2 and C_1 - C_5 gaseous hydrocarbons (Fig. 5). C_2 - C_5 are the main gases generated after heating at 338°C, whereas C_1 is mainly produced after heating at 390°C for 104 days. The concentration of C_1 - C_5 is greater for samples M31, E38 and E40, which have higher HI values (Fig. 5, Tab. 1). On the contrary, the concentration of CO_2 is higher for samples M2, M23 and E19, which have lower HI values (Fig. 5, Tab. 1).

Tab. 1. Mean calculated vitrinite reflectance (R_c), total extractible organic matter (EOM), total organic carbon (TOC), Hydrogen index (HI) and biomarker ratios of Kimmeridge clay samples after 104 days of maturation.

Sample	T (C°)	TOC (wt. %)	HI (mgHC/g TOC)	EOM (mg/g TOC)	Mean R_c ¹ (%)	Dev	Pr/Ph	Pr/n-C ₁₇	Ph/n-C ₁₈	20S/(20S+R) ²	22S/(22S+R) ²
M2		3.0	327	150	*	*	1.07	1.02	0.63	0.29	0.50
M23		5.4	471	249	0.70	0.07	1.03	1.25	0.85	0.35	0.51
M31	283	6.8	501	365	0.71	0.09	1.42	0.75	0.98	0.29	0.53
E19		10.0	469	201	0.72	0.10	1.08	0.82	0.52	0.27	0.49
E38		14.8	525	257	0.67	0.06	1.37	0.65	0.60	0.36	0.48
E40		15.4	530	366	0.66	0.03	1.53	0.74	1.02	0.42	0.51
Mean					0.69	0.07	1.25	0.87	0.77	0.33	0.50

M2		2.03	101	122	1.20	0.11	0.57	0.13	0.09
M23		2.22	220	181	1.27	0.14	0.68	0.08	0.08
M31	338	4.05	162	172	1.22	0.09	0.76	0.09	0.13
E19		4.02	190	111	1.21	0.09	0.50	0.07	0.06
E38		13.27	102	116	1.24	0.12	0.59	0.04	0.07
E40		13.18	96	142	1.19	0.04	0.94	0.08	0.14
Mean					1.22	0.10	0.67	0.08	0.10
M2		2.35	9	35	*	*	0		
M23		3.72	19	12	*	*	0		
M31	390	3.93	23	87	*	*	0		
E19		8.47	34	27	*	*	0		
E38		5.44	46	19	2.15	0.13	0		
E40		5.02	72	76	2.19	0.10	0		
Mean					2.17	0.12			

Pr: pristane. *Ph*: phytane. *Unquantifiable ratios (very low concentrations). ¹see appendix 1. ²20S and 20R are epimers at C-20 in the C₂₉ 5 α ,14 α ,17 α ; 22S is C₃₂ 17 α 21 β 22S hopane and 22R is C₃₂ 17 α 21 β 22R hopane.

C₁₅-C₃₈ n-alkanes, accompanied by a few isoprenoids, methylated and alkylcycloalkanes (alkylcyclohexanes and pentanes), are predominant in samples up to their total degradation after heating at 390°C for 104 days (Fig. 5). Pr/n-C₁₇, Ph/n-C₁₈ and Pr/Ph ratios show a progressive decrease with increasing thermal maturity (Tab .1, Fig. 6A, B). Hopanes and steranes as well as triaromatic (TAS) and ring-C monoaromatic steroids (MAS) are quickly degraded during maturation and completely disappear after heating 104 days at 338°C (Fig. 5). 22S/(22S+R) 17 α 21 β 22R hopane and 20S/(20S+R) C₂₉ 5 α ,14 α ,17 α sterane ratios are > 0.30 after heating 104 days at 283°C corresponding to a maturity just below the peak of oil generation (Tab.1, Fig. 6C). Present in lower amounts, di-aromatic hydrocarbons dibenzothiophenes, naphthalenes and biphenyles) and heavier polyaromatic hydrocarbons (phenanthrenes, pyrenes and fluorethenes) and their methylated derivatives are present in the ARO (Fig. 5). The relative proportion of some of them with increasing maturity has allowed the calculation of the mean Rc (Tab. 1, appendix 1). The n-alkanes and the di-aromatic hydrocarbon concentrations are higher for samples with the higher OM oil generation potential (M31, E40 and E38) while the concentrations of the heavier polyaromatic components are higher for the less oil-prone samples (M2, M23 and E19, Fig. 5).

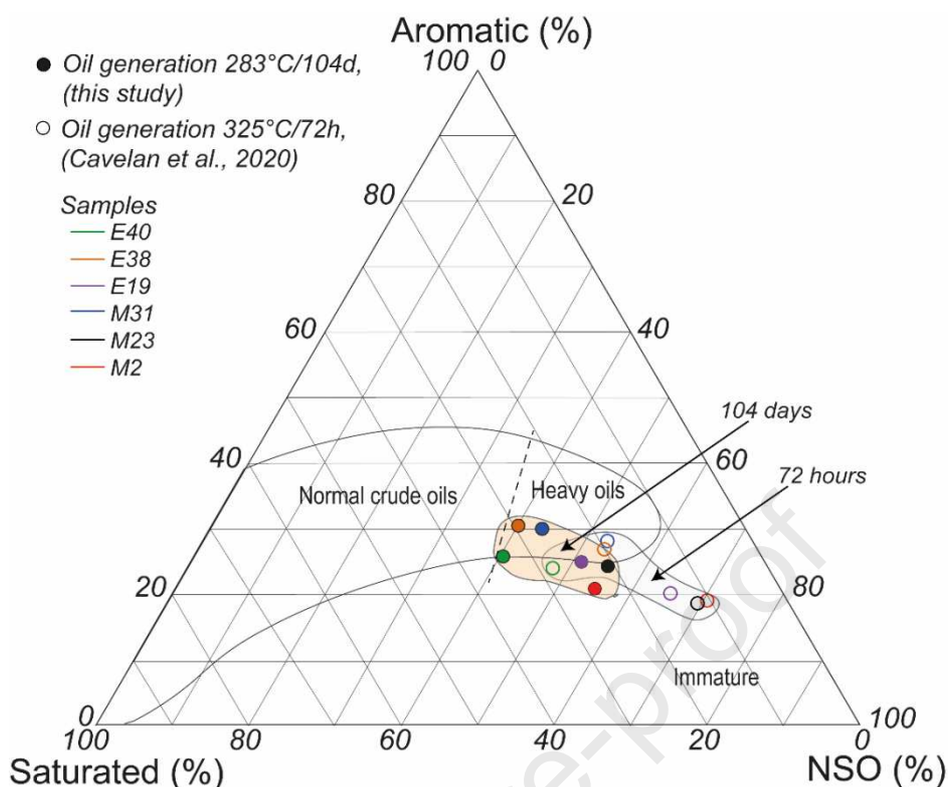
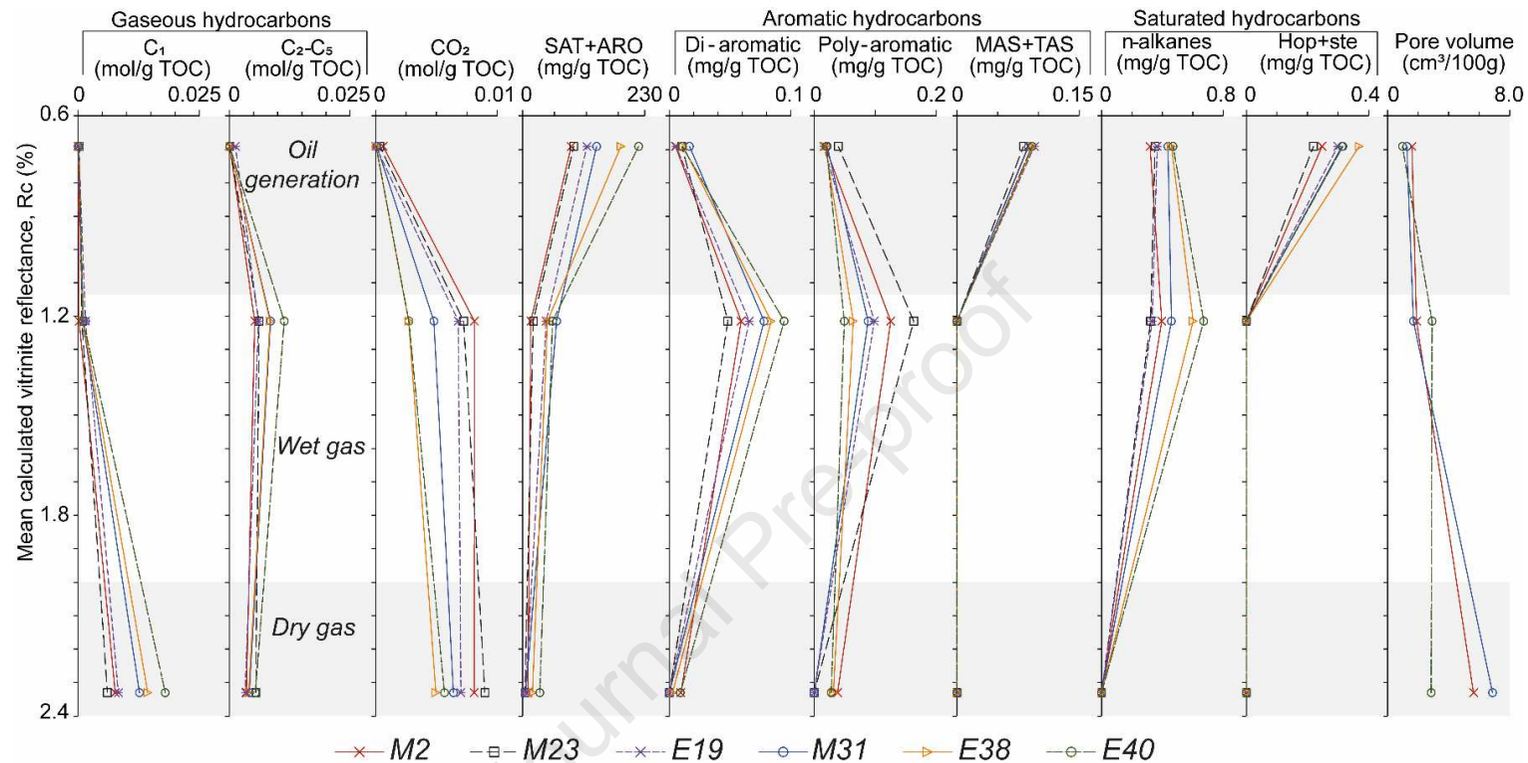


Fig. 4. Comparison of the relative proportion of saturated, aromatic hydrocarbons and NSO during oil generation (R_c of ca. 0.70%) after 104 days (filled symbols) and 72 hours (open symbols) of thermal maturation. Data of normal crude-oil from Tissot and Welte, (1984).

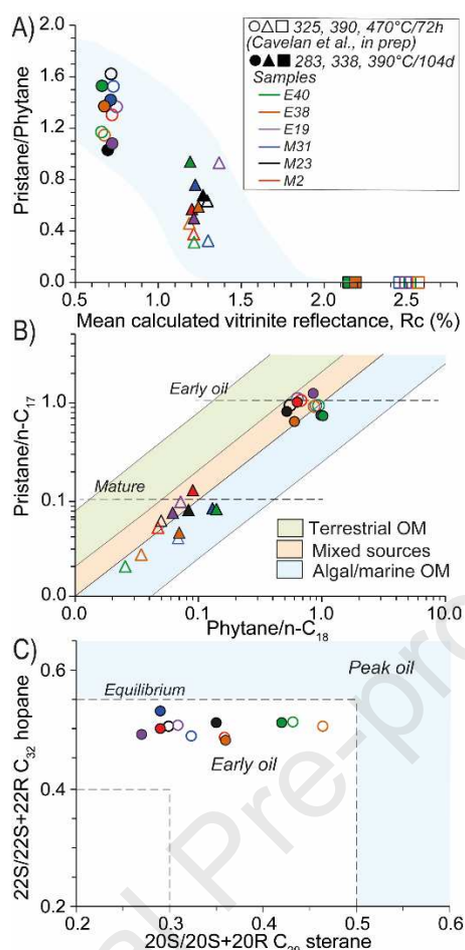
1.1. Comparison with shorter duration experiments

Comparable Rock Eval[®] parameters and mean R_c are obtained after maturation for 104 days and 72 hours (Fig. 3, 7A), indicating that similar stages of thermal maturity have been reached. The R_c obtained after maturation at 470°C for 72 hours (ca. 2.50%) is slightly higher than the mean R_c of 2.17% reached after maturation at 390°C for 104 days (Fig. 6A). However, the uncertainty of R_c values at this stage of maturity is greater because the concentrations of the aromatic components which served for its calculation are lower. This difference may be related only to this uncertainty attached to the calculation of the R_c . Note that the measured vitrinite reflectance values are similar between the two series: 2.12% \pm 0.10 for the short-duration experiments and 2.14% \pm 0.05 for the long-duration experiments. Similar biomarker ratios, including 20S/(20S+R), 22S/(22S+R), pristane/phytane (Pr/Ph), Pr/n-C₁₇ and Ph/n-C₁₈ ratios are obtained after maturation for 104 days and 72 hours (Fig. 6A-C).



1 —x— M2 —□— M23 —x— E19 —○— M31 —▶— E38 —○— E40

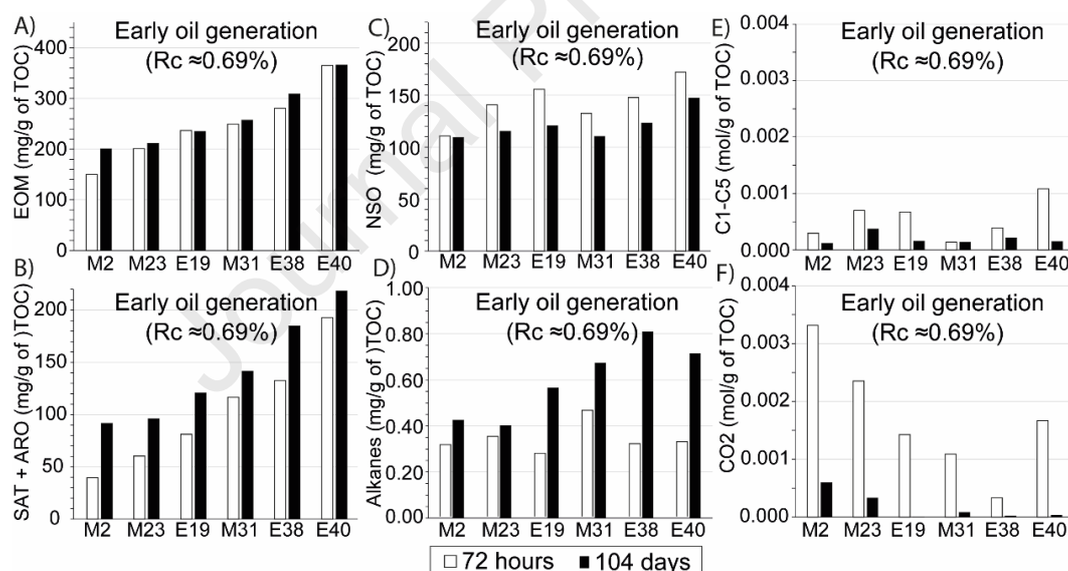
2 Fig. 5. Evolution of the concentrations of gas, aromatic (ARO), saturated (SAT) hydrocarbons, n-alkanes (including alkylcycloalkanes and isoalkanes),
 3 hopanes (Hop), sterane (ste), monoaromatic (MAS), triaromatic steroids (TAS), di-aromatic (dibenzothiophenes, naphthalenes, biphenyles) and poly-aromatic
 4 hydrocarbons (phenanthrenes, pyrenes, fluorethenes) after 104 days of maturation at 283°C, 338°C and 390°C. The division of hydrocarbons generation
 5 stages (in black) is based on Rc values, Rock Eval parameters, biomarker ratios and corresponding data from the literature (Fig. 2, Landais et al., 1994).



6
 7 Fig. 6. Evolution of A) pristane/phytane as a function of the mean calculated vitrinite reflectance, B)
 8 pristane/ $n-C_{17}$ versus phytane/ $n-C_{18}$, C) $22S/(22S+22R)$ $17\alpha 21\beta$ C_{32} hopane versus $20S/(20S+20R)$
 9 $5\alpha,14\alpha,17\alpha$ C_{29} sterane after 104 days (in red) and 72 hours of thermal maturation (black).

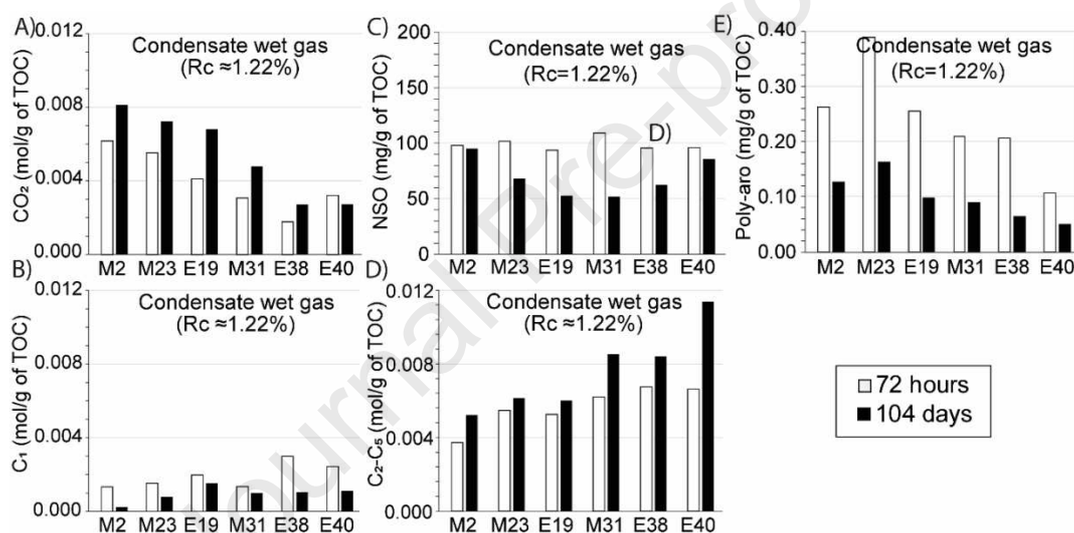
10 In accordance with the R_c values, the similarity of the biomarker ratios between the two series of
 11 laboratory thermal maturations suggests that similar stages of maturity have been reached after the
 12 long and the short-duration experiments. The $20S/(20S+R)$ value of ca.0.33 and $22S/(22S+R)$ value of
 13 ca.0.50 obtained after thermal maturation at 283°C for 104 days are typical values of the early oil
 14 generation stage (Fig. 6C) (Mackenzie et al., 1980; Peters et al., 2007; Seifert and Moldowan, 1981).
 15 Whatever the duration of the maturation, samples M31, E40 and E38, with the higher oil generation
 16 potential ($HI > 500$ mgHC/g of TOC, Tab. 1), still contain slightly lower Pr/ $n-C_{17}$ ratios for relatively
 17 high Ph/ $n-C_{18}$ ratios. This indicates the presence of a less oxygenated OM (Tab. 1, Fig. 6B). As for the
 18 shorter experiments (Cavelan et al., 2019a), samples M31, E38 and E40 (higher HI values) generated
 19 higher amounts of SAT+ARO hydrocarbons enriched in alkanes, di-aromatic hydrocarbons and C_1-C_5

20 gaseous hydrocarbons, while M2, M23 and E19 (lower HI values) generated the highest amount of
 21 CO₂, lower contents of SAT+ARO richer in heavier polyaromatic components (Fig. 5). However,
 22 some differences are observable between the longer and the shorter maturation experiments. While the
 23 total concentrations of EOM generated in the early oil generation stage are similar between the two
 24 maturation series, samples matured for 104 days exhibit lower NSO, CO₂, C₁-C₅ and slightly higher
 25 SAT, ARO, and alkane concentrations (including n-alkanes but also branched alkanes, methylalkanes,
 26 isoprenoids and alkylcycloalkanes) than samples matured for 72 hours (Figs. 4, 7). The relative
 27 proportions of SAT, ARO and NSO after 104 days of maturation are closer to the composition of the
 28 natural crude oils from Tissot and Welte, (1984) (Fig. 4). The difference in alkane concentrations is
 29 more pronounced for samples with the higher OM oil generation potential (IH >500 mgHC/g TOC):
 30 M31, E38 and E40 (Fig. 7). Similar concentrations of poly- and di-aromatic hydrocarbons are
 31 obtained.

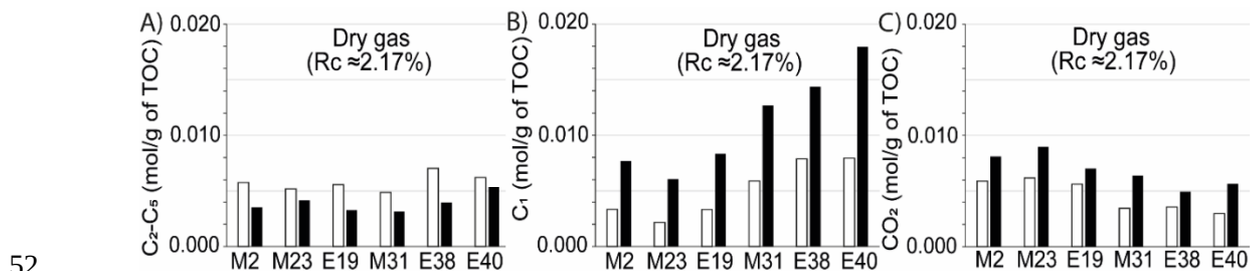


32
 33 Fig. 7. Comparison of A) extractible organic matter (EOM), B) saturated (SAT), aromatic (ARO)
 34 hydrocarbons, C) NSO, D) alkanes (hopanes, steranes, n-alkanes, branched alkanes, isoprenoids and
 35 alkylcycloalkanes), E) C₁-C₅ gaseous hydrocarbons and F) CO₂ concentrations in the early oil
 36 generation stage after 104 days and 72 hours of thermal maturation. The division of hydrocarbons
 37 generation stages is based on Rc values, Rock Eval parameters, biomarker ratios and Fig.2.

38 In the condensate wet gas zone, the increase in the maturation duration resulted in the
 39 production of higher C_2 - C_5 concentrations for samples with the higher oil generation potential (M31-
 40 E40), lower C_1 and lower CO_2 concentrations (Fig. 8). Comparable alkane and di-aromatic
 41 hydrocarbon concentrations are obtained in the condensate wet gas zone after maturation for 104 days
 42 and 72 hours. However, significantly lower concentrations of poly-aromatic hydrocarbons are
 43 observed after maturation for 104 days (Fig. 8E). The concentration of polyaromatic hydrocarbons
 44 decreases from the sample M2 and E40 as a function of the initial oil generation potential of the
 45 samples (Fig.8E). Lower NSO concentrations are obtained for most of the samples from the longer
 46 duration experiment (Fig. 8C).



47
 48 Fig.8. Comparison of the concentrations of A) CO_2 , B) C_1 , C) NSO, D) C_2 - C_5 gases and E) selected
 49 polyaromatic hydrocarbons (phenanthrenes, pyrenes and fluorethenes) in the condensate wet gas
 50 zone after 104 days and 72 hours of thermal maturation. The division of hydrocarbons generation
 51 stages is based on Rc values, Rock Eval parameters, biomarker ratios and Fig.2.



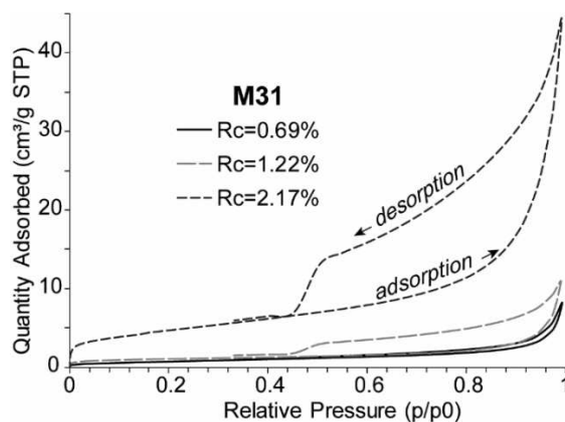
52
53 Fig. 9. Comparison of A) C₂-C₅, B) C₁ and C) CO₂ concentrations in the dry gas zone after 72 hours
54 (Cavelan et al., 2019a) and 104 days (this study) of thermal maturation. The hydrogen index increases
55 from samples B2 to E40. The division of hydrocarbons generation stages is based on Rc, Rock Eval
56 parameters, biomarker ratios and corresponding data from the literature.

57 In the dry gas zone, the increase in the duration of maturation allows the production of higher C₁
58 concentrations and still higher concentrations of CO₂ (Fig. 9). C₂-C₅ are, however, present in slightly
59 lower amounts after maturation for 104 days in the dry gas zone (Fig. 9A). The concentration of
60 residual NSO, poly- and di-aromatic hydrocarbons are too low at this stage of maturity to allow a good
61 comparison between the two series. The concentration of C₁ increases from M31 to E40 as a function
62 of the initial oil generation potential of the OM (Fig.9B) while the concentration of CO₂ decreases
63 (Fig.9C).

64 1.2. Evolution of pore volume and surface area after 104 days of maturation

65 1.2.1. Isotherms

66 An example of the evolution of nitrogen adsorption and desorption isotherms at 77K with
67 increasing maturity is presented in Fig. 10. Whatever their initial composition and maturity, all
68 samples show type IV isotherms, indicating the presence of mainly mesoporous materials (Sing,
69 1985). The shape of the hysteresis loops indicates the presence of pores with slit-shaped associated
70 with aggregated plate-like particles (Sing, 1985). However, in natural materials such as shale rocks,
71 previous studies (Clarkson et al., 2013; Schmitt et al., 2013) revealed that the assumption of slit-
72 shaped pores could be subjected to error, and thus has to be considered with caution.



73

74 Fig. 101. Example of the evolution of KCF samples' nitrogen gas adsorption and desorption isotherms
 75 (at 77K) with increasing thermal maturity.

76 The volume of gas adsorbed around $p/p_0 = 0.992$ is low after maturation at 283°C for 104 days
 77 ($R_c=0.69\%$, Fig. 10). After maturation at 338°C and 390°C for 104 days, the volume of gas adsorbed
 78 increases progressively, indicating the increase in sample pore volumes.

79 1.2.2. Pore volumes, specific surface areas and pore size

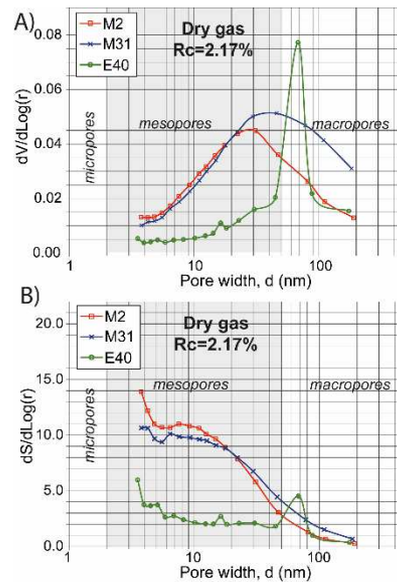
80 The total pore volumes (V_p) and the BET specific surface areas (S_{BET}) of samples are low after heating
 81 at 283°C for 104 days (Tab. 2). At this maturation stage, the V_p (≤ 1.59 cm³/100g) and the S_{BET} (≤ 9.31
 82 m²/g) decrease from M2 to E40, and thus decrease with the increase in the initial OM oil generation
 83 potential. The PSD determined using the DFT and the BJH methods give similar results (Figs. 11, 12)
 84 and show that all KCF samples exhibit a large pore size distribution mainly ranging from meso to
 85 macropores (Fig. 12) after maturation at 283°C for 104 days. The BJH method was used to
 86 characterize the mesoporosity, the DFT was used to determine the mesoporosity and part of the
 87 microporosity, and the DR was used to assess the microporosity. A comparison of the surface areas
 88 and the pore volume calculated with these different models with the S_{BET} and the V_p enabled the
 89 relative contribution of the meso and the micropore surface area and pore volume of each sample to be
 90 estimated.

91 Tab. 2. Total pore volumes (V_p), BET specific surface areas (S_{BET}), surface areas (S_{BJH}) and pore
 92 volumes (V_{BJH}) determined using the Barrett, Johner and Halenda method (BJH) (adsorption

93 isotherms) and the density functional theory (DFT), micropore volumes (V_{micro}) and micropore surface
 94 areas (S_{micro}) determined using the Dubinin-Radushkevich models (DR) for three representative
 95 samples after 104 days of thermal maturation.

Sample	T (°C)	V_p (cm ³ /100g)	S_{BET} (m ² /g)	S_{BJH} (m ² /g)	V_{BJH} (cm ³ /100g)	$S_{\text{BJH}}/$ S_{BET}	S_{DFT} (m ² /g)	V_{DFT} (cm ³ /100g)	$S_{\text{DFT}}/$ S_{BET}	$S_{\text{micro, DR}}$ (m ² /g)	$V_{\text{micro, DR}}$ (cm ³ /100g)	$S_{\text{DR}}/$ S_{BET}	$V_{\text{DR}}/$ V_t (%)
M2		1.59	9.31	3.1	1.2	0.35	5.92	1.49	0.64	9.99	0.35	1.13	22.3
M31	283	1.26	2.64	1.7	1.19	0.64	1.55	1.12	0.59	2.02	0.07	0.77	5.7
E40		0.97	1.89	1.24	0.92	0.69	1.05	0.95	0.55	1.18	0.04	0.65	4.3
M2		1.89	3.76	2.63	1.82	0.7	2.46	1.76	0.65	4.18	0.15	1.11	7.9
M31	338	1.69	3.6	2.04	1.62	0.57	2.23	1.55	0.62	4.02	0.14	1.12	8.5
E40		2.91	6.61	3.33	2.72	0.5	3.9	2.74	0.59	5.57	0.2	0.84	6.8
M2		5.62	19.5	18.22	5.08	0.94	14.36	5.25	0.74	20.92	0.74	1.33	13.2
M31	390	6.87	16.9	11.21	6.49	0.67	11.36	6.47	0.67	16.56	0.59	0.97	8.6
E40		2.85	5.36	3.83	2.75	0.72	3.04	2.81	0.57	3.97	0.14	0.74	4.9

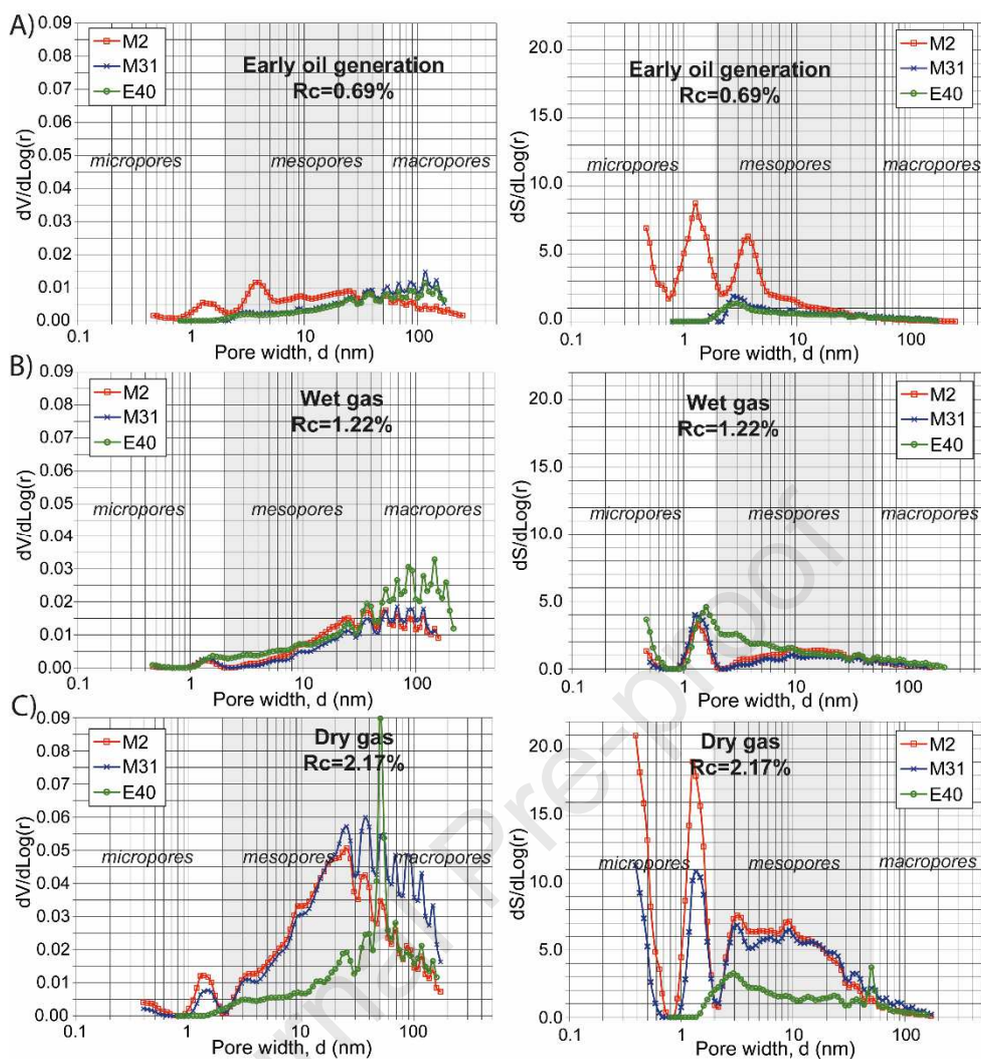
96
 97 As shown by the pore volume determined using the BJH (V_{BJH}) close to the V_p , most of the pore
 98 volume is provided by meso and macropores in the studied samples (Tab. 2). However, contrary to the
 99 other two samples, the lower $S_{\text{BJH}}/S_{\text{BET}}$ ratio, far from 1.0, the slightly higher $S_{\text{DFT}}/S_{\text{BET}}$ ratio and
 100 especially the higher $S_{\text{DR}}/S_{\text{BET}}$ ratio close to 1 after heating at 283°C indicate that most of the specific
 101 surface area of the sample M2 is provided by micropores (Tab. 2). The V_{DR}/V_t ratio indicates that
 102 micropores account for about 22% of the total pore volume in this sample and less than 6% in the
 103 other two samples. This is well substantiated by the PSD from the DFT methods, which showed the
 104 presence of two controlled-size porosities about 1.3 and 4 nm in diameter in M2 (Fig. 12). These
 105 porosities have a significant contribution to the total specific surface area after maturation at 283°C for
 106 104 days.



107
 108 Fig. 11. BJH A) pore size and B) surface area distributions of samples M2, M31 and E40 after thermal
 109 maturation at 390°C for 104 days.

110 After heating at 338°C for 104 days, the V_p , the $V_{BJH, ads}$ and the V_{DFT} increase (Tab. 2, Fig. 5). While
 111 the S_{BET} increases for M31 and E40, S_{BET} strongly decreases for M2 due to the decrease in its number
 112 of micropores around 1.3 nm in diameter after maturation at 338°C (Tab. 2, Fig. 12). E40, the highly
 113 oil-prone sample, exhibits higher V_p and S_{BET} than M2 and M31, which show similar features (Tab. 2).
 114 The differences between samples are less significant at this thermal maturity. All samples exhibit a
 115 similar PSD, with a predominance of mesopores ranging mainly from 4 to 150 nm in diameter (Fig.
 116 12).

117 As shown by the V_{BJH} close to the V_p , these mesopores account for most of the pore volume
 118 after maturation at 338°C. The PSD from the DFT, however, shows the presence of a controlled size
 119 microporosity around 1.3 nm with a relatively low contribution to the pore volume (V_{DR}/V_t of ca. 8%,
 120 Tab. 2, Fig. 12) but a significant contribution to the specific surface area. This feature may explain the
 121 S_{DR} / S_{BET} ratio close to 1 in all samples and the S_{DFT} / S_{BET} and S_{BJH} / S_{BET} ratios far from 1.0 (Tab. 2).
 122 As shown by the V_{DR}/V_t (%), these micropores account for about 8% of the pore volume of KCF
 123 samples after maturation at 338°C for 104 days. After maturation at 390°C for 104 days, the V_p and
 124 the S_{BET} increase strongly except for E40 for which these values remain relatively stable (Tab. 2, Fig.
 125 5).



126

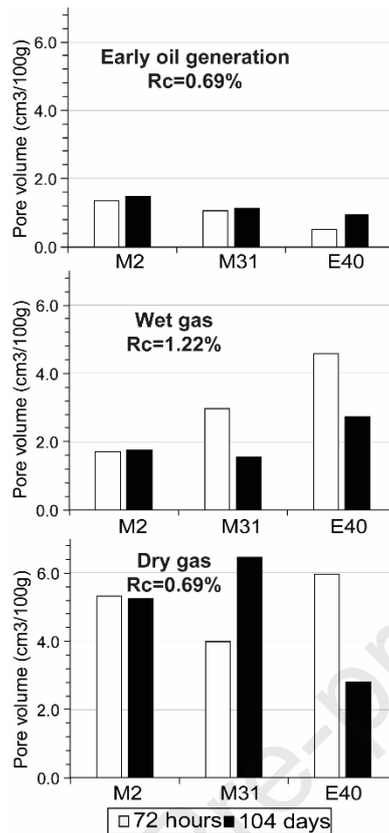
127 Fig. 12. DFT pore size (left) and surface area distribution (right) of samples M2, M31 and E40 after
 128 thermal maturation at A) 283°C, B) 338°C and C) 390°C for 104 days.

129 The V_p and the S_{BET} thus decrease with the increase in the initial TOC and OM oil generation
 130 potential. The V_{BJH} still close to the V_p indicate that the total pore volume of samples is still mainly
 131 provided by meso and macropores (Tab. 2), widely distributed between 2 and 150 nm after maturation
 132 at 390°C for 104 days (Figs. 11, 12). The PSD from the DFT, however, shows the increase in the
 133 number of pores at ≈ 1.3 nm and 4 nm for M2 and M31 (Fig. 12). While these micro and small
 134 mesopores make a relatively low contribution to the total pore volume (V_{DR}/V_t ratio between 8.6% and
 135 13.2%), they provide a significant part of the specific surface area marked by a high DR micropore
 136 surface area (S_{DR}) (Fig. 12, Tab. 2). The PSD from the DFT method suggest that micropores less than

137 0.4 nm in diameter with a high contribution to the specific surface area are present after heating at
138 390°C for 104 days but are not quantifiable by nitrogen adsorption.

139 **1.2.3. Comparison of porosity evolution with shorter duration experiments.**

140 Similar trends in pore volume evolution are observed after 104 days (Fig. 5, this study) and 72 hours
141 (Cavelan et al., 2019a) of thermal maturation. The total pore volume, very low in the early oil
142 generation stage, increases progressively to the dry gas zone (Fig. 5). While no real differences are
143 observed during oil generation, some slight differences are, however, visible between the two series
144 during gas generation (Fig. 13). Slightly lower pore volumes are observed in the condensate wet gas
145 zone. Increasing the duration of thermal maturation does not allow the formation of a greater pore
146 volume at this stage of maturity. In the dry gas zone, the relation between the pore volume and the
147 thermal maturation duration is not clear. While M2 exhibits a similar pore volume after maturation for
148 72 hours and 104 days, M31 exhibits a higher pore volume and E40 a lower pore volume after
149 maturation for 104 days (Fig. 13). Despite the fact that all samples exhibit a similar porosity mainly
150 composed of widely distributed mesopores and macropores, differences in PSD are observed after 104
151 days (Fig. 12) and 72 hours of thermal maturation (Cavelan et al., 2019a), especially in the condensate
152 wet gas zone. Contrary to the results from Cavelan et al. (2019a), where highly oil prone samples
153 contain mainly pores < 10 nm and less oil prone samples contain pores > 10 nm, less important
154 differences of PSD are observed between our samples after 104 days of maturation (Fig. 12). The
155 pores are mainly comprised between 1 and 80 nm in diameter.



156

157 Fig. 13. Comparison of total pore volumes after 104 days (this study) and 72 hours (Cavelan et al.,
 158 2019a) of thermal maturation at different maturities.

159 2. Discussion

160 2.1. How useful is it to combine different models for nitrogen adsorption measurements?

161 The concomitant use of different models for nitrogen adsorption measurements, such as the DFT and
 162 the DR, enables to learn more about the pore structure of KCF rocks. In accordance with BJH pore
 163 size distribution and previous work, DFT pore size distributions show that most of the remaining pore
 164 volume in KCF rocks is provided by meso and macropores which vary widely in diameter from 2 to
 165 160 nm in the early oil generation stage. However, the PSD curves from the DFT, which are more
 166 precise for microporosity and small mesoporosity assessment, reveal the presence of two controlled-
 167 size porosities at about 1.3 and 4 nm in diameter, only visible in the sample with the lowest oil
 168 generation potential in the early oil generation stage (M2, Fig. 12). These two controlled-size
 169 porosities are also observed in the condensate wet gas zone and are clearly visible in M2 and M31 in
 170 the dry gas zone (Fig. 12). According to the shape of the isotherms hysteresis loops (Fig. 10), these

171 micro and small regular mesoporosity could correspond to slit-shaped pores associated with the plate-
172 like clay particle aggregates previously observed in KCF rocks using SEM imaging (Cavelan et al.,
173 2019b). While these small meso and micropores generally make a limited contribution to the pore
174 volume, they provide a high specific surface area (Fig. 12) at the origin of the higher total BET and
175 DR specific surface area of M2. Their role in gas storage and adsorption phenomena should thus not
176 be neglected. In previous SEM images, the phytoplanktonic orange AOM, abundant in oil-prone
177 rocks, appeared as a 'gel' which seems to fill the interlayer space between minerals, reducing the
178 mineral associated pore volume of certain samples before maturation (Cavelan et al., 2019b). We can
179 thus suggest that M31 and E40, which initially contain significantly higher amounts of orange AOM
180 than M2 (Cavelan et al., 2019a), exhibit initially a lower micro and mesoporosity associated with clay
181 sheet aggregates. Then, the filling of the porosity of these two higher oil prone samples by effluents
182 during oil generation makes the rest of these pores inaccessible to nitrogen adsorption. Hence their
183 lower porosity at 1.3 and 4 nm in diameter during oil generation (Fig. 12). During the progressive
184 conversion of residual OM and oil into gaseous hydrocarbons and the ensuing generation of secondary
185 pores in OM, some micro and mesopores associated with clay minerals may be progressively released
186 and/or become more accessible to nitrogen adsorption. This probably explains their increasing
187 contribution to the pore volume and the specific surface area (increase in $S_{\text{micro, DR}}$ and $V_{\text{micro, DR}}$) from
188 the condensate wet gas to the dry gas zone in samples M2 and M31 (Tab. 2, Fig. 12). Indeed, the $S_{\text{DR}} /$
189 $S_{\text{BET}} \geq 1$ in M2 and M31 indicates that most of the specific surface area is provided by micropores
190 (Tab. 2). In the sample E40, a more extensive collapses/alteration of the pore structure in response to
191 its important gas generation may have limited the accessibility of these pores to nitrogen adsorption,
192 explaining the absence of pores 1.3 nm in diameter in the dry gas zone (Fig. 12). Although this
193 relatively scarce porosity associated with clay sheet aggregates makes a small contribution to the total
194 pore volume, it is a major contributor to the specific surface area and may have a significant ability to
195 adsorb and retain gas. These pores could thus also contribute to the gas storage capacity of these
196 source rocks.

197 **2.2. Effect of the duration of confined thermal maturation on OM thermal degradation**

198 As previously demonstrated (Landais et al., 1994; Wood, 1988), different time-temperature pairs can
199 thus be used to reach the same stage of maturity. As expected, comparable Rc and thermal maturity
200 biomarker ratios are obtained after thermal maturation for 72 hours (Cavelan et al., 2020) and 104
201 days, indicating indeed that similar stages of maturity have been reached: the early oil generation (Rc
202 ca. 0.70%), the condensate/wet gas (Rc ca. 1.23%) and the dry gas zone (Rc: 2.2-2.5%) (Tab. 1, Fig.
203 6). It is thus possible to compare the bulk, the molecular composition of OM and the porosity between
204 the two series as a function of the duration of maturation experiments. Our results show that Pr/Ph,
205 Pr/n-C₁₇, Ph/n-C₁₈ ratios and the relative proportion of the different hopane and sterane configurations
206 do not depend on the duration of the maturation (Tab. 1, Fig. 6). The Rock Eval[®] parameters do not
207 exhibit variations related to the duration-temperature pairs chosen for the experiments (Fig. 3). This
208 well substantiates the results of Landais et al. (1994) on type II source rocks and OM isolated from the
209 Paris basin. Similar evolution of EOM, CO₂ and C₁-C₅ concentrations and comparable OM
210 components with similar distributions are observed in SAT and ARO fractions for long (104 days)
211 (Fig. 5) and short (72 hours) maturation experiments. Furthermore, the choice of the time-temperature
212 pairs does not influence the differences observed between samples with various initial oil generation
213 potential. In both cases, the concentration of EOM (especially of SAT and ARO), alkanes and di-
214 aromatic hydrocarbons in the early oil generation stage and thus the concentration of C₁-C₅
215 hydrocarbon gases during gas generation increase with the OM oil generation potential (Fig. 5). These
216 longer maturation experiments thus confirm the influence of the slight variations in the initial OM
217 composition of KCF mudstones on the amount and the composition of EOM generated, as previously
218 discussed (Cavelan et al., 2020, 2019a).

219 However, some differences are visible between the two series during oil and gas production.
220 Shorter duration experiments (72 hours) appear to favor the production of EOM enriched in NSO,
221 while long-duration experiments (104 days) seem to favor the production of EOM enriched in SAT
222 and, to a lesser extent, ARO hydrocarbons in the early oil generation stage (Figs. 4, 7). The relative
223 proportion in EOM of SAT and ARO hydrocarbons appears thus closer to the composition of natural
224 crude oils (Fig. 4). Hence, residual EOM exhibit lower NSO and especially lower polyaromatic

225 concentrations such as phenanthrenes, pyrenes or fluorethenes during gas generation stages (Fig. 8).
226 The increase in the maturation duration, and thus the increase in the residence time in the thermal
227 maturation system, appears to enhance the degradation of heavy NSO components into SAT and ARO
228 hydrocarbons. This probably explains the higher alkane concentrations in the SAT fraction after the
229 long-duration experiment in the early oil generation stage (Fig. 7). Similar results were previously
230 found on type II Paris Basin source rocks and type III Mahakam coal (Landais, 1991; Landais et al.,
231 1994). On the contrary, lower C₁-C₅ concentrations are generated in the early oil generation stage after
232 the long- duration experiment (Fig. 7). As previously suggested by Landais et al. (1994), the lower
233 temperature used for the long-duration experiment (283°C for 104 days) probably led to a less
234 pronounced early cracking of OM into light components by preventing low energy bonds from being
235 broken during the early stage of maturation. This may also explain the slightly lower C₁ concentration
236 for the longer duration experiment in the condensate/wet gas (Fig. 8). This lower early transformation
237 of OM then leads to a more efficient secondary cracking of OM at higher maturity stages. This results
238 in higher C₂-C₅ concentrations in the condensate wet gas zone and to a more efficient conversion of
239 C₂-C₅ into C₁ in the dry gas zone responsible for the lower remaining C₂-C₅ concentrations and the
240 significantly higher C₁ concentration of the longer maturation experiment (Figs. 8, 9). Note, however,
241 that contrary to the Paris basin source rocks, where CO₂ concentrations did not appear to be influenced
242 by these processes (Landais et al., 1994), increasing the duration of the experiments in the KCF also
243 seems to lead to the production of lower CO₂ concentrations in the early oil generation stage but
244 higher concentrations during gas generation (Figs. 7-9). Nevertheless, the difference in CO₂
245 concentrations between the two series seems to depend on the initial OM composition of KCF
246 samples, especially during condensate wet gas generation. This difference, which is significant for the
247 less oil-prone samples (M2 to E19), does not exist in the two highly oil-prone samples (E38 and E40,
248 Fig. 8). This suggests that the effect of the duration-temperature pair choice on OM thermal
249 degradation may also slightly depend on the original OM composition and oil generation potential.

250 After the long-duration experiments, the concentrations of C₂-C₅ are still very high in the dry gas
251 zone compared to the concentrations of C₁ (Fig.5). All biomarker ratios and Rc values (verified by

252 punctual vitrinite reflectance measurements) indicate that the dry gas zone has been reached after
253 maturation at 390°C for 204 days. However, based on gas concentrations alone, the production and the
254 composition of gaseous hydrocarbons are closer to those of the middle/end of the condensate/wet gas
255 zone. These differences were also visible for short-duration experiments, but were more pronounced
256 (Cavelan et al., 2020). The conversion of C₂-C₅ gaseous hydrocarbons into methane was delayed
257 during the short and the long-duration experiments, leading to a discrepancy between the predicted
258 maturity (R_c, biomarkers) and the real thermal maturity of the OM. Despite a significant increase in
259 thermal maturation duration, the composition of the gaseous hydrocarbons generated by laboratory
260 experiments remains somewhat different from those obtain under low temperature, after long-duration
261 geologic maturations. Moreover, artificial thermal maturations were carried out on powdered samples
262 and under an external hydrostatic confining pressure. This cannot accurately reproduce the effect of
263 the lithostatic pressure that affect the structure of the rock in sedimentary basins. During laboratory
264 thermal maturations, the pressure is transmitted into the rock by the effluents (water, hydrocarbons and
265 gases). The relation between porosity, oil expulsion and retention processes is thus not accurately
266 simulated. This may impact both the OM thermal degradation and the development of pores.
267 Nevertheless, the pore volumes obtained during artificial maturations (ranging from 0.97 cm³/g to
268 6.87 cm³/g) are very consistent with those of naturally-matured organic-rich shales. For example,
269 Mastalerz et al. (2013), Milliken et al. (2013), Pan et al. (2015), Song et al. (2019), among many
270 others, found similar porosity and pore volumes in their naturally-matured samples for similar
271 maturities. Artificial thermal maturation remains therefore a good technique to study the porosity of
272 shales at different thermal maturities, despite the differences with regard to natural sedimentary basins.

273 **2.3. Does the duration of confined thermal maturation affect the evolution of porosity?**

274 Similar variations in pore volume are observed from the early oil generation to the dry gas zone after
275 104 days (this study, Fig. 5) and 72 hours of thermal maturation (Cavelan et al., 2020, 2019a). Total
276 pore volumes and specific surface areas are very low during early oil generation (probably
277 filled/occluded by oil) and then exhibit a progressive increase (probably due to the formation of pores
278 in OM), which depends on the initial OM oil generation potential (Fig. 5). The more a sample has

279 produced hydrocarbons (especially light saturated compounds) able to migrate and fill the adjacent
280 porosity, the smaller the pore volume is during oil generation. This was observed during the short-
281 duration and the long-duration experiments. Indeed, samples with a higher OM oil-generation
282 potential (such as E40) have lost a higher volume of pores during oil generation, occluded by the
283 larger quantity of SAT and ARO generated (Tab. 2). The increase in maturation duration allows the
284 production of less viscous EOM enriched in SAT and ARO hydrocarbons, but this does not influence
285 the total amount of EOM generated during oil generation (Fig. 7). This probably explains why similar
286 BJH pore size distributions and similar pore volumes are observed after 72 hours and 104 days of
287 maturation in the early oil generation stage (Fig. 13). The slight differences of SAT, ARO and NSO
288 relative proportions between the two series appear thus to have a limited effect on pore occlusion
289 processes during the early stages of maturation. This well substantiates the hypothesis that a link exists
290 between the OM composition, its capacity to generate hydrocarbons and the evolution of porosity.

291 In most of the studies carried out on organic-rich shales, gas generation coincides with the
292 development of abundant pores in OM, in response to the production and the expulsion of gaseous
293 hydrocarbons (Bernard et al., 2012b; Curtis et al., 2012; Han et al., 2017; Ko et al., 2018; Loucks et
294 al., 2012; Song et al., 2019; Y. Wang et al., 2020; Wu et al., 2019, among many others). This
295 formation of pores was also observed in the KCF after short-duration thermal maturation experiments
296 (Cavelan et al., 2020, 2019a). However, these results showed that the massive production of gaseous
297 hydrocarbons may lead to deformations of the OM structure that may cause the decrease of the pore
298 volume at the end of the condensate/wet gas zone, especially for the highly oil-prone samples (such as
299 E40) which generated higher concentrations of gaseous hydrocarbons. We can suggest that a more
300 unstable pore network, particularly sensitive to pressure-induced deformations (tied to the presence of
301 more abundant and well-connected gas devolatilization vacuoles), formed in the highly oil-prone rocks
302 in response to the massive generation of gaseous hydrocarbons. The increase in maturation duration
303 causes an increase in the amount of C₂-C₅ generated in the condensate wet gas zone, especially for
304 samples with a higher OM oil generation potential (such as E40, Fig. 8). These samples may thus have
305 suffered earlier or more pronounced alteration of their pore network in response to the more important

306 generation of gaseous hydrocarbons, leading to slightly lower pore volumes than for the shorter
307 duration experiments (Fig. 13). The less oil-prone sample M2 generates relatively similar amounts of
308 gaseous hydrocarbons after heating for 72 hours and 104 days (Figs. 8) and exhibits comparable pore
309 volume and PSD (pores widely distributed from 8 to 160 nm in diameter) in the condensate wet gas
310 zone. This provides support for this hypothesis (Fig. 13). A different timing and amplitude of the
311 alteration of the pore network in response to the different amounts of gaseous hydrocarbons generated
312 between the two series of maturation may explain the different PSD observed for the highly oil-prone
313 samples M31 and E40. After maturation for 72 hours, M31 and E40 are composed of meso and
314 micropores less than 40 nm (Cavelan et al., 2019a), while they contain mainly meso and macropores
315 more than 8 nm after heating for 104 days (Fig. 12). We can suggest that the lower temperature used
316 by the longer maturation experiments, which seems to prevent the lower energy bonds from being
317 broken early on, may result in a slower and more homogeneous gas generation throughout the OM
318 particles forming a better interconnected and distributed OM-hosted porosity in the condensate wet
319 gas zone. Despite frequent collapse/alteration events, this may have enhanced early pore coalescence
320 at the origin of the larger pore sizes after heating for 104 days.

321 In the dry gas zone, as previously observed after maturation for 72 hours, the specific surface
322 area and the pore volume of samples increase more or less as a function of the initial OM composition
323 and gas concentrations (Tab. 2). Moreover, the PSD indicates a comparable porosity mainly composed
324 of pores larger than 10 nm in the two series. As observed during the short-duration experiments, the
325 sample with the highest OM oil generation potential (E40) generated higher amounts of SAT and
326 ARO enriched in hydrogenated components (such as n-alkanes) and exhibits lower pore volume than
327 less oil prone samples in the dry gas zone (Fig. 13). This confirms that the pore volume formed during
328 gas generation is not linearly correlated with the OM oil generation potential. However, the slightly
329 different timing of generation and concentrations of C_1 - C_5 gaseous hydrocarbon has probably
330 influenced the ability of the rock to preserve its pore network from collapse or deformation events that
331 have affected the pore volume, probably causing the differences in pore volume between samples of
332 the two series of maturation. Our results show that increasing the duration of thermal maturation

333 experiments influences the composition of EOM, the amount and the timing at which gaseous
334 hydrocarbons are generated. Nevertheless, these differences, relatively small between 72h and 104
335 days of maturation, do not significantly affect the development and the evolution of KCF mudstones'
336 porosity with increasing maturity. Similar trends of pore volume and specific surface area are observed
337 with increasing maturity and similar variations of porosity are observed between samples with a
338 slightly different initial OM composition. The choice of duration-temperature pairs for thermal
339 maturation experiments seems thus to have a limited impact on the porosity of organic-rich source
340 rocks.

341 **Conclusions**

342 The comparison between short-duration (72h) and long-duration thermal maturations of organic-rich
343 mudstones shows that, for a same thermal maturity, increasing the duration of thermal maturation
344 experiments: (i) does not influence the amount of EOM generated; (ii) increases the residence time of
345 the effluents in the thermal maturation system, enhancing the degradation of heavy polar OM
346 components into saturated and aromatic hydrocarbons and leading to the production of EOM enriched
347 in saturated and aromatic components; (iii) has no significant impact on the evolution of KCF rocks'
348 porosity at the early oil generation stage, which is controlled more by the amount of effluents filling
349 the interparticle mineral associated porosity than by the composition of EOM.

350 In preventing the early breaking of low energy bonds, the lower temperature used for the
351 longer duration experiments results in a less pronounced early cracking of OM, resulting in a different
352 timing of gas generation, more gradual throughout the OM particles. This process does not influence
353 the global evolution of the pore volume during maturation. However, it seems to influence the
354 amplitude and the timing of pore alteration/deformation events that occur during gas generation,
355 leading to slightly different pore size distributions and pore volumes, especially in highly oil-prone
356 samples. The impact of the duration-temperature pairs of thermal maturation experiments on the
357 development of KCF mudstones porosity remains, however, limited, suggesting that results from
358 different time-duration experiments and natural samples can be easily compared.

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366 **References**

- 367 Barrett, E.P., Joyner, L.G., Halenda, P.P., 1951. The Determination of Pore Volume and Area
368 Distributions in Porous Substances. I. Computations from Nitrogen Isotherms. *Journal of the*
369 *American Chemical Society* 73, 373–380. <https://doi.org/10.1021/ja01145a126>
- 370 Behar, F., Lewan, M.D., Lorant, F., Vandenbroucke, M., 2003. Comparison of artificial maturation of
371 lignite in hydrous and nonhydrous conditions. *Organic Geochemistry* 34, 575–600.
372 [https://doi.org/10.1016/S0146-6380\(02\)00241-3](https://doi.org/10.1016/S0146-6380(02)00241-3)
- 373 Bernard, S., Horsfield, B., Schulz, H.-M., Wirth, R., Schreiber, A., Sherwood, N., 2012a. Geochemical
374 evolution of organic-rich shales with increasing maturity: A STXM and TEM study of the
375 Posidonia Shale (Lower Toarcian, northern Germany). *Marine and Petroleum Geology* 31,
376 70–89. <https://doi.org/10.1016/j.marpetgeo.2011.05.010>
- 377 Bernard, S., Wirth, R., Schreiber, A., Schulz, H.-M., Horsfield, B., 2012b. Formation of nanoporous
378 pyrobitumen residues during maturation of the Barnett Shale (Fort Worth Basin). *International*
379 *Journal of Coal Geology* 103, 3–11. <https://doi.org/10.1016/j.coal.2012.04.010>
- 380 Boreham, C.J., Crick, I.H., Powell, T.G., 1988. Alternative calibration of the Methylphenanthrene
381 Index against vitrinite reflectance: Application to maturity measurements on oils and
382 sediments. *Organic Geochemistry* 12, 289–294. [https://doi.org/10.1016/0146-6380\(88\)90266-](https://doi.org/10.1016/0146-6380(88)90266-5)
383 5
- 384 Boussafir, M., Gelin, F., Lallier-Vergès, E., Derenne, S., Bertrand, P., Largeau, C., 1995a. Electron
385 microscopy and pyrolysis of kerogens from the Kimmeridge Clay Formation, UK: source
386 organisms, preservation processes, and origin of microcycles. *Geochimica et Cosmochimica*
387 *Acta* 59, 3731–3747.
- 388 Boussafir, M., Lallier-Vergès, E., 1997. Accumulation of organic matter in the Kimmeridge Clay
389 Formation (KCF): an update fossilisation model for marine petroleum source-rocks. *Marine*
390 *and Petroleum Geology* 14, 75–83.
- 391 Boussafir, M., Lallier-Vergès, E., Bertrand, P., Badaut-Trauth, D., 1995b. SEM and TEM studies on
392 isolated organic matter and rock microfacies from a short-term organic cycle of the
393 Kimmeridge Clay Formation (Yorkshire, GB), in: *Organic Matter Accumulation*. Berlin,
394 Heidelberg, pp. 15–30.
- 395 Brunauer, S., Emmett, P.H., Teller, E., 1938. Adsorption of Gases in Multimolecular Layers. *Journal*
396 *of the American Chemical Society* 60, 309–319. <https://doi.org/10.1021/ja01269a023>
- 397 Cavelan, A., Boussafir, M., Le Milbeau, C., Laggoun-Défarge, F., 2020. Impact of Oil-Prone
398 Sedimentary Organic Matter Quality and Hydrocarbon Generation on Source Rock Porosity:
399 Artificial Thermal Maturation Approach. *ACS Omega*.
400 <https://doi.org/10.1021/acsomega.0c01432>
- 401 Cavelan, A., Boussafir, M., Le Milbeau, C., Rozenbaum, O., Laggoun-Défarge, F., 2019a. Effect of
402 organic matter composition on source rock porosity during confined anhydrous thermal

- 403 maturation: Example of Kimmeridge-clay mudstones. *International Journal of Coal Geology*
404 212. <https://doi.org/10.1016/j.coal.2019.103236>
- 405 Cavelan, A., Boussafir, M., Rozenbaum, O., Laggoun-Défarage, F., 2019b. Organic petrography and
406 pore structure characterization of low-mature and gas-mature marine organic-rich mudstones:
407 Insights into porosity controls in gas shale systems. *Marine and Petroleum Geology* 103, 331–
408 350.
- 409 Chalmers, G.R., Bustin, R.M., Power, I.M., 2012. Characterization of gas shale pore systems by
410 porosimetry, pycnometry, surface area, and field emission scanning electron
411 microscopy/transmission electron microscopy image analyses: Examples from the Barnett,
412 Woodford, Haynesville, Marcellus, and Doig units. *AAPG Bulletin* 96, 1099–1119.
413 <https://doi.org/10.1306/10171111052>
- 414 Chalmers, G.R.L., Bustin, R.M., 2008. Lower Cretaceous gas shales in northeastern British Columbia,
415 Part I: geological controls on methane sorption capacity. *Bulletin of Canadian Petroleum*
416 *Geology* 56, 1–21. <https://doi.org/10.2113/gscpgbull.56.1.1>
- 417 Chen, J., Xiao, X., 2014. Evolution of nanoporosity in organic-rich shales during thermal maturation.
418 *Fuel* 129, 173–181. <https://doi.org/10.1016/j.fuel.2014.03.058>
- 419 Clarkson, C.R., Solano, N., Bustin, R.M., Bustin, A.M.M., Chalmers, G.R.L., He, L., Melnichenko,
420 Y.B., Radliński, A.P., Blach, T.P., 2013. Pore structure characterization of North American
421 shale gas reservoirs using USANS/SANS, gas adsorption, and mercury intrusion. *Fuel* 103,
422 606–616. <https://doi.org/10.1016/j.fuel.2012.06.119>
- 423 Curtis, M.E., Cardott, B.J., Sondergeld, C.H., Rai, C.S., 2012. Development of organic porosity in the
424 Woodford Shale with increasing thermal maturity. *International Journal of Coal Geology* 103,
425 26–31. <https://doi.org/10.1016/j.coal.2012.08.004>
- 426 Desprairies, A., Bachaoui, M., Ramdani, A., Tribovillard, N., 1995. Clay diagenesis in organic-rich
427 cycles from the Kimmeridge Clay Formation of Yorkshire (G.B.): implication for
428 palaeoclimatic interpretations, in: *Organic Matter Accumulation*. Springer Berlin Heidelberg,
429 Berlin, Heidelberg, pp. 63–91. <https://doi.org/10.1007/BFb0117668>
- 430 DiStefano, V.H., McFarlane, J., Anovitz, L.M., Stack, A.G., Gordon, A.D., Littrell, K.C., Chipera,
431 S.J., Hunt, R.D., Lewis, S.A., Hale, R.E., Perfect, E., 2016. Extraction of organic compounds
432 from representative shales and the effect on porosity. *Journal of Natural Gas Science and*
433 *Engineering* 35, 646–660. <https://doi.org/10.1016/j.jngse.2016.08.064>
- 434 Espitalie, J., Deroo, G., Marquis, F., 1985a. Rock-Eval pyrolysis and its applications. *Revue De L*
435 *Institut Francais Du Petrole* 40, 563–579.
- 436 Espitalie, J., Deroo, G., Marquis, F., 1985b. Rock-Eval pyrolysis and its application. 2. *Revue De*
437 *L'Institut Francais Du Petrole* 40, 755–784.
- 438 Furmann, A., Mastalerz, M., Bish, D., Schimmelmann, A., Pedersen, P.K., 2016. Porosity and pore
439 size distribution in mudrocks from the Belle Fourche and Second White Specks Formations in
440 Alberta, Canada. *Bulletin* 100, 1265–1288. <https://doi.org/10.1306/02191615118>
- 441 Gallois, R.W., 2004. The Kimmeridge Clay: the most intensively studied formation in Britain. *Open*
442 *University Geological Journal* 25.
- 443 Guo, H., He, R., Jia, W., Peng, P., Lei, Y., Luo, X., Wang, X., Zhang, L., Jiang, C., 2018. Pore
444 characteristics of lacustrine shale within the oil window in the Upper Triassic Yanchang
445 Formation, southeastern Ordos Basin, China. *Marine and Petroleum Geology* 91, 279–296.
446 <https://doi.org/10.1016/j.marpetgeo.2018.01.013>
- 447 Guo, H., Jia, W., Peng, P., Zeng, J., He, R., 2017. Evolution of organic matter and nanometer-scale
448 pores in an artificially matured shale undergoing two distinct types of pyrolysis: A study of
449 the Yanchang Shale with Type II kerogen. *Organic Geochemistry* 105, 56–66.
450 <https://doi.org/10.1016/j.orggeochem.2017.01.004>
- 451 Han, H., Pang, P., Li, Z., Shi, P., Guo, C., Liu, Y., Chen, S., Lu, J., Gao, Y., 2019. Controls of organic
452 and inorganic compositions on pore structure of lacustrine shales of Chang 7 member from
453 Triassic Yanchang Formation in the Ordos Basin, China. *Marine and Petroleum Geology* 100,
454 270–284. <https://doi.org/10.1016/j.marpetgeo.2018.10.038>
- 455 Han, Y., Horsfield, B., Wirth, R., Mahlstedt, N., Bernard, S., 2017. Oil retention and porosity
456 evolution in organic-rich shales. *Bulletin* 101, 807–827. <https://doi.org/10.1306/09221616069>

- 457 Herbin, J.P., Fernandez-Martinez, J.L., Geysant, J.R., Albani, A.El., Deconinck, J.F., Proust, J.N.,
458 Colbeaux, J.P., Vidier, J.P., 1995. Sequence stratigraphy of source rocks applied to the study
459 of the Kim meridgian/Tithonian in the north-west European shelf (Dorset/UK, Yorkshire/UK
460 and Boulonnais/France). *Marine and Petroleum Geology* 12, 177–194.
461 [https://doi.org/10.1016/0264-8172\(95\)92838-N](https://doi.org/10.1016/0264-8172(95)92838-N)
- 462 Hu, H., Zhang, T., Wiggins-Camacho, J.D., Ellis, G.S., Lewan, M.D., Zhang, X., 2015. Experimental
463 investigation of changes in methane adsorption of bitumen-free Woodford Shale with thermal
464 maturation induced by hydrous pyrolysis. *Marine and Petroleum Geology* 59, 114–128.
- 465 Jarvie, D.M., Hill, R.J., Ruble, T.E., Pollastro, R.M., 2007. Unconventional shale-gas systems: The
466 Mississippian Barnett Shale of north-central Texas as one model for thermogenic shale-gas
467 assessment. *Bulletin* 91, 475–499. <https://doi.org/10.1306/12190606068>
- 468 Juliao, T., Suárez-Ruiz, I., Marquez, R., Ruiz, B., 2015. The role of solid bitumen in the development
469 of porosity in shale oil reservoir rocks of the Upper Cretaceous in Colombia. *International*
470 *Journal of Coal Geology* 147–148, 126–144. <https://doi.org/10.1016/j.coal.2015.07.001>
- 471 Katz, B.J., Arango, I., 2018. Organic porosity: A geochemist's view of the current state of
472 understanding. *Organic Geochemistry* 123, 1–16.
473 <https://doi.org/10.1016/j.orggeochem.2018.05.015>
- 474 Ko, L.T., Loucks, R.G., Zhang, T., Ruppel, S.C., Shao, D., 2016. Pore and pore network evolution of
475 Upper Cretaceous Boquillas (Eagle Ford–equivalent) mudrocks: Results from gold tube
476 pyrolysis experiments. *AAPG Bulletin* 100, 1693–1722. <https://doi.org/10.1306/04151615092>
- 477 Ko, L.T., Ruppel, S.C., Loucks, R.G., Hackley, P.C., Zhang, T., Shao, D., 2018. Pore-types and pore-
478 network evolution in Upper Devonian-Lower Mississippian Woodford and Mississippian
479 Barnett mudstones: Insights from laboratory thermal maturation and organic petrology.
480 *International Journal of Coal Geology* 190, 3–28. <https://doi.org/10.1016/j.coal.2017.10.001>
- 481 Kuila, U., McCarty, D.K., Derkowski, A., Fischer, T.B., Topór, T., Prasad, M., 2014. Nano-scale
482 texture and porosity of organic matter and clay minerals in organic-rich mudrocks. *Fuel* 135,
483 359–373. <https://doi.org/10.1016/j.fuel.2014.06.036>
- 484 Lafargue, E., Marquis, F., Pillot, D., 1998. Rock-Eval 6 Applications in Hydrocarbon Exploration,
485 Production, and Soil Contamination Studies. *Revue de l'Institut Français du Pétrole* 53, 421–
486 437. <https://doi.org/10.2516/ogst:1998036>
- 487 Landais, P., 1991. Assessment of coal potential evolution by experimental simulation of natural
488 coalification. *Organic Geochemistry* 17, 705–710. [https://doi.org/10.1016/0146-6380\(91\)90014-B](https://doi.org/10.1016/0146-6380(91)90014-B)
- 490 Landais, P., Michels, R., Elie, M., 1994. Are time and temperature the only constraints to the
491 simulation of organic matter maturation? *Organic Geochemistry* 22, 617–630.
492 [https://doi.org/10.1016/0146-6380\(94\)90128-7](https://doi.org/10.1016/0146-6380(94)90128-7)
- 493 Loucks, R.G., Reed, R.M., Ruppel, S.C., Hammes, U., 2012. Spectrum of pore types and networks in
494 mudrocks and a descriptive classification for matrix-related mudrock pores. *Bulletin* 96,
495 1071–1098. <https://doi.org/10.1306/08171111061>
- 496 Loucks, R.G., Reed, R.M., Ruppel, S.C., Jarvie, D.M., 2009. Morphology, Genesis, and Distribution
497 of Nanometer-Scale Pores in Siliceous Mudstones of the Mississippian Barnett Shale. *Journal*
498 *of Sedimentary Research* 79, 848–861. <https://doi.org/10.2110/jsr.2009.092>
- 499 Mackenzie, A.S., Patience, R.L., Maxwell, J.R., Vandenbroucke, M., Durand, B., 1980. Molecular
500 parameters of maturation in the Toarcian shales, Paris Basin, France—I. Changes in the
501 configurations of acyclic isoprenoid alkanes, steranes and triterpanes. *Geochimica et*
502 *Cosmochimica Acta* 44, 1709–1721.
- 503 Mastalerz, M., Schimmelmann, A., Drobniak, A., Chen, Y., 2013. Porosity of Devonian and
504 Mississippian New Albany Shale across a maturation gradient: Insights from organic
505 petrology, gas adsorption, and mercury intrusion. *AAPG Bulletin* 97, 1621–1643.
506 <https://doi.org/10.1306/04011312194>
- 507 Michels, R., Landais, P., Elie, M., Gerard, L., Mansuy, L., 1992. Evaluation of factors influencing the
508 thermal maturation of organic matter during confined pyrolysis experiments. *ABSTRACTS*
509 *OF PAPERS OF THE AMERICAN CHEMICAL SOCIETY* 204, 76.

- 510 Michels, R., Landais, P., Torkelson, B.E., Philp, R.P., 1995. Effects of effluents and water pressure on
511 oil generation during confined pyrolysis and high-pressure hydrous pyrolysis. *Geochimica et*
512 *Cosmochimica Acta* 59, 1589–1604. [https://doi.org/10.1016/0016-7037\(95\)00065-8](https://doi.org/10.1016/0016-7037(95)00065-8)
- 513 Milliken, K.L., Rudnicki, M., Awwiller, D.N., Zhang, T., 2013. Organic matter-hosted pore system,
514 Marcellus Formation (Devonian), Pennsylvania. *Bulletin* 97, 177–200.
515 <https://doi.org/10.1306/07231212048>
- 516 Modica, C.J., Lapiere, S.G., 2012. Estimation of kerogen porosity in source rocks as a function of
517 thermal transformation: Example from the Mowry Shale in the Powder River Basin of
518 Wyoming. *Bulletin* 96, 87–108. <https://doi.org/10.1306/04111110201>
- 519 Pan, L., Xiao, X., Tian, H., Zhou, Q., Chen, J., Li, T., Wei, Q., 2015. A preliminary study on the
520 characterization and controlling factors of porosity and pore structure of the Permian shales in
521 Lower Yangtze region, Eastern China. *International Journal of Coal Geology* 146, 68–78.
522 <https://doi.org/10.1016/j.coal.2015.05.005>
- 523 Peters, K.E., Walters, C.C., Moldowan, J.M., Peters, K.E., 2007. Biomarkers and isotopes in the
524 environment and human history, 2. ed., reprinted with corrections, digitally printed version
525 2007. ed, *The biomarker guide*. Cambridge Univ. Press, Cambridge, UK.
- 526 Powell, J.H., 2010. Jurassic sedimentation in the Cleveland Basin: a review. *Proceedings of the*
527 *Yorkshire Geological Society* 58, 21–72.
528 <https://doi.org/10.1144/pygs.58.1.278>
- 529 Radke, M., 1988. Application of aromatic compounds as maturity indicators in source rocks and crude
530 oils. *Marine and Petroleum Geology* 5, 224–236. [https://doi.org/10.1016/0264-](https://doi.org/10.1016/0264-8172(88)90003-7)
531 [8172\(88\)90003-7](https://doi.org/10.1016/0264-8172(88)90003-7)
- 532 Radke, M., Welte, D.H., Willsch, H., 1986. Maturity parameters based on aromatic hydrocarbons:
533 Influence of the organic matter type. *Organic Geochemistry* 10, 51–63.
534 [https://doi.org/10.1016/0146-6380\(86\)90008-2](https://doi.org/10.1016/0146-6380(86)90008-2)
- 535 Ramanampisoa, L., Disnar, J.R., 1994. Primary control of paleoproduction on organic matter
536 preservation and accumulation in the Kimmeridge rocks of Yorkshire (UK). *Organic*
537 *Geochemistry* 21, 1153–1167. [https://doi.org/10.1016/0146-6380\(94\)90160-0](https://doi.org/10.1016/0146-6380(94)90160-0)
- 538 Ramanampisoa, L.R., Radke, M., 1995. Extractable aromatic hydrocarbons in a short-term organic
539 cycle of the Kimmeridge Clay formation, Yorkshire (U.K.): relationship to primary production
540 and thermal maturity. *Organic Geochemistry* 23, 803–817. [https://doi.org/10.1016/0146-](https://doi.org/10.1016/0146-6380(95)80002-9)
541 [6380\(95\)80002-9](https://doi.org/10.1016/0146-6380(95)80002-9)
- 542 Rawson, P.F., Riley, L.A., 1982. Latest Jurassic - Early Cretaceous Events and the “Late Cimmerian
543 Unconformity” in North Sea Area. *AAPG Bulletin*, *AAPG Bulletin* 66, 2628–2648.
- 544 Schmitt, M., Fernandes, C.P., da Cunha Neto, J.A.B., Wolf, F.G., dos Santos, V.S.S., 2013.
545 Characterization of pore systems in seal rocks using Nitrogen Gas Adsorption combined with
546 Mercury Injection Capillary Pressure techniques. *Marine and Petroleum Geology* 39, 138–
547 149.
- 548 Seifert, W.R., Moldowan, J.M., 1981. Paleoreconstruction by biological markers. *Geochimica et*
549 *Cosmochimica Acta* 45, 783–794.
- 550 Sing, K.S., 1985. Reporting physisorption data for gas/solid systems with special reference to the
551 determination of surface area and porosity (Recommendations 1984). *Pure and applied*
552 *chemistry* 57, 603–619.
- 553 Song, L., Martin, K., Carr, T.R., Ghahfarokhi, P.K., 2019. Porosity and storage capacity of Middle
554 Devonian shale: A function of thermal maturity, total organic carbon, and clay content. *Fuel*
555 241, 1036–1044. <https://doi.org/10.1016/j.fuel.2018.12.106>
- 556 Tissot, Welte, D.H., 1984. *Petroleum formation and occurrence*, 2. rev. and enl. ed. ed. Springer,
557 Berlin, Heidelberg [usw.].
- 558 Wang, X., Jiang, Z., Jiang, S., Chang, J., Li, X., Wang, Xin, Zhu, L., 2020. Pore Evolution and
559 Formation Mechanism of Organic-Rich Shales in the Whole Process of Hydrocarbon
560 Generation: Study of Artificial and Natural Shale Samples. *Energy Fuels* 34, 332–347.
561 <https://doi.org/10.1021/acs.energyfuels.9b03789>
- 562 Wang, Y., Liu, L., Hu, Q., Hao, L., Wang, X., Sheng, Y., 2020. Nanoscale Pore Network Evolution of
563 Xiamaling Marine Shale during Organic Matter Maturation by Hydrous Pyrolysis. *Energy*
564 *Fuels* 34, 1548–1563. <https://doi.org/10.1021/acs.energyfuels.9b03686>

- 565 Wang, Y., Liu, L., Zheng, S., Luo, Z., Sheng, Y., Wang, X., 2019. Full-scale pore structure and its
 566 controlling factors of the Wufeng-Longmaxi shale, southern Sichuan Basin, China:
 567 Implications for pore evolution of highly overmature marine shale. *Journal of Natural Gas
 568 Science and Engineering* 67, 134–146. <https://doi.org/10.1016/j.jngse.2019.04.020>
- 569 Wood, D.A., 1988. Relationships between thermal maturity indices calculated using Arrhenius
 570 equation and Lopatin method: implications for petroleum exploration. *Assoc. Pet. Geol. Bull.*
 571 72, 115–134.
- 572 Wu, S., Yang, Z., Zhai, X., Cui, Jingwei, Bai, L., Pan, S., Cui, Jinggang, 2019. An experimental study
 573 of organic matter, minerals and porosity evolution in shales within high-temperature and high-
 574 pressure constraints. *Marine and Petroleum Geology* 102, 377–390.
 575 <https://doi.org/10.1016/j.marpetgeo.2018.12.014>
- 576 Zhang, T., Ellis, G.S., Ruppel, S.C., Milliken, K.L., Lewan, M.D., Sun, X., 2013. Effect of organic
 577 matter properties, clay mineral type and thermal maturity on gas adsorption in organic-rich
 578 shale systems. Presented at the Unconventional Resources Technology Conference, Society of
 579 Exploration Geophysicists, American Association of Petroleum Geologists, Society of
 580 Petroleum Engineers, pp. 1996–2001.
 581

582 Appendix 1

583 Calculation of biomarker ratios

- 584 • MDBT: methyl dibenzothiophene ratio = 4-MDBT/1-MDBT (Radke, 1988).
- 585 • $R_{\text{MDBT}} (\%) = 0.40 + 0.30 \cdot \text{MDR} - 0.094 \cdot \text{MDR}^2 + 0.011 \cdot \text{MDR}^3$ (Radke, 1988).
- 586 • DPR: dimethylphenanthrene ratio = (2,6-DMP + 2,7-DMP + 3,5-DMP) / (1,3-DMP + 1,6-DMP
 587 + 2,5-DMP + 2,9-DMP + 2,10-DMP + 3,9-DMP + 3,10-DMP) (Radke, 1988).
- 588 • $R_{\text{DPR}} (\%) = 1.50 + 1.34 \cdot \log_{10} \text{DPR}$ (Radke, 1988).
- 589 • MPI-1: methylphenanthrene index = $1.5 \cdot (2\text{-MP} + 3\text{MP}) / (\text{Ph} + 1\text{-MP} + 9\text{-MP})$ (Radke et al.,
 590 1986).
- 591 • $R_{\text{MPI-1}} (\%)$: for $R_o \leq 1.4\%$, $R_o = 0.7 \cdot \text{MPI-1} + 0.22$ (Boreham et al., 1988). For $R_o \geq 1.4\%$, R_c
 592 $= -0.55 \cdot \text{MPI-1} + 3.0$ (Boreham et al., 1988).
- 593 • Mean calculated vitrinite reflectance, $R_c (\%) = (R_{\text{MDBT}} + R_{\text{DPR}} + R_{\text{MPI-1}}) / 3$.
- 594

Highlights

- Long and short-duration thermal maturation experiments are compared.
- Longer duration experiments enhance the degradation of polar organic components.
- Longer duration experiments exhibit slightly different gas concentrations.
- Duration-temperature pairs do not greatly influence the evolution trend of porosity.
- Duration-temperature pairs influence pore size distributions and pore volumes.

Journal Pre-proof

Declaration of interests

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

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