

Origin of compositional differences in organic matter abundance and oil potential of cherty and clayey Cenomanian black levels in the Umbria-Marche basin (Italy)

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Abstract

Rock-Eval pyrolysis of a large set of Cenomanian samples, collected from the black levels (clayey, cherty and mixed) in three sections of the Umbria-Marche basin, showed large differences in organic matter (OM) quantity and quality. The chert samples systematically exhibit much lower TOC contents, markedly lower HI and higher OI. This reflects the extensive oxidative destruction of the initial kerogen that occurred upon the chertification of some clayey sediments. A comparative study, by a combination of microscopic, spectroscopic and pyrolytic methods, was performed on kerogens of the chert and clay layers of a representative mixed level. The various fractions of the initial kerogen underwent differential destruction or alteration during chertification, resulting in (i) relative enrichments of microfossils and woody debris although lignin was altered by demethoxylation and (ii) extensive destruction of the amorphous fraction while it remained predominant. The amorphous fraction retained in the chert kerogen showed large changes in composition related to oxygen incorporation and probably escaped complete destruction owing to oxidative reticulation. The above features account for the pronounced systematic differences in OM abundance and oil potential between the chert and clay layers in the black levels.

1. Introduction

A number of black levels (laterally extensive, dark coloured, clay- or chert-rich horizons intercalated with pale-coloured pelagic limestones) occur just below the Cenomanian/Turonian boundary (ca. 93.3 Ma) in the Umbria-Marche basin, Central Italy ([Arthur and Premoli, 1982 and Beaudoin et al]). Approximately 40 of the levels have been identified and correlated in numerous sections over the entire basin ([Beaudoin et al]). These levels are relatively thin (<1 cm up to ca. 15 cm) and their deposition spanned about 2.5 million years ([Arthur and Premoli, 1982, M'ban, 1994 and Beaudoin et al]). The black levels exhibit rhythmic deposition whose observed cyclicity can be related to precession ([Beaudoin et al]) as previously observed in other deposits (e.g. [Klemme and Ulmishek, 1991 and de Boer and Smith, 1994]). Such levels occur within predominantly organic-lean carbonate sediments and exhibit a clayey or siliceous lithology. Some mixed levels showing both types of lithology are also observed, with the siliceous (chert) layers always overlaying the clay ones.

Sedimentological field observations and morphological studies on whole rock samples ([Salmon, 1999]) indicate that the chert levels (layers) result from the complete (partial) chertification of clay sediments. Accordingly, the organic matter (OM) in the chert and clay layers of a mixed level share a common origin. It also appeared that this transformation into chert took place at a relatively early stage of sediment compaction. As observed in other deposits, oxygen and silica for chertification could be supplied by clay ([Behl, 1990]); radiolarian tests were probably another important potential source of silica through dissolution and recrystallisation. Cherts generally exhibit low contents of OM, as observed for example in Cretaceous deposits from South America ([Marcellari and de Vries, 1987]) and laboratory studies on simulated silicification of blue green algae ([Oehler, 1976]) showed that extensive OM destruction took place due to the intense oxidation processes associated with chert formation. Indeed, (i) examination of a set of thirteen samples from black levels from the Furlo section in the Umbria-Marche Basin showed large differences in OM quantity and quality, with total organic carbon (TOC) ranging from ca. 0.5 to 14% and Rock-Eval Hydrogen Index (HI) from ca. 120 to 450 mg HC/g TOC ([M'ban, 1994]) and (ii) these observations pointed to correlations between the above features and the lithology of the black levels, the cherts generally exhibiting markedly lower TOC contents (<1%) and HI values (commonly <200 mgHC/g TOC) when compared to the clays.

In the present work a larger set of samples, collected from the black levels (clay, chert, mixed) of three sections in the Umbria-Marche Basin, was examined by Rock-Eval pyrolysis. These bulk analyses confirmed the occurrence of conspicuous differences in OM abundance and oil potential between the clay and chert samples. Such differences are well illustrated by the samples from the chert and clay layers from a mixed black level, termed NN8, from the Sta Maria di Burano (SMB) section. A detailed study was, therefore, performed on the kerogen isolated from the chert layer (NN8si) of this mixed level, via a combination of microscopic (transmitted light microscopy, transmission electron microscopy), spectroscopic (FTIR and solid state ¹³C NMR) and pyrolytic methods, to determine its composition. Comparison with previous observations ([Salmon et al and Salmon et al]) on the kerogen from the associated clay layer (NN8c) afforded information on the origin of the compositional differences observed between the clay and chert samples in the black levels of the basin.

2. Materials and methods

2.1. Rock-Eval analysis of whole rock samples

Forty-eight samples, from 34 black levels (including 14 mixed levels), were collected from three sections located north of Perugia (Italy): the SMB section (ca. 40 km), the Furlo section (ca. 65 km) and the Monte Petrano section (ca. 90 km). Rock-Eval pyrolyses were performed on ca. 90 mg of ground rock with a temperature program from 300 °C (20 min) up to 600 °C at 25 °C/min under a He flow, followed by oxidation at 600 °C for 7 min under an oxygen flow.

2.2. Study of the chert kerogen

The NN8 level in the SMB section consists of a 3 cm thick clay layer overlain by a 2 cm thick chert layer. This black level spans a large area in the Umbria-Marche basin and was identified in all the known outcrops over the whole basin ([Beaudoin et al]). Vitrinite reflectance (0.55%) indicated a relatively low maturity corresponding to the onset of the oil window (

[Salmon, 1999]). The NN8si sample was taken from a fresh surface of the section in an active quarry. This siliceous sample was collected in the first centimeter of the base of the cherty layer of the NN8 black level just above the previously studied clayey sample (NN8c). The kerogen concentrate was prepared via the classical HF/HCl treatment ([Durand and Nicaise, 1980]) and was immediately analysed to avoid possible alteration upon storage ([Salmon et al]).

Rock-Eval pyrolysis was performed on ca. 5 mg of NN8si and NN8c kerogen concentrates using the same conditions as above for whole rock samples. Elemental composition was determined at the Service Central d'analyse of the CNRS (Vernaison) by combustion at 1050 °C under O₂.

X-ray diffraction powder patterns were recorded at room temperature, with a SIEMENS D5000 diffractometer (CoK_α radiation), equipped with a graphite back monochromator set on the diffracted beam. X-ray data collection (scanning step 0.02°, counting time 2.4 s) and data treatment were monitored by the Siemens Diffract-AT routines.

The FTIR spectra were recorded with a Bruker IFS48 spectrometer as KBr pellets and the cross polarization/magic angle spinning (CP/MAS) ¹³C NMR spectra were obtained on a Bruker MSL 400 under the same conditions as previously described ([Salmon et al]).

A Leitz MPVII microscope was used for transmitted light microscopy observations. The relative abundance of the main types of particles in the clay and chert kerogens was determined through area estimation on 500 particles per slide. Transmission electron microscopy (TEM) observations were performed as previously described ([Boussafir et al]) using a Philips CM 20 microscope.

Off-line pyrolysis under a helium flow and pyrolysate separation by column chromatography were performed as described in [Largeau et al]. The methanol-eluted fraction was esterified to improve fatty acid detection. Each fraction was analysed by combined gas chromatography/mass spectrometry (GC/MS) as described in [Salmon et al].

3. Results and discussion

3.1. Rock-Eval analysis of whole rocks

The Rock-Eval data obtained for the 48 samples, collected from homogeneous or mixed black levels, reveal pronounced differences between the clay and chert rocks (Table 1, Fig. 1). Thus, TOC is markedly lower in the latter (systematically below 1%), whereas most of the clays exhibit values above 10%. The HI is markedly higher for the clays, whereas the reverse is observed for the oxygen index (OI). HI and OI values of kerogens are controlled by different factors, including kerogen maturity, type and level of oxidation ([Tissot and Welte, 1984]). In the present study the former factor can be ruled out since the black levels experienced close (identical for the mixed levels) thermal histories in terms of burial. The second factor can also be straightforwardly eliminated, at least for the mixed levels since the OM in the chert and clay layers has a common origin. Accordingly, in agreement with general observations on cherts and laboratory silicification mentioned in the Introduction, the large differences observed for the HI and OI values between the chert and clay samples must reflect differences in post-depositional oxidation associated with chert formation.

Table 1. Rock-Eval data for whole samples from black levels of the Sta maria di Burano (SMB), Furlo (FU) and Monte Petrano (MP) sections

Sample ^a	TOC ^b	HI ^b	OI ^b
<i>SMB section</i>			
(NN1si)	0.5	107	60
(NN2si)	0.7	152	54
(NN3c)	11.2	418	7
<i>(NN3si)</i>	<i>0.1</i>	<i>nd</i>	<i>nd</i>
(NN4si)	0.3	nd	nd
(NN5c)	8.8	375	9
(NN6c)	12.2	427	12
(NN7c)	13.2	452	7
(NN8c)	13.6	449	7
<i>(NN8si)</i>	<i>0.2</i>	<i>nd</i>	<i>nd</i>
(NN9c)	10.7	362	11
(NN10c)	10.2	436	10
<i>(NN10si)</i>	<i>0.6</i>	<i>138</i>	<i>48</i>
(NN15si)	0.4	109	72
(NN18c)	12.7	359	7
<i>(NN18si)</i>	<i>0.1</i>	<i>nd</i>	<i>nd</i>
(NN19c)	10.9	424	10
(NN20c)	11.6	487	5
(NN21c)	13.2	469	11
<i>(NN21si)</i>	<i>0.4</i>	<i>156</i>	<i>68</i>
(NN22c)	13.8	445	17
<i>(NN22si)</i>	<i>0.1</i>	<i>nd</i>	<i>nd</i>
(NN24c)	9.6	337	13
<i>(NN24si)</i>	0.4	97	53
(NN25si)	0.2	nd	nd
(NN26si)	0.1	nd	nd
(NN27si)	0.1	nd	nd
<i>FU section</i>			
(NN1si)	0.8	179	39
(NN3c)	10.7	442	6
<i>(NN3si)</i>	0.2	nd	nd
(NN5c)	13.5	397	10
(NN7c)	9.2	379	8
<i>(NN7si)</i>	<i>0.2</i>	<i>nd</i>	<i>nd</i>
(NN8c)	11.2	439	7
<i>(NN8si)</i>	<i>0.1</i>	<i>nd</i>	<i>nd</i>
(NN10c)	12.8	409	13
(NN20c)	8.9	349	12
(NN21c)	13.2	447	8
(NN24c)	14.8	485	6
<i>(NN24si)</i>	<i>0.5</i>	<i>122</i>	<i>57</i>
<i>MP section</i>			
(NN1si)	0.2	nd	nd
(NN3c)	8.2	357	5
<i>(NN3si)</i>	0.1	nd	nd
(NN5c)	12.2	427	21
<i>(NN5si)</i>	<i>0.8</i>	<i>113</i>	<i>61</i>
(NN7si)	0.4	97	49
(NN8c)	11.9	478	7
<i>(NN8si)</i>	<i>0.1</i>	<i>nd</i>	<i>nd</i>

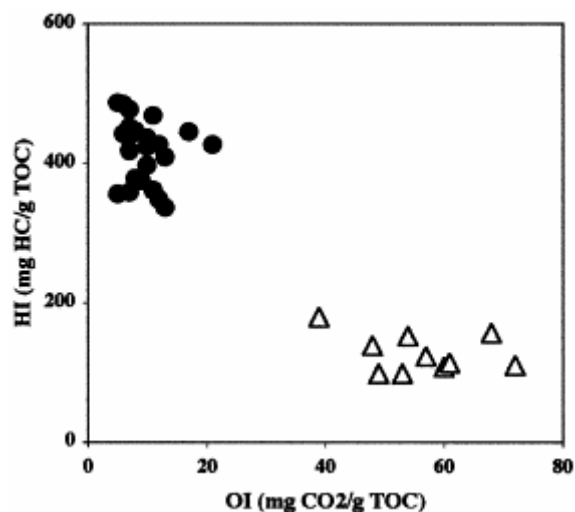


Fig. 1. HI versus OI diagram for whole samples from black levels of the Sta maria di Burano, Furlo and Monte Petrano sections (circles: clayey samples, triangles: cherty samples; empty symbols: TOC <1%, filled symbols: TOC around 10–15%).

The representative NN8 black level was selected for further study so as to derive detailed information on the differences between the chert and clay kerogens and on the consequences of chertification on kerogen composition.

3.2. Detailed study of the chert kerogen and comparison with the clay kerogen

3.2.1. Bulk geochemical features

Rock-Eval pyrolysis was performed on the kerogen concentrates since significant HI and OI values cannot reliably be obtained for rocks with a TOC <0.3% ([Espitali and Espitali]). The chert kerogen shows a markedly lower oil potential, whereas OI is much higher (Table 2). Elemental analysis also showed large differences: the chert kerogen exhibits a lower H/C atomic ratio, while its O/C ratio is ca. two times higher when compared to the clay kerogen (Table 2). In addition, the elemental analysis of the concentrates indicated a markedly lower ash content for the chert. It is well documented that some minerals, including pyrite and various sulfates survive HF/HCl treatment ([Tissot and Welte, 1984]). Pyrite indeed occurs in both concentrates as shown by microscopy and by the presence of a sharp band at 427 cm^{-1} in their FTIR spectra. Organic sulfur content of kerogens is classically calculated from elemental analysis data on concentrates, by assuming that mineral sulfur only occurs as pyrite and that all the Fe corresponds to pyrite, so that $S_{\text{org}} = S_{\text{total}} (\text{measured}) - S_{\text{ash}} (\text{pyritic, calculated from Fe content})$. Such a calculation showed for the clay kerogen a low sulfur content and a S_{org}/C atomic ratio of 0.02, i.e. in the range commonly observed for low maturity type II kerogen. In contrast, this calculation gives a much higher "S_{org}" content for the chert kerogen, corresponding to an unusually high S_{org}/C atomic ratio of 0.19. In fact (i) XANES spectroscopy (G. Sarret, personal communication) showed that sulfur in the chert concentrate occurs almost exclusively as sulfate and (ii) X-ray diffraction showed the presence of jarosite, an inorganic sulfate of general formula $\text{KFe}_3(\text{SO}_4)_2(\text{OH})_6$: main peaks (2θ Co K_α) at 20.12, 20.95, 22.52, 24.89, 27.60, 28.29, 30.23, 33.98, 38.40, 43.57, 48.45, 51.74 and 55.44). This sulfate can survive HF/HCl treatment but is decomposed upon heating around $600\text{ }^\circ\text{C}$, with SO_2 production ([Mackenzie, 1970]). Indeed, the trace for the Curie-point flash pyrolysate at $610\text{ }^\circ\text{C}$ of the chert concentrate (not shown) exhibits a prominent peak corresponding to sulfur

dioxide that overwhelms the pyrolysis products of the kerogen. Due to the presence of jarosite in the concentrate, a significant value cannot be obtained for S_{org} content in the chert kerogen, using the elemental data and the classical calculation. In fact, the sulfur of jarosite is then taken into account in the total sulfur but not in the ash sulfur, due to jarosite decomposition and SO_2 release upon heating during ash isolation. Hence, in that case, the value calculated for S_{org} , by subtracting ash sulfur from total sulfur, is largely overestimated. Finally, it should be noted that jarosite can be formed by oxidation of pyrite. Therefore, the presence of jarosite in the chert concentrate must reflect the extensive oxidation that took place during chertification.

Table 2. Rock-Eval data and elemental composition for NN8c kerogen concentrate, NN8si kerogen concentrate and the 400 °C pyrolysis residue of the latter

Sample	TOC ^a	HI ^a	OI ^a	H/C ^b	O/C ^b	Sorg/C ^b	Ash ^e
SMB (NN8c)	13.6 ^d	472 ^c	9 ^e	1.07 ^d	0.20 ^d	0.02 ^d	20.5 ^d
SMB (NN8si)	0.2	185 ^e	42 ^e	0.70	0.41	ns ^f	5.8
R400. NN8si	nd	nd	nd	0.40	0.49	ns ^f	11.0

nd: pyrolysis residue not examined by Rock-Eval pyrolysis.

3.2.2. Morphological features

Light microscopy showed that the chert kerogen is mostly composed of amorphous OM (ca. 63%) dominated by tiny (ca. 1 μm), brown, particles. A substantial contribution (ca. 31%) of woody debris and a few microfossils, including some pollen and spores, was also detected. TEM observations did not reveal any structure in the tiny particles, which are composed of nanoscopically-amorphous OM. The clay kerogen showed a higher contribution (ca. 88%) and much larger size (ca. 10 μm) for the amorphous particles, whereas the woody debris (ca. 9%) and microfossils were less abundant.

3.2.3. Spectroscopic features

The solid state CP/MAS ^{13}C NMR spectrum of the chert kerogen (Fig. 2A) shows the presence of aliphatic carbons (0–60 ppm range, maximum around 30 ppm corresponding to CH_2 groups in polymethylenic chains), olefinic and/or aromatic unsaturations (peak centered at 130 ppm), C–O groups (peak centered at 86 ppm) and C=O in esters and/or amides (peak centered at 170 ppm). Comparison with the spectrum of the clay kerogen (Fig. 2B); ([Salmon et al]) showed some quantitative differences for the chert kerogen: (i) an increase in the relative intensity of the peak at 170 ppm corresponding to carboxyl groups, (ii) the occurrence of two maxima for the C–O groups, including the peak at 86 ppm probably related to aromatic carbons linked to oxygen and (iii) a larger shoulder around 40 ppm probably corresponding to carbons adjacent to carbons bearing oxygen functions. Comparison of the FTIR spectra (not shown) showed a substantial shift in the maximum of the C=C band (1605 versus 1615 cm^{-1}) for the chert kerogen, indicating a relatively higher contribution of aromatic unsaturations with respect to olefinic ones. The spectroscopic features therefore indicate a higher contribution of carboxyl groups and of aromatic units (including oxygen-substituted units) for the chert kerogen, in agreement with its higher O/C and lower H/C ratios (Table 2).

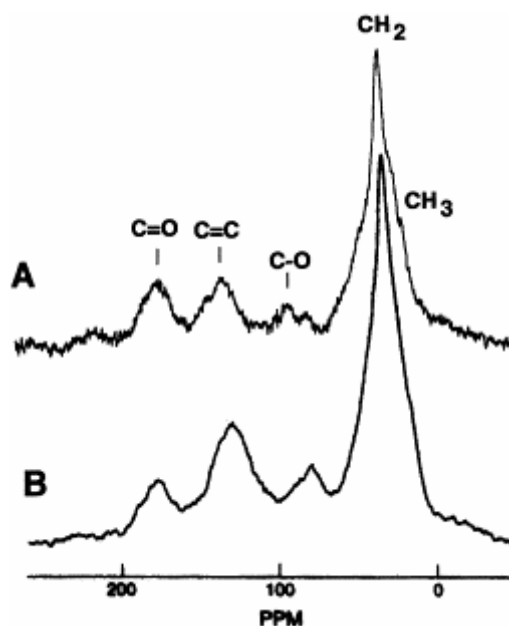


Fig. 2. Solid state CP/MAS ^{13}C NMR spectra of (A) NN8si and (B) NN8c kerogen.

3.2.4. Pyrolysis

Off-line pyrolysis at 400 °C resulted in a total weight loss of ca. 25% for the chert kerogen. The pyrolysis residue shows a decrease in the H/C atomic ratio (Table 2), reflecting the release of the aliphatic moieties and the aromatization processes that led to the formation of this residual material. A higher weight loss (ca. 40%) was previously observed, under the same pyrolysis conditions, for the clay kerogen ([Salmon et al]) and the pyrolysis residue showed a higher H/C atomic ratio (0.64 against 0.40). Such differences should reflect the higher level of aromatic moieties in the chert kerogen, hindering the formation of volatile pyrolysis products and promoting the formation of residual polyaromatic material.

The GC trace of the crude pyrolysate of the chert kerogen showed a large unresolved complex mixture (UCM) of co-eluting compounds. The pyrolysate was, therefore, separated by column chromatography, into three fractions eluted with heptane, toluene and methanol, accounting for ca. 20, 15 and 50 wt.% of the total pyrolysate, respectively. A large number of compounds, commonly found in kerogen pyrolysates were identified by GC/MS analysis of these three fractions, such as series of alkanes and alkenes, alkylated benzenes and alkylated polyaromatic hydrocarbons, hopanes and hopenes, *n*-alkanones and mid-chain ketones, fatty acids and phenols. Only the main differences in pyrolysate composition compared to the clay kerogen, previously examined by [Salmon et al] are presented below. These differences are chiefly related to the distribution of some pyrolysis products with long alkyl chains and of the phenols:

The abundant *n*-alkan-2-ones in the toluene-eluted fraction exhibit a bimodal distribution with maxima at C_{17} and C_{29} (Fig. 3) and the odd-carbon-numbered compounds predominate, whereas a unimodal distribution with a maximum at C_{13} followed by a rather smooth decrease in intensity and no odd-over-even predominance, was observed for the clay kerogen. This trend for a higher relative contribution of long-chain compounds exhibiting a marked predominance for the chert kerogen was also observed for the fatty acids, the C_{20+} homologues accounting for ca. 20% of the total fatty acids and exclusively corresponding to even-carbon-numbered compounds.

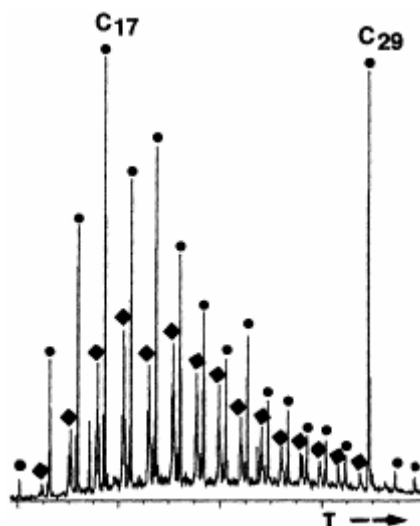


Fig. 3. Mass chromatogram at $m/z=58$ showing the distribution of the n -alkan-2-ones (filled circles) and of the co-eluting series of mid-chain ketones (filled diamonds) in the toluene-eluted fraction of the pyrolysate at 400 °C of NN8si kerogen (numbers refer to chain length).

The phenolic compounds in the pyrolysate of the chert kerogen correspond only to alkyl phenols, the C_1 – C_3 homologues being predominant. In contrast, these short-chain phenols occur along with methoxy-substituted counterparts in the case of the clay kerogen.

The significance of these differences is discussed in the next section, in relation to the differences in the bulk, microscopic and spectroscopic features of the chert and clay kerogens.

3.2.5. Comparison of chert and clay kerogens

The OM in the clay and chert layers of mixed levels shares a common origin. Therefore, the compositional quantitative and qualitative differences observed between the clay and chert kerogens should reflect differences in post-depositional diagenesis associated with chert formation. Chertification resulted in extensive destruction of a large part of the initial OM in the silicified zone. This oxidative destruction is reflected in the chert kerogen by a sharp drop in the size of the brown amorphous particles, by the increase in O/C atomic ratio and OI and by the marked decrease in HI and TOC. The latter decrease also reflects dilution resulting from chert production. In agreement with the generally higher resistance of terrestrial OM relative to marine OM and of palynomorphs relative to amorphous OM (e.g. [Tyson, 1995]), the woody debris and the microfossils were less affected by oxidative degradation than the brown amorphous material. As discussed below, this differential degradation that took place during chertification accounts for the marked morphological, spectroscopic and pyrolytic differences observed between the chert and clay kerogens.

3.2.5.1. Microfossils

The microfossils in both kerogens mostly correspond to pollen and spore walls. It is well documented that such microfossils contribute to kerogen formation through the "selective preservation" pathway, based on the existence of biomacromolecules exhibiting a very high resistance to diagenetic degradation (reviewed in [de Leeuw and Largeau, 1993 and Largeau and de Leeuw, 1995]). This type of biomacromolecule, when submitted to laboratory degradation experiments, also exhibits a high resistance to oxidative treatment (e.g. [Berkaloff

et al]). Thus, the pollen and spore walls that occurred in the kerogen of the clayey zone that was transformed into chert were not highly affected by the oxidation processes associated with chertification; hence the enrichment observed for these microfossils in the chert kerogen.

Generally, these resistant biomacromolecules are highly aliphatic and their pyrolysates show the abundant presence of *n*-alkanones, derived from the thermal cleavage of ether cross-links. Such ketones show a typical distribution that reflects the location of the ether links on specific positions of the alkyl chains, this location being biosynthetically controlled (e.g. [Derenne et al, Derenne et al, Gillaizeau et al, Gelin et al, Gelin et al, Schouten et al and Blokker et al]). The *n*-alkan-2-ones that abundantly occur in the pyrolysate of the chert kerogen exhibit a bimodal (C₁₇, C₂₉) distribution (Fig. 3), with a sharp maximum at C₂₉. This bimodal distribution reflects a marked preferential location of the ether linkages on the alkyl chains for the kerogen moieties from which these ketones are generated by pyrolysis. The distribution of these *n*-alkan-2-ones thus probably reflects, in agreement with microfossil enrichment, the substantial contribution of resistant aliphatic biomacromolecules in the chert kerogen. These components afforded a much lower contribution to the clay kerogen so that such features were not observed due to large dilution by the pyrolysis products of the other fractions. In contrast, as a result of the extensive destruction that took place during chertification, the aliphatic biomacromolecules were enriched and their typical signature could now be observed. Moreover, the preferential location of the ether bridges suggests that such components did not undergo substantial alteration during chertification since, as discussed below, oxidative alteration is reflected in a random location of the oxygen-containing bridges.

3.2.5.2. Woody debris

Complete degradation of lignin into soluble compounds can hardly be achieved through strong oxidative laboratory treatments ([Isherwood, 1965]) and lignin also shows a high resistance to diagenetic degradation. Short chain (C₁–C₃) alkyl phenols and methoxy-substituted alkyl phenols are typical pyrolysis products of lignin (e.g. [Saiz-Jimenez and de Leeuw, 1986 and Hatcher and Spiker, 1988]). C₁–C₃ Alkyl phenols and methoxy-substituted alkyl phenols were observed in pyrolysate of the clay kerogen, in agreement with the contribution of woody debris. The relative abundance of such debris is substantially higher in the chert kerogen and the isomers of methylphenol and several isomers of dimethylphenol, probably related to lignin, were detected in its pyrolysate. In contrast to the clay kerogen, however, no methoxy-substituted phenols were observed. It is well documented (e.g. [Hatcher and Clifford, 1997]) that lignin alteration results in demethoxylation reactions. Thus, it appears that lignin was enriched during chertification but the retained lignin underwent some alteration through elimination of the methoxy groups.

3.2.5.3. Amorphous OM

Despite the increase in the relative contribution of microfossils and woody debris, amorphous particles still dominate the chert kerogen. As previously shown ([Salmon et al]), the bulk of the amorphous particles in the clay kerogen originated from the condensation of potentially labile compounds that survived diagenesis owing to mineral protection provided by alternation of organic and clay nanolayers. The kerogen fractions thus formed are highly sensitive to oxidative degradation, once this physical protection is removed by destruction of the mineral nanolayers. Indeed, the isolated clay kerogen undergoes pronounced and fast oxidative alteration upon storage, when exposed to air at room temperature ([Salmon et al]). Such a high sensitivity accounts for the extensive degradation that took place, for the

amorphous fraction of the clay kerogen, when submitted to the oxidative conditions associated with chertification. Moreover, the low amount of this amorphous material that escaped destruction probably underwent major alteration in chemical structure, as reflected by the high level of oxygen-containing groups occurring in the chert kerogen. This alteration possibly resulted in the formation of resistant macromolecular compounds with a high degree of oxygen cross-linking. As recently shown ([Riboulleau et al and Riboulleau et al]), such cross-linking was implicated in the formation of some kerogens and "oxidative reticulation" was thus put forward as a new pathway of OM preservation in sediments. A similar process may therefore account for the lack of complete destruction of the "oxidation-sensitive" amorphous material in the chert kerogen. The occurrence of oxidative cross-linking is supported by the presence in the pyrolysate of several series of "mid-chain" *n*-alkanones with the keto group located on any carbon atom from C(3) to C(12). Contrary to the *n*-alkan-2-ones these series showed a smooth distribution (Fig. 3), indicating the lack of preferential location of the corresponding ether bridges on the alkyl chains. Such a random location of the bridges would be typical of the materials formed by oxidative reticulation ([Riboulleau et al and Riboulleau et al]). The preservation of this amorphous fraction by oxidative reticulation is consistent with the relatively high OI and the relatively low HI values observed for the chert sample, as typical for all samples of this type in the black levels of the basin.

4. Conclusion

Rock-Eval analysis of a large set of samples from three sections confirmed the occurrence of compositional differences in OM abundance and quality between clay and chert samples. The low TOC and the low oil potential typical of the latter must reflect the large oxidative destruction of the initial kerogen that took place during chertification.

Different levels of destruction occurred between the various fractions of the initial kerogen in the silicified NN8 layer, as reflected by an increase in the relative contribution of the microfossils and woody debris. Moreover, pyrolytic signatures suggest that the microfossil components did not undergo significant alteration during chertification, whereas the chemical structure of lignin in the retained woody debris was substantially altered due to demethoxylation.

The amorphous particles, physically protected by clay mineral nanolayers in the initial kerogen, were extensively destroyed during chertification so that their relative contribution decreased while remaining predominant. The amorphous OM retained in the chert kerogen showed major changes in chemical structure due to substantial incorporation of oxygen. This probably reflects the oxidative reticulation that allowed a small part of the amorphous OM to escape destruction during chertification.

The differential processes that extensively affected the initial, clayey, material during chertification account for the morphological and chemical features of the chert kerogen and the differences in OM abundance and oil potential, which are systematically observed between the chert and clay samples in the black levels of the Umbria-Marche Basin.

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