# ORGANIC GEOCHEMICAL STUDY OF THE LOWER MIOCENE KREMNA BASIN, SERBIA

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Perunović, T., Stojanović, K., Simić, V., Kašanin-Grubin, M., Šajnović, A., Erić, V., Schwarzbauer, J., Vasić, N., Jovančićević, B. & Brčeski, I., 2014. Organic geochemical study of the Lower Miocene Kremna Basin, Serbia. *Annales Societatis Geologorum Poloniae*, 84: 185–212.

**Abstract:** The Kremna Basin is located in southwest Serbia, in the Zlatibor area, which is part of the Internal Dinarides. This basin is noteworthy because of the type of bedrock drainage, which it represents. It was formed on ultrabasic rocks and volcanic materials that influenced the occurrence of organic matter (OM) in the basin fill. The objective of the study was to determine the organic geochemical characteristics of sediments from the central part of the Kremna Basin.

The sediments studied belong to an intrabasinal facies, in which two sequences were distinguished. The lower sequence occurs at depths of 216–343 m, while the upper sequence is found from 13.5–216 m.

At the start of basin development (265–343 m) sedimentation took place in shallow alkaline water, rich in Mg ions. Through time, a slight deepening of the basin occurred. This was followed by chemical deposition of carbonates (216–265 m). The most important change in the sedimentary environment occurred with the formation of sediments marking the transition between the sequences (at about 200 m).

Sediments from the lower sequence are characterized by the dominance of dolomite and magnesite. There are indications of volcanic activity, i.e. tuff layers and the presence of searlesite. The upper sequence is characterized by the prevalence of calcite and dolomite. The amounts of MgO, Na<sub>2</sub>O and B are higher in the lower sequence, whereas the CaO content is higher in the upper sequence.

The sediments contain different amounts of immature OM (late diagenesis). Biomarker analysis shows diverse precursors of the sedimentary OM: methanogenic archaea, photosynthetic green sulfur bacteria (*Chlorobiaceae*), bacterivorous ciliates, various bacteria, both photosynthetic and non-photosynthetic, the green unicellular microalga, *Botryococcus braunii* race A (exclusively in the upper sequence) and terrestrial plants. The lower sequence contains lower amount OM, composed primarily of kerogen II/III and III types, indicating a higher contribution of the allochtonous biomass of land plants from the lake catchment, particularly in the lower part. The sediments of the upper sequence are enriched in autochthonous aquatic OM, which comprises mostly kerogen I, I/II and II types. The transition from the lower sequence to the upper one is associated with a decrease in pristane to phytane ratio, gammacerane index and content of  $C_{28}$  steroids, absence or significantly lower amount of squalane, absence of  $C_{24}$  and  $C_{25}$  regular isoprenoids, 8-methyl-2-methyl-2-(4,8,12-trimethyltridecyl) chroman and  $C_{30}$  hop-17(21)-ene.

Pyrolytic experiments showed that the sediments of the upper sequence, rich in aquatic OM, at a catagenetic stage could be a source of liquid hydrocarbons. The values of hopane, sterane and phenanthrene maturation parameters indicate that through pyrolysis at 400°C the samples investigated reached a value of vitrinite reflectance equivalent of approximately 0.70%. It was estimated that the sediments should be found at depths of 2300–2900 m in order to become active source rocks. The calculated minimum temperature, necessary for catagenetic hydrocarbon generation, is between 103 and 107°C.

Key words: Kremna Basin, lacustrine sediments, organic matter, mineralogy, biomarkers, pyrolysis.

Manuscript received 6 December 2013, accepted 17 September 2014



**Fig. 1.** Location and simplified geological map of Kremna Basin (modified after Basic Geologic Map of SFRJ, Užice Sheet, Mojsilović *et al.*, 1973), showing location of sampled ZLT-2 borehole.

## **INTRODUCTION**

Hydrology (water output and input of surface water, rainfall, and groundwater), sediment input, and temperature changes represent three major factors which control sedimentation patterns, and thus carbonate deposition in lake basins (Tucker and Wright, 1990; Platt and Wright, 1991). These factors are influenced by both climate and tectonics (Bohacs *et al.*, 2000, 2003). Considering the complexity of sedimentation in lake basins, an understanding of the sedimentation history requires a variety of sedimentological, palaeontological and geochemical analyses.

Bioproductivity in lacustrine environments can be as high as ten times, than in marine environments. Under such conditions, preservation of the organic matter can be favoured as well (Peters *et al.*, 2005). These two factors contribute to formation of sediments rich in organic matter (OM).

Despite of the influence of oxidation processes, which favour the OM degradation, organic carbon content (Corg) is often used as a preliminary palaeoproductivty indicator (Meyers, 1997). Rock-Eval pyrolysis is used as a rapid, preliminary method for examining the origin/type and maturity of the organic matter (Espitalié *et al.*, 1985).

The conditions of sedimentation determine the carbon to nitrogen ratio (C/N) in OM-rich sediments (Meyers, 1994; Mackie *et al.*, 2005). The C/N ratio lower than 10 is typical for algal material, which is depleted in cellulose in comparison to OM originated from higher plants (Meyers and Ishiwatari, 1993). Values of C/N ratio over 20 can be considered an indicator of a greater contribution of higher terrestrial plant material as a precursor for the OM (Meyers, 1994). However, an elevated C/N ratio also may be a consequence of the easier degradation of algal OM, rich in labile nitrogen, during diagenesis (Meyers, 1994).

Analysis of the extractable organic matter (bitumen) taken from sediments has been proven to be a promising tool for reconstruction of the sedimentary environment as well as the precursor biomass and its transformation during diagenesis. The analysis of bitumen is based on identification of finger print compounds (biomarkers) which are by structure strongly related to precursor biomolecules from organisms. Numerous parameters for biomarkers are used in the assessment of the origin, depositional environment and maturity of the OM (Peters et al., 2005, and references therein). Apart from the analyses mentioned, a full characterisation of the OM in sedimentary rocks also requires the investigation of the kerogen. In the case of an immature sample, its potential might be estimated by simulation of changes in organic matter maturity under laboratory conditions, using different pyrolytic experiments (Huizinga et al., 1988; Yoshioka and Ishiwatari, 2002; Parsi et al., 2007; Budinova et al., 2014, and references therein).

The objective of the study was to determine the organic-geochemical characteristics of sediments from the central part of the Kremna Basin. This basin is noteworthy, because of the specific palaeorelief, type of bedrock drainage and volcanic input that influenced the development of organic matter. For this purpose, the major-element content, bulk and specific organic-geochemical parameters, and qualitative mineralogical composition were investigated. Pyrolytic experiments were performed on bitumen-free samples in a detailed study of the *l*iquid-hydrocarbon potential of the sediments for prediction of the conditions necessary for them to become active oil-generating source rocks.

# **GEOLOGICAL SETTING**

The Zlatibor area is located in SW Serbia, about 200 km SW of Belgrade (Fig. 1). As a part of the Internal Dinarides, it consists mostly of a Dinaric ophiolite nappe, which was emplaced during the Jurrasic (Ilić and Neubauer, 2005). During the middle early Miocene, the Zlatibor area was affected by the opening of Dinaride Lake system (Krstić *et al.*, 2001). The relics of Miocene sediments are preserved only locally (the Kremna, Bioska, Mačkat, Kačer and Braneš basins), usually fault-bounded and interbedded with fine-grained volcanic tuffs and rarely with coal.

According to recent palaeontological investigations, the age of the sedimentary units is early Miocene, between 19 and 17 Ma (Prysjazhnjuk *et al.*, 2000; Krstić *et al.*, 2001). The only known fossil flora remains in the Kremna Basin were interpreted as early Miocene (Pantić, 1956). A rare fossil fauna (*Planorbis, Unio, Bithinia*) and ostracoda also were found, indicating the same age (Eremija, 1977). The whole Dinarides subsequently were affected by late Neogene and Quaternary uplift and denudation (Marović *et al.*, 1999, 2002).

Sedimentation in the Kačer and Bioska basins was dominated by clastic rocks, while in the Kremna Basin, carbonate rocks also were deposited in addition to clastics (Obradović and Vasić, 2007). This study is focused on the Kremna Basin, covering an area of approximately 15 km<sup>2</sup> (Fig. 1). The Kremna Basin was previously investigated for potential deposits of boron and magnesite as well as sepiolite and palygorskite clay (Živković and Stojanović, 1976; Dedić, 1978; Ilić and Rubežanin, 1978; Obradović *et al.*, 1994, 1995; Kovačević, 1998).

The bedrock that underwent erosion during basin evolution consisted primarily of Jurassic ultrabasic rocks, serpentinite and ophiolitic mélange, and Triassic carbonate rocks (Fig. 1). Intense weathering of ultramafic rocks during the Palaeogene formed a thick weathering crust of smectite (Maksimović, 1996). The weathering processes yielded intense leaching of MgO, which consequently influenced the formation of magnesite rich sedimentary rocks.

Obradović and Vasić (2007) summarized previous studies of the Kremna Basin and on the basis of the results presented by Dedić (1978), they distinguished alluvial and lacustrine series, and in them marginal and intrabasinal facies. The alluvial series consists of conglomerates and sand-stones made up of ultramafic rock fragments (Obradović and Vasić, 2007). Thin intercalations of coal (2–5 cm thick) are found in these sediments (Dedić, 1978). Marginal lacustrine and intrabasinal lacustrine facies consist of carbonate sediments, namely dolomite and/or magnesite, marlstones and rarely limestone (Obradović *et al.*, 1996).

## **MATERIALS**

For the purpose of this study, 43 core samples (5–10 cm in length) were taken from the ZLT-2 borehole at different depths. Sample preparation for chemical analyses was performed at the Institute for Technology of Nuclear and Mineral Raw Materials (ITNMS), Belgrade. The sediment samples were dried at 105°C. In the next step, the samples were

successively crushed to 2.36 mm in three stages, using a jaw crusher, a cone crusher and a roller crusher, respectively. Then the samples were homogenized. A rough mill-fragmented sample was subsequently finely pulverized and sifted through a 63  $\mu$ m sieve. Whole rock samples for thin sections and scanning electron microscopy (SEM) analyses were taken separately from the core.

#### **METHODS**

#### X-Ray diffraction analysis

X-ray diffraction analyses were performed in the Laboratory for Crystallography, Faculty of Mining and Geology, University of Belgrade, Serbia. For determination of the qualitative composition of the minerals, an X-ray generator PHILIPS Type PW 1729, and a diffractometer from the same manufacturer, type PW 1710 with genuine software processing (Philips APD), were used. As a source of radiation, an X-ray LLF-type tube with a copper anticathode and arched graphite monochromator placed between the sample and the detector were used, so that the radiation was  $CuK_{a1}$ = 1.5405 Å, thus avoiding possible X-ray fluorescence. The anode tube load was 40 kV and 35 mA. The gaps (slits) were fixed at 1.0 and 0.1 mm. Samples were pressed into standard aluminum frames and recorded in the range of  $2\Theta$ from 5° to 60°. Data were collected by measuring each of 1/50° (0.02°) for a duration of 0.5 sec. Identification of minerals was performed by comparing d<sub>hlk</sub> values with those of the standards from an electronic database (Joint Committee on Powder Diffraction Standards, JCPDS-International Centre for Diffraction Data).

#### Determination of contents of major and trace elements

The contents of major and trace elements were determined at AcmeLabs, in Canada. Prepared samples were mixed with an LiBO<sub>2</sub>/Li<sub>2</sub>B<sub>4</sub>O<sub>7</sub> flux. The crucibles were fused in a furnace. The cooled bead was dissolved in ACS-grade nitric acid. The amounts of Al<sub>2</sub>O<sub>3</sub>, CaO, Cr<sub>2</sub>O<sub>3</sub>, Fe<sub>2</sub>O<sub>3</sub>, K<sub>2</sub>O, MgO, MnO, Na<sub>2</sub>O, P<sub>2</sub>O<sub>5</sub>, SiO<sub>2</sub> and TiO<sub>2</sub> were determined by inductively coupled plasma-optical emission spectroscopy (ICP-OES). The amounts of Ba, Co, Cs, Rb, Sb, Sr, Th, U, V, Y and Zr were determined by inductively coupled plasma mass spectrometry (ICP-MS). The ICP-OES analysis was performed, using a SPECTRO ARCOS instrument, while for ICP-MS analysis an ELAN 9000 from Perkin Elmer was employed. A rack of 40 samples on either instrument comprised 36 samples, one analytical blank, one sample replicate, one internal reference material (IRM) and one of the Certified Reference Materials (CRM). Loss on ignition (LOI) was determined by igniting a sample split, then measuring the weight loss. Additionally, for the screening of possible occurrences of B and Li, the amounts of these two elements, together with the percentages of major elements and Sr (due to correlation) in core samples at 10 m intervals, were also determined. These analyses were performed in the Société Générale de Surveillance (SGS) Laboratory, Canada, following procedures that were described in previous papers (Šajnović et al., 2009, 2012).

#### Scanning electron microscopy (SEM)

SEM analyses were performed on gold-coated, polished thin sections at the Faculty of Mining and Geology, University of Belgrade, Serbia. Scanning electron microscopy was used for imaging and collecting chemical data, using a JEOL JSM–6610 LV Scanning Electron Microscope, equipped with an energy-dispersive spectrometer (EDS). Analyses were run at an accelerating voltage of 20 kV, and a working distance of 10 mm.

#### **Rock Eval pyrolysis**

Rock-Eval pyrolysis was performed at the Institute of Geology and Geochemistry of Petroleum and Coal, Aachen University, Germany. Depending on mineral composition, the amounts of macro-elements, the organic carbon content and the biomarker composition, twenty-five samples were selected for Rock-Eval pyrolysis. Measurements were carried out, using a DELSI INC Rock-Eval VI instrument, according to guidelines published by Espitalié *et al.* (1985) and Lafargue *et al.* (1998). About 100 mg of each powdered sample were pyrolyzed.

#### Determination of organic carbon, sulphur and nitrogen contents and bitumen analysis

Analyses were performed at the Faculty of Chemistry, University of Belgrade, Serbia. Elemental analysis was applied to determine the amounts of carbon, sulphur and nitrogen. Organic carbon (Corg) was determined after the removal of carbonates with diluted hydrochloric acid (1:3 v/v). The measurements were performed, using a Vario EL III, CHNOS Elemental Analyser, Elementar Analysensysteme GmbH (CHNS operation mode: furnace temperature 1150°C with a thermoconductivity detector, TCD).

Extractable organic matter (bitumen) was removed from the sediments by Soxhlet's extraction with an azeotrope mixture of dichloromethane and methanol (88:12 volume %), for 42 h (Šajnović *et al.*, 2009, 2012). Separation of the bitumen into the saturated and aromatic hydrocarbon fractions was achieved, using column chromatography (adsorbents: SiO<sub>2</sub>, 2.88 g and Al<sub>2</sub>O<sub>3</sub>, 2.12 g; eluents, including a "dead volume" of 1.2 cm<sup>3</sup>: *n*-hexane, 50 cm<sup>3</sup> and benzene, 25 cm<sup>3</sup>; per 10 mg of bitumen).

The saturated and aromatic fractions, isolated from the initial bitumen and liquid pyrolysates, were analyzed by means of the gas chromatography-mass spectrometry (GC- MS) techique. The GC-MS was performed using an Agilent 7890A gas chromatograph (HP-5MS column, 30 m × 0.25 mm, 0.25  $\mu$ m film thickness, He carrier gas 1.5 cm<sup>3</sup>/min), coupled to an Agilent 5975C mass selective detector (70 eV). The column was heated from 80 to 310°C, at a rate of 2°C/min., and the final temperature of 310°C was maintained for an additional 25 minutes. The individual peaks were identified on the basis of comparison of the mass spectrum obtained with those from the literature (Risatti *et al.*, 1984; Philp, 1985; Radke, 1987; Sinninghe Damsté *et al.*, 1987; Koopmans *et al.*, 1996a; Peters *et al.*, 2005) and the mass spectra library (NIST5a). Quantification of amounts of the com-

pounds for calculating the molecular parameters was performed by integration of peak areas (software GCMS Data Analysis) in the appropriate mass chromatograms [m/z 71 for *n*-alkanes; m/z 217 for steranes; m/z 215 for sterenes; m/z for 191 hopanes and m/z 121, 135 and 149 for mono-, di- and trimethylated 2-methyl-2-(4,8,12-trimethyltridecyl) chromans, respectively]. Methyl-, dimethyl- and trimethylnaphthalenes in the aromatic fractions of liquid pyrolysates were identified from the m/z 142, 156, and 170 ion fragmentograms, whereas phenanthrene, methyl-, and dimethylphenanthrene isomers were analyzed from the m/z 178, 192, and 206 ion fragmentograms.

#### Pyrolysis of the samples rich in organic matter

Pyrolysis was performed on samples free of soluble organic matter (bitumen), which contained kerogen with a native mineral matrix. The initial mass of a bitumen-free sample was ~5 g. Pyrolyses were performed in an autoclave under nitrogen for 4 hours at 400°C. Liquid pyrolysis products were extracted with hot chloroform. The gaseous products were not analyzed, although the production of gaseous products was indicated by the pressure change in the autoclave. Liquid pyrolysates were separated into saturated hydrocarbon, aromatic hydrocarbon, and NSO fractions using the same method as applied in the fractionation of extracted bitumen. The saturated and aromatic fractions were analyzed by GC-MS, as described in the previous section.

## RESULTS

#### Mineral composition and content of major elements

The chemical composition of the sediments from the Kremna Basin was determined for 43 core samples (Tables 1–3). Samples for mineralogical and petrological analyses (thin section and SEM) were selected with reference to the analyses obtained.

Generally, the mineral composition of the Kremna Basin sediments is represented by dolomite, quartz, calcite, magnesite, and clay minerals. Different lacustrine facies could not be clearly distinguished from the results obtained. However, it was evident that the sediments from the core analyzed belong to an intrabasinal facies (Figs 2, 3).

Sediments from the lower sequence in the core (216– 343 m) contain more Mg-rich minerals (Fig. 3B) and the Mg originated from the underlying ultramafites. This section is characterised by pronounced volcanic activity (tuff, searlesite; Fig. 3C). The spherulitic structure, common in silica-rich glassy rocks, is visible in thin (20–30 cm) potassium-rich unaltered volcanic material (depth 262 m; Figs 2, 4A, B). The glass is occasionally devitrified and the spherulites are recrystallised. Clay minerals are poorly represented in the entire column, except in intervals where sepiolite and smectite occur by paragenesis with dolomite and magnesite (Fig. 3D). Ca-rich sediments are dominant in the upper sequence of the core (13.5–216 m, Fig. 3A). Volcanic material was also determined in a thin layer at a depth of 77.5 m (Figs 2, 4C, D).

Amounts of major elements (weight %) in sediments

Sequence	Descrip- tion	No	Depth (m)	$\mathrm{SiO}_2$	Al <sub>2</sub> O <sub>3</sub>	Fe <sub>2</sub> O <sub>3</sub>	MgO	CaO	Na <sub>2</sub> O	K <sub>2</sub> O	TiO <sub>2</sub>	P <sub>2</sub> O <sub>5</sub>	MnO	Cr <sub>2</sub> O <sub>3</sub>	LOI*
Clayey c	arbonates	1	11.5	55.57	0.94	1.03	4.97	8.88	0.12	0.47	0.04	0.01	0.01	0.02	28.6
		2	13.5	8.40	1.02	0.81	16.19	22.19	0.09	0.56	0.03	0.01	0.02	0.02	50.7
		3	27	17.44	3.03	1.31	14.46	21.58	0.11	1.42	0.07	0.02	0.03	0.02	40.5
		4	32	12.62	1.87	0.69	19.4	22.43	0.11	1.13	0.04	0.01	0.02	0.01	41.8
	Marly dolomite	5	42.5	15.02	3.04	1.44	2.15	37.64	0.07	1.51	0.06	0.02	0.05	0.01	39.4
	uoronnie	6	54	34.79	6.80	2.33	11.15	9.24	0.08	2.63	0.12	0.01	0.03	0.02	34.3
		7	55.5	26.09	1.97	0.81	15.63	18.86	0.08	0.67	0.04	0.03	0.03	0.01	36.6
		8	64.5	17.58	0.61	0.21	17.1	23.15	0.08	0.18	0.01	0.01	0.01	0	41
		9	70	30.34	6.04	1.80	2.27	27.7	0.06	1.47	0.09	0.01	0.04	0.02	31.7
		10	78	10.24	2.09	0.69	11.21	32.77	0.13	0.47	0.04	0.01	0.02	0	42.5
Upper		11	80	10.68	2.38	0.53	6.08	36.77	0.11	0.36	0.05	0.02	0.02	0	43.6
(US)		12	83	11.99	3.31	0.88	2.43	41.81	0.15	0.41	0.07	0.04	0.02	0.01	39.9
		13	96	10.98	3.18	1.44	1.50	42.84	0.13	0.55	0.08	0.11	0.03	0.01	39.9
	Marlstone,	14	111	12.45	2.59	0.80	2.52	36.33	0.14	0.53	0.05	0.15	0.02	0.01	45
	calcite	15	113	22.39	5.46	2.68	1.47	35.35	0.20	1.19	0.28	0.04	0.06	0.04	31.9
	dominated	16	127	18.28	5.18	2.57	1.69	38.01	0.25	1.12	0.14	0.04	0.04	0.03	33.6
		17	137.5	6.98	1.18	0.63	2.66	44.58	0.15	0.24	0.03	0.06	0.01	0.01	43.6
		18	150	27.69	1.87	0.87	1.09	32.43	0.16	0.39	0.13	0.01	0.02	0.02	36.6
		19	164	13.99	1.82	1.16	1.21	43.88	0.20	0.41	0.07	0.12	0.03	0.02	37.7
		20	185	17.16	1.04	0.76	7.51	29.38	0.23	0.36	0.03	0.07	0.02	0.01	43.8
		21	189.5	23	1.39	1.20	8.88	26.89	0.22	0.71	0.04	0.05	0.03	0.02	37.7
		22	216	36.81	0.31	0.96	21.41	13.46	0.41	0.10	0.02	0.01	0.01	0.02	28.3
		23	219	12.91	0.85	1.24	35.35	5.99	0.29	0.29	0.02	0.01	0.02	0.03	42.8
		24	224	24.05	1.79	2.51	20.39	17.10	0.45	0.80	0.06	0.01	0.05	0.05	33.8
		25	238	46.93	4.89	1.17	24.44	2.36	1.24	2.79	0.03	0.01	0.02	0.01	20.4
	Marly	26	243.5	16.61	0.94	1.19	32.75	6.29	1.19	0.48	0.04	0.01	0.02	0.02	40.6
	magnesite	27	245	16.44	0.73	2.66	35.98	4.79	0.69	0.27	0.03	0.01	0.04	0.08	39.1
		28	248.3	19.39	0.66	1.34	30.28	6.50	2.28	0.29	0.03	0.01	0.02	0.03	39.4
		29	255	12.26	0.60	1.20	36.68	4.58	0.70	0.22	0.03	0.01	0.03	0.05	43.8
		30	258	40.14	3.40	8.85	20.92	6.15	1.71	1.60	0.11	0.01	0.11	0.19	21
		31	265	15.93	1.30	1.95	33.62	5.84	1.40	0.62	0.05	0.01	0.03	0.03	39.9
Lower		32	283	20.75	1.35	1.89	26.3	13.79	1.16	0.45	0.05	0.01	0.03	0.04	35.7
(LS)		33	286	29.18	2.27	3.46	24.87	11.49	1.56	0.66	0.08	0.02	0.04	0.06	29
		34	297.5	23.76	2.09	1.92	28.62	9.19	1.53	0.44	0.07	0.01	0.03	0.03	34.3
		35	309	33.83	4.62	5.93	21.84	9.25	2.23	1.23	0.19	0.02	0.07	0.08	24.1
	dolomite.	36	317.5	22.68	2.19	2.72	25.39	12.59	1.45	0.53	0.08	0.01	0.04	0.05	33.7
	Silty	37	324	34.6	3.65	5.08	23.11	7.70	1.98	1.07	0.12	0.02	0.06	0.1	25.7
	Mg-marls-	38	329	34.22	4.05	5.51	19.05	10.76	1.83	1.61	0.13	0.02	0.06	0.11	25.6
	Mg-clay	39	335	27.27	0.06	0.15	23.9	13.48	2.11	0.07	0.01	0.02	0.07	0.01	35.8
		40	336	23.11	0.28	0.19	22.58	14.3	2.10	0.07	0.01	0.02	0.04	0.01	39.6
		41	340	43.08	1.51	5.53	22.88	2.96	1.9	0.34	0.04	0.02	0.03	0.1	27.2
		42	341	46.15	2.37	8.17	20.53	2.93	2.22	0.53	0.07	0.02	0.04	0.15	22.8
		43	343	52.4	4.39	11.55	16.66	0.68	2.12	2.39	0.12	0.02	0.03	0.2	15.6
Minimum	US			6.98	0.61	0.21	1.09	9.24	0.06	0.18	0.01	0.01	0.01	0	31.7
Maximun	n US			34.79	6.8	2.68	19.4	44.58	0.25	2.63	0.28	0.15	0.06	0.04	50.7
Average US			17.41	2.79	1.18	7.33	31.19	0.14	0.82	0.07	0.04	0.03	0.01	39.59	
Standard deviation US			7.71	1.77	0.69	6.37	9.57	0.06	0.6	0.06	0.04	0.01	0.01	4.75	
Minimum LS			12.26	0.06	0.15	16.66	0.68	0.29	0.07	0.01	0.01	0.02	0.01	15.6	
Maximum LS			52.4	4.89	11.55	36.68	17.1	2.28	2.79	0.19	0.02	0.11	0.2	43.8	
Average LS				28.75	2.01	3.42	25.8	8.28	1.48	0.77	0.06	0.01	0.04	0.07	31.74
Standard	deviation L	s		11.83	1.52	3.05	5.89	4.55	0.64	0.74	0.05	0.01	0.02	0.06	8.14

\*-Loss on ignition.

Amounts of trace elements (mg/kg) in sediments

Sequence	Descrip- tion	No	Depth (m)	Ba	Со	Cs	Rb	Sc	Sr	Th	U	V	Zr
Clayey ca	arbonates	1	11.5	109.2	8.90	1.11	12.74	2.02	190.7	0.71	0.61	13.14	12.84
		2	13.5	108.89	12.10	2.42	19.66	2.02	906.53	0.50	1.11	34.28	6.15
		3	27	451.39	10.88	11.79	70.33	3.02	815.82	1.41	0.81	20.15	24.08
		4	32	233.55	5.26	2.63	29.22	2.02	1227.88	1.72	0.81	12.13	6.77
	Marly	5	42.5	171.73	8.79	6.26	60.31	3.03	875.04	2.42	1.21	18.18	19.6
	doioinne	6	54	201.46	14.49	10.79	116.76	4.11	984.89	2.06	0.51	39.06	31.86
		7	55.5	153.83	5.91	4.28	35.35	2.04	773.63	2.55	2.24	16.30	10.59
		8	64.5	93.51	1.21	1.41	9.85	0.50	794.57	0.40	0.80	8.04	9.15
		9	70	307.76	10.57	16.82	82.17	4.10	1521.65	3.49	1.85	18.47	36.32
		10	78	515.36	3.44	7.68	25.97	2.02	1232.52	3.13	3.13	10.11	23.65
Upper		11	80	388.01	2.23	6.09	24.58	1.02	926.26	1.93	2.03	14.22	32.61
sequence (US)		12	83	277.61	3.78	13.47	33.68	2.04	1404.16	2.55	1.33	15.31	14.9
(05)		13	96	437.44	10.27	19.43	37.13	3.05	1291.05	2.75	1.73	19.33	14.34
	Marlstone	14	111	258.83	6.60	13.20	32.07	2.03	1144.74	2.33	2.84	16.24	14.41
	calcite	15	113	415.01	19.58	32.83	79.74	7.14	1258.59	7.85	3.26	61.18	35.28
	dominated	16	127	371.24	20.40	47.42	78.53	6.12	1707.29	3.47	1.84	41.82	27.03
		17	137.5	226.72	5.95	16.22	16.02	1.01	1193.97	0.91	1.11	12.09	7.76
		18	150	263.97	11.56	3.17	20.46	2.05	701.04	1.64	0.61	18.42	19.85
		19	164	247.16	11.75	1.72	19.65	3.04	944.89	1.72	1.32	17.22	16.00
		20	185	171.79	8.08	2.83	16.98	1.01	659.36	1.52	2.22	17.18	8.19
		21	189.5	149.99	11.88	2.92	31.11	2.01	600.56	1.31	1.51	17.11	10.57
		22	216	82.59	10.94	0.31	4.96	1.03	1316.13	0.10	0.31	8.26	5.27
		23	219	134 59	15.69	0.81	12.85	2.02	2197 53	0.30	0.40	9.11	5.67
		23	219	338.81	29.88	1 44	26.49	4 11	2510.78	1 33	0.92	21.56	13.96
		25	238	108.22	7 75	0.53	19.31	1.06	861.64	0.74	0.42	4 24	21.96
	Marly	25	243.5	121.83	15.63	1.02	12 79	2.03	1484.67	0.51	0.51	12.18	8 32
	laminated	20	245.5	69.89	32.99	0.82	10.69	3.08	1276 57	0.21	0.51	18.50	5.45
	magnesite	28	248.3	117.96	20.24	1.12	11.80	2.03	1869.33	0.21	0.61	10.17	4 47
		20	255	105.73	16.98	0.92	9.86	1.02	1514.95	0.20	0.01	11.18	4 27
		30	255	110.46	05.17	5.10	50.03	8.50	720.34	0.51	0.41	33.00	21.77
		31	258	82.85	21.70	2.56	17.80	2.05	11/7 90	0.04	0.04	11.25	0.41
Lower		32	203	176.30	18.00	10.69	18.16	3.11	2002.01	0.83	0.83	14.53	10.27
sequence		32	285	153.10	20.78	21.20	20.05	4.10	2392.01	0.83	0.63	20.07	18.67
(LS)		33	200	133.10	12.70	24.24	10.77	2.08	1557.05	0.04	0.03	16.65	20.44
		25	297.5	174.03	13.42	60.01	19.77	6.22	2201 12	1.48	0.51	20.50	27.44
	Marly Mg-	36	217.5	228.64	25.25	24.25	25.07	4.14	2301.13	0.82	0.53	21.72	15.62
	dolomite,	30	324	80.44	54.02	02.42	61.66	5.26	1174 56	0.85	0.32	25.77	24.21
	Silty Mg-marl-	28	324	02.25	50.44	93.43	51.58	7.24	1244.30	0.74	0.42	38.70	24.51
	stone,	20	225	92.23	1 4 2	94.24	1 16	0.52	1244.10	0.75	0.75	4.22	14.74
	Mg-clay	39	226	183.34	4.43	0.93	0.72	0.53	1062.28	0.84	0.55	4.22	14.30 9.47
		40	240	185.02	40.22	0.42	0.73	0.52 5.42	1005.58	1.30	0.42	4.18	0.11
		41	240	39.02	40.33	4.44	8.02	5.42	123.58	0.87	0.43	/6.96	9.11
		42	341	32.58	54.75	5.54	11.84	8.69	105.12	1.41	0.43	83.61	12.16
NC 1 1		43	343	17.29	64.4	12.75	52.62	11.89	24.53	1.19	0.22	106.97	19.45
Minimum	05			93.51	1.21	1.41	9.85	0.50	000.56	0.40	0.51	8.04	0.15
Maximum	08			515.36	20.40	47.42	116.76	/.14	1/0/.29	7.85	3.26	61.18	36.32
Average U	5			272.26	9.24	11.17	41.98	2.67	1048.22	2.28	1.61	21.34	18.46
Standard deviation US			121.53	5.21	11.62	29.00	1.67	301.81	1.57	0.81	12.95	9.97	
Minimum LS			17.29	2.20	0.31	0.73	0.52	24.53	0.10	0.22	4.18	4.27	
Maximum LS			338.81	95.17	94.24	61.66	11.89	3252.12	1.48	1.57	106.97	37.23	
Average LS				126.06	31.04	17.13	23.00	3.93	1460.72	0.76	0.56	27.06	14.75
Standard d	eviation LS			71.25	23.29	28.74	18.29	3.05	866.2	0.42	0.28	27.63	8.98

# Amounts of major (weight %) and trace elements B, Li, Sr (mg/kg) in core samples at 10-m intervals

													-			
Sequence	Descrip- tion	Depth (m)	$SiO_2$	Al <sub>2</sub> O <sub>3</sub>	Fe <sub>2</sub> O <sub>3</sub>	MgO	CaO	Na <sub>2</sub> O	K <sub>2</sub> O	TiO <sub>2</sub>	$P_2O_5$	MnO	Cr <sub>2</sub> O <sub>3</sub>	В	Li	Sr
		10-20	12.10	0.62	0.60	18.40	25.50	0.16	0.24	0.02	0.01	0.02	0.02	95	290	1200
		20-30	9.61	1.60	1.05	9.19	34.50	0.10	0.73	0.05	0.01	0.03	0.03	87	84	1300
	Marly	30-40	22.20	2.47	1.14	17.80	18.80	0.16	1.47	0.04	0.01	0.03	0.01	73	460	1200
	dolomite	40-50	12.80	1.64	0.97	11.90	30.80	0.11	0.75	0.04	0.02	0.04	0.02	140	130	1000
		50-60	18.00	2.37	0.94	12.50	25.80	0.12	1.11	0.05	0.01	0.03	0.03	96	210	860
		60-70	14.30	2.21	0.98	7.48	33.00	0.12	0.63	0.05	0.01	0.05	0.02	110	110	1300
		70-80	24.40	2.11	0.60	10.40	25.10	0.12	1.18	0.03	0.01	0.02	0.03	71	81	740
		80-90	7.82	1.46	0.46	2.97	43.40	0.25	0.24	0.03	0.05	0.03	0.01	69	34	1700
		90-100	8.64	0.83	1.03	13.60	32.10	0.32	0.25	0.03	0.04	0.03	0.02	280	85	1400
Upper		100-110	5.74	0.91	0.44	2.68	45.80	0.20	0.18	0.03	0.12	0.02	0.01	100	20	1600
sequence (US)		110-120	19.00	3.10	1.05	3.75	34.30	0.19	0.61	0.07	0.08	0.03	0.02	81	27	1400
(05)		120-130	9.15	1.54	0.94	4.13	40.40	0.25	0.37	0.04	0.11	0.03	0.01	62	20	1300
	Marlstone,	130-140	10.00	0.99	0.78	5.40	39.00	0.14	0.23	0.03	0.08	0.04	0.02	57	20	1200
	calcite dominated	140-150	12.40	1.55	0.96	2.57	41.10	0.23	0.35	0.05	0.08	0.04	0.02	72	20	1400
	uommuteu	150-160	12.20	1.79	0.96	6.71	35.80	0.19	0.41	0.05	0.05	0.04	0.02	39	20	580
		160-170	14.00	1.35	0.86	6.94	35.00	0.17	0.32	0.04	0.06	0.04	0.01	52	21	610
		170-180	13.50	1.67	0.94	6.41	35.90	0.35	0.46	0.04	0.04	0.02	0.02	76	33	630
		180-190	16.70	1.84	1.24	7.68	32.30	0.27	0.54	0.06	0.03	0.04	0.03	76	50	620
		190-200	44.90	7.33	1.63	4.42	14.00	1.78	1.90	0.12	0.05	0.06	0.03	77	20	320
		200-210	12.60	1.47	1.42	7.38	36.30	0.19	0.57	0.06	0.01	0.04	0.03	74	62	670
		210-220	21.70	1.92	1.64	20.80	14.80	2.04	0.71	0.07	0.01	0.03	0.02	4400	390	3300
		220-230	13.40	0.78	1.22	34.10	6.95	0.31	0.30	0.03	0.01	0.03	0.04	300	190	1500
		230-235	13.10	1.02	1.23	33.30	7.22	0.36	0.44	0.03	0.01	0.03	0.03	190	170	1300
	Morly	235-240	12.10	0.66	0.75	36.30	5.58	0.35	0.24	0.02	0.01	0.02	0.03	320	140	1600
	laminated	240-245	14.50	0.93	2.08	32.80	6.31	0.52	0.39	0.03	0.01	0.03	0.05	680	160	1300
	magnesite	245-250	14.90	0.88	1.89	29.90	8.90	0.99	0.41	0.04	0.01	0.03	0.05	2100	130	1200
		250-255	13.60	0.91	1.33	31.60	8.50	0.82	0.36	0.04	0.01	0.03	0.04	1600	110	1200
Tanan		255-260	16.70	1.28	1.64	29.10	8.39	1.39	0.59	0.04	0.01	0.04	0.04	2900	130	1300
sequence		260-265	18.30	1.96	1.64	26.80	9.22	1.48	1.15	0.06	0.01	0.04	0.04	3700	140	1100
(LS)		265-271	10.80	0.75	0.90	28.70	12.70	1.10	0.37	0.03	0.01	0.03	0.03	2600	150	2300
		271-281	15.00	0.86	1.08	24.70	16.60	0.99	0.32	0.04	0.02	0.03	0.02	1600	170	3400
	Marly Mg-	281-291	18.70	1.13	1.57	25.20	14.50	1.02	0.34	0.05	0.01	0.03	0.03	1300	270	2500
	dolomite, Silty	291-301	18.70	1.34	1.40	26.70	12.30	1.15	0.32	0.06	0.01	0.03	0.03	1000	440	1900
	Mg-marl-	301-311	12.50	0.88	1.15	23.20	18.80	0.17	0.37	0.03	0.02	0.03	0.03	140	120	1200
	stone,	311-321	16.80	1.20	1.49	23.50	17.00	0.99	0.31	0.04	0.01	0.04	0.04	810	400	4200
	Mg-clay	321-331	16.30	0.90	1.17	22.60	18.10	1.04	0.31	0.03	0.01	0.03	0.03	1400	420	2500
		331-342	19.60	0.90	2.17	31.40	5.62	0.97	0.31	0.03	0.01	0.03	0.05	830	220	480
Minimum	US		5.74	0.62	0.44	2.57	14.00	0.10	0.18	0.02	0.01	0.02	0.01	39	20	320
Maximun	n US		44.90	7.33	1.63	18.40	39.00	1.78	1.90	0.12	0.12	0.06	0.03	280	460	1700
Average I	US		15.00	1.94	0.95	8.12	30.25	0.27	0.63	0.05	0.04	0.03	0.02	89	90	1052
Standard	deviation		8.48	1.40	0.29	4.70	7.96	0.36	0.46	0.02	0.04	0.01	0.01	50	112	393
Minimum	LS		10.80	0.66	0.75	20.80	5.58	0.17	0.24	0.02	0.01	0.02	0.02	140	110	480
Maximun	n LS		21.70	1.96	2.17	36.30	18.80	2.04	1.15	0.07	0.02	0.04	0.05	4400	440	4200
Average I	LS		15.69	1.08	1.43	28.28	11.26	0.92	0.43	0.04	0.01	0.03	0.04	1522	221	1899
Standard LS	deviation		3.01	0.37	0.39	4.55	4.61	0.48	0.22	0.01	0.00	0.00	0.01	1252	117	995

	Fac	ies	Lithology	Description	Reconstruction of s	edimentation history
,	Allu	vial	~~~~~~~	Clayey carbonates		
		<sup>G</sup> Depth (m)		Thin-bedded marly dolomite		
Lacustrine intrabasinal	Upper sequence	100		Thin tuff layer at 77.5 m Thin-bedded marlstone, calcite-dominated	Dominance of calcite and dolomite, more humid conditions, freshwater environment, palaeoproductivity increased, better preservation of organic matter	Higher water level, dominance of the aquatic organic matter, seasonal algal blooms
é		200		Manha Mar dalamata		
	uence	250		Marly laminated magne- site with searlesite veinlets and, rarely, tuff layers (0.03–0.06 m) rich in searlesite	Dominance of magnesite	Sedimentation in shallow water, chemical precipitation of carbonates, dominance of the aquatic organic matter
	Lower seq	300 -		Marly Mg-dolomite with thin layers of silty Mg-marlstone Mg-clay and silty Mg-marlstone Weathered serpentinite	volcanic activity, presence of searlesite, low palaeoproductivity	Sedimentation in shallow water, dominance of the allochtonous organic matter from the lake catchment

Fig. 2. Lithological column of ZLT-2 borehole.

The mineral content of the sediments from the Kremna Basin is reflected in their chemical composition. The lower sequence (216-343 m) has higher MgO, Na<sub>2</sub>O, Fe<sub>2</sub>O<sub>3</sub>, SiO<sub>2</sub>, B and Li contents, while the upper sequence (13.5-216 m) has a higher CaO content (Tables 1, 3). The entire core has a very low Al<sub>2</sub>O<sub>3</sub> content (0.06-6.80%; Table 1), owing to the low clay content and a specific clay mineralogy with Mg clays predominant.

## Bulk organic geochemical parameters and Rock Eval data

The OM content, expressed as total organic carbon (Corg), varies across a wide range of 0.32 to 12.38% (Table 4). The same observation is applicable to the amounts of extrac- table organic matter (bitumen) and both free (S1) and pyrolysis-derived bound hydrocarbons (S2), obtained by Rock-Eval pyrolysis (Tables 4, 5). Samples from the upper sequence are characterised by enhanced Corg, S1, S2 and nitrogen contents (Tables 4, 5). These results indicate that variations



Fig. 3. Characteristic X-ray diffractograms of sediments from the Kremna Basin.



**Fig. 4.** Petrographic analyses of volcanic material from the Kremna Basin: (**A**) thin-section image of volcanic material containing biotite and glass, 262 m; (**B**) thin-section image of the same sample as under (A) showing sulphide minerals as centers of spherulites; (**C**) thin-section image of volcanic material with lamination and silica minerals in lenses, 77.5 m; (**D**) SEM image of the same sample as under (C).

in mineral proportions are followed by a notable change in the OM content. The elevated concentration of OM in the lower sequence, detected only in the depth interval 329 to 336 m (Table 4) can be related to coal layers (allochtonous OM, derived from the biomass of land plants), observed on a macroscopic scale during the sampling.



**Fig. 5.** Modified van Krevelen diagram Hydrogen Index (HI) *vs.* Oxygen Index (OI) (Langford and Blanc-Valleron, 1990).

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Values of bulk organic geochemical parameters

# Table 4

					2	0	1				
Sequence	Description	No	Depth (m)	Corg (%) <sup>1</sup>	N (%)	S (%)	$C/S^2$	C/N <sup>3</sup>	Bitumen (mg/g Corg)	HC <sup>4</sup> (mg/g Corg)	Asphaltenes + NSO <sup>5</sup> (mg/g Corg)
Clayey ca	arbonates	1	11.5	10.64	0.31	0.97	29.28	40.03	117.11	3.44	113.67
		2	13.5	4.13	0.16	0.38	29.02	30.10	334.79	5.69	329.10
		3	27	12.07	0.35	1.20	26.85	40.22	59.93	1.20	58.73
		4	32	4.01	0.16	0.37	28.93	29.23	112.89	2.61	110.29
	Marly	5	42.5	5.71	0.28	0.77	19.80	23.78	95.02	5.63	89.40
	doioinite	6	54	11.04	0.44	1.05	28.07	29.26	134.26	7.56	126.70
		7	55.5	5.52	0.20	0.47	31.36	32.19	122.10	6.85	115.25
		8	64.5	3.03	0.07	0.24	33.71	50.48	78.35	4.67	73.68
		9	70	4.72	0.14	0.78	16.16	39.32	111.41	5.90	105.51
		10	78	3.24	0.11	0.37	23.38	34.35	93.18	5.19	87.99
Upper		11	80	5.78	0.17	0.57	27.07	39.65	124.6	5.69	118.91
(US)		12	83	2.65	0.12	0.50	14.15	25.75	90.2	4.78	85.42
		13	96	3.62	0.16	0.76	12.72	26.38	80.93	3.70	77.23
	Marlstone,	14	111	10.23	0.33	0.95	28.75	36.15	135.51	5.76	129.75
	calcite	15	113	1.58	0.11	0.70	6.03	16.75	136.27	5.44	130.83
	dominated	16	127	1.45	0.08	0.57	6.79	21.14	94.19	4.08	90.11
		17	137.5	6.30	0.20	0.75	22.43	36.73	74.38	3.17	71.21
		18	150	5.71	0.26	0.51	29.89	25.61	33.58	1.35	32.23
		19	164	1.71	0.07	0.31	14.73	28.49	88.03	4.14	83.89
		20	185	8.86	0.27	0.71	33.32	38.27	108.82	5.88	102.95
		21	189.5	5.81	0.23	0.61	25.43	29.46	31.87	1.37	30.51
	-	22	216	1.42	0.06	0.15	25.27	27.60	183.71	8.08	175.63
		23	219	2.07	0.08	0.30	18.42	30.17	213.71	8.29	205.42
		24	224	0.32	0.10	0.56	1.53	3.73	276.69	13.56	263.13
		25	238	0.46	0.06	0.54	2.27	8.94	270.49	11.33	259.16
	Marly laminated	26	243.5	2.14	0.08	0.50	11.43	31.2	213.75	10.58	203.17
	magnesite	27	245	1.04	0.03	0.55	5.05	40.43	266.2	5.14	261.06
		28	248.3	3.02	0.09	0.48	16.80	39.13	329.45	10.51	318.94
		29	255	2.40	0.06	0.42	15.26	46.65	323.46	11.39	312.08
		30	258	0.81	0.07	1.83	1.18	13.49	158.19	5.33	152.86
		31	265	1.37	0.04	0.58	6.31	39.94	218.77	16.39	202.39
Lower		32	283	1.23	0.04	0.44	7.46	35.86	270.12	7.89	262.23
(LS)		33	286	0.88	0.03	0.60	3.92	34.21	262.38	12.10	250.28
		34	297.5	1.69	0.06	0.41	11.00	32.85	326.03	25.46	300.57
		35	309	0.51	0.04	0.97	1.40	14.87	251.39	10.26	241.13
	dolomite.	36	317.5	1.48	0.06	0.48	8.23	28.77	389.65	13.13	376.52
	Silty	37	324	2.08	0.08	0.83	6.69	30.32	336.19	17.25	318.95
	Mg-marl-	38	329	12.38	0.48	3.88	8.51	29.74	40.41	2.55	37.86
	Mg-clay	39	335	4.50	0.20	0.30	40.05	26.24	81.17	4.69	76.48
		40	336	5.88	0.26	0.37	42.43	26.37	80.37	2.61	77.76
		41	340	2.69	0.12	1.37	5.24	26.14	89.33	4.63	84.70
		42	341	1.89	0.08	1.81	2.79	27.55	89.27	6.65	82.62
		43	343	0.77	0.06	2.50	0.82	14.97	81.95	3.96	77.99
Minimum U	JS			1.45	0.07	0.24	6.03	16.75	31.87	1.20	30.51
Maximum U	JS			12.07	0.44	1.20	33.71	50.48	334.79	7.56	329.1
Average US	8			5.36	0.20	0.63	22.93	31.67	107.02	4.53	102.48
Standard de	viation US			3.09	0.10	0.25	8.48	7.85	61.50	1.82	60.54
Minimum L	.S			0.32	0.03	0.15	0.82	3.73	40.41	2.55	37.86
Maximum I	LS			12.38	0.48	3.88	42.43	46.65	389.65	25.46	376.52
Average LS	5			2.32	0.10	0.90	11.00	27.69	216.03	9.63	206.40
Standard de	viation LS			2.61	0.10	0.89	11.66	10.76	102.01	5.49	98.04

 $^{1}$  Corg – Organic carbon content;  $^{2}$  C/S is given as molar ratio;  $^{3}$  C/N is given as molar ratio;  $^{4}$  HC – Content of saturated and aromatic hydrocarbons;  $^{5}$  NSO – Polar compounds containing N, S and O.

s

Sequence	Description	No	Depth (m)	S1 <sup>1</sup> (mg HC <sup>2</sup> /g sample)	S2 <sup>3</sup> (mg HC/g sample)	$S3^4$ (mg $CO_2/g$ sample)	HI <sup>5</sup> (mg HC/ (g Corg)	OI <sup>6</sup> (mg CO <sub>2</sub> / g Corg) <sup>5</sup>	S2/S3	PI <sup>7</sup>	Tmax (°C) <sup>8</sup>
		3	27	2.35	53.42	2.69	442.58	22.29	19.86	0.04	430
	Marla dalamita	6	54	7.78	112.31	3.93	1017.3	35.60	28.58	0.06	425
	Marly dolomite	7	55.5	3.51	41.96	1.73	760.14	31.34	24.25	0.08	430
		8	64.5	1.01	25.69	1.31	847.85	43.23	19.61	0.04	428
		9	70	1.35	33.64	1.67	712.71	35.38	20.14	0.04	427
Upper		10	78	0.63	24.21	1.57	747.22	48.46	15.42	0.03	429
(US)		12	83	0.47	12.18	1.92	459.62	72.45	6.34	0.04	424
	Marlstone,	14	111	2.84	77.74	3.81	759.92	37.24	20.40	0.04	430
	dominated	18	150	1.29	17.19	3.53	301.05	61.82	4.87	0.07	424
		19	164	0.38	9.44	1.41	552.05	82.46	6.70	0.04	427
		20	185	2.84	84.25	2.79	950.9	31.49	30.20	0.03	425
		21	189.5	0.95	46.21	1.73	795.35	29.78	26.71	0.02	429
		22	216	0.47	7.08	1.10	498.59	77.46	6.44	0.06	452
		23	219	0.89	13.56	1.15	655.07	55.56	11.79	0.06	442
	Marly laminated	25	238	0.24	1.41	0.95	306.52	206.52	1.48	0.15	423
	magnesite	28	248.3	0.59	21.01	1.46	695.70	48.34	14.39	0.03	428
	magnesite	30	258	0.16	0.93	1.14	114.81	140.74	0.82	0.15	434
Lower		31	265	0.23	6.05	1.21	441.61	88.32	5.00	0.04	425
sequence		32	283	0.14	5.76	1.19	468.29	96.75	4.84	0.02	437
(LS)		33	286	0.10	2.62	1.04	297.73	118.18	2.52	0.04	445
	Marly Mg	35	309	0.10	0.77	1.08	150.98	211.76	0.71	0.11	425
	dolomite, Silty	37	324	0.26	8.98	1.47	432.77	70.84	6.11	0.03	439
	stone, Mg-clay	38	329	0.31	8.54	2.30	68.98	18.58	3.71	0.04	436
		39	335	0.17	16.72	3.00	371.56	66.67	5.57	0.01	442
		41	340	0.10	5.53	1.98	205.58	73.61	2.79	0.02	438
Minimum U	JS			0.38	9.44	1.31	301.05	22.29	4.87	0.02	424
Maximum U	US			7.78	112.31	3.93	1017.3	82.46	30.20	0.08	430
Average US			2.12	44.85	2.34	695.56	44.29	18.59	0.04	427.33	
Standard deviation US			2.07	32.01	0.97	215.12	18.60	8.69	0.02	2.35	
Minimum LS			0.10	0.77	0.95	68.98	18.58	0.71	0.01	423	
Maximum LS			0.89	21.01	300	695.7	211.76	14.39	0.15	452	
Average LS			0.29	7.61	1.47	362.17	97.95	5.09	0.06	435.85	
Standard deviation LS			0.23	6.23	0.60	196.17	57.99	4.07	0.05	8.67	

 $^{1}$ S1 – Free hydrocarbons;  $^{2}$ HC – Hydrocarbons;  $^{3}$ S2 – Pyrolysate hydrocarbons;  $^{4}$ S3 – Amount of CO<sub>2</sub> generated from oxygenated functional groups;  $^{5}$ HI – Hydrogen Index = S2 × 100/Corg;  $^{6}$ OI – Oxygen Index = S3 × 100/Corg;  $^{7}$ PI – Production Index = S1/(S1+S2);  $^{8}$ Tmax – Temperature corresponding to S2 peak maximum.

Modified van Krevelen, Hydrogen Index (HI) vs. Oxygen Index (OI) diagram (Langford and Blanc-Valleron, 1990) shows that samples from the upper and lower sequence of the borehole ZLT-2, in addition to the OM content, also differ with regard to the OM type. The OM of sediments from the upper sequence comprises kerogen I/II, II, I and II/III types, whereas the OM of samples from the lower sequence consists of kerogen II/III, III and II types (Fig. 5).

There is no significant difference in sulphur content between samples (Table 4), although sediments from the lower sequence are generally enriched in this element. Higher concentrations of total organic carbon, accompanied by lower concentrations of sulphur, resulted in a notably higher C/S ratio in the upper sequence (Table 4).

Preferred utilization of carbon and bacterial fixation of

nitrogen result in lowering of C/N values with increasing depth (Table 4), as it is reported for lacustrine sediments (Meyers and Ishiwatari, 1993).

The bitumen of all samples investigated is mainly represented by asphaltenes and polar-NSO compounds, whereas the relative amounts of saturated and aromatic hydrocarbons are low (Table 4), indicating low maturity. Values of Production Index (PI) lower than 0.1 confirms the low OM maturity (Table 5). Two samples have values of PI = 0.15. However, these samples contain low amounts of Corg (0.46% and 0.81%, respectively; Tables 4, 5) and therefore the Rock Eval parameters should be treated with caution (Kluska *et al.*, 2013). The maximal temperature (T<sub>max</sub>), which corresponds to S2 peak maximum, varies in a wide range (Table 5). However, this variation in T<sub>max</sub> is related to different kerogen types, rather than to thermal maturity (Peters *et al.*, 2005).

Values of bulk	organic	geochemical	parameters in	1 liquid	pyrolysates
	$\mathcal{O}$	0	1	1	1 2 2

	Viald of liquid	Yield of liquid	Vialda of UC <sup>2</sup>	Violda of UC	Group composition of	of liquid pyrolysates		
Sample	pyrolysates <sup>1</sup> (ppm)	pyrolysates (mg/g Corg)	(ppm)	(mg/g Corg)	Saturated + aromatic HC (%)	NSO + asphaltenes (%)	p <sub>o</sub> <sup>3</sup> (bar)	P <sup>4</sup> (bar)
3	4980	41.26	1796	14.88	36.07	63.93	5.0	5.9
6	13114	118.79	5996	54.31	45.72	54.28	4.9	5.5

 $^{1}$  Relative to the bitumen-free sample;  $^{2}$  HC – Hydrocarbons;  $^{3}$  p<sub>o</sub> – Initial pressure;  $^{4}$  p – Pressure at the end of pyrolysis.

# Characteristics of liquid pyrolysates – Bulk organic geochemical parameters

The generative potential of samples from the upper sequence, rich in organic matter containing kerogen I and II types, was investigated using conventional pyrolysis (Stojanović *et al.*, 2010). Pyrolysis was performed on bitumenfree samples 3 and 6. Heated at 400°C for 4 hours, the bitumen-free samples generated a total liquid pyrolysate of 4980 ppm and 13114 ppm, and hydrocarbons of 1796 and 5996 ppm, respectively (Table 6). The yields are consistent with those of a source rock with good potential (Peters *et*  *al.*, 2005) and support the assumption, derived from elemental analysis and Rock-Eval pyrolysis of the initial samples (Tables 4, 5). Sample 6 showed a higher liquid hydrocarbons potential (Table 6), consistent with the Rock-Eval data (Table 5). Apart from the liquid pyrolysate, the pyrolytic experiments also produced gaseous products that may be generated by direct degradation of kerogen or as secondary products of the degradation of liquid hydrocarbons. The gaseous products were not analyzed. However, their presence is proved by measuring pressure in the autoclave at the end of pyrolysis, in relation to the initial pressure, which typically was  $\sim$ 5 bar (Table 6).



**Fig. 6.** Characteristic Total Ion Currents (TICs) of the saturated fractions in sedimentary rocks from the upper sequence (A, B) and the lower sequence (C, D). *n*-alkanes are labelled according to their carbon number; Pr - pristane; Ph - phytane; PMI - 2,6,10,15, 19-pentamethylicosane; *i*-24 - C<sub>24</sub> regular isoprenoid; *i*-25 - C<sub>25</sub> regular isoprenoid; Sq - squalane; S - sulphur;  $\beta$  designates configurations at C<sub>17</sub> in hopanes;  $\beta\beta$  and  $\beta\alpha$  designate configurations at C<sub>17</sub> and C<sub>21</sub> in hopanes.



**Fig. 7.** Characteristic Total Ion Current (TIC) of the saturated fraction (**A**) and characteristic GC–MS chromatogram of *n*-al-kanes (m/z 71) (**B**) in liquid pyrolysate.  $\Delta - n$ -alkenes with same number of carbon atoms as *n*-alkanes; NPr – norpristane; \* – phthalate contamination; for other peak assignments, see Fig. 6 legend.

#### Molecular composition of the organic matter

The total ion currents (TICs) of saturated fractions of the samples investigated are shown in Figure 6. The main constituents are *n*-alkanes, isoprenoids, and polycyclic alkanes with sterane and hopane skeleton. The *n*-alkanes are the predominant biomarkers in all the samples. However, samples showed differences with respect to *n*-alkane patterns and abundances of isoprenoids and steroids (Fig. 6), which were consistent with different OM types (Fig. 5).

The total ion currents (TICs) of saturated fractions of liquid pyrolysates were dominated by n-alkanes, typical of oil distribution, whereas n-alkenes were observed in low amounts (Fig. 7A).

#### n-Alkanes and acyclic isoprenoids

On the basis of m/z 71 mass chromatograms (Fig. 8), in almost all of the samples, *n*-alkanes are identified in the range C<sub>15</sub> to C<sub>35</sub> (Table 7). In all samples, the CPI values for the full range of *n*-alkanes (C<sub>16</sub>–C<sub>34</sub>) are higher than 2.4, indicating the significant predominance of odd-chain homologues (Table 7). CPI values for short-chain *n*-alkanes



**Fig. 8.** Characteristic GC–MS chromatograms of *n*-alkanes (m/z 71) in sedimentary rocks from the upper sequence  $(\mathbf{A}, \mathbf{B})$  and the lower sequence  $(\mathbf{C}, \mathbf{D})$ . For peak assignments, see Fig. 6 legend.

Values of organic geochemical parameters calculated from the distributions and abundances of *n*-alkanes and isoprenoids

Sequence	Description	No	Depth (m)	<i>n</i> -Alkane range	$\begin{array}{c} {\rm CPI} \\ {\rm (C_{16}\text{-}C_{34})^1} \end{array}$	CPI (C <sub>16</sub> -C <sub>24</sub> ) <sup>2</sup>	CPI (C <sub>24</sub> -C <sub>34</sub> ) <sup>3</sup>	<i>n</i> -Alkane maximum	$\begin{array}{c} (C_{15}\text{-}C_{24})/\ (C_{25}\text{-}C_{35})^4 \end{array}$	LAD <sup>5</sup>	Pr <sup>6</sup> /Ph <sup>7</sup>	Pr/ <i>n</i> -C <sub>17</sub>	Ph/n-C <sub>18</sub>
Clayey c	arbonates	1	11.5	16 - 35	4.92	2.15	5.61	C <sub>29</sub>	0.34	1.21	0.20	0.54	1.06
		2	13.5	15 - 35	4.53	2.05	5.08	C <sub>29</sub> , C <sub>27</sub>	0.47	1.15	0.33	0.26	0.60
		3	27	15 - 35	3.92	1.29	5.70	C <sub>31</sub> , C <sub>29</sub>	0.70	1.96	0.15	0.51	1.58
	M. 1	4	32	16 - 37	2.94	1.28	3.11	C <sub>31</sub> , C <sub>29</sub>	0.29	3.28	0.35	0.52	0.60
	Marly dolomite	5	42.5	16 - 35	4.53	2.40	4.96	C <sub>27</sub> , C <sub>29</sub>	0.61	0.95	0.16	0.43	1.83
	doronnite	6	54	15 - 35	5.58	2.86	6.21	C <sub>27</sub>	0.40	0.98	0.25	0.37	1.38
		7	55.5	15 - 35	4.79	2.21	5.76	C <sub>31</sub> , C <sub>23</sub>	0.87	1.05	0.32	0.24	1.07
		8	64.5	15 - 35	2.58	1.34	3.20	C <sub>22</sub>	2.47	0.67	0.66	0.48	0.39
		9	70	15 - 35	3.47	1.97	3.68	C <sub>29</sub> , C <sub>22</sub>	0.77	1.02	0.18	0.40	1.60
		10	78	17 - 33	2.95	1.42	4.08	C <sub>22</sub>	2.51	0.67	0.44	0.35	0.32
Upper		11	80	14 - 33	2.79	1.23	3.91	C <sub>21</sub>	1.39	0.91	0.40	0.41	0.68
(US)		12	83	14 - 35	3.17	1.39	4.19	C <sub>22</sub> , C <sub>21</sub>	1.53	1.16	0.36	0.20	0.67
		13	96	14 - 35	3.33	1.26	5.56	C <sub>21</sub> , C <sub>22</sub>	1.63	1.64	0.23	0.15	0.78
	Marlstone.	14	111	14 - 35	5.99	1.83	8.02	C <sub>31</sub>	0.34	2.20	0.18	0.30	3.65
	calcite	15	113	15 - 35	2.57	1.29	2.83	C <sub>22</sub> , C <sub>21</sub>	3.43	0.65	0.33	0.22	0.59
	dominated	16	127	14 - 33	2.88	1.40	3.44	C <sub>22</sub> , C <sub>21</sub>	1.84	0.90	0.38	0.19	1.25
		17	137.5	16 - 35	6.16	1.93	7.32	C <sub>31</sub>	0.28	2.68	0.18	0.77	3.47
		18	150	16 - 35	2.43	1.37	2.54	C <sub>22</sub> , C <sub>21</sub>	2.14	0.62	0.52	0.65	1.07
		19	164	15 - 35	3.86	2.48	3.98	C <sub>22</sub> , C <sub>21</sub>	2.56	0.44	0.30	0.37	1.07
		20	185	16 - 35	6.19	2.66	7.10	C <sub>29</sub> , C <sub>31</sub>	0.28	1.89	0.07	0.42	7.21
		21	189.5	16 - 35	4.85	1.78	5.92	C <sub>31</sub> , C <sub>29</sub>	0.44	1.91	0.08	0.32	3.76
		22	216	15 - 35	2.47	1.33	2.81	C <sub>29</sub> , C <sub>31</sub>	0.79	1.12	0.49	0.94	2.41
		23	219	15 - 35	4.25	1.54	5.42	C <sub>31</sub> , C <sub>29</sub>	0.47	1.62	0.06	0.69	14.16
		24	224	15 - 35	3.72	1.80	4.27	C <sub>21</sub> , C <sub>22</sub>	1.02	1.00	0.11	0.25	3.15
		25	238	15 - 35	2.86	1.70	2.57	C <sub>22</sub>	4.17	0.32	1.72	0.62	0.64
	Marly	26	243.5	15 - 35	4.32	2.69	3.88	C <sub>17</sub>	1.62	1.13	0.14	0.22	25.72
	laminated magnesite	27	245	15 - 35	3.01	1.36	3.70	C <sub>22</sub> , C <sub>21</sub>	1.32	1.20	0.35	0.50	1.91
	magneone	28	248.3	15 - 35	2.91	2.72	2.37	C <sub>17</sub>	0.99	1.71	0.25	0.17	4.04
		29	255	15 - 35	6.49	7.48	3.50	C <sub>17</sub>	2.27	1.62	0.28	0.06	3.62
		30	258	15 - 35	4.66	2.15	5.64	C <sub>29</sub>	0.67	1.44	1.24	0.38	1.99
		31	265	15 - 35	3.53	1.96	3.49	C <sub>22</sub> , C <sub>21</sub>	1.90	0.88	0.17	0.46	24.87
Lower		32	283	15 - 35	3.79	1.57	4.55	C <sub>29</sub> , C <sub>31</sub>	0.75	1.61	0.40	0.53	3.18
sequence (LS)		33	286	16 - 35	3.91	1.84	4.56	C <sub>29</sub> , C <sub>22</sub>	0.92	1.32	0.43	0.14	1.29
(25)		34	297.5	15 - 33	3.40	2.12	3.29	C <sub>17</sub>	1.38	1.40	2.17	1.39	3.87
		35	309	16 - 35	3.38	1.98	4.13	C <sub>17</sub>	1.34	0.93	1.44	0.24	0.63
	Marly Mg	36	317.5	16 - 35	2.50	1.41	3.21	C <sub>21</sub> , C <sub>29</sub>	1.27	0.92	0.52	0.82	2.20
	Silty	37	324	16 - 34	3.32	1.48	4.79	C <sub>29</sub>	0.79	1.36	0.22	0.66	4.74
	Mg-marl-	38	329	16 - 35	3.54	1.51	5.10	C <sub>29</sub> , C <sub>31</sub>	0.90	1.04	0.10	0.56	6.89
	stone,	39	335	16 - 33	2.43	1.09	3.06	C <sub>29</sub> , C <sub>31</sub>	0.84	1.50	0.54	0.33	0.55
	wig-ciay	40	336	16 - 33	5.10	1.80	8.45	C <sub>29</sub> , C <sub>27</sub>	0.75	1.02	0.6	0.31	0.45
		41	340	16 - 35	4.19	1.13	6.42	C <sub>31</sub> , C <sub>29</sub>	0.44	2.71	0.88	0.19	0.13
		42	341	15 - 35	4.06	1.27	5.68	C <sub>31</sub> , C <sub>29</sub>	0.54	2.25	0.33	0.45	0.39
		43	343	15 - 35	3.83	1.25	4.91	C <sub>31</sub> , C <sub>29</sub>	0.43	2.08	0.20	0.42	0.55
Minimum	US				2.43	1.23	2.54		0.28	0.44	0.07	0.15	0.32
Maximum	US				6.19	2.86	8.02		3.43	3.28	0.66	0.77	7.21
Maximum US Average US					3.98	1.77	4.83		1.25	1.34	0.29	0.38	1.68
Average US Standard deviation US					1.27	0.52	1.58		0.96	0.76	0.15	0.16	1.68
Standard deviation US Minimum LS					2.43	1.09	2.37		0.43	0.32	0.06	0.06	0.13
Minimum LS Maximum LS					6.49	7.48	8.45		4.17	2.71	2.17	1.39	25.72
Maximum LS Average LS					3.71	1.96	4.36		1.16	1.37	0.57	0.47	4.88
Standard d	eviation LS				0.94	1.31	1.42		0.82	0.52	0.57	0.31	7.28

Sample	<i>n</i> -Alkane range	CPI (C <sub>16</sub> -C <sub>34</sub> )	CPI (C <sub>16</sub> -C <sub>24</sub> )	CPI (C <sub>24</sub> -C <sub>34</sub> )	<i>n</i> -Alkane maximum	(C <sub>15</sub> -C <sub>24</sub> )/ (C <sub>25</sub> -C <sub>35</sub> )	LAD	Pr/Ph	Pr/n-C17	Ph/n-C18
3	13 - 35	1.14	0.96	1.08	C16, C17	1.39	0.99	0.93	0.88	0.98
6	13 - 35	1.16	1.07	1.01	$C_{16}, C_{17}$	1.11	0.97	1.04	0.77	0.94

Values of organic geochemical parameters calculated from the distributions and abundances of *n*-alkanes and isoprenoids in liquid pyrolysates

For abbreviations see Table 7 legend.

(C<sub>16</sub>–C<sub>24</sub>) range from 1.09 to 7.48, most probably resulting from relatively high contents of n-C<sub>17</sub> and n-C<sub>21</sub> (Fig. 8, Table 7). The CPI values for long-chain *n*-alkanes (C<sub>24</sub>–C<sub>34</sub>) are higher than 2.3 (Table 7). Average CPI values for three different ranges (full, short and long) are comparable, indicating uniform, low maturity level (Table 7). The samples investigated differ significantly according to *n*-alkane maxima and the distributions of *n*-alkanes, i.e. the abundance of short- and mid-chain homologues (C<sub>15</sub>–C<sub>24</sub>) *vs.* abundance of long-chain homologues (C<sub>25</sub>–C<sub>35</sub>; Table 7; Fig. 8).

Both pyrolysates have similar *n*-alkane distributions, in which the *n*-alkanes  $C_{16}$ – $C_{20}$  are predominant, as is typical for aquatic organic matter (Fig. 7B). The CPI values for the three different ranges (full, short and long) for the pyrolysates are close to 1, which is typical of a mature oil distribution (Table 8).

The isoprenoids, pristane (Pr) and phytane (Ph), were identified in all samples (Figs 6, 8). Generally, higher values of Pr/Ph ratio, followed by more common variations of this parameter were observed in the lower sequence (Table 7). In samples from the lower sequence, the non-regular isoprenoid, squalane,  $C_{30}$ , was identified (Figs 6C, D, 8C, D). In the upper sequence, squalane is absent or present in very low amounts. The regular  $C_{24}$  isoprenoid was observed exclusively in sediments from depth interval 219–265 m (lower sequence, Figs 6C, 8C).

The C<sub>25</sub> isoprenoid was observed only in samples from depths above 265 m, with values of the Pr/Ph ratio  $\leq 0.35$ , (Figs 6A, C, 8A, C). Careful checking of the mass spectra (ratio of fragmentations ions, 239 and 253 as well as the abundance of fragmentation ion 113) of the corresponding peak (Vink et al., 1998; Peters et al., 2005) showed that in the upper sequence, the peak represents the irregular  $C_{25}$ isoprenoid, 2,6,10,15,19-pentamethylicosane (PMI). In the sediments of the lower sequence, the peak is a mixture of regular and irregular C25 isoprenoids, with a dominance of the latter. The irregular PMI is related to methanogenic archaea (Schouten et al., 1997; Vink et al., 1998). The regular C<sub>25</sub> isoprenoid is interpreted as an indicator of high salinity (Waples et al., 1974; Wang and Fu, 1997; Grice et al., 1998a, b; Yangming et al., 2005). However, the elevated content of the regular C25 isoprenoid was also identified in alkaline environments (Šajnović et al., 2008b). In addition it was reported that some families of methanogenic archaea (*Methanobacterium thermoautotrophicum*) also synthesize the precursor of the regular  $C_{25}$  isoprenoid (Risatti *et al.*, 1984).

Unlike bitumen in the initial samples, squalane and PMI were absent from the liquid pyrolysates (Fig. 7). The values of Pr/Ph ratio were higher than in the initial bitumen (Tables 7, 8), which may be explained by the fact that the degradation of kerogen during laboratory simulations results in the uniform formation of both pristane and phytane (Stojanović *et al.*, 2009, 2010).

#### Chromans

Alkylated 2-methyl-2-(4,8,12-trimethyltridecyl)chromans (MTTCs) were detected in aromatic fractions by ion chromatograms of m/z = 121 + 135 + 149 (Fig. 9). Samples from the lower sequence contained 8-methyl-MTTC, 5,8-dimethyl-MTTC, 7,8-dimethyl-MTTC and 5,7,8-trimethyl-MTTC (Fig. 9B), whereas in samples from the upper sequence only dimethyl- and trimethyl- MTTCs derivatives were observed (Fig. 9A). In all samples, 5,7,8-trimethyl-MTTC was the most abundant compound. On the basis of empirical observations, it was suggested by Sinninghe Damsté et al. (1989) that in sediments from non-hypersaline environments, 5,7,8trimethyl-MTTC dominates and 8-methyl-MTTC is completely missing. Therefore, it might be assumed that samples from the upper sequence were deposited under non-saline conditions. For samples from the lower sequence, 5,7,8-trimethyl-MTTC was the most prominent compound, but 8-methyl-MTTC also occurred, which indicates certain differences in depositional environment between the sequences. For a more elaborate characterisation of palaeosalinities, the MTTC ratio was defined by Sinninghe Damsté et al. (1987, 1993) as MTTC ratio = 5,7,8-trimethyl-MTTC/total MTTCs (Table 9).

Chromans were not identified in the liquid pyrolysates.

#### Arylisoprenoids

In samples 3, 4, 14, 23, 26, 27, 28, 31 and 32, isoreniratane (I) and several other digenetic products of isoreniratene (II–X) are identified (Fig. 10).

Arylisoprenoids are absent from the liquid pyrolysates. This result is expected, owing to the sensitivity of their side chain to thermal stress.

 $<sup>\</sup>begin{array}{l} \text{CPI} - \text{Carbon Preference Index;} \ ^{1}\text{CPI} \left(\text{C}_{16}\text{-}\text{C}_{34}\right) = 1/2 \times \left[ \Sigma \text{odd}(n\text{-}\text{C}_{17} - n\text{-}\text{C}_{33}) / \Sigma \text{even}(n\text{-}\text{C}_{16} - n\text{-}\text{C}_{32}) + \Sigma \text{odd}(n\text{-}\text{C}_{17} - n\text{-}\text{C}_{33}) / \Sigma \text{even}(n\text{-}\text{C}_{18} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left(\text{C}_{16}\text{-}\text{C}_{24}\right) = 1/2 \times \left[ \Sigma \text{odd}(n\text{-}\text{C}_{17} - n\text{-}\text{C}_{23}) / \Sigma \text{even}(n\text{-}\text{C}_{16} - n\text{-}\text{C}_{22}) + \Sigma \text{odd}(n\text{-}\text{C}_{17} - n\text{-}\text{C}_{23}) / \Sigma \text{even}(n\text{-}\text{C}_{18} - n\text{-}\text{C}_{24}) \right]; \ ^{3}\text{CPI} \left(\text{C}_{24}\text{-}\text{C}_{34}\right) = 1/2 \times \left[ \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) + \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) + \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) + \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) + \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) + \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) + \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) + \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) + \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) + \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25} \text{-}n\text{-}\text{C}_{34} \right) + \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) + \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34}) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) + \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34} \right) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) = \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34} \right) \right]; \ ^{2}\text{CPI} \left( \text{C}_{25}\text{-}n\text{-}\text{C}_{34} \right) = \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34} \right) = \Sigma \text{odd}(n\text{-}\text{C}_{25} - n\text{-}\text{C}_{34} \right) \right]; \ ^{2}\text{$ 



**Fig. 9.** Characteristic GC–MS chromatograms of MTTCs (m/z 121+135+149) in sedimentary rocks from the upper sequence (**A**) and the lower sequence (**B**).

#### Steroids and hopanoids

The distributions of steroid biomarkers are characterised by the presence of  $C_{27}$ – $C_{29}$  saturated steranes with  $5\alpha(H)14\alpha(H)17\alpha(H)$  and  $5\beta(H)14\alpha(H)17\alpha(H)$  20R configurations, unsaturated sterenes in the same range with double bonds in the  $\Delta^2$ ,  $\Delta^4$  and  $\Delta^5$  positions, respectively, as well as steradienes (Fig. 11). A similar distribution of steroids was reported in Malm Zeta laminated carbonates from the Franconian Alb, SW Germany (Schwark *et al.*, 1998). Thermodynamically more stable sterane isomers, with  $5\alpha(H)14\alpha(H)17\alpha(H)$  20S and  $5\alpha(H)14\beta(H)17\beta(H)$  20R or 20S configurations, are not present. Such steroid distributions indicate a low thermal maturity of the OM, consistent with the bitumen composition, PI and CPI discussed above (Tables 4, 5, 7). The sediments investigated are characteri-



**Fig. 10.** Characteristic GC–MS chromatograms of arylisoprenoids (m/z 133) in sedimentary rocks from the upper sequence (**A**) and the lower sequence (**B**). **C.** The structures of arylisoprenoids identified in the samples. \* Unidentified compound, M<sup>+</sup> 544?, basic ion, m/z 133.

sed by the dominance of  $C_{29}$  compounds or  $C_{27}$  compounds (Table 9), indicating a prevalence of terrestrial, or aquatic OM, respectively (Volkman, 1986), although elevated amounts of  $C_{29}$  steroids in samples containing kerogen I and I/II types (Fig. 5) also can be related to a contribution from green algae (Volkman, 2003). The main difference between sediments from the upper and lower sequences with regard to steroid biomarkers is expressed as elevated contents of  $C_{28}$  steroids in the lower sequence (Table 9).

<sup>&</sup>lt;sup>1</sup>% C<sub>27</sub> = 100 × C<sub>27</sub>5α(H)14α(H)17α(H)-Sterane/[ΣC<sub>27</sub>-C<sub>29</sub> 5α(H)14α(H)17α(H)-Steranes]; <sup>2</sup>% C<sub>28</sub> = 100 × C<sub>28</sub>5α(H)14α(H)17α(H)-Sterane/[ΣC<sub>27</sub>-C<sub>29</sub> 5α(H)14α(H)17α(H)-Steranes]; <sup>4</sup>ΔC<sub>27</sub> = 100 × C<sub>27</sub>(Δ<sup>2</sup> + Δ<sup>4</sup> + Δ<sup>5</sup>)-Sterenes]; <sup>5</sup>ΔC<sub>28</sub> Sterenes = 100 × C<sub>28</sub>(Δ<sup>2</sup> + Δ<sup>4</sup> + Δ<sup>5</sup>)-Sterenes]; <sup>4</sup>ΔC<sub>27</sub> = 100 × C<sub>27</sub>(Δ<sup>2</sup> + Δ<sup>4</sup> + Δ<sup>5</sup>)-Sterenes]; <sup>5</sup>ΔC<sub>28</sub> Sterenes = 100 × C<sub>28</sub>(Δ<sup>2</sup> + Δ<sup>4</sup> + Δ<sup>5</sup>)-Sterenes]; <sup>6</sup>ΔC<sub>29</sub> Sterenes = 100 × C<sub>29</sub>(Δ<sup>2</sup> + Δ<sup>4</sup> + Δ<sup>5</sup>)-Sterenes]; <sup>6</sup>ΔC<sub>29</sub> Sterenes = 100 × C<sub>29</sub>(Δ<sup>2</sup> + Δ<sup>4</sup> + Δ<sup>5</sup>)-Sterenes]; <sup>6</sup>ΔC<sub>29</sub> Sterenes = 100 × C<sub>29</sub>(Δ<sup>2</sup> + Δ<sup>4</sup> + Δ<sup>5</sup>)-Sterenes]; <sup>6</sup>ΔC<sub>29</sub> Sterenes = 100 × C<sub>29</sub>(Δ<sup>2</sup> + Δ<sup>4</sup> + Δ<sup>5</sup>)-Sterenes]; <sup>6</sup>ΔC<sub>29</sub> Sterenes]; <sup>7</sup> Ster/Hop = [Σ(C<sub>27</sub>-C<sub>29</sub>)(Δ<sup>2</sup> + Δ<sup>4</sup> + Δ<sup>5</sup>)-Sterenes]; <sup>6</sup>ΔC<sub>29</sub> Sterenes]; <sup>6</sup>ΔC<sub>29</sub> Sterenes]; <sup>7</sup> Ster/Hop = [Σ(C<sub>27</sub>-C<sub>29</sub>)(Δ<sup>2</sup> + Δ<sup>4</sup> + Δ<sup>5</sup>)-Sterenes]; <sup>6</sup>ΔC<sub>29</sub> C<sub>30</sub>(H)14α(H)17α(H) + 5β(H)14α(H)17α(H))-Steranes]/[Σ(C<sub>29</sub>-C<sub>32</sub>)17α(H)21β(H)-Hopanes + Σ(C<sub>29</sub>-C<sub>31</sub>)17β(H)21α(H)-Hopanes + Σ(C<sub>29</sub>-C<sub>31</sub>)17β(H)21β(H)-Hopanes + C<sub>27</sub>17α(H)-Hopane + C<sub>27</sub>17β(H)-Hopane + C<sub>30</sub>Hop-17(21)-ene + C<sub>27</sub>Hop-17(21)-ene + C<sub>27</sub>Hop-13(18)-ene + C<sub>30</sub>Hop-13(18)-ene + C<sub>30</sub>Hop-17(21)-ene + C<sub>30</sub>Hop-17(21)-ene + C<sub>30</sub>Hop-13(18)-ene + C<sub>30</sub>Hop-13(18)-ene + C<sub>30</sub>Hop-17(21)-ene + C<sub>31</sub>Hop-13(18)-ene + C<sub>30</sub>Hop-13(18)-ene + C<sub>31</sub>17α(H)21β(H)-Hopane + C<sub>31</sub>17α(H)21β(H)-Hopane + C<sub>31</sub>17α(H)21β(H)-Hopane); <sup>10</sup> GI – Gammacerane Index = 100 × G/(G + C<sub>30</sub>17α(H)21β(H)-102(S)-Hopane); <sup>11</sup> MTTC - 2-Methyl-2-(4,8,12-trimethyltridecyl)chromans (MTTC) ratio = 5,7,8-Trimethyl-MTTC/(8-Methyl-MTTC + 5,8-Dimethyl-MTTC); 12 N.D. - Not determined due to the absence of steranes, sterenes, C<sub>31</sub>17α(H)21β(H)22(S)-hopane and gammacerane. For sample 34 parameters calculated from the distributions of steroids and hopanoids are missing.

 $C_{30}\beta\beta/$ C31(S)/ Descrip Depth % C<sub>27</sub> % C28 2 % C<sub>29</sub> <sup>3</sup>  $\% \Delta C_{27}^{4}$  $\% \Delta C_{28}^{5}$  $\% \Delta C_{29}^{6}$ Ster/Hop GI 10 MTTC 11 Sequence No  $C_{30}(\beta\beta)$ C<sub>31</sub>(S+R) tion (m)  $+\alpha\beta)^{8}$ 11.5 27.32 29.16 43.52 23.45 27.81 48.75 0.11 0.82 N.D. N.D. 0.79 Clayey carbonates 1 2 13.5 34.78 20.72 44.50 49.89 18.24 31.87 0.06 0.75 N.D. N.D. 0.75 27 35.52 3 34.12 16.69 49.19 14.82 49.66 0.24 0.76 N.D. N.D. 0.82 4 51.96 15.13 57.97 0.79 32 32.36 15.68 26.90 0.20 0.83 N.D. N.D. Marly 5 20.39 52.82 32.08 46.76 0.14 0.82 0.06 42.5 26.78 21.17 N.D 0.81 dolomite N.D. <sup>12</sup> 6 54 N.D. N.D. 27.93 15.28 56.80 0.20 0.84 N.D. N.D. 0.84 7 55.5 N.D. N.D. N.D. 30.04 9.46 60.50 0.32 0.80 0.10 N.D 0.82 64.5 9.23 69.57 20.26 7.80 71.94 0.76 8 21.21 0.13 0.75 0.12 N.D 9 70 N.D. N.D. N.D. 26.14 14.60 59.26 0.09 0.79 0.15 N.D. 0.82 10 78 32.73 19.78 47.49 27.00 17.17 55.83 0.17 0.74 0.17 N.D 0.84 Upper 51.54 0.72 19.77 45.50 32.84 0.34 0.04 11 80 34.72 15.63 N.D. 0.83 sequence 12 83 42.55 18.26 39.20 50.62 16.42 32.96 0.30 0.47 N.D. 17.14 0.63 (US) 47.47 33.52 13 96 40.14 12.40 59.47 13.23 27.30 0.31 0.46 N.D. 0.79 14 111 51.45 7.86 40.69 42.48 10.62 46.90 0.30 0.56 N.D. 23.68 0.75 Marlstone 15 113 25.52 18.08 56.40 38.34 1.82 59.84 0.46 0.39 N.D. 5.95 0.86 calcite lominated 16 127 52.57 15.42 32.00 N.D. N.D. N.D. 0.18 0.39 N.D. N.D. 0.67 13.71 0.74 0.71 17 137.5 N.D. N.D. N.D 32.63 53.66 0.18 N.D N.D 9.85 0.44 0.50 18 150 N.D. N.D. N.D 32.63 57.51 0.71 N.D N.D. 19 164 43.24 15.76 41.00 48.92 12.01 39.07 0.30 0.75 N.D. N.D. 0.74 20 185 N.D. N.D. 48 28 10.11 0.80 ND 41.62 0.56 0.87 0.17 N.D. 21 189.5 N.D. N.D. N.D. 40.90 12.68 46.42 0.47 0.77 N.D. N.D. 0.81 216 51.92 21.56 36.88 40.49 0.58 0.51 13.73 22 26.52 22.63 0.25 0.7223 219 N.D. N.D. N.D. 66.29 13.04 0.35 0.84 N.D. 67.08 20.67 0.82 24 224 N.D. N.D. N.D. 53.83 15.83 30.34 0.31 0.80 N.D. 63.09 0.78 25 238 25.98 29.68 44.35 N.D. N.D. N.D. 0.48 0.11 0.15 12.31 0.61 Marly N.D. N.D. 41.79 0.14 45.00 0.77 26 243.5 N.D. 25.56 32.65 3.64 0.68 laminated 27 245 48.78 25.20 26.02 47.15 30.62 22.22 0.27 0.86 N.D. 30.44 0.78 magnesite 28 248.3 51.35 24.54 24.11 36.22 14.45 49.33 0.55 0.79 N.D. 37.10 0.81 25.36 55.76 29 255 42.10 32.55 29.84 35.87 34.30 0.71 0.81 N.D. 0.78 0.71 30 258 29.52 36.50 33.98 27.65 31.32 41.03 0.60 0.69 0.30 24.95 27 51 17 54 54 95 5.02 0.57 24 52 0.79 31 265 0.31 Lower 32 283 37.80 26.96 35.24 47.12 26.54 26.33 0.48 0.73 N.D. 40.45 0.69 sequence 33 286 31.60 22.15 46.24 27.11 42.68 30.21 0.51 0.66 N.D. 35.04 0.58 (LS) 34 297.5 N.D. N.D N.D. N.D N.D. N.D N.D. N.D. N.D. N.D 0.73 35 309 32.40 33.47 34.13 N.D. N.D. N.D. 0.23 0.44 N.D. 8.77 0.61 Marly Mg 317.5 30.62 26.37 53.44 22.50 0.53 N.D. 36 43.01 24.06 0.64 N.D 0.60 dolomite, 37 324 42.69 29.45 27.86 43.45 27.73 28.81 0.74 0.72 N.D. 62.01 0.70 Silty Mg-marl-40.20 38 329 34.90 24.91 41.40 30.77 27.84 1.44 0.66 N.D. 37.85 0.71 stone. 39 335 11.91 9.91 78.18 17.08 9.37 73.55 0.18 0.83 N.D. N.D. 0.63 Mg-clay 40 0.74 336 8 28 8.41 83 31 9 55 8 04 82.41 0.45 ND 23 55 0.61 41 340 7.29 21.98 70.73 5.67 11.84 82.49 0.19 0.83 N.D N.D. 0.67 42 341 9.72 19.87 70.41 N.D. N.D. N.D. 0.11 0.78 N.D. 22.32 0.57 343 12.55 15 90 71 55 0.43 43 11 46 6 90 81.64 0.08 0.74 0.16 ND Minimum US 7.86 32.00 20.26 27.30 0.06 0.39 0.00 0.00 0.50 21.21 1.82 Maximum US 52.57 20.72 69.57 59.47 21.17 71.94 0.56 0.87 0.21 33.52 0.86 36.99 49.86 0.27 36.32 47.52 0.70 4.01 Average US 16.16 13.14 0.05 0.77 Standard deviation US 9.44 10.35 11.28 10.22 7.42 12.67 0.15 0.15 0.07 9.43 0.09 Minimum LS 7.29 8.41 21.56 5.67 6.90 20.67 0.08 0.00 0.00 0.43 0.11 Maximum LS 51.35 51.92 83.31 66.29 42.68 82.49 5.02 0.86 0.31 67.08 0.82 Average LS 28.22 27.58 44.20 34.64 22.84 42.53 0.83 0.69 0.06 28.76 0.69

#### Values of organic geochemical parameters calculated from the distributions and abundances of steroids, hopanoids and chromans



Fig. 11. Characteristic GC–MS chromatograms of sterenes, m/z 215 (A, B) and steranes, m/z 217 (C, D) in sedimentary rocks from the upper sequence and the lower sequence.  $\Delta^2$ ,  $\Delta^4$  and  $\Delta^5$  designate double bonds in positions 2, 4 and 5 in sterenes;  $\alpha\alpha\alpha$  and  $\beta\alpha\alpha$  designate configurations at C<sub>5</sub>, C<sub>14</sub> and C<sub>17</sub> in 20(R) steranes.

The sterane distributions in pyrolysates obtained at 400°C are typical for oils (Fig. 12A), which confirms the good potential of the sediments investigated and shows that catagenesis was simulated successfully by pyrolysis. Apart from the regular  $5\alpha(H)14\alpha(H)17\alpha(H)20(R)$ -steranes,  $C_{27}$ - $C_{29}$  isomers with thermodynamically more stable  $5\alpha(H)14\alpha(H)17\alpha(H)$ 20(S)-,  $5\alpha(H)14\beta(H)17\beta(H)20(R)$ -, and  $5\alpha(H)14\beta(H)17\beta(H)$ 20(S)-configurations, as well as the typical geoisomers,  $\beta\alpha$ and  $\alpha\beta$ -diasteranes, were present (Fig. 12A). As in bitumen of initial samples 3 and 6,  $C_{27}$  and  $C_{29}$  steranes are more abundant in liquid pyrolysates than  $C_{28}$  homologues (Tables 9 and 10). Values of the most used sterane maturation parameters based on the ratios of  $C_{29}$  sterane isomers,  $C_{29}\alpha\beta\beta(R)/(C_{29}\alpha\alpha\alpha(S)/C_{29}\alpha\alpha\alpha(S+R))$  in all pyrolysates are lower than equilibrium values (Peters *et al.*, 2005; Table 10).

On the basis of a mass chromatogram, m/z 191 of the saturated fractions, the hopane composition is characterised by the presence of C<sub>27</sub>–C<sub>32</sub> 17 $\alpha$ (H)21 $\beta$ (H), C<sub>29</sub>–C<sub>31</sub> 17 $\beta$ (H) 21 $\alpha$ (H), and C<sub>27</sub>–C<sub>33</sub> 17 $\beta$ (H)21 $\beta$ (H) compounds with the exception of C<sub>28</sub> homologues in all three series (Fig. 13). Other hopanoid-type constituents of the saturated fraction are C<sub>27</sub> hop-13(18)-ene and C<sub>27</sub> hop-17(21)-ene, whereas C<sub>30</sub> hop-17(21)-ene was observed in samples from the lower sequence (depth interval 219–265 m). The presence of unsaturated hopenes, the dominance of  $\beta\beta$ -isomers over  $\alpha\beta$ -hopanes and the complete absence of  $\alpha\beta$ -22(S) isomers or low values of C<sub>31</sub>(S)/(S+R) ratio (< 0.3) confirm an immature

## Table 10

Values of organic geochemical parameters calculated from the distributions and abundances of steranes and hopanes in liquid pyrolysates

Sample	% C <sub>27</sub>	% C <sub>28</sub>	% C <sub>29</sub>	C <sub>29</sub> ααα(S)/ C <sub>29</sub> ααα(S+R)	$\begin{array}{c} C_{29}\alpha\beta\beta(R) / \\ (C_{29}\alpha\beta\beta(R) + \alpha\alpha\alpha(R)) \end{array}$	C <sub>31</sub> (S)/ C <sub>31</sub> (S+R)	$C_{29}\beta\alpha/C_{29}\alpha\beta$	$C_{30}\beta\alpha/C_{30}\alpha\beta$	Ts/(Ts+Tm) <sup>2</sup>
3	33.78	29.68	36.54	0.50	0.55	0.58	0.29	0.35	0.42
6	31.83	29.96	38.21	0.49	0.54	0.57	0.36	0.44	0.41
E.V.				0.52-0.55	0.67-0.71	0.57-0.62		0.15 <sup>1</sup>	

<sup>1</sup> – For samples of Tertiary age;  ${}^{2}\text{Tm} - \text{C}_{27}17\alpha$ (H)-22,29,30-trisnorneohopane; for other abbreviations see Table 9 and Figure 12 legend.



Fig. 12. Characteristic GC–MS chromatograms of steranes, m/z217 (A) and hopanes, m/z 191 (B) in liquid pyrolysate.  $1 - C_{27}5\alpha$ (H)14 $\alpha$ (H)17 $\alpha$ (H)20(S)-sterane + C<sub>28</sub>13 $\alpha$ (H)17 $\beta$ (H)20(S)-diasterane;  $2 - C_{27}5\alpha(H)14\beta(H)17\beta(H)20(R)$ -sterane +  $C_{29}13\alpha(H)$  $17\alpha(H)20(S)$ -diasterane;  $3 - C_{27}5\alpha(H)14\beta(H)17\beta(H)20(S)$ -sterane + C<sub>28</sub>13 $\alpha$ (H)17 $\beta$ (H)20(R)-diasterane; 4 - C<sub>27</sub>5 $\alpha$ (H)14 $\alpha$ (H)  $17\alpha(H)20(R)$ -sterane;  $5 - C_{29}13\alpha(H)17\alpha(H)20(R)$ -diasterane;  $6 - C_{29}13\alpha(H)17\alpha(H)20(R)$ -diasterane;  $C_{29}13\alpha(H)17\alpha(H)20(R)$ -diasterane;  $C_{29}13\alpha(H)17\alpha(H)17\alpha(H)20(R)$ -diasterane;  $C_{29}13\alpha(H)17\alpha(H)17\alpha(H)17\alpha(H)17\alpha(H)17\alpha$  $C_{28}5\alpha(H)14\alpha(H)17\alpha(H)20(S)$ -sterane; 7 –  $C_{29}13\alpha(H)17\beta(H)20$ (S)-diasterane;  $8 - C_{28}5\alpha(H)14\beta(H)17\beta(H)20(R)$ -sterane + C<sub>29</sub>13  $\alpha(H)17\beta(H)20(R)$ -diasterane; 9 – C<sub>28</sub>5 $\alpha(H)14\beta(H)17\beta(H)20(S)$ sterane;  $10 - C_{28}5\alpha(H)14\alpha(H)17\alpha(H)20(R)$ -sterane;  $11 - C_{29}$  $5\alpha(H)14\alpha(H)17\alpha(H)20(S)$ -sterane;  $12 - C_{29}5\alpha(H)14\beta(H)17\beta(H)$ 20(R)-sterane; 13 - C<sub>29</sub>5α(H)14β(H)17β(H)20(S)-sterane; 14 - $C_{295\alpha}(H)14\alpha(H)17\alpha(H)20(R)$ -sterane;  $\beta$  and  $\alpha$  designate configurations at  $C_{17}$  in hopanes;  $\beta\beta$ ,  $\beta\alpha$  and  $\alpha\beta$  designate configurations at  $C_{17}$  and  $C_{21}$  in hopanes; Ts –  $C_{27}18\alpha$ (H)-22,29,30-trisnorneohopane; (S) and (R) designate configuration at C<sub>22</sub> in hopanes.

stage for the OM (Table 9; Fig. 13). The absence of  $C_{34}$  and  $C_{35}$  hopanes could be explained by the high sensitivity of the side chains from biologic hopanoids, which were readily degraded in the upper, oxygenated part of the stratified water column and/or to the lower contribution of those bacteria that synthesise bacteriohopanetetrol.

Gammacerane was identified in the majority of samples from the lower sequence and in several samples from the upper sequence. Its precursor, 2-gammacerene, formed by the dehydration of tetrahymanol (Peters *et al.*, 2005), is present in almost all of the samples (Fig. 13).

Both pyrolysates contain a greater quantity of thermodynamically more stable  $C_{29}$  and  $C_{30} \alpha\beta$ -hopanes, compa-



**Fig. 13.** Characteristic GC–MS chromatograms of hopanoids  $(m/z \ 191)$  in sedimentary rocks from the upper sequence (**A**) and the lower sequence (**B**).  $\Delta^2$ G – 2-gammacerene; G – gammacerane; for other peak assignments, see Fig. 12 legend.

red to the corresponding  $\beta\alpha$ -moretanes (C<sub>29</sub> $\beta\alpha$ /C<sub>29</sub> $\alpha\beta$  and  $C_{30}\beta\alpha/C_{30}\alpha\beta$  below 1; Table 10), whereas unstable  $\beta\beta$ -hopanes and unsaturated hopenes were not identified (Fig. 12B). On the basis of the mass spectra of the corresponding peaks, Ts and the 22R and 22S epimers of C<sub>31</sub>-C<sub>35</sub> homohopanes were determined in the pyrolysates (Fig. 12B). Values for  $C_{31}(S)/C_{31}(S+R)$ -homohopanes indicated that in isomerisation  $22(R) \rightarrow 22(S)$  equilibria had been achieved in both pyrolysates; which is established in the earliest phase of catagenesis, at vitrinite reflectance, Ro  $\approx 0.60$  (Peters et al., 2005; Table 10). This result, together with the appearance of the hopane distribution in the liquid pyrolysates (ion fragmentogram m/z 191; Fig. 12B), which is typical for mature source rocks and oils, provide proof that catagenesis using pyrolysis at 400°C was simulated successfully and confirms the good potential of the samples investigated.

#### Alkylnaphthalenes and alkylphenanthrenes

The main constituents of the aromatic fractions of the liquid pyrolysates are alkylnaphthalenes and alkylphenanthrenes, which showed distributions typical for mature source rocks and oils (Fig. 14). The values of the corresponding maturity ratios are given in Table 11.



**Fig. 14.** Characteristic GC-MS chromatograms of (**A**) MN, DMN and TMN (m/z 142 + 156 + 170), and (**B**) P, MP and DMP (m/z 178 + 192 + 206) in liquid pyrolysate. MN – methylnaphthalene; DMN – dimethylnaphthalene; TMN – trimethylnaphthalene; PrN – propylnaphthalene; EMN – ethylmethylnaphthalene; P – phenanthrene; A – anthracene; MP – methylphenanthrene; DMP – dimethylphenanthrene; EP – ethylphenanthrene.

## DISCUSSION

#### Mineral composition and geochemistry

The entire section studied belongs to an intrabasinal lacustrine facies, although Mg-clays at the bottom of the basin fill indicate a marginal sedimentary environment. The lower sequence of the intrabasinal facies (216-343 m) is most probably more marginal, but that cannot be strictly defined on the basis of only one borehole core. In general, in this lower sequence, sedimentation occurred in shallow water with the predominant Mg ions derived from ultramafic rocks, while sedimentation of the upper sequence occurred in slightly deeper water, with predominant Ca ions (Fig. 15; Table 1). The higher variations of the Sr and Na<sub>2</sub>O contents in the lower sequence, compared to the upper sequence of the core studied (Fig. 16; Tables 1, 2), indicates more variable sedimentary conditions at the beginning of basin evolution, i.e. the formation of Mg-clays and marly magnesite, and rare thin (up to 2 mm) layers with dissolved, readily soluble minerals in the lower sequence of the intrabasinal facies (Fig. 17). Pronounced volcanic activity, evidenced by several tuff layers, influenced lower the palaeoproductivity and/or preservation of the organic matter (Fig. 2).

The significant positive correlation between B and Na2O contents (Fig. 18) indicates that the high boron concentrations in the lower sequence of the intrabasinal facies (Table 3) are most probably related to searlesite, identified by XRD analyses (Fig. 3C). This result is consistent with an earlier study of the Kremna Basin (Živković and Stojanović, 1976). which reported the presence of searlesite. The presence of boron minerals in the Kremna Basin was interpreted as hydrothermal in origin (Ilić, 1969; Obradović et al., 1996). However, boron rich sediments also can originate from diagenetic processes in tuffaceous sediments in saline-alkaline lakes (Sheppard and Gude, 1973; Stamatakis, 1989; Šajnović et al., 2008b). In arid areas, boron is likely to be co-precipitated with Mg and Ca hydroxides as coatings on the sediment particles and it also may occur as Na-metaborate (Floyd et al., 1998; Alonso, 1999).

The upper sequence of the intrabasinal facies (13.5–216 m) was deposited in slightly deeper water, with predominant Ca ions (Table 1), resulting in the formation of more

#### Table 11

Values of organic geochemical parameters calculated from distributions and abundances of naphthalene and phenanthrene hydrocarbons in liquid pyrolysates

Sample	MNR <sup>1</sup>	DMNR <sup>2</sup>	DNx <sup>3</sup>	$\alpha/\beta DN 1^4$	TNR 1 <sup>5</sup>	TNR 2 <sup>6</sup>	TMNR <sup>7</sup>	TNy <sup>8</sup>	MPI 1 <sup>9</sup>	MPI 3 <sup>10</sup>	$Rc^{11}$	DMPI 1 <sup>12</sup>	DMPI 2 <sup>13</sup>	PAI 1 <sup>14</sup>	PAI 2 <sup>15</sup>	DBT/P <sup>16</sup>
3	0.93	0.32	2.79	1.04	0.63	0.56	0.35	2.05	0.55	1.06	0.73	0.45	0.35	1.08	0.71	0.16
6	1.11	0.40	2.57	0.93	0.61	0.71	0.61	1.71	0.56	1.06	0.74	0.65	0.38	1.14	0.86	0.14

<sup>&</sup>lt;sup>1</sup>MNR = 2-MN/1-MN (Radke *et al.*, 1982b); <sup>2</sup>DMNR = (2,6-+2,7-DMN)/(1,4-+1,5-+1,6-+2,3-+2,6-+2,7-DMN) (Yawanarajah and Kruge, 1994); <sup>3</sup>DNx = (1,3-+1,6-+1,7-DMN)/(1,4-+1,5-+2,3-DMN) (Stojanović *et al.*, 2007); <sup>4</sup>α/ DN 1 = (1,4-+1,5-+2,3-DMN)/(2,6-+2,7-DMN) (Golovko, 1997); <sup>5</sup>TNR 1 = 2,3,6-TMN/(1,3,5-+1,4,6-TMN) (Alexander *et al.*, 1985); <sup>6</sup>TNR 2 = (1,3,7-+2,3,6-TMN)/(1,3,5-+1,3,6-+1,4,6-TMN) (Radke, 1987); <sup>7</sup>TMNR = 1,3,7-TMN/(1,3,7-+1,2,5-TMN) (van Aarssen *et al.*, 1999); <sup>8</sup>TNy = (1,3,6++1,3,7-TMN)/(1,3,5-+1,4,6-TMN) (Stojanović *et al.*, 2007); <sup>9</sup>MPI 1 = 1.5 × (2-+3-MP)/(P+1-+9-MP) (Radke *et al.*, 1982a); <sup>10</sup>MPI 3 = (2-+3-MP)/(1-+9-MP) (Radke, 1987); <sup>11</sup>Rc = 0.6 MPI 1 + 0.37 (Radke and Welte, 1983); <sup>12</sup>DMPI 1 =  $4 \times (2,6++2,7-+3,5-+3,6-DMP+1-+2-+9-EP)/(P+1,3-+1,6-+1,7-+2,5-+2,9-+2,10-+3,9-+3,10-DMP)$  (Radke *et al.*, 1982a); <sup>13</sup>DMPI 2 = (2,6++2,7-+3,5-DMP)/(1,3-+1,6-+2,5-+2,9-+2,10-+3,9-+3,10-DMP) (Radke *et al.*, 1982a); <sup>13</sup>DMPI 2 = (2,6-+2,7-+3,5-DMP)/(1,3-+1,6-+2,5-+2,9-+2,10-+3,9-+3,10-DMP) (Radke *et al.*, 1982b); <sup>14</sup>PAI 1 = (1-+2-+3-+9-MP)/P (Ishiwatari and Fukushima, 1979); <sup>15</sup>PAI 2 = ΣDMP/P (Ishiwatari and Fukushima, 1979); <sup>16</sup>DBT/P = Dibenzothiophene/Phenanthrene; DMP – Dimethylphenanthrene; EP – Ethylphenanthrene.



Fig. 15. Correlation diagram MgO/CaO vs. Na<sub>2</sub>O.



Fig. 16. Correlation diagram Sr vs. Na<sub>2</sub>O.

uniform marlstone and dolomite (Figs 2, 3A). The rare volcanic input and stable sedimentary conditions were responsible for the higher productivity and better preservation of the organic matter.

#### Content and precursors of the organic matter

**Lower sequence.** The lower sequence contains lower amounts of total and soluble OM (Tables 4, 5). The OM of these sediments is composed of kerogen II/III, III and II types (Fig. 5). This result indicates a significant contribution of the allochtonous biomass of land plants from the lake catchment to the OM of the sediments, particularly in the lower part of the sequence (265–343 m). The relatively low OM content could be attributed to a lower palaeoproductivity (i.e. *in situ* OM production) and/or more intense degradation (i.e. a higher redox potential in the environment).

Samples from the lower part of the lower sequence are characterised by the dominance of odd long-chain *n*-alkane homologues  $C_{27}-C_{31}$ , a  $(C_{15}-C_{24})/(C_{25}-C_{35})$  ratio < 1 and the prevalence of  $C_{29}$  homologues in the  $C_{27}-C_{29}$  sterane and sterene distributions (Tables 7, 9; Figs 8D, 11D), indicating significant input of terrestrial OM (Bray and Evans, 1961; Volkman, 1986; Cranwell *et al.*, 1987). Samples from upper part of the lower sequence (219–265 m, an interval of chemical precipitation of carbonates) are characterised by *n*-alkane maxima at  $C_{17}$ ,  $C_{21}$  or  $C_{22}$ ,  $(C_{15}-C_{24})/(C_{25}-C_{35}) > 1$  and a ratio of  $C_{27}/C_{29}$  steranes and sterenes > 1 (Tables 7, 9; Figs 8C, 11B), implying dominance of aquatic OM (Neto *et al.*, 1998; Peters *et al.*, 2005).

The identification of PMI in samples with  $Pr/Ph \le 0.35$  (Figs 6C, 8C) indicates methanogenic archaea as the precursor of the OM (Rissati *et al.*, 1984; Schouten *et al.*, 1997).



**Fig. 17.** Thin-section image of layers with dissolved, readily soluble minerals in the lower sequence of the intrabasinal facies (sample 26).



Fig. 18. Correlation diagram B vs. Na<sub>2</sub>O.

The regular  $C_{25}$  isoprenoid is a common indicator of elevated salinity (Waples *et al.*, 1974; Wang and Fu, 1997; Grice *et al.*, 1998a, b; Yangming *et al.*, 2005). However, the presence of this biomarker in the lower sequence in part also may be related to the methanogenic archaea *Methanobacterium thermoautotrophicum*, which synthesize  $C_{25}$  alkenes, the hydrogenation of which results in the formation of the regular  $C_{25}$  isoprenoid. The presence of phytenes in these samples (Figs 6C, 8C) corroborates the previous assumption, as it was reported that *Methanobacterium thermoautotrophicum* also synthesizes  $C_{20}$  alkenes with a phytene skeleton (Rissati *et al.*, 1984).

Isorenieratene (I; see Fig. 10C for structures) is a carotenoid, uniquely biosynthesized by the brown-coloured strains of the photosynthetic green sulphur bacterium (*Chlorobiaceae*). In samples (23, 26, 27, 28, 31 and 32), isorenieratane, the hydrogenated counterpart of isorenieratene and other diagenetic products of isorenieratene were identified (II–X, Fig. 10B, C). Although isorenieratane also can originate from  $\beta$ -carotene (Koopmans *et al.*, 1996b), a widespread carotenoid synthesized by algae, bacteria and land plants (Peters *et al.*, 2005), the presence of other diagenetic products of isorenieratene (II–X; Fig. 10B) uniquely indicates the contribution of a photosynthetic green sulphur bacterium (*Chlorobiaceae*) to the precursor OM. It also should be noted that the diagenetic products of  $\beta$ -carotene, such as  $\beta$ -carotane and other derivatives with the prominent typical fragmentation ions m/z 119 and 120 (Koopmans *et al.*, 1997), were absent.

The main precursor of 2-gammacerene and gammacerane is tetrahymanol. The principal source of tetrahymanol appears to be bacterivorous ciliates, which occur at the interface between oxic and anoxic zones in stratified water columns. Moreover, tetrahymanol was reported in ferns (Zander *et al.*, 1969), an anaerobic rumen fungus (Kemp *et al.*, 1984) and photosynthetic bacteria (Kleeman *et al.*, 1990). Considering the previous discussion, the presence of 2-gammacerene and gammacerane in the samples investigated partly can be related to bacterivorous ciliates. In several ciliates,  $C_{30}$  hopan-3- $\beta$ -ol was also identified. Therefore, it is reasonable to speculate that the abundant  $C_{27}$ ,  $C_{29}$ and  $C_{30}$  hopanoids with  $\beta\beta$ -configuration in the samples studied (Fig. 13), could be attributed at least to some extent to ciliate sources.

Abundant hopanoid biomarkers (ster/hop ratio < 1 in the majority of the samples; Table 9) indicate a significant contribution of prokaryotic organisms to the sedimentary OM. This result is consistent with the significant proportion of kerogen II type in the OM (Fig. 5) and the conclusions drawn by Bohacs *et al.* (2000, 2003).

Bacteriohopanetetrol with minor contribution of 3-desoxyhopanes from prokaryotes and fungi, are considered as the main biological precursors of geohopanoids (Ourisson *et al.*, 1979; Rohmer *et al.*, 1992). In addition, Chaffee *et al.* (1986) reported that hopanes containing less than 30 carbon atoms could originate from ferns, lichens and mosses. Elevated amounts of hopanoids with less than 30 carbon atoms (Fig. 13), observed in some samples, may also signify methanotrophic bacteria (e.g., *Methylococcus capsulatus* or *Methylomonas methanica*, which synthesized aminobacteriohopanepentol; Neunlist and Rohmer, 1985), as a source of the OM. These bacteria had suitable conditions, owing to the presence of methanogenic archaea (see the discussion related to the irregular C<sub>25</sub> isoprenoid, PMI), which produce methane as a source of methanotrophics.

C<sub>30</sub> Hop-17(21)-ene was detected in samples 20–32, with the exception of sample 22 (Fig. 13B). According to Bottari *et al.* (1972) and Volkman *et al.* (1986), C<sub>30</sub>hop-17(21)ene is introduced into sediments by bacteria or in some cases of ferns and mosses, whereas Brassell *et al.* (1980) suggest a diagenetic origin from the transformation of hop-22(29)-ene (diploptene). Wolff *et al.* (1992) suggest sulphate-reducing bacteria as a probable source of C<sub>30</sub>hop- 17(21)-ene.

**Upper sequence.** The upper sequence is relatively rich in OM that comprises kerogen I/II, II, I and II/III types (Tables, 4, 5). Samples from this sequence are characterised by the dominance of  $C_{21}$  and  $C_{22}$  *n*-alkanes or long-chain odd homologues in the range  $C_{27}$  to  $C_{33}$ , and the predominance of either  $C_{27}$  or  $C_{29}$  homologues in the  $C_{27}$ – $C_{29}$  steroid distribution (Tables 7, 9; Figs 8A, B, 11A, C). *n*-Alkane (C<sub>15</sub>–  $C_{24}$ )/(C<sub>25</sub>–C<sub>35</sub>) ratio vary in a wide range of 0.28–3.43 (Table 7). These data indicate mixed aquatic/terrestrial OM sources (Peters *et al.*, 2005).

Although the *n*-alkane distributions of samples 3, 6, 14, 20 and 21 show maxima at odd long-chain homologues and

have CPI and  $(C_{15}-C_{24})/(C_{25}-C_{35})$  values > 1 (Table 7), Rock Eval data suggest kerogen I and I/II types (Fig. 5; Table 5). In addition to an origin from higher land plants, there are reports of  $C_{27}$ – $C_{31}$  *n*-alkanes originating from strains of Botryococcus braunii race A (e.g., Moldowan et al., 1985; Derenne et al., 1988) by the reduction of the n-alkadienes and trienes they biosynthesized (e.g., Banerjee et al., 2002, and references therein). Therefore, it can be assumed that the green unicellular microalga, Botryococcus braunii race A was the source of the OM in the sediments of the upper sequence. For purpose of quantification of the variability of the long chain n-alkanes, Kluska et al. (2013) proposed novel parameter, the LAD (long *n*-alkane distribution) ratio. This ratio includes range of *n*-alkanes which are considered typical for Botryococcus braunii race A. For the samples mentioned (3, 6, 14, 20 and 21) this ratio varies in the range 0.98 to 1.96 (Table 7). These values are similar to those observed during an investigation of sediments from the Werra cyclothem (Upper Permian, Fore-Sudetic Monocline, Poland), for which Botryococcus braunii race A also was proposed as a source of the OM (Kluska et al., 2013). The significant contribution of a green alga is further supported by the high relative abundance of C29 steroid homologues, observed in these samples (Fig. 11A, Table 9).

The identification of PMI in samples with values of the Pr/Ph ratio  $\leq 0.35$ , indicates methanogenic archaea as precursors of the OM (Risatti *et al.*, 1984; Schouten *et al.*, 1997; Vink *et al.*, 1998). In samples 3, 4 and 14, isorenieratane and other diagenetic products of isorenieratene were identified (Fig. 10A), indicating a photosynthetic green sulphur bacterium (*Chlorobiaceae*) as an OM source.

The presence of gammacerane and 2-gammacerene (Fig. 13) partly can be related to bacterivorous ciliates (Peters *et al.*, 2005), whereas abundant hopanoid biomarkers indicate a significant contribution from prokaryotic organisms (possibly including methanotrophic bacteria, owing to elevated amounts of  $C_{27}$  to  $C_{30}$  hopanes, Fig. 13A; Neunlist and Rohmer, 1985) to the sedimentary OM.

#### Characteristics of depositional environment

Lower sequence. The pristane to phytane ratio is widely used as indicator of redox settings (Didyk et al., 1978). However, it also depends on thermal maturity and increases with thermal alteration of the OM. Low values of this ratio (< 0.8) were also reported in hypersaline environments (ten Haven et al., 1987; Peters et al., 2005). On the basis of the amounts of NSO compounds in bitumen, values of PI, CPI, and biomarker maturity parameters (Tables 4, 5, 7 and 9), the influence of thermal maturity on this sample set could be ruled out. On the other hand, the distributions of MTTCs (Fig. 9B) and the values of the MTTC ratio (Table 9) show that there are no indications of hypersalinity. Therefore, a Pr/ Ph ratio lower than 1 (with exception of 4 samples, Table 7) could imply sedimentation under reducing conditions. The plot Pr/n-C<sub>17</sub> vs. Ph/n-C<sub>18</sub> ratio (Shanmugam, 1985) corroborates the previous assumption (Fig. 19).

The higher amount of  $C_{28}$  homologue in the distribution of  $C_{27}$ – $C_{29}$  steroids (Fig. 11B; Table 9; Volkman 1986; Wang and Fu, 1997), the presence of squalane,  $C_{24}$  and  $C_{25}$  regular isoprenoids (Fig. 6C; Waples, 1974; Grice *et al.*, 1998a, b), the higher values of GI (Table 9; Peters *et al.*, 2005) and the distribution of MTTCs (Fig. 9B; Sinninghe Damsté *et al.*, 1987, 1993), followed by elevated MgO/CaO ratio and elevated content of Na<sub>2</sub>O (Fig. 15; Table 1), indicate that sediments from lower sequence were deposited under alkaline conditions. Similar conditions were established during the investigation of the Lower Miocene lacustrine Valjevo-Mionica Basin, located close to the Kremna Basin (Šajnović *et al.*, 2008a, 2009), and the Piskanja borate deposit, Jarandol Basin, Serbia (Szabó *et al.*, 2009).

The presence of gammacerane, 2-gammacerene (Fig. 13), MTTCs and isoreniratene diagenetic products (Figs 9B, 10B) implies stratification of the water column (Sinninghe Damsté *et al.*, 1995). The presence of isoreniratene diagenetic products (Fig. 10B) indicates a photic zone of anoxia (Koopmans *et al.*, 1996a; Kluska *et al.*, 2013), characterised by periods when the water column was highly stratified and anoxic waters extended up into the photic zone.

**Upper sequence.** Considering that there was no influence of hypersalinity and maturity, the Pr/Ph ratio < 1 (Table 7) and plot Pr/n-C<sub>17</sub> vs. Ph/n-C<sub>18</sub> (Fig. 19; Shanmugam, 1985) implies reducing conditions.

Generally, the lower values of Pr/Ph ratio in the upper sequence, followed by less variation of this parameter, indicate more reductive settings, i.e. deposition under a higher water column by comparison with sediments from the lower sequence (Table 7). This result could imply higher precipitation that is consistent with the palaeoclimate reconstruction obtained from an investigation of the palaeoflora in Serbia, which showed that during the Lower Miocene, precipitation rates increased, peaking in the Eggenburgian/Ottnangian (Utescher *et al.*, 2007). Therefore, a higher amount of OM (Tables 4, 5) in sediments from the upper sequence to some extent could be related to more reducing conditions (resulting from a deepening of the water column), which contributed to better OM preservation.

The low amounts of Na<sub>2</sub>O (Fig. 15; Table 1) and the  $C_{28}$  homologue in the distribution of  $C_{27}$ – $C_{29}$  steranes and sterenes (Table 9; Fig. 11A, C), the absence of or trace amounts of squalane, the absence of  $C_{24}$  and  $C_{25}$  regular isoprenoids (Fig. 6A, B), the distribution of MTTCs (Fig. 9A) along with the dominance of calcite as the primary carbonate and the low MgO/CaO ratio (Figs 2A, 15) show that sediments from the upper sequence were formed in a freshwater environment.

The presence of 2-gammacerene (Fig. 13A), MTTCs (Fig. 9A) and isoreniratene diagenetic products (Fig. 10A) implies that stratification of the water column (Sinninghe Damsté *et al.*, 1995; Kluska *et al.*, 2013) resulted from temperature differences within the higher water column. A photic zone of anoxia is also indicated.

## Investigation of the liquid hydrocarbon generation potential and assessment of the conditions for achieving early catagenesis

The yields of total liquid pyrolysate of 4980 and 13114, and hydrocarbons of 1796 ppm and 5996 ppm, respectively (Table 6) indicate a good generation potential for the orga-



**Fig. 19.** Plot of  $Pr/n-C_{17}$  vs.  $Ph/n-C_{18}$  (Shanmugam, 1985). I – Algal OM, reducing environments; II – Mixed OM, reducing and transitional environments; III – Peat-coal environments; IV – Terrestrial OM, oxic environments.

nic-rich sediments of the upper sequence (samples 3 and 6). Liquid pyrolysates have typical oil distributions of biomarkers (*n*-alkanes, steranes, hopanes; Figs 7, 12) and alkylaromatics (Fig. 14), which confirms the good potential of the sediments investigated and shows that catagenesis was simulated successfully by pyrolysis.

Values for C<sub>31</sub>(S)/C<sub>31</sub>(S+R)-homohopanes indicate that in isomerisation  $22(R) \rightarrow 22(S)$  the equilibria was achieved in liquid pyrolysates; established in the earliest phase of catagenesis, at vitrinite reflectance  $\approx 0.60$  (Peters et al., 2005; Table 10). On the other hand, the values of sterane maturation parameters,  $C_{29}\alpha\beta\beta(R)/(C_{29}\alpha\beta\beta(R)+\alpha\alpha\alpha(R))$  and  $C_{29}\alpha\alpha\alpha$  $(S)/C_{29}\alpha\alpha\alpha(S+R)$  in pyrolysates are lower than the equilibrium values (Peters et al., 2005; Table 10). Bearing in mind that in the previous studies it was noticed that equilibria in the sterane isomerisations are established at vitrinite reflectance value of approximately 0.80% and taking into consideration the fact that equilibria were attained in homohopane isomerisations  $22(R) \rightarrow 22(S)$  in both pyrolysates (Table 10), it may be assumed that during pyrolysis at 400°C the sample investigated reached an equivalence value of vitrinite reflectance between 0.60 and 0.80% (Peters et al., 2005).

The low dibenzothiophene/phenanthrene ratio (DBT/P; Table 11) confirms formation of the sediments of the upper sequence in a freshwater environment. The values of naphthalene and phenanthrene maturity ratios in pyrolysates are similar and in a range that is typical for oils (Table 11). Applying the equation Rc = 0.6 MPI 1 + 0.37 (Radke and Welte, 1983), the vitrinite reflectance equivalent (Rc) of 0.70% for pyrolysates of at 400°C was calculated (Table 11). This Rc value is in full agreement with the results, obtained in the interpretation of the terpane and sterane biomarkers (Table 10).

Therefore, it can be assumed that pyrolysis of the samples investigated at 400°C achieved oil generation at a vitrinite reflectance equivalent of ~ 0.70%. Applying a generalized diagram that relates vitrinite reflectance, depth and a



regional geothermal gradient (Suggate, 1998) of between 40 and 50°C/km (Kostić, 2010), the minimum depth of 2300– 2900 m was estimated, as that at which the sediments would become a thermally mature source rock (Fig. 20). The minimum temperature necessary for catagenetic generation of hydrocarbons (temperature = depth x geothermal gradient + annual mean surface temperature; Suggate, 1998) was calculated at 103°C (t =  $2.3 \times 40 + 11 = 103$ °C). Using the basin-independent equation T = (lnRo+1.68)/0.0124 (Barker & Pawlewicz, 1994) and a vitrinite reflectance, Ro, value of 0.70%, the temperature is estimated to be 107°C. The estimated temperature for hydrocarbon generation and the necessary depth are in good agreement with corresponding data for the active source rocks in the region (Dragaš *et al.*, 1991; Jovančićević *et al.*, 2002; Kostić, 2010; Mrkić *et al.*, 2011).

# **CONCLUSIONS**

Lacustrine sediments of the Kremna Basin (borehole ZLT-2, at depths of up to 343 m) originated from an ultramafic source. One main intrabasinal facies with two sequences was distinguished. The lower sequence occurs at depths of 216–343 m, while the upper sequence is encountered from 13.5–216 m. The sediments contain variable amounts and types of immature OM.

A comprehensive analysis of the OM indicates diverse precursors: bacteria, algae and, terrestrial plants. A higher contribution of allochtonous biomass of land plants from the lake catchment is observed in the sediments of the lower part of the lower sequence (265–343 m). Biomarker distributions reveal the following possible sources of the OM in the stratified water column: methanogenic archaea, a photosynthetic green sulphur bacterium (*Chlorobiaceae*), bacterivorous ciliates, various bacteria (photosynthetic and nonphotosynthetic), and the green unicellular microalga, *Botryococcus braunii* race A (strictly in the upper sequence).

At the beginning of basin development (216–343 m), sedimentation took place in shallow water. This lower sequence consists of Mg-rich sediments with lower amounts of OM. Indications of volcanic activity include tuff layers and the presence of searlesite. The OM of these sediments is composed of kerogen II/III, III and II types. Through time, a slight deepening of basin occurred. This was followed by deposition of Ca-rich sediments (depth interval 13.5–216 m). These sediments are richer in OM, which comprises kerogen I/II, II, I and II/III types.

A higher water level (more reducing conditions) contributed to better OM preservation and also resulted in an elevated content of autochthonous aquatic OM in the sediments of the upper sequence. The transition between the lower and upper sequences is associated with a decrease of MgO/CaO ratio, a decrease of Na<sub>2</sub>O and B contents, and an increase in the OM content. The main differences between the sequences with regard to biomarker distribution are expressed in the more uniform and lower Pr/Ph ratio, the lower amounts of  $C_{28}$  steroids and gammacerane index, the absence or significantly lower amount of squalane, the absence of 8-methyl-MTTC,  $C_{24}$  and  $C_{25}$  regular isoprenoids, as well as  $C_{30}$  hop-17(21)-ene in the sediments of the upper sequence.

The liquid pyrolysate and hydrocarbons yields obtained in the pyrolitic experiments, the distributions of saturated biomarkers and the alkylaromatics in pyrolysates supported the assumption, derived on the basis of the Rock-Eval data and the analysis of the initial bitumen, that the samples of the upper sequence, rich in OM in its catagenetic phase, could be a source of liquid hydrocarbons. The values of hopane, sterane and phenanthrene maturation parameters indicate that through pyrolysis at 400°C the samples investigated reached a value of vitrinite reflectance, equivalent to approximately 0.70%. It was estimated that the sediments investigated should occur at depths of 2300–2900 m to become active source rocks. The calculated minimum temperature, necessary for catagenetic hydrocarbon generation, is between 103 and 107°C.

#### Acknowledgments

The investigations that formed this study were done in cooperation with the company, Rio Tinto – Rio Sava Exploration, from Serbia. The study was partly financed by the Ministry of Education, Science and Technological Development of the Republic of Serbia (Projects 176006 and 176016). The authors are also grateful to Bartosz Budzyń (Polish Academy of Sciences, Kraków, Poland), Ladislav Palinkaš (University of Zagreb, Zagreb, Croatia) and two anonymous reviewers, whose helpful suggestions and comments greatly benefited this paper.

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