Graptolite turnover and $\delta^{13}C_{org}$ excursion in the upper Wenlock shales (Silurian) of the Holy Cross Mountains (Poland)

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Abstract: The mid–late Homerian Age of the Silurian Period was a time of intense changes in biota, oceanic chemistry, and sea level and is known as the *lundgreni* extinction (for the graptolite extinctions), the Mulde bioevent (for the condont turnover event) or the Homerian carbon isotope excursion (CIE) probably related to glacially influenced climate perturbation. New information on this interval from the deep water sedimentary and graptolite succession of the Kielce Region (Holy Cross Mountains, Poland) of the northern margin of the Małopolska Block is presented here based on analysis of the Pragowiec Ravine section. The *lundgreni–nilssoni* graptolite biozones interval have been recognized there. This interval is composed by dark shales with very rare benthic fauna, which indicate the deep open-marine (pelagic) paleoenvironment. Ten samples were taken for the $\delta^{13}C_{org}$ analysis from the *lundgreni* (2 samples), *parvus* (2 samples), *praedeubeli–deubeli* (1 sample), *ludensis* (2 samples) and *nilssoni* (1 sample) biozones. According to the $\delta^{13}C_{org}$ results, the first positive $\delta^{13}C_{org}$ excursion of the Mulde Bioevent is well recognized. The $\delta^{13}C_{org}$ values rise from -30.7 - 30.1 ‰ in the *lundgreni* Biozone to -29.3 - -28.7 ‰ in the *parvus* Biozone and fall below -30 ‰ in the *praedeubeli–deubeli* interval. The second positive $\delta^{13}C_{org}$ peak of the Mulde Event was not recognized in the Pragowiec Ravine. Based on the numerical comparisons using Raup-Crick metric of co-occurrences of graptolite species, the upper Homerian was characterized by significant between-biozone turnover of these taxa at the given locality.

Keywords: Poland, Holy Cross Mountains, Silurian, Mulde Event, geochemistry, 813Corro-

Introduction

The Silurian was one of the most unstable periods in the Paleozoic, marked by significant environmental changes and biotic perturbations (Crampton et al. 2016). One such episode is in the late Wenlock Epoch (mid-late Homerian). Graptolite workers refer to the onset of this episode as the lundgreni extinction (Koren' 1987) or the "Große Krise" (Jaeger 1991), and conodont workers refer to the immediately preceding changes in conodont assemblages as the Mulde Event (e.g., Jeppsson et al. 1995) or Homerian carbon isotope excursion (CIE; Calner 2008). Although it is debatable whether the Mulde Event had a significant impact on conodont communities (e.g., Radzevičius et al. 2014c; Jarochowska et al. 2018), it has been determined that, at least in Baltica, the community structure and abundance fluctuation patterns radically changed as a consequence of the events (Spiridonov 2017; Spiridonov et al. 2017a). The impact on the microphytoplankton communities of the Mulde Event is not so clear and the changes in their taxonomic composition and the size distribution probably were more related to sea level changes (Porebska et al. 2004; Venckutė-Aleksienė et al. 2016; Spiridonov et al. 2017b).

A twin-peaked positive carbon isotope excursion has been documented globally in the upper Wenlock. Such carbon isotope excursions of this age are documented: in Baltica (Samtleben et al. 1996; Wenzel & Joachimski 1996; Bickert et al. 1997; Kaljo et al. 1997, 1998, 2007; Samtleben et al. 2000; Porebska et al. 2004; Martma et al. 2005; Calner et al. 2006b, 2012; Jarochowska et al. 2014, 2016a, 2016b; Radzevičius et al. 2014c, 2016; Jarochowska & Munnecke 2016; Makhnach et al. 2018); Laurentia (Saltzman 2001; Noble et al. 2005; Cramer et al. 2006; Lenz et al. 2006; Sullivan et al. 2016); Avalonia (Corfield et al. 1992; Marshall et al. 2012; Blain et al. 2016; Fry et al. 2017); Timan (Shebolkin & Männik 2014); Perunica (Frýda & Frýdová 2014, 2016); and Gondwana (Vecoli et al. 2009). Thus, most of the stable carbon isotopic data are from the Baltica, Laurentia and Avalonia paleocontinents.

The purpose of the investigation is to document biostratigraphy, constrain the stable carbon isotopic trends of the middle–upper Homerian, and to enlighten the patterns of graptolite community change in the Kielce Region of the Holy Cross Mountains, Poland. Here we present the first stable carbon isotope ($\delta^{13}C_{ore}$) data linked to a biostratigraphical framework for the Kielce Region (the Małopolska Block), and the Homerian biogeochemical event in the deep water open marine facies environments.

Geological background

The Holy Cross Mountains (HCM) are located in central Poland (Fig. 1A) and expose Paleozoic rocks in the central part of the Teisseyre–Tornquist Zone (TTZ) which extends from the North Sea in the NW to the Black Sea in the SE. According to the differences in the stratigraphic, lithological, facies, and tectonic evolution of the Lower Paleozoic, the HCM are divided into the Lysogóry Region or Unit in the north (Fig. 1B) and the Kielce Region (part of Małopolska Block) in the south (Fig. 1B) (Dadlez et al. 1994).

The Łysogóry Unit is considered to be a passive margin of Baltica (Dadlez et al. 1994; Narkiewicz 2002). The Silurian succession of the Łysogóry Unit is without significant gaps (Modliński & Szymański 2001) and comprises lower Llandovery deep open-marine (pelagic) to uppermost Přídolí lagoonal/fluvial (continental) deposits (Kowalczewski et al. 1998; Kozłowski 2003). Most natural outcrops of Silurian strata are exposed in the "Silurian Zone" (Fig. 1B) of the Łysogóry Unit (Kozłowski 2008). The upper Ludlow positive carbon isotope excursion associated with the Lau Event was first documented from the "Silurian Zone" (Fig. 1B) of the Łysogóry Unit by Kozłowski & Munnecke (2010) but no data from the upper Wenlock were presented.

The Kielce Region is on the northern margin of the Małopolska Block (Fig. 1A). There are different opinions regarding the development of the Małopolska Block: (1) it has a peri-Gondwanan origin, rifted in the Cambrian and amalgamated with the margin of SW Baltica during the Cambrian-Early Ordovician (Bełka et al. 2002; Walczak & Belka 2017); and (2) the Małopolska Block originated near the present SW margin of Baltica (Cocks 2002; Nawrocki et al. 2007). However, the Kielce and the Łysogóry regions were paleogeographically separate sub-basins in the Silurian (Kozłowski 2008). The Łysogóry Region was located relatively distally and the Kielce Region more proximally in relation to the same orogen, in other words the Łysogóry Region was closer to Baltica then the Kielce Region (Kozłowski et al. 2004, 2014). The Silurian sequence in the Kielce Region has numerous stratigraphic gaps and is represented by lowermost Llandovery deep open-marine deposits through to upper Ludlow turbidites (Kozłowski & Tomczykowa 1999). Perhaps because of the dominance of clastic sedimentation, stable carbon isotopes have not been investigated in the Kielce Region.

The exposed upper Wenlock and Ludlow succession is about 600 m thick (Tomczyk 1962) and is divided into the upper Bardo, Prągowiec, and Niewachlów beds (Fig. 1C). The upper Bardo beds are dark yellow and brown clayey shales belonging to the *Cyrtograptus lundgreni* and probably *Pristiograptus parvus* (see below) graptolite biozones. The Prągowiec beds are composed of dark grey silty shale with rare limestone

concretions and abundant graptolites from the *Gothograptus* nassa to Saetograptus leintwardinensis biozones (Tomczyk 1962). The Niewachlów beds are composed of mediumgrained greywackes with mudstone interbeds (Malec 2001) with *Bohemograptus bohemicus* (Barrande), *B. bohemicus* tenuis (Bouček) (Tomczyk 1962) and trilobites of Ludfordian age (Tomczykowa 1993). The Bardo Diabase occurs between the Prągowiec and Niewachlów beds. Using ⁴⁰Ar-³⁹Ar isotope dating its age is either 424±6 Ma-415±2 Ma latest Ludlow and earliest Lochkovian (Nawrocki et al. 2013). The igneous intrusion is spatially separated from the sampled outcrops. Good preservation of organic skeletons of graptolites points to the absence of significant contact metamorphism in the studied part of the section.

Material and methods

About 50 samples from the upper Wenlock and lower Ludlow (*lundgreni–nilssoni* biozones) were collected from the Prągowiec Ravine (50°44'46.04" N, 21°01'46.77" E) located in the Kielce region (Małopolska Block) of the HCM (Fig. 1B) about 2 km north of Bardo village on the northern limb of the Bardo Syncline (Kozłowski et al. 2017).

Material for geochemical analysis has been selected from five small outcrops in the Pragowiec Ravine (Fig. 1C). The height of outcrops varies from 0.5 to 2 m. Ten samples with well-preserved graptolites were selected from the upper Bardo and the Pragowiec shales for stable carbon isotope analysis. Graptolites from these shales are important for precise biostratigraphy. Sampling at high-resolution (e.g., each 0.1 to 1 m) was not possible because the ravine is overgrown and is covered by recent mudflow deposits and anthropogenic debris and at the present is far away from H. Tomczyk's (1962) interpretation (Fig. 1C). Due to these factors and also to the complex folding and faulting of the shales, the exact superposition of layers below biozonal level is impossible to determine. Due to these severe constraints on the availability of the material, the resolution of carbon isotopic sampling in this study is kept approximately at two samples per biozone (see below), where biozones were determined based on abundant graptolite material. All the material is stored in the Geological Museum of Vilnius University, Lithuania.

Stable carbon isotope ($\delta^{13}C_{org}$)

Samples are mostly composed of terrigenous (shale) material with different carbonate content. For the purpose of the $\delta^{13}C_{org}$ analysis, the samples that were powdered were the same as those from which fossils were identified (total 10 samples). Approximately 0.9 mg of sample powder was used from each sample. Powder was dissolved using 5 N (mass equivalents) HCl acid for 24 hours at the room temperature to remove carbonate minerals. After that, the powder residue was washed with distilled water and dried.



Fig. 1. A — Simplified structural map of Central Europe (Bełka et al. 2002). HCM — The Holy Cross Mountains; MGCH — Mid German Crystalline High; OZ — Odra Zone; TBT — Tepla–Barrandian Terrane; USM — Upper Silesian Massif. B — Distribution of Silurian rocks in the Holy Cross Mountains area (Kozłowski et al. 2014) and the Prągowiec Ravine location. C — stratigraphical interpretation of the Silurian succession in the Prągowiec Ravine (Tomczyk 1962, fig. 9). D — Possible correlation of the Prągowiec Ravine Upper Wenlock graptolite biozones (Tomczyk 1962) with those of revised Lithuanian (Radzevičius 2006).

After the pre-treatment removal of carbonates, all samples (at least two per sediment sample) were weighed and wrapped in tin capsules. The prepared samples were combusted with combustion module (Costech Analytical Technologies, Inc.) connected via Picarro Liaison Interface A0301 to the laser-based Picarro Cavity Ring-Down Spectrometer G2121-i. The stable carbon isotope ratio was measured in CO₂ and presented as per mil deviations from internationally accepted standards with the reproducibility of $\pm 0.3\%$ for δ^{13} C. The international standards (IAEA-600, IAEA C1, IAEAC2 and SRM 4990C) were used for the calibration of the reference gas (CO₂).

Multivariate comparison of graptolite assemblages

In order to compare the compositional changes in graptolite assemblages in the Pragowiec Ravine, two methods were employed: non-metric multidimensional scaling and the comparison of turnover within the zones and among them. We employed a non-metric multidimensional scaling technique, a multivariate dimension reduction technique which is robust in revealing non-linear gradients in species composition (Patzkowsky & Holland 2012). We used the PAST program and the Raup-Crick metric (Hammer & Harper 2008), which is suitable for comparison of samples with differing and unknown abundances (Raup & Crick 1979). In our case, the latter property is especially desirable, since the abundance of graptolites is very difficult to measure, especially if there are variations in preservation (e.g., many fragmented rhabdosomes) and in the sizes of rock samples. In order to test the level of between-biozonal turnovers to inside-biozonal turnovers of graptolites in the Pragowiec Ravine, we performed all possible pair-wise comparisons between samples in a given biozone $c=(N_1^2-N_1)/2$, where N₁ is the number of samples in the first biozone) and between all the samples between two biozones $c=(N_1 \times N_2)$, where N₂ is the number of samples in the second biozone). For this purpose, as in the previous case, we used Raup-Crick compositional distance, which was calculated in the *Vegan* package (Oksanen et al. 2018) for the R programming environment (R Development Core Team 2015). Later on, the yielded distributions of compositional distances were compared with each other using the non-parametric Mann-Whitney test.

The analysis was performed on the graptolite occurrence data from the *praedeubeli*, *deubeli* and *ludensis* biozones (spanning most of the upper Homerian), since only those biozones had more than two collected samples each and were represented by sufficiently abundant graptolite material. The *praedeubeli* Biozone was represented by 15 samples, the *deubeli* Biozone by 12 samples, and the *ludensis* Biozone by 18 unambiguously assigned samples with abundant graptolite material. The single occurrence of *Semigothograptus* cf. *meganassa* was not used in these multivariate analyses because of the ambiguity of its species assignment.

Graptolite biozones

H. Tomczyk (1962) distinguished the graptolite biozones in the Prągowiec Ravine (Fig. 1C), describing six graptolite biozones in the Homerian–lower Gorstian interval (Fig. 1D). Some of these biozones are not used at present. The Prągowiec Ravine graptolite revision and new graptolite biozones correlation of Tomczyk's biozones were given by E. Porębska (in Masiak 2010, fig. 20). At the same time as the above biozonations were being developed, S. Radzevičius (2006) distinguished graptolite biozones (Fig. 1D) in the *lundgreni–nilssoni* interval of the Prągowiec ravine.

Cyrtograptus lundgreni Tullberg (Fig. 2A), *Monograptus flemingii* (Salter) (Fig. 2B); *Monoclimacis flumendosae* (Gortani) (Fig. 2C) and *Pristiograptus pseudodubius* Bouček occur in the Bardo beds (sample VU-U-10). *M. flemingii*, *Mcl. flumendosae* and *P. pseudodubius* are long ranging species, which appeared in the middle Sheinwoodian above the *riccartonensis* Biozone (Zalasiewicz et al. 2009) and disappeared in the middle Homerian during the *lundgreni* Event (Koren' 1987). *C. lundgreni* is the index species and its range defines the *lundgreni* Biozone (Zalasiewicz & Williams 1999).

The only *Testograptus testis* (Barrande) (Fig. 2D) identified was in the Bardo bed sample VU-U-6. A *testis* Biozone is recognized between the *lundgreni* and *nassa* biozones in the HCM (Tomczyk 1958; Ryka & Tomczyk 1959) as well in the Prągowiec ravine (Tomzcyk 1962). *T. testis* appeared later than *Cyr. lundgreni*, which accompanies *T. testis* until the *lundgreni* extinction. Therefore, the interval with *T. testis* represents the upper part of the *lundgreni* Biozone (Zalasiewicz et al. 2009) and is sometimes referred to it as the upper subzone (Jaeger 1991; Štorch 1994).

Pristiograptus parvus Ulst (Fig. 2E) and fragments of *Gothograptus nassa* (Holm) (Fig. 2F) occur in sample VU-U-7 which is composed of dark yellow clayey shales, probably of the Bardo beds. *G. nassa* ranges from the lower part of *parvus* Biozone to the middle part of the *praedeubeli* Biozone

(Kozłowska et al. 2009) or the *parvus–nassa* interval (Maletz 2010). The short-ranging *P. parvus* appears after the *lundgreni* extinction and defines the *parvus* range Biozone (Ulst 1974).

Gothograptus nassa (Fig. 2H) and a trilobite pygidium of Odontopleura cf. ovata Emmrich (Fig. 2G) occur in Bardo beds sample VU-U-9. G. nassa ranges through the parvus– praedeubeli interval. A mass occurrence of benthic fauna (e.g., Odontopleura) marks the lower part of the nassa Biozone in the Prągowiec ravine (Tomczykowa 1957) and Bartoszyce IG 1 borehole (Porębska et al. 2004) in Poland. According to Calner et al.'s (2006a) data, O. ovata was recovered from just above the Grötlingbo Bentonite on Gotland. The Grötlingbo Bentonite is widespread in Laurussia at a stratigraphic level corresponding to the parvus Biozone (Kiipli et al. 2008).

Colonograptus praedeubeli (Jaeger) (Fig. 2K) and *G. nassa* (Fig. 2L) occur in the Prągowiec beds (sample VU-U-2). *C. praedeubeli* ranges from the *praedeubeli* Biozone to the middle of the *ludensis* Biozone (Koren' 1991), but is most common in the *praedeubeli* Biozone. *Col. praedeubeli* (Fig. 2M) also occurs in sample VU-U-8 (Prągowiec beds).

Colonograptus praedeubeli (Fig. 2O) occurs in sample VU-U-5 (Pragowiec beds). There is Col. cf. deubeli (Jaeger) (Fig. 2N) identified in the same sample. Col. deubeli is characterized by funnel- or trumpet-like sicula with distinct dorsal process and rapid increase in rhabdosome width (Koren' & Suyarkova 1994). Our Col. cf. deubeli specimen has a moderately expanded sicular aperture, but the width of rhabdosome increases gradually. Col. cf. deubeli is similar in terms of rhabdosome width to Pristiograptus idoneus Koren', but differs in the form of the sicula. The sicula of P. idoneus is strongly ventrally curved (Koren' 1992). Some fragments of retiolitids with the nassa type of apertural hoods have been found in the same sample (Fig. 2P). There is G. nassa, Semigothograptus meganassa (Rickards & Palmer 2002) and Neogothograptus eximinassa Maletz with genicular hoods of the nassa type in the parvus-ludensis interval (Kozłowska 2016). The flattened and poor preservation specimens complicate identification. Retiolitids are distinguished based mostly on isolated, 3D material (Lenz & Kozłowska 2007). However, N. eximinassa marks the ludensis Biozone (Maletz 2008), S. meganassa and G. nassa have similar stratigraphic ranges, and are confined to the parvus-deubeli interval (Kozłowska et al. 2009; Kozłowska 2016). The main difference that can be distinguished in the poorly preserved material between S. meganassa and G. nassa is in the width of rhabdosome. S. meganassa is wider than G. nassa (Kozłowska-Dawidziuk et al. 2001; Rickards & Palmer 2002) and N. eximinassa as well. Our specimen is relatively wide (approximately 2 mm) but the structure of apertural hoods is invisible and the proximal end of rhabdosome is absent. Based on this, this specimen has been identified as Semigothograptus cf. meganassa. However, the index species Col. deubeli, the first appearance of which marks the base of the deubeli and ranges to the middle of the ludensis Biozone (Jaeger 1991; Koren' 1991), is not found in the sample. Therefore, the graptolite assemblage of VU-U-5 sample is interpreted as belonging to the praedeubeli-deubeli interval.

Pristiograptus virbalensis Paškevičius and Col. ludensis (Murchison) (Fig. 2J) occur in sample VU- U-3 (Prągowiec beds). *P. virbalensis* is known from the *praedeubeli–nilssoni* biozones in Lithuania (Radzevičius et al. 2014a). Col. ludensis is the index species for the uppermost Homerian *ludensis* Biozone (Zalasiewicz et al. 2009).

P. frequens Jaekel and *Col. gerhardi* (Kühne) (Fig. 21) occur in sample VU-U-1 (Prągowiec beds). The long-ranging



Fig. 2. The main fauna from the Prągowiec Ravine. **A–C:** Sample VU-U-10, upper Bardo beds, *lundgreni* Biozone; A — *Cyrtograptus lundgreni* Tullberg, SV-PER-027; B — *Monograptus flemingii* (Salter), SV-PER-026; C — *Monoclimacis flumendosae* (Gortani), SV-PER-026a. **D** – Sample VU-U-6, upper Bardo beds, *lundgreni* Biozone, *Testograptus testis* (Barrande), SV-PER-018. **E**, **F:** Sample VU-U-7, upper Bardo beds, *parvus* Biozone; E — *Pristiograptus parvus* Ulst, SV-PER-023; F — *Gothograptus nassa* (Holm), SV-PER-023a. **G**, **H:** Sample VU-U-9, upper Bardo beds, *parvus* Biozone; G — trilobite pygidium of *Odontopleura* cf. *ovata* Emmrich SV-PER-001a; H — *Gothograptus nassa* (Holm) SV-PER-001. **I** — Sample VU-U-1, Prągowiec beds, *ludensis* Biozone, *Colonograptus gerhardi* (Kühne) SV-06-8. **J** — Sample VU-U-3, Prągowiec beds, *ludensis* Biozone, *Colonograptus nassa* (Holm) SV-69-9. **M** — Sample VU-U-8, Prągowiec beds, *praedeubeli* (Jaeger) SV-69-6; L — *Gothograptus nassa* (Holm) SV-69-9. **M** — Sample VU-U-8, Prągowiec beds, *praedeubeli* (Jaeger) SV-A04-6. **N–P:** Sample VU-U-5, Prągowiec beds, *praedeubeli* (Jaeger) SV-A07-1; O — *Colonograptus praedeubeli* (Jaeger) SV-A07-1; P — *Semigothograptus* cf. *meganassa* (Rickards Palmer) SV-A07-1b. **R** — Sample VU-U-4, Prągowiec beds, *nilssoni* Biozone, *Neodiversograptus nilssoni* (Barrande) SV-A06-4. Scale bars are 1mm.

P. frequens appears in the *nassa* Biozone and disappears in the *leintwardinensis* Biozone (Urbanek et al. 2012) or above the *tenuis* Biozone (Štorch et al. 2014). *Col. gerhardi* appears in the upper part of the *ludensis* Biozone and disappears in the lower part of the *nilssoni* Biozone (Kozłowska-Dawidziuk et al. 2001; Radzevičius & Paškevičius 2005) but dominated in the upper part of the *ludensis* Biozone (Štorch et al. 2016).

An approximately 10 cm long mesial rhabdosome fragment of *Neodiversograptus nilssoni* (Barrande) (Fig. 2R) was found in the sample VU-U-4 (Pragowiec beds). *N. nilssoni* indicates the lower part of the Ludlow and is the index species for the *nilssoni* Biozone (Zalasiewicz et al. 2009).

In summary, the samples for the $\delta^{13}C_{org}$ analyses come from *lundgreni*, *parvus*, *praedeubeli–deubeli*, *ludensis* and *nilssoni* biozones (Table 1). There are no findings of typical *Col. Deubeli*, the index species of the *deubeli* Biozone, in the studied isotopic samples. Graptolites described in the samples VU-U-2, VU-U-8 and VUU-5 have long range and can be discovering in both *praedeubeli*, *deubeli* biozones. According to that, the *praedeubeli–deubeli* interval is not split into the separate biozones there.

Results and discussion

Turnover in graptolite assemblages

Although some indications of the community turnover can be distinguished from the range charts of the graptolite species, the ecological significance of those changes is not especially obvious. The composition of ecological communities is expressed as the identity of species and their abundance. Therefore, even if we observe a very similar set of species in two consecutive zones, if the relative frequencies of species in these zones are very different the final compositions in these two zones will be very ecologically different. As a result, numerical comparisons of sets of assemblages between zones can be used in statistical testing of significance of temporal changes. If the turnover between zones is continuous (when

Table 1	1:	Isotopic	data	for	the	Prągow	iec	Ravine	e sam	ples
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compositions change continuously without breaks), we should expect the fossil assemblages to significantly overlap in the multivariate compositional spaces. On the other hand if there are sharp changes in assemblage composition between zones we should expect clear separation.

The comparison of distances between samples in the praedeubeli Biozone Median=0.21, SD=0.21 with distances between samples of praedeubeli and deubeli biozones Median=0.658, SD=0.21 revealed their highly significant difference, p<0.01 (Fig. 3A). Similarly, there is a highly significant difference between sample distances in the deubeli Biozone Median=0.211, SD=0.16 and between distribution of distances between samples from the deubeli and ludensis biozones Median=0.817, SD=0.11, p<0.01 (Fig. 3B). A similar pattern of divergence can be seen between assemblage distances in the ludensis Biozone Median=0.249, SD=0.27, and the assemblage compositional distances between deubeli and *ludensis* biozones, Median=0.817, SD=0.11, p<0.01 (Fig. 3C). Although an overlap in distances can be observed between consecutive biozones, there is strong separation between assemblages. Moreover, compositional distances between biozones show strong modality, which points to the conclusion that biozonal boundaries represent genuine turnover episodes in development of graptolite faunas in the area represented by samples from the Pragowiec Ravine. Additionally, it appears that the differences in graptolite assemblages were on average greater between the deubeli and ludensis biozones (difference between medians of distances=0.448) than between the praedeubeli and deubeli biozones (difference between medians of distances=0.547).

The same conclusion of between-zonal differentiations in graptolite assemblages can be drawn based on the results of the non-metric multidimensional scaling analysis of the composition of graptolite assemblages (Fig. 4). The assemblages from all three analysed biozones are very clearly distinguished, with graptolite communities from the *praedeubeli* and *deubeli* biozones being more closely related to each other, as was suggested by the results of the previously presented pair-wise comparative analyses. Based on the taxonomic compilation of

	Sample No.	Sampling sites	GPS coordinates	Formation	Biozone	$\delta^{13}C_{\rm org}$	Fauna
1	VU-U-10	A00	50°44'46.2" N, 21°02'16.0" E	Bardo beds	lundgreni	-30.7	M. flemingii, Mcl. flumendosae, P. pseudodubius, Cyr. lundgreni
2	VU-U-6	A00	50°44'46.2" N, 21°02'16.0" E	Bardo beds	lundgreni	-30.1	T. testis
3	VU-U-7	A00	50°44'46.2" N, 21°02'16.0" E	Bardo beds	parvus	-29.3	P. parvus, G. nassa
4	VU-U-9	A00	50°44'46.2" N, 21°02'16.0" E	Bardo beds	parvus	-28.7	G. nassa, Odontopleura cf. ovata
5	VU-U-2	A08	50°44'46.8" N, 21°02'10.8" E	Prągowiec beds	praedeubeli	-30.2	Col. praedeubeli, G. nassa
6	VU-U-8	A04	50°44'48.7" N, 21°02'01.9" E	Prągowiec beds	praedeubeli	-30.8	Col. praedeubeli
7	VU-U-5	A07	50°44'47.6" N, 21°02'08.1" E	Prągowiec beds	praedeubeli–deubeli	-30.4	Col. praedeubeli, Col. cf. deubeli, S. cf. meganassa
8	VU-U-3	A04	50°44'48.7" N, 21°02'01.9" E	Prągowiec beds	ludensis	-31.2	P. virbalensis, Col. ludensis
9	VU-U-1	A06	50°44'49.3" N, 21°01'58.1" E	Prągowiec beds	ludensis	-30.2	P. frequens, Col. gerhardi
10	VU-U-4	A06	50°44'49.3" N, 21°01'58.1" E	Prągowiec beds	nilssoni	-30.5	N. nilssoni



Fig. 3. Raup-Crick distances of graptolite assemblages from the Prągowiec Ravine (p-values show probabilities of null hypothesis for equality of medians): A — between samples of the praedeubeli Biozone (c=105), and between samples of the praedeubeli and deubeli (c=180) biozones (p<0.01); B — between samples of the deubeli Biozone (c=66), and between samples of the deubeli and ludensis (c=216) biozones (p<0.01); C — between samples of the ludensis Biozone (c=153), and between samples of the deubeli and ludensis (c=216) biozones (p<0.01). Arrows point to the medians for each distribution of between sample distances. Here c=the number of pairwise comparisons in each category.

Koren' (1991), the *ludensis* Biozone witnessed significant restructuring in diversity of graptolites-retiolitids increased in species richness and monograptids a decline in species richness. Similarly, in the Canadian sections described by Lenz (1994, 1995), there was a significant turnover of monograptids between the *deubeli* and *ludensis* biozones. According to the global sequencing of the graptoloid clade, the *ludensis* Biozone marked an interval of significant increase in graptolite diversity which ended in the early Ludlow (Cooper et al. 2014). At least two large extinction events separated by hundreds of thousands of years occurred in the late Homerian, not including the *lundgreni* extinction event (Crampton et al. 2016). Thus the intensive graptolite turnover described in the Pragowiec Ravine is part of the wider pattern.

Trend in $\delta^{13}C_{org}$ values

The $\delta^{13}C_{org}$ values are low and vary from -31.2 ‰ to -28.7 ‰; (Table 1). Similar variations in upper Homerian $\delta^{13}C_{org}$ values are recorded from the part of Poland corresponding to the West of Baltica (Porębska et al. 2004; Sullivan et al. 2018). The highest variation of $\delta^{13}C_{org}$ values is found in the upper Bardo beds. The values rise from -30.7 ‰ and -30.1 ‰ in the *lundgreni* Biozone to -29.3-28.7 ‰ in the *parvus* Biozone (Fig. 5) and drop in the Prągowiec beds and fluctuate overall by 1 ‰ between -31.2 ‰ and -30.2 ‰ through the *praedeubeli* – *nilssoni* Biozone interval (Fig. 5).

There are two positive $\delta^{13}C_{carb}$ peaks in the mid-upper Homerian (Cramer et al. 2011; Melchin et al. 2012). The first $\delta^{13}C_{carb}$ peak is found in the *parvus–nassa* biozones and second one in the *ludensis* Biozone. Such interpretation is based on material from the West Midlands (England) and Gotland (Cramer et al. 2012). The West Midlands sections are dominated by shallow marine facies with very rare graptolites and the correlation of these localities is based only on conodont biostratigraphy, sequence stratigraphy and high-precision zircon (U–Pb) dating of bentonites. Therefore, bearing in mind all the possible uncertainties in correlation, the incorporation of $\delta^{13}C_{carb}$ excursion with graptolite biozones should be imprecise. In the material from the Viduklė–61 borehole (Lithuania)



Fig. 4. Non-metric multidimensional scaling of graptolite assemblages from the Pragowiec Ravine (Stress=0.26).

with a continuous graptolite sequence, the first positive excursion represents the *parvus* Biozone to lower *praedeubeli* Biozone interval and the second positive excursion represents the *deubeli* Biozone to lower *ludensis* Biozone (Radzevičius et al. 2014a) for the Gėluva Regional Stage in Lithuania (Fig. 5). After graptolite revision from the West Midlands, the same conclusions have been reached by Fry et al (2017).

In the gamma log data from the Viduklė–61 well, two cycles of ~16.7 m and five with ~6.7 m period lengths were determined in the Gėluva interval (Fig. 5) (Radzevičius et al. 2014b). Similar cyclicities were found in other mid-upper Homerian geological sections of West Lithuania (Radzevičius et al. 2017). The Gėluva age corresponds to the mid-later Homerian, namely from 428.45±0.35 Ma to 427.86±0.32 Ma (total ~0.59 Ma) (Cramer et al. 2015). Based on these dates, long cycles are about 0.3 Ma and short about 0.12 Ma. The duration of cycles alone could be tentatively interpreted as the 4th and 5th order cycles which are close to the Milankovitch eccentricity cycles generated by the orbital forcing (Miall 2010).

Two stage slices (Ho2 and Ho3) have been distinguished in the mid-upper Homerian (Cramer et al. 2011) or the Geluva



Fig. 5. Mid–upper Homerian global stratigraphical scale (Melchin et al. 2012); graptolite biozones (Koren' et al. 1996); stage slices (Cramer et al. 2011); regional stages of the East Baltic (Paškevičius et al. 1994); Conodont turnover stages (Radzevičius et al. 2016); cyclostratigraphy in the Viduklė-61 borehole and situation of the Ančia Member (Radzevičius et al. 2017); generalized $\delta^{13}C_{carb}$ curve (Cramer et al. 2011); lithology (Tomczyk 1962); $\delta^{13}C_{org}$ data (this paper) and generalized distribution of graptolites (Radzevičius 2006) of the Prągowiec Ravine (HCM, Poland).

Regional Stage in Lithuania. A stage slice is an informal stratigraphic unit that is defined on the basis of biochemostratigraphy. According to Cramer et al. (2011), the base of the Ho2 slice lies within the *parvus* Biozone and includes the first peak of the Mulde δ^{13} C excursion. The Ho3 slice ranges from the base of the *ludensis* Biozone to the base of the *nilssoni* Biozone with the second peak of the Mulde δ^{13} C excursion. *P. parvus* are found from the upper Bardo Beds in the Prągowiec Ravine. According to that, the lower Ho2 slice boundary doesn't directly corresponding to the Bardo and Prągowiec beds boundary. In the cyclostratigraphic data from West Lithuania (Radzevičius et al. 2017), the boundary between Ho2 and Ho3 is coincident with the boundary between Gėluva 4,1 and Gėluva 4,2 (4th order) sedimentary cycles.

It has previously been hypothesized that the evolutionary turnover of conodonts was spread through the upper Homerian (Jeppsson et al. 1995), although more detailed integrated analysis of interregional data revealed that the first appearances of zonal taxa, *O. bohemica longa*, *K. ortus absidata* and *Ctenognathodus murchisoni* converge to the beginning of the Mulde event as indicated by the integrated stratigraphy (Radzevičius et al. 2016). Therefore appearances of those taxa in different regions should be highly diachronous (see in fig. 5 "delayed dispersal phase model", which shows an initial phase of evolutionary changes which is followed by the longer dispersal phase of conodont taxa). On the other hand, based on the analyses of local communities, it was shown that there was no significant community turnover in conodonts at the proposed boundary of the Mulde event (Jarochowska et al. 2018). Since there is positive evidence on evolutionary taxic disappearance and appearance (Jeppsson et al. 1995; Radzevičius et al. 2016), this lack of congruence could be explained by disproportionate effects of the Mulde turnover event on rare taxa. On the other hand, although the composition was similar in conodonts before and after the Mulde event, their communities changed to a more even and simple abundance distribution, and there was a shift to lower abundance and higher autocorrelation of abundance fluctuations which points to the transition to differing community states in the upper Homerian in the conodont clade (Spiridonov 2017; Spiridonov et al. 2017a).

On the other hand, range data for graptolites (including those presented here) and the multivariate analyses of their cooccurrences point to the possibility that the late Homerian

experienced several significant turnover events between biozones. Those are possibly related to the periodic sea level perturbations associated with the 4th and 5th order cycles which were clearly distinguished in the Baltic data (Fig. 5, and also Radzevičius et al. 2017) and briefly described in this paper. The same physical changes in environment are highly congruent with Cramer et al.'s (2011) proposed stage slices of the Homerian which are conceptually similar to assemblage biozones (Fig. 5). Moreover, recent cyclostratigraphic-spectral analytical studies revealed that external forcing due to Milankovitch forcing was a significant factor in driving changes in community compositions, and abundance of conodonts (Spiridonov et al. 2016, 2017a), as well as macroevolutionary diversity of graptolites during the Ordovician and Silurian (Crampton et al. 2018). The confluence of evidence points toward the dominance of the "common geological cause" (sensu Peters & Foote 2002) mechanisms of change in stratigraphy, ecology and macroevolution of biota in the Silurian.

The first $\delta^{13}C_{org}$ excursion is well represented in the *parvus* Biozone in the Pragowiec Ravine and related to the trilobite mass occurrence interval in the upper part of the Bardo beds (Fig. 5). This interval could be correlated with upper part of microlaminated, varve-like marlstone of the Ančia Member in the Lithuania (Radzevičius et al. 2014a) and with the lower part of Ho2. The second $\delta^{13}C_{\text{org}}$ excursion is very indistinct. This could be related to the deep marine facies which dominate the sedimentary record at the Pragowiec Ravine. A similar situation is found in the Zwierzyniec-1 borehole from deep marine facies of South-East Poland (Sullivan et al. 2018). The magnitudes of $\delta^{13}C$ excursions are higher in shallow water than in deeper water settings (Noble et al. 2005), which makes it more difficult to distinguish them in offshore shales. On the other hand, it can be related to low sampling resolution. However, the present results are consistent with previous observations and show that the $\delta^{13}C_{\mbox{\scriptsize org}}$ data can be used as a supporting data for stratigraphic correlation even in terrigenous deep water facies.

Conclusions

The first positive Homerian $\delta^{13}C_{org}$ excursion peak is well represented in the Prągowiec Ravine and is in the *parvus* Biozone (upper part of Bardo beds). The upper Bardo beds can be correlated with the lower part of the Ho2 stage slice and the Ančia Member in the Baltic Silurian Basin as well. Nevertheless, the $\delta^{13}C_{org}$ curve shows a broadly similar trend to that represented in the global Homerian $\delta^{13}C_{carb}$ curve and can be used for the correlation between these datasets.

Additionally, the quantitative analyses of graptolite samples revealed that there was a significant turnover in this planktic group between biozones in the late Homerian *(post lundgreni* Event) at the studied site. The greatest turnover occurred between the *deubeli* and *ludensis* biozones in the Prągowiec Ravine area. Acknowledgements: We thank colleagues J. Mažeika and R. Paškauskas from the Nature Research Centre (Lithuania) for undertaking isotope analyses. We sincerely thank M. Whittingham for the English correction A.S.'s research is supported by the Research Council of Lithuania grant No. 09.3.3-LMT-K-712-02-0036. This research was supported by the Open Access to research infrastructure of the Nature Research Centre under the Lithuanian open access network initiative. This is a contribution to "IGCP 652: Reading geological time in Paleozoic sedimentary rocks: the need for an integrated stratigraphy" and to "Event Stratigraphy in the Silurian Sedimentary Basin of Lithuania", a Vilnius University project.

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Appendix

Graptolite data used in multivariate analysis

Zone/taxon	P. ludensis	P. frequens	C. gerhardi	P. virbalensis	C. deubeli	Gothograptus sp.	P. praedeubeli
praedeubeli	0	0	0	1	0	0	0
praedeubeli	0	0	0	1	0	0	1
praedeubeli	0	0	0	0	0	0	1
praedeubeli	0	0	0	0	0	0	1
praedeubeli	0	0	0	1	0	0	1
praedeubeli	0	0	0	1	0	0	1
praedeubeli	0	0	0	1	0	0	0
praedeubeli	0	0	0	0	0	0	1
praedeubeli	0	0	0	0	0	0	1
praedeubeli	0	0	0	0	0	0	1
praedeubeli	0	1	0	1	0	1	1
praedeubeli	0	0	0	0	0	0	1
praedeubeli	0	0	0	0	0	0	1
praedeubeli	0	0	0	0	0	0	1
praedeubeli	0	0	0	0	0	1	1
deubeli	0	0	0	0	1	1	0
deubeli	0	0	0	0	1	0	0
deubeli	0	0	0	0	1	1	0
deubeli	0	0	0	0	1	0	0
deubeli	0	0	0	0	1	0	1
deubeli	0	1	0	0	1	0	1
deubeli	0	0	0	0	1	0	0
deubeli	0	0	0	1	1	0	0
deubeli	0	0	0	0	1	0	1
deubeli	0	0	0	0	1	0	0
deubeli	0	0	0	0	1	0	0
deubeli	0	0	0	0	1	1	1
ludensis	1	1	0	0	0	0	0
ludensis	0	1	1	0	0	0	0
ludensis	1	0	0	0	0	0	0
ludensis	0	0	1	0	0	0	0
ludensis	0	0	1	0	0	0	0
ludensis	1	1	0	0	0	0	0
ludensis	1	0	0	1	0	0	0
ludensis	1	1	0	1	0	0	0
ludensis	1	1	0	0	0	0	0
ludensis	1	0	0	0	0	0	0
ludensis	1	0	0	0	0	0	0
ludensis	1	0	0	0	0	0	0
ludensis	1	0	0	0	0	0	0
ludensis	1	0	1	1	0	0	0
ludensis	1	1	0	0	0	0	0
ludensis	1	0	0	1	0	0	0
ludensis	1	1	0	0	0	0	0
ludensis	1	0	0	1	0	0	0