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## Faunal dynamics across the Silurian–Devonian positive isotope excursions ( $\delta^{13}$ C, $\delta^{18}$ O) in Podolia, Ukraine: Comparative analysis of the Ireviken and Klonk events

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Two global isotopic events, the early Sheinwoodian (early Wenlock) and that at the Silurian-Devonian transition, have been comprehensively studied in representative carbonate successions at Kytayhorod and Dnistrove, respectively, in Podolia, Ukraine, to compare geochemistry and biotic changes related correspondingly to the Ireviken and Klonk events. These two large-scale isotope excursions reveal different regional ecosystem tendencies. The well-defined increasing trend across the Llandovery-Wenlock boundary in siliciclastic input, redox states and, supposedly, bioproductivity, was without strict correlative relations to the major <sup>13</sup>C enrichment event. The environmental and biotic evolution was forced by eustatic sea-level fluctuations and two-step climate change toward a glaciation episode, but strongly modified by regional epeirogeny movements due to location near the mobile Teisseyre-Törnquist Fault Zone. Thus, the global early Sheinwoodian biogeochemical perturbation was of minor depositional significance in this epeiric sea, as in many other Laurussian domains. Conversely, the Podolian sedimentary record of the Klonk Event exhibits temporal links to the abrupt  $\delta^{13}$ C anomaly, overprinted by a tectonically driven deepening pulse in the crucial S–D boundary interval. This carbon cycling turnover was reflected in the regional carbonate crisis and cooling episodes, paired with a tendency towards eutrophication and recurrent oxygen deficiency, but also with major storms and possible upwelling. Faunal responses in both Podolian sections follow some characters of the Silurian pattern worldwide, as manifested by conodont changeover prior to the major early Sheinwoodian isotopic/climatic anomaly. This contrasts with the relative brachiopod and chitinozoan resistances in the course of the Ireviken Event. Also, during the Klonk Event, a moderate faunal turnover, both in benthic and pelagic groups, occurred only near the very beginning of the prolonged <sup>13</sup>C-enriched timespan across the system boundary, possibly due to progressive dysoxia and temperature drop. The characters point to a peculiarity of the Klonk Event by comparison with the Silurian global events, and some similarity already to the succeeding Devonian transgressive/anoxic episodes.

Key words: Brachiopoda, Conodonta, Chitinozoa, carbon isotopes, oxygen isotopes, geochemistry, faunal dynamics, Ireviken Event, Klonk Event, Silurian, Podolia.

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## Introduction

In contrast to previous views, the Silurian Period has recently been shown as a short mid-Palaeozoic time slice characterized by recurring changes of the global ecosystem following

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recovery from the end-Ordovician extinction (Calner 2008). Surprisingly large-amplitude positive carbon and oxygen stable isotope anomalies (up to 10% in the late Ludlow; Fig. 1) were episodes of profound environmental change ("oceanic events"; Jeppsson 1990, 1998) manifested by distinctive fa796

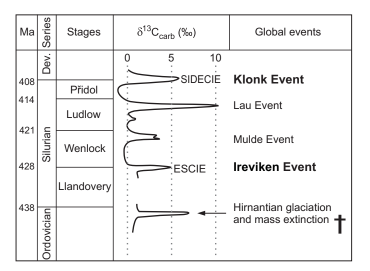


Fig. 1. Schematic compilation of positive  $\delta^{13}$ C excursions and global events in the Late Ordovician to Early Devonian interval (adapted from Stricanne et al. 2006: fig. 1; for more refined composite isotopic curve see Cramer et al. 2011a: fig. 3; for discussion of the Silurian timescale see Cramer et al. 2011b and Melchin et al. 2012). Abbreviations: Dev., Devonian; ESCIE, early Sheinwoodian carbon isotope excursion; SIDECIE, Silurian–Devonian carbon isotope excursion.

cies variation and associated stepwise extinctions both preceding plus coincident with the isotopic excursions (see summary in Munnecke et al. 2003, 2010; Calner 2008; and Melchin et al. 2012). Regular occurrences of teratological acritarchs record these high stress conditions (Munnecke et al. 2012). In general, pelagic communities were more strongly affected than shallow-water biota in the repeatedly pronounced oceanographic and climatic changes (see overviews in Kaljo et al. 1996; Jeppsson 1998; and Calner 2008; also e.g., Boucot 1990; Brett and Baird 1995; Porebska et al. 2004; Johnson 2006; Stricanne et al. 2006; Lehnert et al. 2007b, 2010; Loydell 2007; Eriksson et al. 2009; Nardin et al. 2011; Molloy and Simpson 2012; Manda et al. 2012; Noble et al. 2012). In fact, Silurian graptolites exhibit a very dynamic evolutionary history (Urbanek 1993; Štorch 1995; Sadler et al. 2011), contrary to the relatively stable Silurian reef biotas (Copper 2002b), and crinoid and brachiopod faunas (Talent et al. 1993; Kaljo et al. 1996; Ruban 2011). On the other hand, synchroneity of mass mortalities in graptolites, conodonts and, especially, the poorly known benthos is uncertain (e.g., Copper 2002b; Loydell 2007; Manda et al. 2012; Noble et al. 2012), and also separate terms have been proposed for the graptolite- and conodont-based crises (see Jeppsson 1990; 1998; Urbanek 1993; Štorch 1995; Kaljo et al. 1996). Conversely, some authors used terms coupling both biotic and geochemical aspects of global events (e.g., the Ireviken isotopic excursion; Cramer et al. 2011a; see also Buggisch and Joachimski 2006), even though the anomalously heavy  $\delta^{13}$ C values are roughly correlated with these medium-order biosphere perturbations (see Melchin et al. 2012: fig. 21.11).

Five distinct  $\delta^{13}$ C excursions have been identified in the "classical" Silurian to Lower Devonian low-latitude car-

bonate successions of Podolia, Ukraine (Azmy et al. 1998; Kaljo et al. 2007, 2012; Małkowski et al. 2009; Fig. 1). Biotic responses and ecological interactions are as yet poorly known from these key stratigraphical levels. Two of these biogeochemical perturbations, the early Sheinwoodian excursion (early Wenlock; ESCIE, cf. Lehnert et al. 2010) and the Silurian-Devonian passage excursion (SIDECIE = SIDE of Kaljo et al. 2012), approximately correlative with the major Ireviken (2<sup>nd</sup> order) and subordinate Klonk (5<sup>th</sup> order) biotic crises, respectively (see Kaljo et al. 1996; Jeppsson 1998), have been comprehensively studied by us in representative Podolian localities to compare geochemical and faunal trends in the epeiric domain. The goal is to present distributional and frequency patterns of benthic (brachiopods) and presumably pelagic (conodonts, chitinozoans) biota across these two  $\delta^{13}C$  anomalies, which are separated by a ca. 14 my interval (Melchin et al. 2012). The integrated event-stratigraphical analysis of biotic signals from very different ecologic groups, against stable isotope and inorganic geochemical proxies, provides an improved insight into concomitant ecosystem relationships in the rapidly evolving Silurian marine realm.

The responsibility of authors is as follows: GR and KM for geochemical data and interpretation; AB and RW for brachiopod and chitinozoan results, respectively; and DD and HS for regional and conodont aspects; the final discussion and conclusions are joint.

Institutional abbreviations.—ZPAL, Institute of Palaeobiology, Polish Academy of Sciences, Warszawa, Poland.

*Other abbreviations.*—ESCIE, early Sheinwoodian carbon isotope excursion; FAD, first appearance datum; GC–MS, Gas Chromatography–Mass Spectrometry; ICP, Inductively Coupled Plasma; S–D, Silurian–Devonian; SIDECIE, Silurian–Devonian carbon isotope excursion; SOIE, Sheinwoodian oxygen isotope excursion; TC, total carbon; TESZ, Trans-European Suture Zone; TIC, total inorganic carbon; TOC, total organic carbon; USGS, U.S. Geological Survey ; V-PDB, Vienna-Pee Dee Belemnite Standard.

### Geological setting

The extended middle Silurian to Lower Devonian marine succession of the Dniester Basin (Fig. 2) is considered as one of the most continuous successions of this age in the world, and have a long history of geological and palaeontological investigations (see Kozłowski 1929; Nikiforova et al. 1972; Tsegel-nyuk et al. 1983; Koren et al. 1989; Uchman et al. 2004; Kaljo et al. 2007; Skompski et al. 2008; Małkowski et al. 2009; Krzemiński et al. 2010). The mostly carbonate deposits are well exposed in the escarpments of the Dniester valley and its tributaries, and form an internally conformable regressive unit with a total thickness of more than 900 m. The essentially undeformed Podolian sequence is strongly controlled by its location in the marginal belt of the East European Platform, near

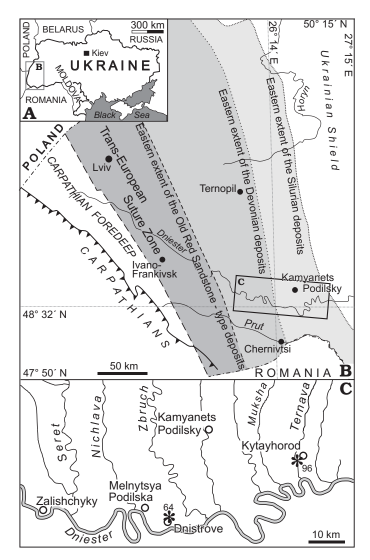


Fig. 2. **A**. General location of the study area in Ukraine. **B**. Distribution of the Silurian and Devonian deposits in Podolia, SW Ukraine, and geological setting of the study area. **C**. Location of two sampled outcrops (arrowed) in the vicinity of Dniester valley (modified from Małkowski et al. 2009: fig. 1). Section numbers after Nikiforova et al. (1972): Kytayhorod (96) and Dnistrove (64).

to the mobile Trans-European Suture Zone—the TESZ or Teisseyre-Törnquist Fault Zone (Fig. 2B; see Drygant 2000; Sliaupa et al. 2006; Krawczyk et al. 2008). The succession thickens toward the suture zone where the sedimentation rate was controlled by platform-margin faults and subsidence.

In the Silurian succession, which is more than 400 m thick, two depositional phases can be distinguished (Drygant 2003). The Wenlock and Ludlow phase of carbonate shelf sedimentation is characterized by numerous bioherms and biostromes (see Koren et al. 1989; Skompski et al. 2008; Łuczyński et al. 2009), and the Pridoli phase is characterized by rapid and differentiated sedimentation rates, especially in the vicinity of the platform margin. The Lower Devonian (Lochkovian) marine deposits, about 530 m thick, can be assigned to a third phase of continuous sedimentation (see Małkowski et al. 2009). The phase is characterized by the flysch-like interbeds of limestone and shale with gradually increasing amounts of terrigenous material resulting for gradual regression and a simultaneous uniform but comparatively strong, subsidence regime (e.g., Uchman et al. 2004).

# Facies evolution and analysed sections

The lower sampled interval (Fig. 3) includes open-shelf carbonate facies, very well naturally exposed in the Dniester River valley (Tarnava tributary) near the village of Kytayhorod (section no. 96 of Nikiforova et al. 1972; see Małkowski et al. 2009: fig. 3A; N48°32'16.9", E26°14'21.4"). The

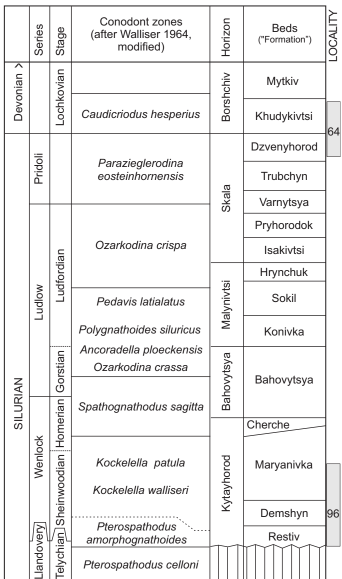


Fig. 3. Stratigraphic scheme of the Silurian deposits of Podolia (after Nikiforova and Predtechensky 1968; Drygant 1984), and stratigraphic position of studied localities (see Fig. 2): Kytayhorod (96) and Dnistrove (64).

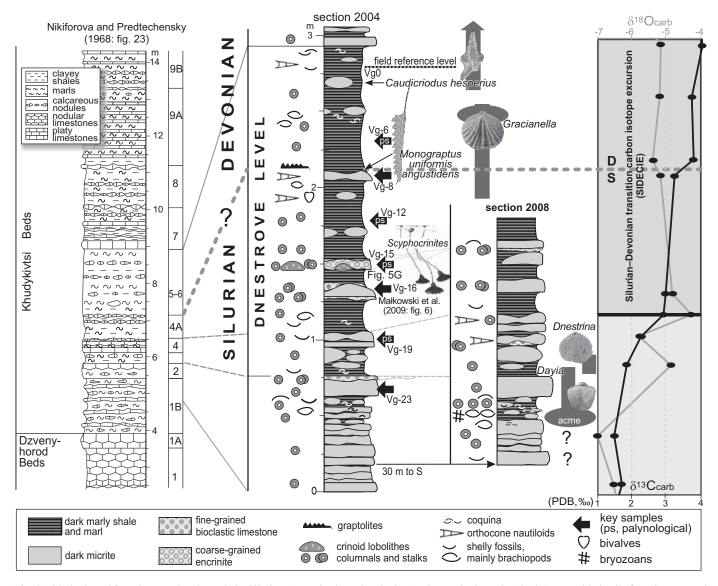


Fig. 4. Lithologic and faunal succession through the Silurian–Devonian boundary beds at Dnistrove in the early Klonk Event, with detail of newly proposed Dnistrove Level, against the secular pattern of  $\delta^{13}$ C (black curve) and  $\delta^{18}$ O (grey curve) values (after Małkowski and Racki 2009: table 2). Note that the system boundary is placed 0.7 m higher than the tentative positioning given in Małkowski et al. (2009: fig. 6). Onset of the carbon isotope anomaly (SIDECIE), first appearances of the index graptolite and conodont species are shown, as well as the key brachiopod distribution (Baliński 2012; see also Figs. 8, 19, 20), rapid lateral changes of crinoid accumulations, and selected samples.

late Llandovery to earliest Wenlock Restiv Beds (rhythmically alternating grey platy limestone and clayey shale beds) are succeeded in the section by Wenlock (Homerian) nodular limestone, with subordinate platy calcareous and clayey sets (Demshyn Beds). These fossiliferous strata pass upward into the more monotonous calcareous-nodular Maryanivka Beds. This succession has been studied in terms of its graptolite (Tsegelnyuk 1976; Tsegelnyuk et al. 1983) and conodont (Drygant 1984) zonation. Koren et al. (1989) claimed that the Llandovery–Wenlock boundary is located 2.2 m above the base of Restiv Beds, and is based on the first and only occurrence of *Cyrtograptus murchisoni* within the *Pterospathodus amorphognathoides* Conodont Zone (see Fig. 3 and below; compare the updated biostratigraphy in Cramer et al. 2010, 2011a; Kleffner and Barrick 2010; and Melchin et al. 2012). The second succession studied is the partly trenched Dnistrove (former Volkovtsy, [64]; N48°38'19.1", E26°47' 18.6") section, which is a regional reference succession across the Silurian–Devonian (S–D) boundary, described in Małkowski et al. (2009; compare the Dnistrove-West locality in Kaljo et al. 2012). The Silurian sequence terminates with the Trubchyn (upper part of the former Rashkiv Beds) and Dzvenyhorod Beds of the upper part of the Skala Horizon (see Skompski et al. 2008: fig. 3). The strata are mostly calcareous, with dark, fossiliferous, nodular limestone predominating in the topmost interval. The S–D boundary is tentatively placed 3.2 m above this interval, in a succession of interbedded greenish-grey argillaceous shale and marl containing dark-grey limestone nodules. This unique, laterally differentiated and clay-enriched part of the lowermost Khudykivtsi Beds (= units 3 to 6 of Nikiforova and Predtechenski 1968: fig. 23) is informally distinguished herein as "Dnistrove Level" (2.2 to 3.1 m thick; Fig. 4).

The boundary is based on the first and only occurrence of the index graptolite species Monograptus uniformis angustidens (Nikiforova et al. 1972; Nikiforova 1977), but some of the conodont species are also important. Ozarkodina typica and Wurmiella excavata curvata do not occur higher than 1.4 m below the boundary while Parazieglerodina eosteinhornensis disappears just below the boundary (Drygant and Szaniawski 2009). Forty cm below the S-D boundary Zieglerodina remsheidensis appears and 60 cm above the boundary is the first representative of the icriodonts, Caudicriodus hesperius. Mashkova (1971) reported first occurence of the icriodonts 1.5 m below the boundary but we can not confirm that. In the recently revised conodont zonation scheme, the Devonian system base is defined by first occurrence of Caudicriodus hesperius (= the bottom of "hesperius Zone" of Corradini and Corriga 2012 and "hesperius-optima Zone" of Slavík et al. 2012; see below). This system boundary can also be fixed by the first appearance of the chitinozoan index species Eisenackitina bohemica, and the last appearance datum of Eisenackitina barrandei and Cingulochitina klonkensis. These chitinozoan faunas provide potential for accurate correlation with the international stratotype at Klonk, Barrandian area, Czech Republic (Chlupač and Hladil 2000; Fatka et al. 2003; see also Wrona 1980; Nestor 1994; Paris et al. 2000).

The lower part of the Lochkovian Stage, the Khudykivtsi Beds of the Borshchiv Horizon (57 m thick), is developed in the form of marly limestone interbedded with shale. The higher fossil-rich, open-marine Lochkovian limestone and shale suite represents an uninterrupted continuation of the Silurian–Devonian deposition (Małkowski et al. 2009).

#### Material and methods

As stressed above, the two intervals selected for quantitative study are characterized by profound carbon and oxygen isotope anomalies (Fig. 1), against well constrained biostratigraphical and facies background information (see also Nikiforova et al. 1972; Kaljo et al. 2007, 2012). The present study utilized a variety of geochemical and quantitative ecological approaches.

**Isotope geochemistry.**—We comprehensively sampled the Kytayhorod succession (38.7 m thick) in 2008 and 2009, whereas the Dnistrove section had already been studied for stable isotopes, based on 28 samples from a 22.35 m thick interval by Małkowski et al. (2009). The samples were collected from fresh surfaces of marly and micritic limestone layers (see microfacies types in Fig. 5, based on 15 thin sections; for generalized mineralogy and petrography see Nikiforova et al. 1972 and Kaljo et al. 2007), with a stratigraphical spacing of decimetres to metres.

A total of 44 bulk-rock samples from Kytayhorod was analysed for carbon and oxygen stable isotopes in the Light Stable Isotopes Laboratory of the Institute of Geological Sciences and Institute of Palaeobiology, Polish Academy of Sciences, Warsaw (see SOM 1 in Supplementary Online Material available at http://app.pan.pl/SOM/app57-Racki\_etal\_SOM.pdf; and Małkowski et al. 2009). The carbonate powder for analysis was extracted with a microdrill and decomposed under vacuum in 100% H<sub>3</sub>PO<sub>4</sub> (ortho-phosphoric acid) for 24 h at 25°C. Released CO<sub>2</sub>, after freezing it out from the separation line, was analyzed on a Finnigan Mat delta plus gas massspectrometer. Every gas sample was introduced into the spectrometer in dual inlet mode and the measurement was repeated eight times, with the mean values treated as results. The standard error of the spectrometer measurements was 0.02%. All carbon isotope values are reported in permil relative to V-PDB. The precision (reproducibility of replicate analyses) of both carbon and oxygen isotope analysis was usually better than  $\pm 0.1\%$ . Data are normalized to the V-PDB scale using National Bureau of Standards NBS-19 ( $\delta^{18}O = -2.20\%$  and  $\delta^{13}C = 1.95\%$ ).

In addition, 11 samples from Kytayhorod and 13 from Dnistrove (SOM 1 and 3) were studied for carbon isotopes in organic carbon in GeoZentrum Nordbayern, Erlangen, Germany. This analysis of organic matter was performed with an elemental analyser (Carlo-Erba 1110) connected online to a ThermoFinnigan Delta Plus mass-spectrometer. Accuracy and reproducibility of the analyses was checked by replicate analyses of international standards (USGS 40). Reproducibility was better than  $\pm 0.07\%$ o.

**Organic geochemistry.**—Twelve samples from Kytayhorod and 16 from Dnistrove were measured for total carbon (TC) and total inorganic carbon (TIC) abundances at the Faculty of Earth Sciences, University of Silesia (SOM 1 and 3). TC and TIC contents were determined using an Eltra CS-500 IR-analyser with a TIC module. Total organic carbon (TOC) was calculated as the difference between total carbon and total inorganic carbon. Analytical precision and accuracy were better than  $\pm 2\%$  for TC and  $\pm 3\%$  for TIC. Four (from six) pilot samples from both sites (SOM 5) were sufficiently rich in extracted organic matter to be analysed by gas chromatography coupled with mass spectrometry (GC–MS) for biomarker characteristics at the Faculty of Earth Sciences, University of Silesia (for method details see Racka et al. 2010).

**Inorganic bulk geochemistry.**—Sixteen rock pulp samples from the Kytayhorod section and 27 samples from the Dnistrove outcrop (SOM 2 and 4); supplemented by extra 7 samples from Zalishchyky (upper Lochkovian Ivanie Beds; Małkowski et al. 2009; SOM 4), were analysed at the Acme Analytical Laboratories (Vancouver) Ltd, Canada. Total abundances of the major oxides and several minor elements were reported from a 0.2 g sample analysed by ICP-emission spectrometry following lithium metaborate/tetraborate fusion and dilute nitric digestion. Rare earth and refractory elements (e.g., Ba, Co, Sr, Th, U, V, Zr) were determined by ICP mass 800

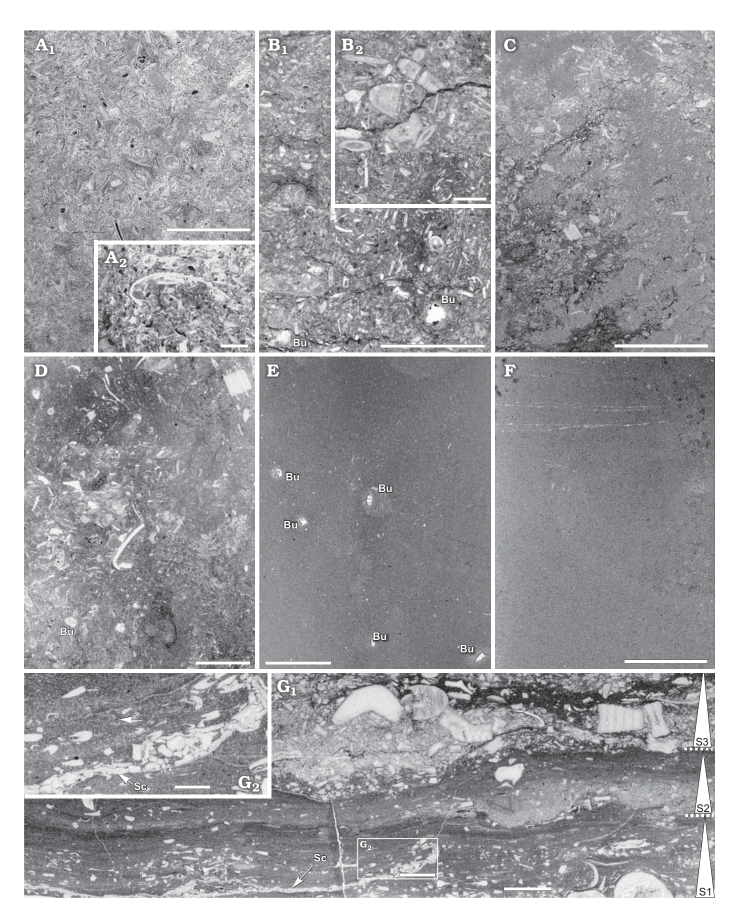


Table 1. Averaged, minimal and maximal Al/normalized concentrations of key elements in the Podolian sections under study against the average limestone and average shale Al/normalized abundance levels (based on Wedepohl 1971, 1991; Taylor and McLennan 1985), and averaged Al abundances (shaded).

Element	(Average limestone/Al) ×10 <sup>4</sup>	(Average shale/Al) ×10 <sup>4</sup>	(Average sample from Kytayhorod/Al) ×10 <sup>4</sup> [minimal value–maximal value]	(Average sample from Dnistrove/Al) ×10 <sup>4</sup> [minimal value–maximal value]
Ba	92.3	65	90 [45–303]	50 [33–165]
Р	230.8	70	310 [20–790]	160 [59–413]
Mn	538.5	90	610 [160–50]	198 [47–469]
Zn	17.7	8.5	25.9 [5.8–170]	7.5* [4.6–19.4]
Pb	6.9	2	5.1 [1.8–13.4]	3.3 [0.6–30.8]
V	15.4	15	12.1 [7.7–17]	15.4** [10.6–23.3]
Cu	3.1	5	2.9 [0.5–10.1]	6.3 [2.4–15.6]
Co	1.5	2.3	2 [1.4–2.7]	2 [1.1–2.9]
Ni	11.5	5.5	5.2 [4.1–9.1]	5.2 [2.7-6.9]
Мо	0.3	0.1	0.26** [≥0.03–1.79]	0.1**[≥0.02–0.6]
U	1.6	0.3	0.8 [0.3–2.1]	0.8 [0.4–3.3]
Zr	14.6	21	29 [23.6–36.9]	26 [16.4–82.2]
Al (%)	1.3	10	2.2 [1.0–6.8]	2.6 [0.7–6.6]

\* Excluding the abnormally enriched sample SD-27 (6543 ppm). \*\*Approximated values based on maximal concentrations in the samples below the detection level.

spectrometry following the same decomposition method. A separate 0.5 g split was digested with 3 ml 2:2:2 HCl-HNO<sub>3</sub>-H<sub>2</sub>O and also analysed by ICP mass spectrometry for the precious and base metals (e.g., Mo, Cu, Pb, Zn, Ni, As, Cd). In addition, total carbon and sulphur analyses by Leco were supplied. The reliability of the results was examined by analyses of some international standard reference materials. Precision and accuracy of the results were better than  $\pm 0.9\%$ (mostly  $\pm 0.4\%$ ) for the 11 major elements, and better than ±15% for the 45 trace elements, considered only in part in this study. When the concentrations of trace metals (especially Mo and Cr) were lower than the detection limits, the highest value below the suitable detection limit value was used in averaged abundance and proxy calculations (Table 1; SOM 2 and 4; see Racka et al. 2010 for further method details).

**Palaeontological data**.—Although various macrofossil groups are present, only the most abundant shelly component, brachiopods, has been quantitatively analyzed in frequency terms (ca. 1800 specimens). Nevertheless, occurrences of other groups were noted as well, and the distribution data have been compared with the faunal lists in Nikiforova and Predtechensky (1968), Nikiforova et al. (1972), and Koren et al. (1989).

Absolute and relative abundances have been estimated for two common acid-resistant microfossil groups in the both sections studied (SOM 6 and 7), derived from 33 samples for conodonts (17 from Kytayhorod and 16 from Dnistrove,) and 47 samples for chitinozoans (18 from Kytayhorod and 29 from Dnistrove; SOM 5). Conodonts were extracted by acetic acid digestion from mostly ca. 500–1000 grams of limestone, whilst about 100–150 grams of each sample were processed for chitinozoans. The samples for chitinozoans were processed using a standard palynological HCI-HF acids technique (see Miller 1996). The residues were sieved using 50 µm screens, but the relatively large amount of resistant residues obtained from the clay- and silt-rich marly samples was reduced about ten times for examination of chitinozoans.

Diversity indices were calculated utilizing a software package for paleontological data analysis PAST Version 1.94b (Hammer et al. 2001). The following statistics are presented (see also Hints et al. 2006):

- Frequency (absolute abundance): the number of specimens (n) per gram or kilogram of rock.
- Taxic diversity (S): the number of species or genera in a sample.
- Margalef's richness index:  $(S-1)/\ln(n)$ , where S is the total

<sup>←</sup> Fig. 5. Variety of microfacies from the Silurian to Lowermost Devonian of Podolia: bioturbated skeletal packstone to grainstone (**A**), bioturbated skeletal packstone (**B**, **C**), bioturbated skeletal wackestone to packstone (**D**), burrowed (**E**) to slightly bioturbated mudstone (**F**), and partly amalgamated crinoid packstone in three bioclastic, allegedly storm-generated intercalations (S1–S3), graded into non-fossiliferous shale (**G**; see also Małkowski et al. 2009: fig. 6). Demshyn (SD-36, D) and Maryanivka Beds at Kytayhorod (SD-50, A; SD-45, B); Dzvenyhorod Beds (SD-21, C), and lower (Vg-15, Dnistrove Level, G; see also Fig. 4) and upper (SD-20, E; SD-3, F) Khudykivtsi Beds at Dnistrove. Differently disarticulated and fragmented skeletal constituents dominated by shelly faunas (brachiopods; molluscs, see bivalve valve on A<sub>2</sub>) and echinoderms (crinoids; rare echinoids), associated with frequent ostracods (A<sub>1</sub>), dasycladacean algae (**B**<sub>1</sub>; close-up in **B**<sub>2</sub>) and trilobites. Note stylo-nodular fabric (C, lower left part), differently preserved burrows (Bu), partly with pellets (E), dispersed fine euhedral pyrite in micrite matrix (C–F), as well as *Scyphocrinites* lobolith arrowed in G<sub>1</sub> (close-up in G<sub>2</sub>). Scale bars 5 mm except A<sub>1</sub>, B<sub>1</sub> and G<sub>1</sub> 1 mm.

number of taxa, and ln(n) is the natural logarithm of the total number of the individuals n.

- Equitability:  $-\text{sum}((n_i/n)\ln(n_i/n))/\ln S$ , where  $n_i$  is number of individuals of taxon *i*, and *n* is the total number of the individuals.
- Dominance: sum  $((n_i/n)^2)$ , where  $n_i$  is number of individuals of taxon *i*, and *n* is the total number of the individuals.

## Geochemical evaluation of environmental factors

As reviewed by Cramer and Saltzman (2005, 2007b), Brand et al. (2006), Loydell (2007), Calner (2008), Vecoli et al. (2009), Munnecke et al. (2003, 2010), Kaljo et al. (2012) and Kozłowski and Sobień (2012), different scenarios have been postulated to explain the Silurian worldwide  $\delta^{13}$ C and  $\delta^{18}$ O positive anomalies. Saltzman (2005) indicated that so striking  $\delta^{13}$ C volatility was sustained by positive feedback between oxygen deficiency and bioproductivity in the usually P-limited and cooler-water Silurian oceans. Thus, one popular explanation invokes elevated primary productivity and/or the increased burial rate of organic matter, visible especially in black shale deposition. The Silurian excursions, however, are usually not synchronous with widespread black shale levels (with exception of the Middle Silurian Mulde Event, Calner 2008; in some regions, also of the ESCIE and SIDECIE, Lüning et al. 2000; Buggisch and Joachimski 2006; Loydell and Frýda 2007; Vecoli et al. 2009; Noble et al. 2012). Therefore, following "a weathering hypothesis" of Kump et al. (1999) for the Hirnantian glaciation as a trigger for the large-scale ecosystem catastrophe, enhanced weathering of exposed Silurian carbonate terrains in a regressive episode has been postulated as well (e.g., Loydell 2007; Noble et al. 2012). Eventually these led to a drop in atmospheric  $pCO_2$  and cooling (anti-greenhouse effect; e.g., Wenzel and Joachimski 1996; Kump and Arthur 1999; Małkowski and Racki 2009), as confirmed by reliable oxygen isotope paleothermometry based on conodont apatite data (Joachimski et al. 2009; Lehnert et al. 2010). In this study, because no data in Podolian successions for substantial changes in redox regimes, organic matter composition and thermal overprint (see discussion in Cramer and Saltzman 2007a), the paired inorganic and organic carbon isotope curves are used to estimate  $\Delta^{13}C (= \delta^{13}C_{carb} - \delta^{13}C_{org})$ , to roughly track changes in atmospheric  $pCO_2$  (Kump and Arthur 1999).

The severe drop in primary productivity and plankton starvation has been considered as a main cause of the trophic web instability and subsequent biotic crises (Jeppsson 1990, 1997, 2005; see also Porębska et al. 2004; Cramer and Saltzman 2007a). During the Ireviken Event, however, phytoplankton data reveal most of the acritarch losses at the final of the biotic overturn, without links to conodont demise steps (Gelsthorpe 2004). This timing of primary production collapse in the Gotland succession, however, is questionable according to Lehnert et al. (2010) because of major facies change. On the other hand, the phytoplankton abundances analysed by Stricanne et al. (2006) indicate, at least for the major Lau (= Kozlowskii) Event, decreased bio-productivities during the isotope excursion peak (see Noble et al. 2012) or perturbations in nutrient composition and availability (Kozłowski and Sobień 2012). If so, the large-scale <sup>12</sup>C sequestration may rather be attributed to black shale deposition in restricted oceanic and/or epeiric regions (e.g., Wenzel and Joachimski 1996; Cramer and Saltzman 2005, 2007a, b), as exemplified by northern Gondwanan hot shales since from Rhuddanian to early Wenlock times (Lüning et al. 2000, Vecoli et al. 2009). Advection of <sup>13</sup>C-enriched near-surface waters in stratified ocean, attributed to onset of an antiestuarine shelf circulation in more arid climate states, is an alternative hypothesis (e.g., Bickert et al. 1997; Munnecke et al. 2003; Jeppson 2005). In fact, the recurring graptolite mass mortalities are seen as triggered by regressive pulses and "oxic events" paired with vigorous thermohaline circulation and retreat of black shale organic-rich facies (e.g., Štorch 1995; Wenzel and Joachimski 1996; Porębska 2005; Johnson 2006). The most hospitable graptolite habitats may have been located near the boundary zone between oxic and denitrified, anoxic waters (= anoxitropic niche sensu Berry et al. 1989).

These conjectural oligotrophic vs. eutrophic photic zone conditions may potentially be tested by combined use of several assumed palaeoproductivity proxies: organic matter abundance (TOC) combined with inorganic tracers (P, Ba, and Si; e.g., Pujol et al. 2006; Tribovillard et al. 2006; Calvert and Pedersen 2007; Jenkyns 2010). As a micronutrient proxy the authigenic fraction of Ni, Zn and Cu is also considered here, as it is associated mostly with organic carbon (e.g., Brumsack 2006; Tribovillard et al. 2006; Śliwiński et al. 2012). Extreme enrichment in some elements (particularly Zn, Cu, Sr, and Ba) requires an additional external source, such as hydrothermal activity (see Brumsack 2006; Tribovillard et al. 2006; Pujol et al. 2006). Limited to a single sample (one-point) overabundances, exemplified by the distinctive sulphide-enriched sample SD-27 from Dnistrove (S-0.38%, Zn-6543 ppm, Cd-20 ppm; SOM 4), may suggest an episodic hydrothermal water influxes to the Podolian sedimentary basin, or rather indicate an imprint of post-depositional diagenetic and/or mineralizing magmatic-hydrothermal fluid signature (see Pujol et al. 2006; Tribovillard et al. 2006; Sindern et al. 2008). However, the studied samples display typically depleted Al-normalized abundances, and in infrequent cases only are distinctly higher relative to the average limestone (e.g., Zr and Cu at Dnestrove, see Table 1).

On the other hand, bottom-water oxygenation depletion is tracked by increased abundances of redox-sensitive (e.g., Mo, U, V, As, Cr,) and/or sulphide-forming trace metals (e.g., Co, Cu, Ni, Pb, Zn; see Brumsack 2006; Tribovillard et al. 2006; Calvert and Pedersen 2007; Schröder and Grotzinger 2007; Ver Straeten et al. 2011; Śliwiński et al. 2012). However, frequently used redox indices often suggest contradictory conclusions (see e.g., Racka et al. 2010). Even if Mo and U contents are mostly low (Table 1), the combined use of Mo/Al and U/Th ratios appears to be the most useful to suggest oxygen-deficient conditions in the Podolian successions. Considering other markers, the time series of V/V+Ni ratio consistently imply far more extensive hypoxia, whereas the Ni/Co ratio variations manifest only oxic regimes in the both sections.

Al is well known as the key lithogenous compositional (i.e., aluminosilicate) standard, and dilution effects by carbonate and quartz were reduced by use of the element/Al ratios as weight-ratios (e.g., Sageman et al. 2003; Pujol et al. 2006; Calvert and Pedersen 2007; Sindern et al. 2008; Ver Straeten et al. 2011). Furthermore, if (i) the non-normalized element concentrations are at the best close to the average limestone contents, and paired with (ii) a good correlation between the element vs Al, this abundance cannot be considered as an environmental proxy; these two characters indicate clearly its dominance by terrigenous provenance and scarcity of the authigenic fraction (Tribovillard et al. 2006; Śliwiński et al. 2012; see Table 1 and examples below). For example, only Al-normalized phosphorus concentrations are more commonly useful in the bioproductivity context in Podolian samples (see below).

The total lithogenous fraction content, as an approximation of the siliciclastic input, is mirrored by Al concentration, whilst aeolian delivery to the outer-shelf Podolian basin can be detected using elemental indicators for sediment grainsize, such as Si/Al, Ti/Al, and Zr/Al ratios (Calvert and Pedersen 2007; Schröder and Grotzinger 2007; Ver Straeten et al. 2011). The applicability of quartz to aluminosilicate (clay) phase ratio is justified by the absence of biogenous silica in Podolian sites. For example, the larger average grainsize is tracked by elevated Zr/Al and Si/Al ratios, recording a proportionally increased content of zircon and quartz grains, respectively. Ver Straeten et al. (2011) designated a Si/Al ratio above 5 as a tracer for sand-dominated clastic supply. Wind-blown dust may also be compositionally influenced by volcanic input, reflected by high Zr abundance levels in fine-grained volcaniclastic admixture (e.g., Pujol et al. 2006), up to 260 ppm in Silurian ash levels of Podolia (Huff et al. 2000: table 1). Elsewhere, Kiipli et al. (2010) have interpreted Silurian sea level variations in a deep-shelf Latvian section by combined SiO<sub>2</sub>/Al<sub>2</sub>O<sub>3</sub> and K<sub>2</sub>O/Al<sub>2</sub>O<sub>3</sub> ratios, reflecting the contribution of quartz, and muscovite and K-feldspar, respectively, in siliciclastic material (see also Sageman et al. 2003; Sindern et al. 2008).

## Geochemical trends across the Ireviken Event

The Kytayhorod succession is characterized by rather monotonous grey carbonate-marly lithologies evolving from planar bedded, rhythmic to unbedded, nodular varieties (Fig. 6A; for microfacies see Figs. 5A, B, D and 11; see also Nikiforova et al. 1972: figs. 31, 32, and 126). Calcium carbonate content (calculated from CaO) is mostly above 70%, with notable exceptions in the lower part, where this abundance drops to 20.8%. The Restiv Beds are distinguished by very high SiO<sub>2</sub> abundance in several layers, above 47% in the basal SD-42 sample, and Al<sub>2</sub>O<sub>3</sub> content above 12% (Fig. 7; compare the basal transgressive sandy set of Nikiforova et al. 1972: 36, 132). In most samples, however, the quartz contribution is negligible (SiO<sub>2</sub> ranges between 9 and 17%), as also is that of the clay minerals (Al<sub>2</sub>O<sub>3</sub> between 2 and 4%). Dolomitisation remains minor in the succession as MgO content is largely below 1.7%.

**Carbon and oxygen isotopes.**—As noted by Kaljo et al. (2007), the distinctive  $\delta^{13}C_{carb}$  increase of 2.6% in the Kytayhorod section begins in the Restiv Beds, and reaches its peak value of 4.0% at the very base of the early Sheinwoodian Demshyn Beds. The isotopic highstand (slice E-2; Fig. 6A) is followed by stable but negative  $\delta^{13}C$  values up to -0.9% in the upper parts (see coeval brachiopod calcite results in Azmy et al. 1998).

Our and Kaljo et al. (2007: fig. 4)  $\delta^{13}$ C time series agree with the worldwide chemostratigraphical pattern in the Llandovery-Wenlock transition (e.g., Brand et al. 2006; Cramer et al. 2010; Noble et al. 2012), especially with Estonian  $\delta^{13}$ C profiles, marked by an obscured slice of transient value decline (isotopic feature 2 of Cramer et al. 2010). The early Sheinwoodian excursion (ESCIE) amplitude approaches 4.5% if a -0.5% regional  $\delta^{13}C_{carb}$  baseline is used (guided by E-4 interval; Fig. 6A). In addition, the C-isotopic event is clearly coupled with a positive  $\delta^{18}O_{carb}$  excursion, evident also in the values in table 3 of Kaljo et al. (2007), but neglected by these authors as a probable diagenetic artifact in whole-rock carbonate samples (see also e.g., Martma et al. 2005). Their overall primary character is demonstrated, however, by comparable brachiopod calcite values from this site in Azmy et al. (1998; see also Wenzel and Joachimski 1996, Bickert et al. 1997, Heath et al. 1998; Munnecke et al. 2003; Cramer and Saltzman 2005; Brand et al. 2006) and, especially by a similar reliable chemostratigraphical  $\delta^{18}$ O pattern (i.e., the SOIE excursion; see below) recognized in conodont apatites by Lehnert et al. (2010, see also discussion on the S-D transition signature in Małkowski et al. 2009). The  $\delta^{18}O_{carb}$  shift begins with values between -5.2 to -6.2%, increases up to -3.9% in the plateau interval of high  $\delta^{13}$ C values, and very gradually diminishes by ca. 2% in the Maryanivka Beds (background E-4 interval; Fig. 5A).

The  $\delta^{13}C_{carb}$  excursion is partly reflected also in the organic carbon data (based on few samples), but marked probably by a distinctly lower amplitude. The Podolian record of the ESCIE is associated in addition with an irregular  $\Delta^{13}C$ rise by ca. 2‰; this highest value corresponds with the uppermost sample from the crucial E-2 slice (Fig. 7A).

**Inorganic bulk geochemistry**.—The detrital admixture exhibits two slightly fluctuating (particularly in the lowest interval;), but contrasting secular tendencies: (i) decreasing clay abundance tied with (ii) increased coarser clastic input indi-

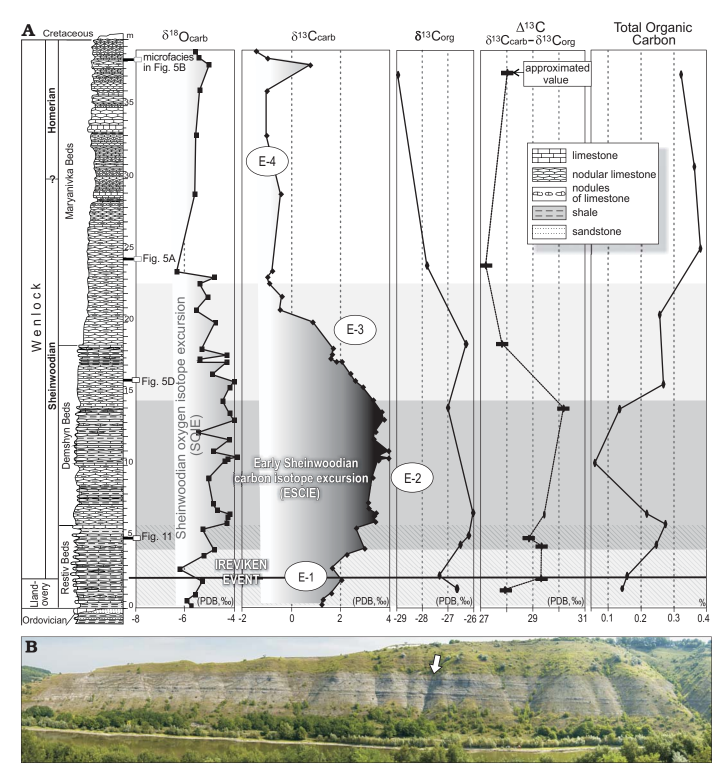


Fig. 6. A. Secular trends of  $\delta^{13}$ C and  $\delta^{18}$ O values in carbonate and organic matter,  $\Delta^{13}$ C and total organic carbon from section 96 near Kytayhorod village (see SOM 1). Lithology after Nikiforova et al. (1972: fig. 82), modified. Three slices E-1 to E-3 (marked by shades of grey) denote of the early Shein-woodian (ESCIE) carbon isotope excursion: the interval E-2 corresponds to the isotopic highstand (see a widened excursion definition in Cramer et al. 2010, also Sadler 2012); E-1 and E-3 to the increasing and decreasing curve segments, respectively, and E-4 to the regional background. Note that the Ireviken biotic event (crossed band) correlates with the initial positive  $\delta^{13}$ C shift slice (E-1); the lower event boundary corresponds the last occurrence of conodont *Pseudooneotodus tricornis*, and the upper boundary is questionably placed above the disappearance of chitinozoan *Eisenackitina* cf. *dolioliformis* (Figs. 15, 16; see Jeppsson 1997; Nestor et al. 2002;). Approximated  $\Delta^{13}$ C levels are calculated for  $\delta^{13}$ C carb values averaged for two stratigraphically adjacent samples, as shown also by a dotted line of the curve. The microfacies illustrated are marked as rectangles. **B**. General view of the natural outcrop in the Dniester River valley (Tarnawa tributary) near Kytayhorod village; arrow indicates the main sampled site.

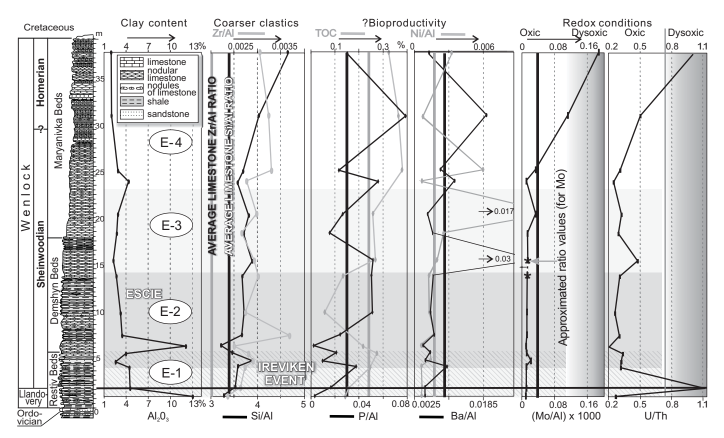


Fig. 7. Stratigraphical distribution of the Al-normalized abundance for selected major and trace elements as well as  $Al_2O_3$  and TOC contents (wt. %) and the U/Th ratio (for redox thresholds see Racka et al. 2010) from the Kytayhorod section (see SOM 2). Thicker vertical lines indicate level of the ratios for average limestone after Wedepohl (1971, 1991). Approximated proxy values, resulted from the content of Mo below the detection levels, are asterisked. Shades of gray denote three slices of the ESCIE.

cated by abruptly increasing Zr/Al and Si/Al ratios, correlated also with the K/Al proxy trend (Fig. 7). Heightened Zr/Al levels are suggestive of some volcaniclastic material in the peak values, even though definite bentonites occur only higher in the Silurian succession of Podolia (Huff et al. 2000).

All presumed bioproductivity tracers show in detail largely dissimilar temporal changes, and are also somewhat questionable because the crucial elements are usually very weakly correlated both with Al and TOC: r = -0.33 and -0.12 for P, respectively, as well as r = 0.12 and -0.52 for Ba, r = 0.99 and -0.24 for Ni, and r = -0.06 and 0.36 for Zn. Thus, their authigenic character is suspected (or organic matter has been differently degraded in the sediments). However, the proxy data analysed collectively exhibit the congruent major increasing-upward trend.

Declining bottom water oxygenation towards at least dysoxic conditions, after the ESCIE (see below), is clearly evidenced by several indices (U/Th, Ni/Co). Normalized Mo contents reveal the most evident chemostratigraphical trend: the long lasting stable low levels are followed by a rapid forty-fold increase during the E-4 timespan (Fig. 7). This tracer is the most reliable as it is unrelated to aluminosilicate phases (r= -0.24 with Al; but r = 0.56 with TOC), by contrast with indices based on V contents (r = 0.97 with Al; but r = -0.41 with TOC). The very high U/Th ratio in transgressive graptolite-bearing horizon of the Restiv Beds (see below) is known from the one sample only, and contrasts with low Mo/Al and most other redox proxy values. This suggests its link rather to a post-depositional enrichment, even if a higherresolution sampling is requested.

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### Geochemical trends across the Klonk Event

Lithofacies variation across the Klonk Event is significant in the Dnistrove section because dark shale set with bioclastic limestone partings (Dnistrove Level; Fig. 4) occurs in the lowermost part of the Khudykivtsi Member, spanning the S–D boundary (Fig. 8). Carbonate content decreased to 22% in the Dnistrove Level (it is largely between 60% and 85%), and this is coupled with SiO<sub>2</sub> values above 45%, and Al<sub>2</sub>O<sub>3</sub> concentrations above 12% (Fig. 9). Otherwise, the marlymicritic lithologies (see Fig. 5C, E, F; Nikiforova et al. 1972: figs. 83, 84, and 141) are enriched overall in terrigenous material, as evidenced by irregularly fluctuating Al and Si contents, varying from 1.3% to 5.4% and 6.2% to 20.7%, respectively. Dolomitization is relatively unimportant at this site even where MgO content is as high as 5.3% as this is notably linked with aluminosilicates (r = 0.82 with Al).

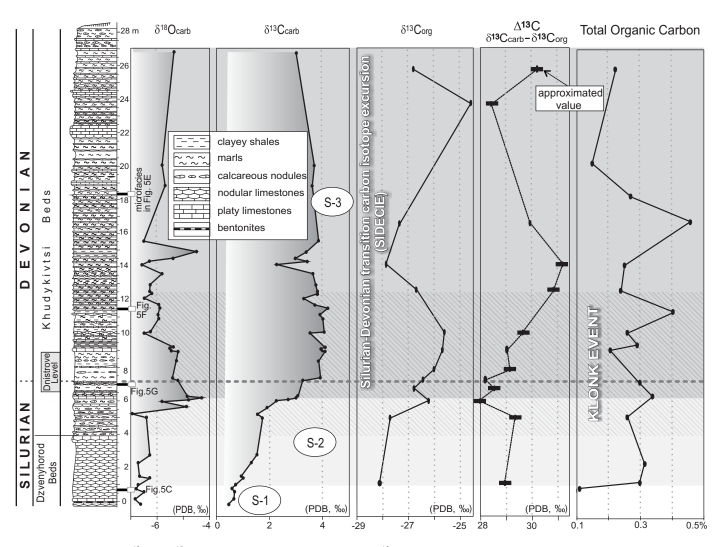


Fig. 8. Secular trends of  $\delta^{13}$ C and  $\delta^{18}$ O values in carbonate and organic matter,  $\Delta^{13}$ C and total organic carbon from section 64 near Dnistrove village (see SOM 3). Two slices S-1 to S-3 (marked by shades of grey) denote of the Silurian–Devonian transition carbon isotope excursion (SIDECIE): the interval S-3 corresponds to the isotopic highstand; S-2 to the increasing curve segment, and S-1 to the regional background. Lithology after Nikiforova et al. (1972: fig. 82), modified. Note that the Klonk biotic event (crossed band) is associated roughly with the final positive shift (S-2) and initial highstand (S-3)  $\delta^{13}$ C curve slices; its lower event boundary corresponds to a collapse of carbonate production and chitinozoan crisis near the base of Dnistrove Level, and the upper boundary is questionably placed below an acme level of brachiopod and conodont faunas (see Figs. 4 and 23).

**Carbon and oxygen isotopes.**—The abrupt and coeval increase of  $\delta^{13}C_{carb}$  and  $\delta^{18}O_{carb}$  values is thoroughly described by Małkowski et al. (2009; see also Kaljo et al. 2012). The  $\delta^{13}C_{carb}$  excursion, with the highest value of 3.8% and 4.5%, is confirmed also in nearby outcrop at Dzvenyhorod and drill core section in the Ternopil area, respectively (Kaljo et al. 2012; see Fig. 2B). The lower part of the S–D excursion (SIDECIE) is recorded in the  $\delta^{13}C_{carb}$  data with an amplitude of ca. 4‰ in the Dnistrove succession (and above 5‰ when considering all the Podolian data).

This trend may be now roughly correlated with a parallel shift in the  $\delta^{13}C_{org}$  levels by 3.5% (Fig. 8), but this is notably reversed higher in the succession. Thus, more high-resolution data are necessary to constrain this co-variation discrepancy, as well as more irregular  $\Delta^{13}C$  secular changes, marked by a distinct increase coeval with the three-point  $\delta^{13}C_{org}$ 

lowstand. The  $\Delta^{13}C$  curve instability notably contrasts with a long-lasting isotopic plateau seen in the  $\delta^{13}C_{carb}$  values.

**Inorganic bulk geochemistry.**—As at Kytayhorod, the terrigenous component reveals two somewhat opposing first-order trends. Clay abundance is distinctly lower in the upper half of the section, whilst the grain-size and possible volcaniclastic admixture abruptly increases up to sand fraction in the uppermost part, as recorded in the very high Zr/Al and Si/Al ratios (Fig. 9).

Considering proxies of bioproductivity, similar serious reservations can be expressed as for the Kytayhorod samples. Irregular secular variations suggest pulses of enhanced fertilization/primary production, but recorded largely in the lower part of the succession, including the Dnistrove Level, and conspicuously absent in its middle part. The pattern is certainly partly biased by the dominantly detrital provenance

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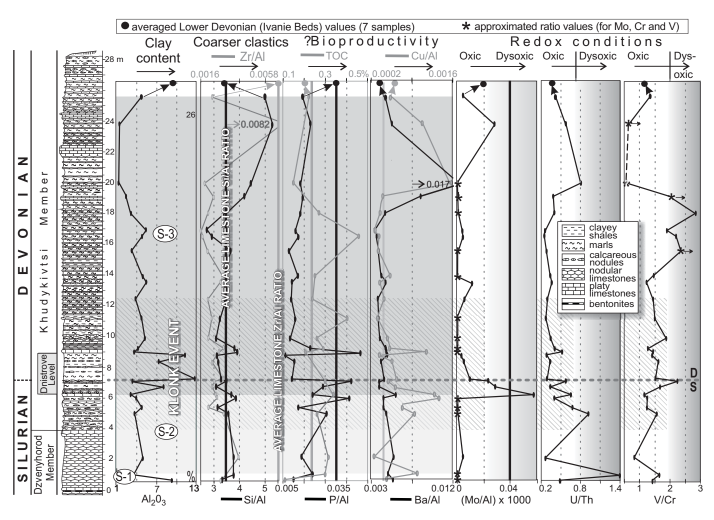


Fig. 9. Stratigraphic distribution of the Al-normalized abundance for selected major and trace elements as well as  $Al_2O_3$  and TOC contents (wt.%) and the U/Th and V/Cr ratios from the Dnistrove section (see SOM 4). Approximated proxy values, resulted from the Mo, V, and Cr content below the detection levels, are asterisked. Averaged data for the higher Lower Devonian strata from Zalishchyky (Ivanie Beds; Małkowski et al. 2009; Fig. 2; see SOM 4) are also shown. Shades of gray denote two slices of the SIDECIE.

of the elements analysed, visible in the adverse relationships between aluminosilicates and organic carbon even for the most trustworthy tracers P (r = 0.39 and -0.10, respectively) and Cu (r = 0.65 and -0.42).

Clearest data are again provided by collective use of redox indicators, because the steady oxic regimes are evidenced exclusively in the middle part of section, just above the S–D boundary, and, at most, dysoxic regimes may be suggested for the remaining SIDECIE portions, marked by fossil-poor, variably bioturbated muddy lithofacies (Fig. 5E, F). Mo is once more predominantly present as an authigenic phase, and therefore the most certain proxy in light of the insignificant correlation with Al, and also with TOC (r = 0.31and 0.25, respectively).

## Organic geochemistry

Organic carbon abundances are invariably low in both sections, as shown by TOC values ranging from 0.06 to 0.38% at

Kytayhorod and from 0.11 to 0.46% at Dnistrove (SOM 1 and 3). The organic matter is rather erratically distributed in the latter succession, and an exact association occurs neither with the major  $\delta^{13}C_{carb}$  shift nor with black shale lithologies around the S–D boundary (Fig. 8). However, the TOC maxima are notably found in the limestone set related to the SIDECIE plateau. By contrast, organic carbon distribution does not show any link with the ESCIE because this is characterized by an upward increasing trend. What is more, the minimal concentration (0.06%) is found in the  $\delta^{13}C_{carb}$  excursion peak (Fig. 6A). Thus, the major secular trends of TOC and inorganic proxies are connected each other at Kytayhorod (Fig. 7), but this correlative link is obscured at Dnistrove, especially in the middle SIDECIE slice (Fig. 9).

Results of the pilot GC-MS study on samples from the Podolian localities (SOM 5) are influenced by thermal maturity: biomarker data indicate at least ca 80–100°C, and of the order of 150°C is indicated by our new observations of the conodont colour alteration (cf. also Drygant 1993 and Huff et al. 2000: 498–499).

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O r d e r STROPHOMENIDA	ORTHOTETIDA PROTORTHIDA	ORTHIDA PENTAMERIDA	RHYNCHONELLIDA ATRYPIDA	ATHYRIDIDA	SPIRIFERIDA	[				
Leptaceoura douvalii	Streptis grayii Skenidioides lewisii	Dolerorthis rustica Dalejina hybrida Resserella canalis Dicoelosia biloba Dicoelosia paralata Anastrophia deflexa Antirhynchonella linguifera	Plagiorhycha depressa Sphaerirhynchia aff. wilsoni Atrypa sp. Oglupes sp. Atrypina barrandii Plectatrypa imbricata Glassia elongata	Meristina bilobata Meristina incrassata Meristina podolica Nucleospira pisum	Cyrtia exporrecta Eospirifer radiatus Howellella globosa	taxa	number of specimens	Margalef's richness	dominance	equitability
						5	29	1.2	0.3	0.9
We n l o c k Maryanivka Beds Maryanivka Beds With the nerian With the nerian W						9		2.5		
Uppermost						14	137	2.6	0.4	0.6
<b>Ε</b> <u>σ</u> 15 D3						4	17	1.1	0.3	0.9
D2						14	54	33	1.2	0.9
1 m above D1						4	8			
								2.5		
A provide the second se				IREVIKEN		1	7	1.8		
				EVENT		13 8	246 17	2.2 2.5	0.2 0.2	0.7 0.9

Fig. 10. Distribution of brachiopod species, diversity, and ecological indices through the Kytayhorod Horizon in Kytayhorod section. Dominating species are marked with thicker lines. Shades of gray denote three slices of the ESCIE.

Cyclic biomarkers like steranes or hopanes have not been found or are present as traces. The most frequently encountered organic compounds are relatively stable n-alkanes and polycyclic aromatic hydrocarbons and their alkyl derivatives. Among n-alkanes, the most abundant are short-chain varieties ([nC17 + nC18 + nC19]/[nC27 + nC28 + nC29]) suggestive of prevalent marine organic matter. However, carbon preference indices (CPI and CPI[25–31]) are above 1 (as high as 2.01 for sample LM3 from Dnistrove) in the long-chain n-alkane sets, characteristic for land-derived organic material due to dominating odd-numbered (over even-numbered) carbon chains (e.g., Peters et al. 2005). Thus, a

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mixed marine-continental source of organic carbon is supposed for the studied shelf successions, and is confirmed by palynofacies at Dnistrove, also grossly obscured by thermal overprint (see below). Furthermore, predominantly oxic sedimentary regimes are suggested by high pristane to phytane ratios (Didyk et al. 1978).

# Faunal dynamics across the Ireviken Event

Quantitative distributional and diversity data on three fossil groups, representing both benthic and most likely largely pelagic biotas, are explored below. The faunal dynamics is considered in the context of environmental variables throughout the two global isotopic events, interpreted from the above geochemical patterns.

**Brachiopods**.—The stratigraphically oldest brachiopod fauna of the Restiv Beds (samples R1 to upper part) is quite diverse taxonomically being represented by up to 14 species (Figs. 10, 11). The most characteristic members of the fauna are species of *Skenidioides* and *Dicoelosia* (see also Modzalevskaya 1968). Other genera represented are *Scamnomena*, *Eoplecto-donta*, *Dalejina*, *Resserella*, *Plagiorhyncha*, *Oglupes*, *Atrypina*, *Glassia*, *Meristina*, and *Cyrtia* (see Figs. 12–14). These generally well preserved brachiopods are mostly characterized by small shell dimensions and thin-ribbed surface sculpture (see also Modzalevskaya and Nikiforova 1980). They were either (i) fixosessile (see Bassett 1984), staying attached to the substrate with their pedicle during all stages of growth

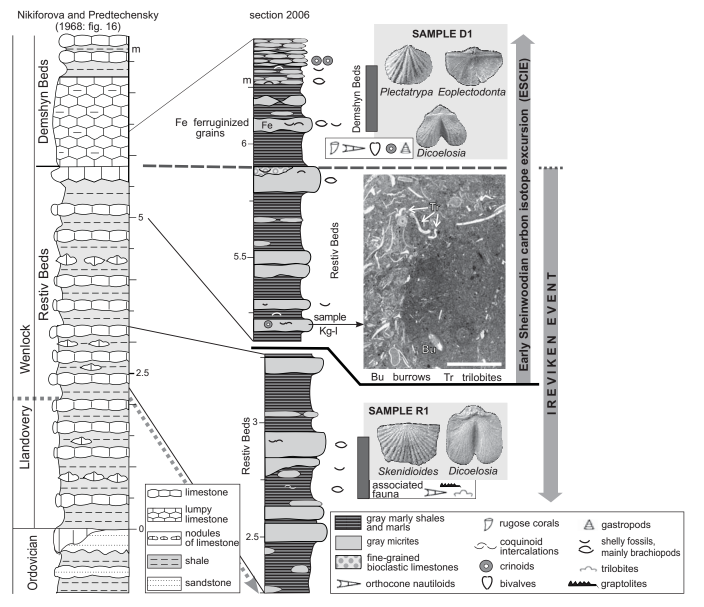


Fig. 11. Lowermost Wenlock lithologic and faunal succession at Kytayhorod, against the key brachiopod assemblages (see Fig. 10) and a representative micritic microfacies of the Restiv Beds, i.e., bioturbated skeletal wackestone to packstone with shelly bioclasts, crinoids, and trilobites (compare Fig. 5). Note onset of the carbon isotope excursion (ESCIE) before the ending of Ireviken Event.

(e.g., Scamnomena, Skenidioides, Dicoelosia, Dolerorthis, Atrypina, Glassia, Cyrtia), or (ii) liberosessile, becoming secondarily free-lying after whole or partial atrophy of the pedicle (e.g., Eoplectodonta, Oglupes). Ambitopic Eoplectodonta and Oglupes developed different adaptations for stabilisation of the shell on soft muddy bottoms. The adult strophomenide Eoplectodonta remained on the surface with the hinge line down partially sunk into the sediment keeping the anterior commissure above a muddy bottom. Dattilo (2004) showed that strophomenid brachiopods were quite mobile and responsive to environmental conditions and that they could live in a convex valve-up position (Lescinsky 1995; Baliński 2010) or assume a near-vertical, semi-infaunal position to escape burial. The adult atrypide Oglupes, after atrophy of the pedicle, became recumbent on the soft substrate and was stabilized by flat ventral valve and widely extended frills (see Fig. 13E). This atrypide, which is the largest species in the association, possessed a highly efficient spirolophe lophophore, which enabled it to flourish in deeper-water settings with an impoverished food supply (see Fürsich and Hurst 1974). The overall taxonomic inventory of brachiopods from the Restiv Beds strongly implies that the fauna represents the globally distributed high diversity Dicoelosia or Dicoelosia-Skenidioides community (see e.g., Boucot 1975, 2005; Gritsenko et al. 1999; Jin and Copper 1999; Watkins et al. 2000; Li and Allen 2008). Both communities have been interpreted as characteristic of outer shelf to slope, deep-water environments indicating a Benthic Assemblage (BA) 4-5 of Boucot (1975). According to Copper (2004), the occurrences of Glassia in Britain and Gotland indicate deeper-water settings, probably equivalent of BA 4 or BA 5. Eoplectodonta, which is a common brachiopod not only in the Restiv Beds but ranges to the top of the Kytayhorod section, is one of the most characteristic members of the deeper-water Clorinda Community (BA 5) during most of the Britain Silurian (Cocks 1970). Johnson (1987; see also Brett et al. 1993; Watkins et al. 2000) estimated the bathymetry of BA 4-5 in the range of 60-120 m. Watkins et al. (2000) demonstrated that brachiopod species diversity of Dicoelosia communities remained unchanged from the early Silurian to the early Devonian on a global basis.

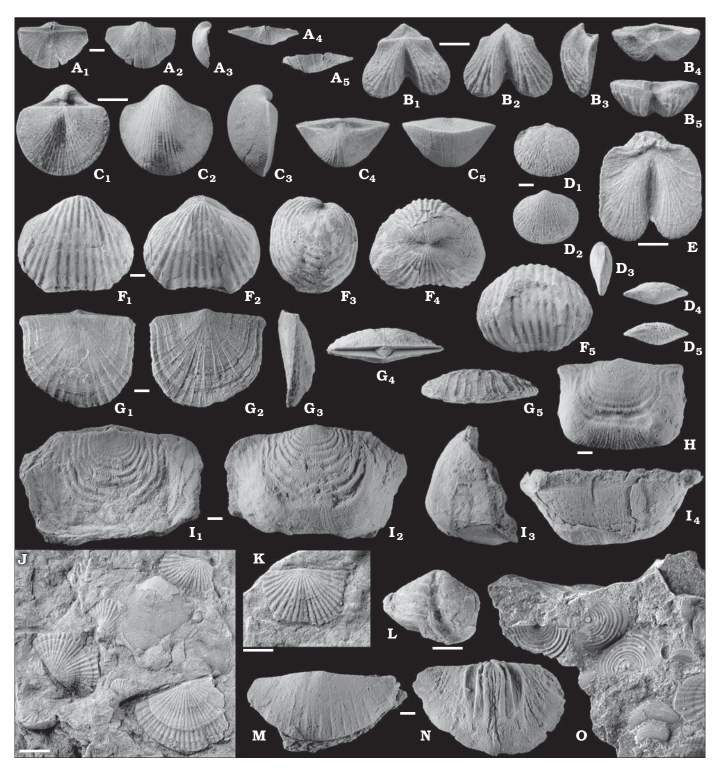
The brachiopods of the overlying Demshyn Beds reveal slightly greater taxonomic diversity than the underlying beds. They are represented by 18 species in total of which 8 continue their ranges from the Restiv beds (Fig. 10). Among them *Dicoelosia* occurs to the top of the interval whereas *Skeni-dioides* was found only in its lowermost part. The brachiopod fauna is dominated by the strophomenide *Eoplectodonta*, atrypides *Atrypa* and *Plectatrypa*, and in the uppermost part of the interval by the mass-occurrence of the rhynchonellide *Sphaerirhynchia*. This genus forms clusters similar to those described by Fürsich and Hurst (1981) for *Sphaerirhynchia* 

wilsoni from the Silurian of England and Gotland. *Kata-strophomena, Leptaena, Streptis, Antirhynchonella, Nucleo-spira, Eospirifer*, and the above-mentioned atrypides appeared in the Demshyn Beds for the first time. Medium- and large-sized forms dominate in this interval. Ambitopic forms with whole or partial atrophy of the pedicle are represented by *Katastrophomena, Leptaena, Eoplectodonta, Antirhynchonella, Sphaerirhynchia, Atrypa*, and *Meristina*.

In general, the brachiopod fauna from the Demshyn Beds suggests continuity of Dicoelosia communities (BA 4-5). This is also supported by the numerous occurrences of Plectatrypa which occurs mostly in deeper, shaly, distal shelf facies on Gotland (Copper 2004). Rare findings of Antirhynchonella, which represents the same morphological type as Clorinda, suggest also some relation to the Clorinda Community of Ziegler et al. (1968) correlated with BA 5 (Boucot 1975). Fürsich and Hurst (1981) concluded that Sphaerirhynchia wilsoni lived typically semi-infaunally on soft muddy substrates in very calm, low energy settings. Some morphological adaptations of brachiopods from the interval, e.g., strongly sulcate anterior commissure, alate hinge margin, and thin shell, indicate a quiet-water environment (Fürsich and Hurst 1974: fig. 6). The brachiopod fauna from the Demshyn Beds reveals also great similarity to the Eoplectdonta douvalii Community described by Hurst (1975) from the Wenlock of Wales and the Welsh Borderland. As many as 78% of the brachiopod species from the Demshyn Beds are also listed by Hurst (1975) in the inventory of the Eoplectdonta douvalii Community. Although the brachiopod communities of the Restiv and Demshyn beds indicate similar settings within BA 4–5, the appearance of generally medium- and large-sized brachiopods in latter and dominance of smaller-sized forms in the Restiv Beds suggest some shallowing efect in the section. However, the general faunal changes at that time indicate that sea-level change appears not to have been substantial.

In the Maryanivka Beds 13 brachiopod species in total have been identified during the present investigations. Medium- and large-sized forms predominate while small species have been not found. Several forms do not continue their range from the preceding Demshyn Beds, with the most important loss being of Dicoelosia. Anastrophia, and Howellella are the only genera which appeared new for this interval (Fig. 10). Besides the absence of Dicoelosia, the total inventory of brachiopod taxa from the Maryanivka Beds did not changed much in comparison with the preceding Demshyn Beds fauna: 10 genera range through both intervals. Two species from the interval are reported at higher levels in other sections. These are Leptaena depressa and Eospirifer radiatus which were revealed in the Muksha and Cherche beds, respectively (Nikiforova et al. 1972). Although it seems probable that the brachiopod fauna of the Maryanivka Beds represents more or less

Fig. 12. Silurian brachiopods from the Kytayhorod section. **A**, **M**, **N**. *Eoplectodonta douvalii* (Davidson, 1847). **A**. ZPAL Bp 71/1, almost complete shell in dorsal (A<sub>1</sub>), ventral (A<sub>2</sub>), lateral (A<sub>3</sub>), posterior (A<sub>4</sub>), and anterior (A<sub>5</sub>) views; Maryanivka beds, sample M1. **M**. ZPAL Bp 71/2, exterior of ventral valve; Demshyn beds. **N**. ZPAL Bp 71/3, interior of dorsal valve; Demshyn beds, sample D2. **B**. *Dicoelosia biloba* (Linnaeus, 1758), ZPAL Bp 71/4, shell in dorsal (B<sub>1</sub>), ventral (B<sub>2</sub>), lateral (B<sub>3</sub>), posterior (B<sub>4</sub>), and anterior (B<sub>5</sub>) views; Demshyn beds, sample D2. **C**. *Resserella canalis* (Sowerby, 1839), ZPAL Bp  $\rightarrow$ 



71/5, shell in dorsal ( $C_1$ ), ventral ( $C_2$ ), lateral ( $C_3$ ), posterior ( $C_4$ ), and anterior ( $C_5$ ) views; upper part of the Restiv beds. **D**. *Dalejina hybrida* (Sowerby, 1839), ZPAL Bp 71/6, shell in dorsal ( $D_1$ ), ventral ( $D_2$ ), lateral ( $D_3$ ), posterior ( $D_4$ ), and anterior ( $D_5$ ) views; Demshyn beds, sample D1. **E**. *Dicoelosia paralata* Bassett, 1972, ZPAL Bp 71/7, slightly damaged shell in dorsal view; Restiv beds, 0.8 m above sample R1. **F**. *Anastrophia deflexa* (Sowerby, 1839), ZPAL Bp 71/8, complete shell in dorsal ( $F_1$ ), ventral ( $F_2$ ), lateral ( $F_3$ ), posterior ( $F_4$ ), and anterior ( $F_5$ ) views; lower part of the Maryanivka beds. **G**. *Katastrophomena antiquata* (Sowerby, 1839), ZPAL Bp 71/9, shell in dorsal ( $G_1$ ), ventral ( $G_2$ ), lateral ( $G_3$ ), posterior ( $G_4$ ), and anterior ( $G_5$ ) views; Demshyn beds. **H**. *I*. *Leptaena depressa* (Sowerby, 1824). **H**. ZPAL Bp 71/10, complete shell in ventral view; Demshyn beds, sample D3. **I**. ZPAL Bp 71/11, slightly damaged shell in dorsal ( $I_1$ ), ventral ( $I_2$ ), lateral ( $I_3$ ), posterior ( $I_4$ ), and anterior ( $I_5$ ) views; lower part of the Maryanivka beds. **J**. K. *Skenidioides lewisii* (Davidson, 1848). **J**. ZPAL Bp 71/12, slab with three well visible dorsal valve; Restiv beds, 0.8 m above sample R1. **K**. ZPAL Bp 71/13, exterior of dorsal valve; Restiv beds, sample R1. **L**. *Streptis grayii* (Davidson, 1848), ZPAL Bp 71/14, slightly incomplete shell in ventral view; lower part of the Demshyn beds. **O**. *Scannomena* sp., ZPAL Bp 71/15, slab with three visible specimens; Restiv beds, slightly above sample R1. Scale bars 2 mm.

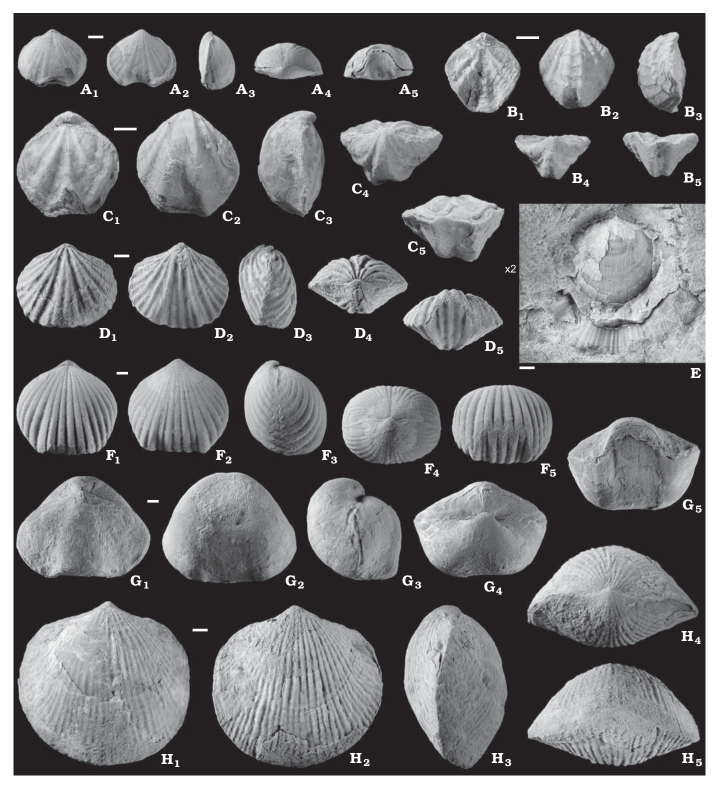


Fig. 13. Silurian brachiopods from the Kytayhorod section. **A**. *Plagiorhycha depressa* (Sowerby, 1839), ZPAL Bp 71/16, shell in dorsal (A<sub>1</sub>), ventral (A<sub>2</sub>), lateral (A<sub>3</sub>), posterior (A<sub>4</sub>), and anterior (A<sub>5</sub>) views; Restiv beds, sample R1. **B**, **C**. *Atrypina barrandii* (Davidson, 1848), ZPAL Bp 71/17 (**B**), ZPAL Bp 71/18 (**C**), two shells in dorsal (B<sub>1</sub>, C<sub>1</sub>), ventral (B<sub>2</sub>, C<sub>2</sub>), lateral (B<sub>3</sub>, C<sub>3</sub>), posterior (B<sub>4</sub>, C<sub>4</sub>), and anterior (B<sub>5</sub>, C<sub>5</sub>) views; upper part of the Restiv beds (B) and upper part of the Demshyn beds (C). **D**. *Plectatrypa imbricata* (Sowerby, 1839), ZPAL Bp 71/19, shell in dorsal (D<sub>1</sub>), ventral (D<sub>2</sub>), lateral (D<sub>3</sub>), posterior (D<sub>4</sub>), and anterior (D<sub>5</sub>) views; Demshyn beds, sample D1. **E**. *Oglupes* sp., ZPAL Bp 71/20, shell with partially preserved frills in ventral view; Restiv beds, sample R2. **F**. *Sphaerirhynchia* aff. *wilsoni* (Sowerby, 1816), ZPAL Bp 71/21, complete shell in dorsal (F<sub>1</sub>), ventral (F<sub>2</sub>), lateral (F<sub>3</sub>), posterior (F<sub>4</sub>), and anterior (F<sub>5</sub>) views; Demshyn beds. **G**. *Antirhynchonella linguifera* (Sowerby, 1839), ZPAL Bp 71/22, large shell in dorsal (G<sub>1</sub>), ventral (G<sub>2</sub>), lateral (G<sub>3</sub>), posterior (G<sub>4</sub>), and anterior (H<sub>5</sub>) views; lower part of the Maryanivka beds. **H**. *Atrypa* sp., ZPAL Bp 71/23, shell in dorsal (H<sub>1</sub>), ventral (H<sub>2</sub>), lateral (H<sub>3</sub>), posterior (H<sub>4</sub>), and anterior (H<sub>5</sub>) views; Demshyn beds. Scale bars 2 mm.

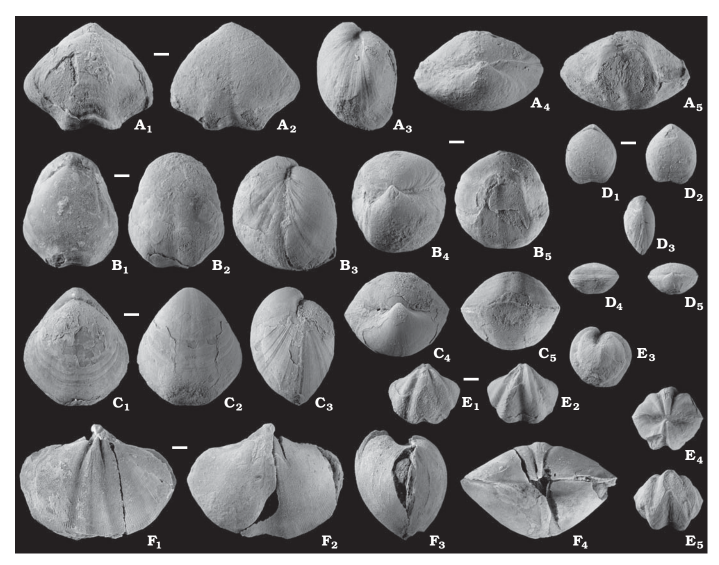


Fig. 14. Silurian brachiopods from the Kytayhorod section. **A**. *Meristina bilobata* Modzalevskaya, 1980, ZPAL Bp 71/24, shell in dorsal (A<sub>1</sub>), ventral (A<sub>2</sub>), lateral (A<sub>3</sub>), posterior (A<sub>4</sub>), and anterior (A<sub>5</sub>) views; Demshyn beds, sample D2. **B**. *Meristina incrassata* Modzalevskaya, 1980, ZPAL Bp 71/25, shell in dorsal (B<sub>1</sub>), ventral (B<sub>2</sub>), lateral (B<sub>3</sub>), posterior (B<sub>4</sub>), and anterior (B<sub>5</sub>) views; Maryanivka beds, sample M1. **C**. *Meristina podolica* (Nikiforova, 1954), ZPAL Bp 71/26, shell in dorsal (C<sub>1</sub>), ventral (C<sub>2</sub>), lateral (C<sub>3</sub>), posterior (C<sub>4</sub>), and anterior (C<sub>5</sub>) views; upper part of the Restiv beds. **D**. *Glassia elongata* Davidson, 1881, ZPAL Bp 71/27, shell in dorsal (D<sub>1</sub>), ventral (D<sub>2</sub>), lateral (D<sub>3</sub>), posterior (D<sub>4</sub>), and anterior (D<sub>5</sub>) views; Restiv beds, 0.8 m above sample R-1. *E*. *Howellella globosa* Tsegelnyuk, 1976, ZPAL Bp 71/28, in dorsal (E<sub>1</sub>), ventral (E<sub>2</sub>), lateral (E<sub>3</sub>), posterior (E<sub>4</sub>), and anterior (E<sub>5</sub>) views; lower part of the Maryanivka beds. **F**. *Eospirifer radiatus* (Sowerby, 1834), ZPAL Bp 71/29, slightly incomplete shell in dorsal (F<sub>1</sub>), ventral (F<sub>2</sub>), lateral (F<sub>3</sub>), posterior (F<sub>4</sub>), and anterior (F<sub>5</sub>) views; Demshyn beds. Scale bars 2 mm.

the same, a slightly impoverished BA 4–5, some changes are noteworthy. Local extinction of the long-lasting *Dicoelosia* combined with first appperance of *Howelella* may be linked to the basin shallowing (the latter spiriferid is a nominative genus for BA 2 in the coeval Appalachian faunas; Brett and Baird 1995; Boucot 2005).

Thus, the succession of the brachiopod fauna in the Kytayhorod section shows rather gradual and minor temporal changes. The most characteristic trend observable in the fauna is a change from stressed assemblages of prevailing small-dimensioned forms in the Restiv Beds (? a reflection of Lilliput effect during the Ireviken crisis; Urbanek 1993) to medium- and large-sized forms in the Demshyn Beds (i.e., during the  $\delta^{13}C \delta^{13}C_{carb}$  highstand; Fig. 6A), and, particularly, Marya-

nivka Beds. Also important seems to be disappearance of *Skenidioides* in the lower part of the Demshyn Beds and, successively, *Dicoelosia* at the top of the interval. Thus, contrary to the diversity changes and extinctions during the Ireviken Event which have been observed among shallow-water brachiopod fauna on Gotland (Kaljo et al. 1996; Erlfeldt 2006; see also data on the Lau Event in Gustavsson et al. 2005), Australia (lingulids only; Valentine et al. 2003) and North America (Boucot 1990; Boucot in Kaljo et al. 1996), the brachiopods from the Kytayhorod section do not reveal any substantial perturbation. This is in agreement with observation of Watkins et al. (2000) that deep-water *Dicoelosia* communities showed long-term stability and were apparently unaffected by Silurian global oceanic events.



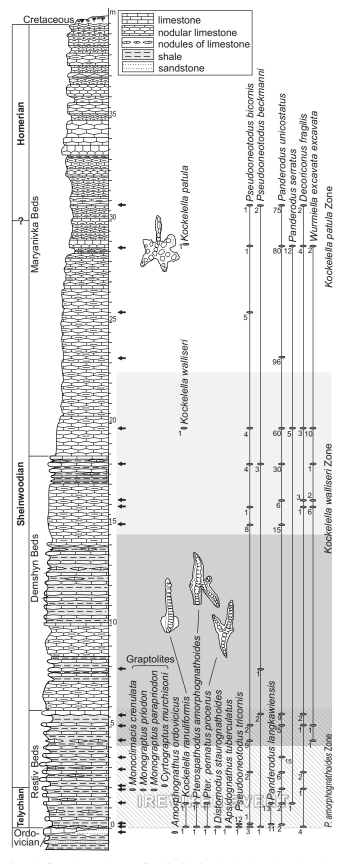


Fig. 15. Conodont and graptolite distribution in the Kytayhorod section. Numbers near the ovals are numbers of specimens in the sample. Shades of gray denote three slices of the ESCIE.

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**Conodonts.**—The conodont assemblages exhibit a two-step distributional pattern, characterized by a low-frequency but relatively high diversity assemblage (up to 9 species) quickly replaced in the succession by a stabilized, but typically species poor (mostly 1–4 species) *Panderodus*-dominated biota of overall increasing abundance (Fig. 15). A similar pattern of abundance was noted for the far richer Estonian assemblages by Hints et al. (2006).

The diverse fauna is limited to two basal samples from the Restiv Beds. The occurrences characterize an initial uppermost Telychian phase of ESCIE, and this regional turnover of the Ireviken Event in the pelagic biota is partly coincident with the oldest "Datum Points" of Jeppsson (1997: fig. 17.2; see also updated data in Cramer et al. 2010). Even if influenced by scarcity of the fauna (mostly single specimens), the First Telychian Datum is clearly suggested by last occurrence of Pseudooneotodus tricornis in the second sample (12 specimens), whilst Panderodus langkawiensis disappeared in the third sample (13 specimens), ca. 1 m higher; this species is considered by Jeppsson (1997, 1998) and Jeppson et al. (2005) to have become extinct at the Sheinwoodian Datum 3.3. Thus, a condensed nature of the lowermost sampled layers in the Ireviken Event may be assumed, especially that coeval condensed sections are known commonly worldwide (Hints et al. 2006; Haq and Schutter 2008: fig. 2; Kleffner and Barrick 2010; Ray and Butcher 2010). A replacement of the series boundary, downward to a ca. 1 m lower level than accepted by Koren et al. (1989), may be therefore considered.

Moreover, even a cryptic hiatus possibly occurs higher in the Ireviken interval (David Loydell, personal communication 2012; see also Kaljo et al. 2007: 216), if the graptolite species *Monoclimacis crenulata* would be confirmed in the Restiv Beds just below entry of the guide Wenlock species, *Cyrtograptus murchisoni* (Fig. 15); *M. crenulata* is unknown worldwide above the middle Telychian *Oktavites spiralis* Biozone (Fig. 2 in Cramer et al. 2010). So, these taxonomic data of Tsegelnyuk et al. (1983) need revision to confirm the implied biostratigraphical or biogeographic peculiarity.

The more widespread faunas from the Demshyn Beds (i.e., in the E-2 highstand) and lower Maryanivka Beds include well-known species that survive the Ireviken Event, such as Decoriconus fragilis and Pseudooneotodus bicornis. In addition, the most common species Panderodus unicostatus also belongs to the survivors, and this ubiquitous species may have had a wide environmental tolerance, as shown by Zhang and Barnes (2002) for the Llandovery of Anticosti Island, Québec. Single occurrences of the index species for the upper, but not uppermost Sheinwoodian Kockelella walliseri Zonal Group, K. walliseri, and K. patula (see Cramer et al. 2010: fig. 3; Sh2 time slice of Cramer et al. 2011a), were found in the Maryanivka Beds. As inferred by Cramer et al. (2010, 211a), this conodont dating, even though under current revision due to a proved biogeographic differentiation in the first occurrence of the both species, coincides with worldwide end of the ESCIE.

	Order	PROSOMATIFERA OPERCULATIFERA		e	colc	rsity and blogical dices			
limestone lumpy limestor nodules of limestone shale sandstone		Ancyrochitina ancyrea Ancyrochitina anserviensis Ancyrochitina primitiva Ancyrochitina sp. Angochitina sp. Angochitina sp. Cingulochitina cingulata Cingulochitina cassa Cingulochitina adiosa Cingulochitina proboscifera Conochitina proboscifera Conochitina aff. proboscifera Conochitina cf. dolioliformis Eisenackitina sp. Margachitina cf. dolioliformis Eisenackitina sp. Margachitina aff. dolioliformis Pterochitina aff. margaritana Pterochitina cf. macroptera	Number of specimens/sample	Number of specimens/gram rock	Diversity -taxa_S	Dominance_D	Margalef's richness index	Events	
Cretaceous	SAMPLES	Ý Ž Ž Ž Ž Ž Ž Ž Ž Ž Ž Ž Ž Ž Ž Ž Ž Ž Ž Ž		NU	D	Do	Ma	Ъ Ш	
▲	Kt-18		118	4	4	0.44	0.63		
Homerian 35 ←	Kt-17		46	3	4	0.25	0.78		
	Kt-16		82	5	6	0.31	1.14	+/-	
Maryanivka Beds	Kt-15		86	5	8	0.16	1.57	_	
↓ ↓	Kt-14		58	5	7	0.2	1.48		
	Kt-13		86	6	7	0.17	1.35		
	Kt-12		15	1	Х	Х	Х		
→ <sup>10</sup>	Kt-11		93	8	7	0.2	1.32		
→ <sup>al</sup> geds	Kt-10		66	6	6	0.22	1.19		
• Demshyn Beds	Kt-09		78	7	4	0.39	0.69		
	Kt-08		46	5	5	0.28	1.05		
0 <b>9 9 9 1 1 1 1</b>	Kt-07		68	7	6	0.20	1.19		
<sup>5</sup> → → → → → → → → → →	Kt-06 Kt-05 Kt-04 Kt-03 Kt-02 Kt-01	IREVIKEN EVENT	72 88 41 69 51 13	7 84551	5 5 4 4 3 X	0.26 0.26 0.31 0.34 0.37 X	0.94 0.92 0.81 0.71 0.51 X	+	

Fig. 16. Distribution of chitinozoans species, diversity, and ecological indices in the Kytayhorod section. Shades of gray denote three slices of the ESCIE.

**Chitinozoans.**—Chitinozoans are organic-walled microfossils of uncertain biological affinity; it is likely that they represent the egg-capsules of soft-bodied marine animals. Their ecological distribution is rather diverse, however, the wide geographical and facies distribution indicate that the chitinozoan biota (chitinozoophorans) may have been either mostly planktic/nekto-pelagic or much more rarely benthic (Miller 1996; Grahn and Paris 2011). The species with thick-walled and poorly ornamented vesicles have been usually interpreted as possibly benthic, high hydrodynamic energy environment occupants, while thin-walled highly ornamented with long processes vesicles have been suggested as pelagic, similar to graptolites in their mode of distribution in the outer shelf environment (Miller 1996; Grahn and Paris 2011).

Chitinozoans in the Kytayhorod section are characterized by a moderately diverse (up to 13 taxa) *Margachitina*-dominated assemblage (SOM 5; Figs. 16, 17). Although their abundance is partly controlled by the facies, with more abundant assemblages from clay-rich rocks, an impoverishment, in frequency and richness terms, has been observed in the basal and upper parts of the succession. This is not a very strong secular differentiation, however, without any obvious

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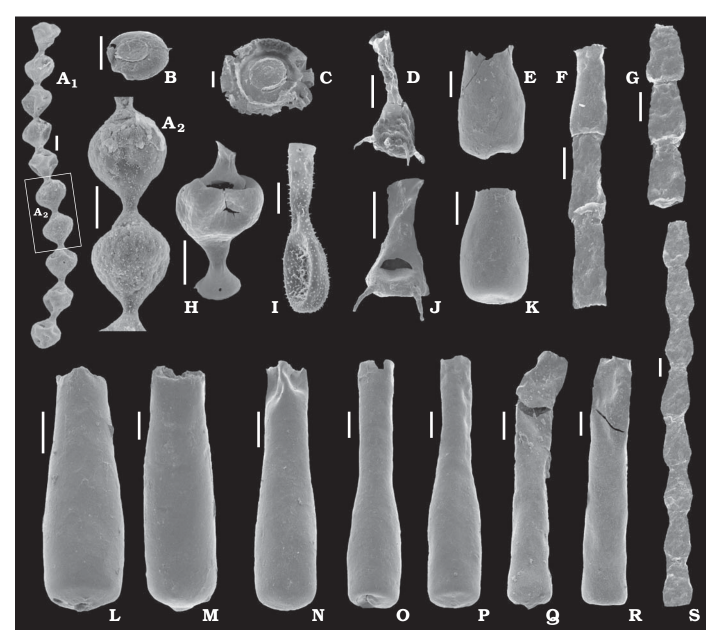


Fig. 17. Stereoscan microphotographs of selected Chitinozoa from the Kytayhorod section. A, H. *Margachitina margaritana* (Eisenack, 1937). A. ZPAL Ch. 8/170S42, sample Kt 06/03; chain of ten vesicles in lateral view (A<sub>1</sub>) and detail of the connection between the vesicle (A<sub>2</sub>). H. ZPAL Ch. 8/170S32, sample Kt 08/11; fragment of a chain of two slightly deformed vesicles showing how aperture is linked to the succeeding vesicle. B. *Calpichitina densa* (Eisenack, 1962), ZPAL Ch 8/176S58, sample Kt 06/15. C. *Pterochitina* cf. *macroptera* Eisenack, 1959, ZPAL Ch. 8/156S128, sample Kt 06/16.
D. *Ancyrochitina* sp., ZPAL Ch 8/176S58, sample Kt 06/06. E. *Eisenackitina dolioliformis* Umnova, 1976, ZPAL Ch 8/176S73, sample Kt 06/05.
F. *Cingulochitina odiosa* (Laufeld, 1974), ZPAL Ch 8/172S46, sample Kt 06/16. G. *Cingulochitina crassa* Nestor, 1994, ZPAL Ch 8/172S48, sample Kt 06/13. I. *Angochitina longicollis* Eisenack, 1959, ZPAL Ch. 8/157S62, sample Kt 08/10. J. *Ancyrochitina primitiva* Eisenack, 1964, ZPAL Ch 8/176S58, sample Kt 06/14. N. *Conochitina proboscifera* Eisenack, 1937.
L. ZPAL Ch 8/176S50, sample Kt 06/17. M. ZPAL Ch 8/176S41, sample Kt 06/14. N. *Conochitina tuba* Eisenack, 1932, ZPAL Ch 8/176S40, sample Kt 06/16.
O. P. *Conochitina pachycephala* Eisenack, 1964. O. ZPAL Ch 8/176S39, sample Kt 06/18. P. ZPAL Ch 8/176S44, sample Kt 06/17. Q. *Conochitina cingulata* (Eisenack, 1937), ZPAL Ch 8/172S55, sample Kt 06/13. R. *Conochitina* sp. ZPAL Ch 8/176S29, sample Kt 06/11. S. *Cingulochitina cingulata* (Eisenack, 1937), ZPAL Ch 8/172S55, sample Kt 06/15.

link with the  $\delta^{13}$ C trends. Two acmes of species representing the genus *Conochitina* were found in the lowermost and topmost samples which were highly-dominated by *C. proboscifera* and *C. pachycephala*, respectively. The chitinozoans suggest a position for the Sheinwoodian–Homerian boundary at ca. 30 m level. This would be above the occurrences of

## *Cingulochitina cingulata* and below the FAD of *Conochitina pachycephala*.

This fairly stabilized dynamics pattern differs from the abrupt changes in chitinozoan faunas at the Llandovery–Wenlock boundary (Nestor et al. 2002; Lehnert et al. 2010), even if partly due to a stratigraphical gap within the *Margachitina* 

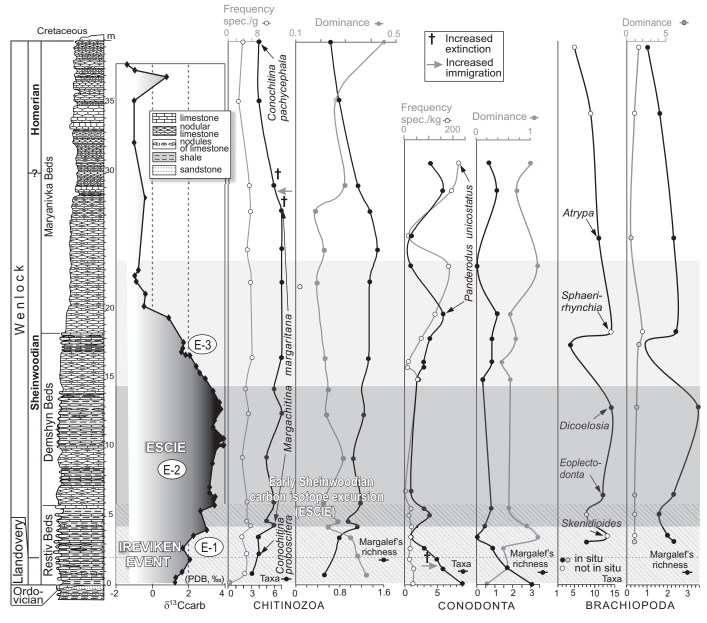


Fig. 18. Faunal dynamics across the Ireviken Event and early Sheinwoodian carbon isotope excursion (ESCIE) in the Kytayhorod section. Shades of gray denote three slices of the ESCIE.

*margaritana* Zone sensu stricto, recognised from Estonia by Loydell and Nestor (2005) and Hints et al. (2006; see also Sadler 2012: fig. 8). Two identified species, *Angochitina longicollis* and, with reservation, *Eisenackitina dolioliformis*, are quoted by Nestor et al. (2002) among chitinozoan victims of the profound late Ireviken crisis in pelagic domain (= level 8 of these authors; or conodont Datum 6; see also Jeppsson et al. 2005).

**Other groups**.—Our field observations, as well literature data on other fossil groups (Nikiforova and Predtechensky 1968: fig. 16; Nikiforova et al. 1972) are in a general agreement with the main distribution patterns outlined above (Fig. 18), controlled primarily by oscillating regressive sea-level trend (Koren et al. 1989: fig. 105).

The Restiv Beds are ecologically distinctive because of the unique occurrences of graptolites (rare in a thin 0.2 m level only; Fig. 15) and macroflora (*Pseudosajania*), and also far more widely distributed benthic groups: crinoids, gastropods, solitary rugose corals (*Orthopaterophyllum*), nautiloids, and trilobites (*Acidaspis*; common in some layers). Also *Tasmanites* (Prasinophyceae, unilocular green algae) was recovered only in the two lowest chitinozoan samples. Assemblages from the lowermost part of the Demshyn Beds include the five above mentioned benthic groups and bivalves, with locally frequent solitary rugose corals (*Orthopaterophyllum*, *Sverigophyllum*), whilst some higher sets (e.g., units 4–6 of Nikiforova et al. 1972) are either unfossiliferous (nodular lithologies) or characterized by infaunal to epifaunal filter-feeding bivalve fauna only, with *Grammysia* and *Mytilarca*. The abundant brachiopod fauna of upper Demshyn Beds is accompanied by a rather impoverished assemblage of trilobites, solitary corals (*Stereotactis*), crinoids, and gastropods.

Poor benthos, in terms of diversity and frequency, is known from the thick nodular limestone units of Maryanivka Beds, where in a few layers, besides brachiopods, merely solitary and colonial corals (*Paleofavosites*), gastropods, crinoids, and trilobites (*Bumastus*) have been found. Intensively bioturbated micritic microfacies varieties of this succession (Figs. 5A, B, D, 11) demonstrate the recurrent development of soft-bodied infauna in the muddy bottom sediments. In addition, there is noteworthy rock-forming abundance of dasycladacean green algae (*Rhabdoporella*) in the shallowing basin (Fig. 5B), up to ca. 50 m water depth (Ischenko 1976, 1985). Also, the Wenlock phytoplankton associations are remarkably diverse in Podolian successions, especially these from the Maryanivka Beds (more than 25 species; Le Hérissé et al. 2009: fig. 8).

## Faunal dynamics across the Klonk Event

Brachiopods.-The stratigraphically oldest brachiopod found in the trench at Dnistrove is Dayia bohemica which occurs in the topmost part of the Dzvenyhorod Beds (see also Kaljo et al. 2012), i.e., about 1.3-1.6 m below the S-D boundary (Fig. 19; see also detailed description of the brachiopod fauna from this section in Baliński 2012). The species occurs mainly in the thin bedded nodular limestone where it forms almost monospecific nest-like clusters of usually articulated shells (Fig. 4). D. bohemica is a characteristic species for the uppermost layers of the Pridoli Series of Bohemia, Estonia, Lithuania, and Latvia (Havlíček and Štorch 1990; Rubel 1977). Tucker (1964) noted that Dayia navicula, a closely related species to D. bohemica, in the Silurian of the Welsh Borderland achieved its widest distribution and greatest abundance in the offshore zones where the sea bottom was below wave base. He also suggested that the incurvature of the ventral umbo in D. navicula prohibited emergence of the pedicle and that the majority of mature individuals rested freely on the sea bottom (see also Kozłowski 1929: 180). Boucot (1975) remarked that Davia constituted a low-diversity, single brachiopod genus community in quiet-water settings within BA 3. Havlíček and Štorch (1990) proposed a separate Dayia bohemica Subcommunity where the nominate species formed brachiopod banks. Johnson (1987) suggested that BA 3 ranges approximately from 30 to 60 m water depth. However, Martma et al. (2005: fig. 3) showed the Ludlow Dayia assemblage in BA4, adjacent to the pelagic biofacies, and this deeper-water position seems to correspond better with the Podolian data (Gritsenko et al. 1999). About 0.3 m above the layer with abundant Dayia bohemica there is a few cm thick shale with numerous specimens of the athyridide *Dnestrina gutta* (Fig. 4). *D. gutta* is a small species, rarely exceeding 6 mm in length, with a plano-convex profile, and open delthyrium. The species is known also from the Pridoli of Moldavia, and Western Europe (Alvarez and Copper 2002). A similar form was described as *D.* cf. *gutta* by Jahnke et al. (1989) from the uppermost Silurian of Yunnan Province, China. The occurrence of *D. gutta* in the Dnistrove Level suggests stressed, impoverished, low diversity, short-ranging assemblages living in somewhat deeper water conditions than *Dayia bohemica*. Besides *D. gutta*, very rare single valves of *D. bohemica* occur in these layers. It seems justified to propose a low diversity, quiet water *Dnestrina gutta* Community which is probably closely analogous to the slightly younger *Gracianella* Community of Boucot (1975) mentioned below.

Although the first appearance of the atrypide Gracianella (Sublepida) paulula at Dnistrove is ca. 20 cm below the S-D boundary, its main occurrence is 40 cm above it. The occurrences are paired with the base of the  $\delta^{13}C_{carb}$  plateau (see Figs. 4 and 19). Other brachiopods constitute 9% of the assemblage being represented by small-sized Resserella elegantuloides, Plectodonta (Plectodonta) mariae pantherae, Pseudoprotathyris infantilis, and Howellella (Howellella) latisinuata. It should be noted that within the range of G. (S.) paulula a layer with the graptolite Monograptus uniformis angustidens has also been recovered. The most characteristic brachiopod G. (S.) paulula is characterized by its minute, up to 4 mm long, weakly biconvex, ribbed shell. It seems that this fauna represents the low-diversity high dominance, quiet-water Gracianella Community of Boucot (1975) recognised in the Silurian non-reef communities of the Uralian-Cordilleran and North Atlantic Regions. According to Boucot (1975) the community relates mainly to BA 4 and 5 but also partly to BA 3 (see also Johnson et al. 1976; Zhang 2001).

In the succeeding 4 m of rhythmic marls and limestones the brachiopods are either very rare, poorly preserved or absent. However, at about 5.5 m above the S–D boundary (beds 47-48) a fossiliferous ca. 30 cm thick layer appears with a high-diversity, well preserved, and distinctive brachiopod assemblage dominated by medium-sized forms. The most characteristic are Septatrypa (Septatrypa) secreta (27% of the assemblage), Plectodonta (Plectodonta) mariae pantherae (13%), Sphaerirhynchia gibbosa (12%), Clorinda pseudolinguifera (11%) and Talentella crassiformis (7%). Thus, the five most common species constitute 70% of the assemblage whereas the remaining 30% of the assemblage comprises 11 brachiopod species. Only Sphaerirhynchia gibbosa occurs more richly higher in the Dnistrove succession. The brachiopod assemblage is represented by a variety of morphological types and ecological adaptations, e.g., small- to large-sized, concavo-convex, biconvex to globose, smooth, ribbed to frilled, with or without functional pedicle (Baliński 2012). The specimens are mostly articulated with the exception of Clorinda pseudolinguifera which possessed a mechanically weaker and more prone to disarticulation hinge structure of deltidiodont type. The presence of numer-

		Order	STROPHOMENIDA	PRODUCTIDA	PROTORTHIDA	ORTHIDA	PENTAMERIDA	RHYNCHONELLIDA	АТКҮРІДА	ATHYRIDIDA	SPIRIFERIDA		Div ec	ersity colog ndice	v and ical es	I
		28 m 26 24 22	Leptaena (L.) aff. altera ?Mesodouvillina sp. Cymostrophia (P.) costatula Plect. (P.) mariae pantherae	<i>Chlupacina</i> sp.	Skenidioides tatyanae	Resserella elegantuloides Dalejina frequens Talentella crassiformis	Clorinda pseudolinguifera	Sphaerirhynchia gibbosa Lanceomyonia borealiformis	Oglu senilis Gracianella (S.) paulula Lissatrypa leprosa Sepatrypa (S.) secreta	Pseudoprotathyris infantilis Camarium sp. Dayia bohemica Dnestrina gutta	Howellella (H.) latisinuata Cyrtina praecedens	taxa	number of specimens	Margalef's richness	dominance	equitability
DEVONIAN	Beds	20										3 2 1 1 2 1 1	7 18 1 5 7 1 1	0.35	0.80	0.50
	Dnistrove Khudykivtsi	112	MLONK EVENT									16 264142221227	1336 4 34 5 3 19 6 6 6 0 130		0.26 0.37 0.81	0.85 0.84 0.49
SILURIAN	Dzvenyhorod Beds	4		marl calca nodul	ar lime limesto	nodules stone						1	130	0	1	0

ous strophomenide *P*. (*P*.) *mariae pantherae* in the assemblage is very distinctive due to its colour pattern performing apparently a protective function through disruptive camouflage against visual systems of potential predators (Baliński

2010). This in turn implies that the brachiopods lived within the photic zone.

The presence of numerous S. (S.) secreta in beds 47-48 suggests that the fauna may be compared to the Silurian

non-reef quiet-water *Dubaria* Community of Boucot (1975) which corresponds with BA 3 (the genus *Dubaria* Termier, 1936 is considered a synonym of *Septatrypa* Kozłowski, 1929; see Copper 2002a: 1465). Jones and Hurst (1984) noted, however, that *Dubaria* in the Silurian of North Greenland inhabited quiet water, cryptic habitats associated with reefs. On the other hand, according to Havlíček and Štorch (1990) in the Pridoli of Bohemia the *Dubaria* Community is related to quiet, shallow-water settings at about BA 2–3. Thus, the inventory and character of the brachiopod and associated fauna from beds 47–48 at Dnistrove (see below) indicate probably a setting near the lower (outer) limit of the photic zone equivalent to the BA 3–4 boundary (see Brett at al. 1993).

In summary, the relations between brachiopod faunal distribution and the isotope excursion are rather complex (see Fig. 19), but progressively improved conditions for the sessile shelly faunas are clearly associated with this event in the earliest Devonian. Far more restricted benthos characterizes the rapid increase phase of  $\delta^{13}C_{carb}$  values prior to the S–D boundary. Only two species became extinct across the period boundary (compare the diversity data in Talent et al. 1993). Interestingly, monospecific acmes of *Dayia minor* distinguish also specific deeper-water habitats through the largescale Ludlow carbon cycle disturbance in the Prague Basin (Lehnert et al. 2007b).

**Conodonts.**—The limestone layers in the Dnistrove succession are mostly rich in conodonts (Figs. 20). Comparatively abundant are: *Ozrkodina typica*, *Parazieglerodina eosteinhornensis*, *Wurmiella excavata*, and *Panderodus unicostatus* (see detailed description of the conodont fauna from this section in Drygant and Szaniawski 2012). Thus, despite somewhat irregular frequency trends, the less abundant faunas occur in the Dnistrove Level and in the uppermost part of the succession (Drygant 2010). In diversity terms, however, more suitable conditions for diverse pelagic biota are found in the Devonian part, i.e., towards the end of the SIDECIE interval (see Fig. 20).

An overturn in the faunas is recognised near the S-D boundary level, when four Silurian-type species have disappeared (e.g., Delotaxis detorta and Parazieglerodina eosteinhornensis), and a more species-rich Devonian assemblage (up to 6 species), with Zieglerodina remscheidensis, Caudicriodus, and Pandorinellina, suddenly flourished in the shallowing epeiric sea. This succession may be overall correlated with the Oulodus elegans detortus to Icriodus hesperius (= Caudicriodus hesperius after Drygant 2010 and Drygant and Szaniawski 2012) zonal passage, as newly redefined by Corradini and Corriga (2012: fig. 5), paired with a radiation of early icriodontids (see also Slavík et al. 2012). Furthermore, a resemblance to the North American faunal step 3 of Barrick et al. (2005), determined by extinction of Delotaxis detorta and approximated with the basal Devonian shift to Decoriconus- and Pseudooneotodus-rich biofacies with Zieglerodina remscheidensis, is noticeable.

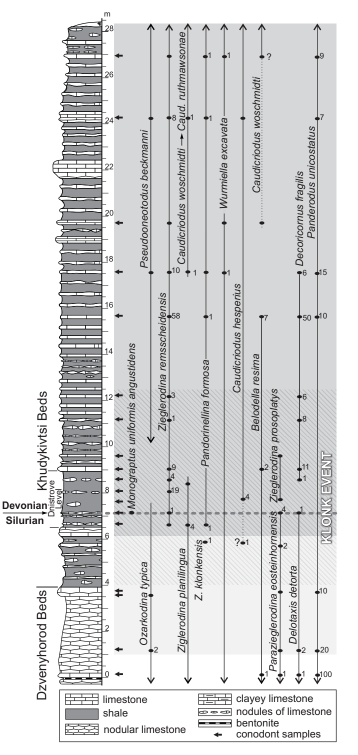


Fig. 20. Conodont and graptolite distribution through the S–D passage beds at Dnistrove. Shades of gray denote two slices of the SIDECIE.

**Chitinozoans.**—The occurrence of characteristic species such as *Urnochitina urna*, *Eisenackitina bohemica*, *Cingulochitina* ex. gr. *ervensis*, *Calpichitina velata*, *Margachitina catenaria* and *Anthochitina* ex. gr. *superba* represents a clear co-occurrence range zone within the S–D boundary interval at Dnistrove (see SOM 6 and Figs. 21, 22). The high abundance of the thick-walled vesicles of *Urnochitina* (Operculatifera)

			PROSOMATIFERA								OPERCULATIFERA									Diversity and ecological indices												
		clayey sha marls calcareou nodular limestone: platy bentonites	s s		Ancyrochitina ancyrea Ancyrochitina cf. aurita	Ancyrochitina desmea	Ancyrochitina cf. lemniscata	Ancyrochitina sp.	Anthochitina superba	Cingulochitina klonkensis	Cingulochitina ex.gr. ervensis	<i>Cinguloochitina</i> sp.	Ramochitina longispina	Kamocnitina ramosus	Bursachitina sp.	Calpichitina velata	Calpichitina aff. gregaria	Eisenackitina barrandei	Eisenackitina bohemica	Margachitina catenaria	Pterochitina sp.	Vinnalochitina pilosa	Vinnalocnitina cr. pilosa Vinnalochitina suchomastvensis	Vinnalochitina sp.	Urnochitina urna	<i>Urnochitina</i> sp.	Number of specimens/samples	Number of specimens/gram rock	Diversity -Taxa_S	Dominance_D	Margalef's richness index	Events
			- ← V-2	29				1	I																		81	8	2	0.51	0.28	
			-26 _ <b></b> V-2	28									1			L								I			96	10	5	0.26	0.88	
			-24 V-2	27																L							217	21	8	0.16	1.30	
			- • <b>-</b> V-2 -22	26																		-					69	7	4	0.29	0.71	
		$\left[ \begin{array}{c} & & & & & \\ & & & & & \\ & & & & & & \\ & & & & & & \\ \end{array} \right] \sim \sim \sim \sim \sim \sim$	<b>←</b> V-2	25																				I			82	8	6	0.18	1.14	
z			-20 . ← V-2	24									1														26	3	2	0.50	0.31	
A - N	Beds		-18 V-2	23														I									45	4	2	0.56	0.26	
N 0			<b>←</b> V-2	22																							18	2	3	0.45	0.69	
D E <	k iv t s		-16 - 🗲 V-2 -	21																							0	0	0			
	hudy		-14 V-2	20																							68	7	3	0.79	0.47	
	¥		<b>←</b> V-1	9																							111	11	5	0.51	0.88	
		~~~~~~		8																							121	12	3	0.41	0.42	
			_10 ▲ V-1	7																	INI						25	2	2	0.51	0.31	
	love		- - <b>*</b> V-1	6																	XEV						23	2	2	0.52	0.32	
	Dnistrove		-	5																	KLONK EVENT			•••			201	20			0.76	
z		20202	🖛 V-1	3														Ì			M						164 5271	16 52		0.83 0.84	0.35	
RIA			← V-1 -4← V-1 -4 V-1	1																							82 53 182	8 5 18	1 5 8	1.00 0.33 0.31	1.00	+
SILU	od Beds		← V-0 ← V-0 ← V-0	)9 )8																					t		28 23 64	18 3 2 6	85 23	0 101	1.20 0.32 0.44	
	Dzwenyhorod Bed		-2	)6 )5 )4																							182 23 64 12 24 47 14 4 3	261 242	NUNNUN	0.40 0.66 0.51 0.50 0.83 0.72 0.34	1.40 1.20 0.32 0.44 0.40 0.42 0.53 0.83	
	Dzw	h	↓ V-0 ↓ V-0	12			ļ																			ļ	4 3	1 1	3	0.34	0.83	

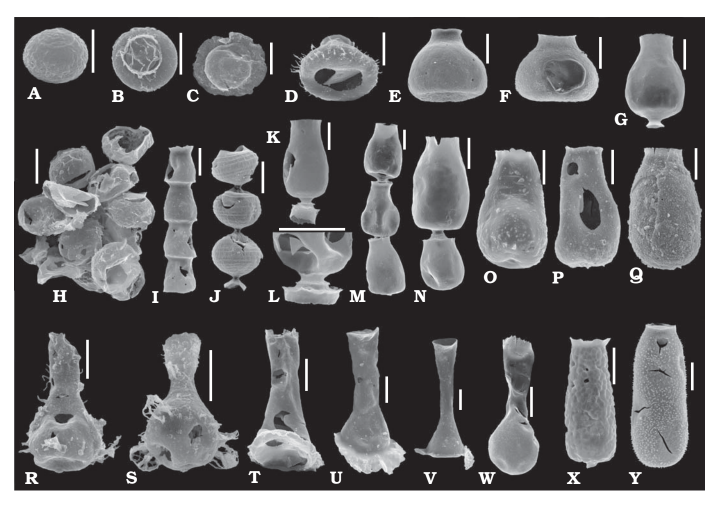


Fig. 22. Stereoscan microphotographs of selected Chitinozoa from the Dnistrove section. **A**, **H**. *Calpichitina* aff. *gregaria* Paris and Křiž, 1984. **A**. Single specimen in oblique lateral view, ZPAL Ch. 8/159S117, sample V-15. **H**. Vesicles aggregated in cluster, ZPAL Ch. 8/161S101, sample V-13. **B**. *Calpichitina velata* (Wrona, 1980), ZPAL Ch 8/160S53, sample V-27. **C**. *Pterochitina* sp., ZPAL Ch. 8/152S099, sample V-28. **D**. *Vinnalochitina pilosa* (Wrona, 1980), ZPAL Ch 8/159S054, sample V-27. **E**–**F**. *Vinnalochitina* cf. *suchomastyensis* Paris and Laufeld, 1981. **E**. ZPAL Ch 8/152S054, sample V-27. **G**, **K**–**N**. *Urnochitina urna* (Eisenack, 1934). **G**. ZPAL Ch 8/160S55, sample V-09. **K**. Specimen with wide copula and operculum attached, ZPAL Ch 8/160S102, sample V-10. **L**. Detail of aboral part showing operculum attachment, ZPAL Ch 8/160S85, sample V-10. **M**. Chain of three closely similar vesicles, ZPAL Ch 8/161S112, sample V-13. **N**. Two marginal vesicles of a chain, showing small terminal vesicle, ZPAL Ch 8/161S104, sample V-15. **I**. *Cingulochitina* ex. gr. *ervensis* (Paris, 1979), Chain of four vesicles, ZPAL Ch 8/159S094, sample V-27. **J**. *Margachitina catenaria* Obut, 1973, Chain of three vesicles, ZPAL Ch 8/159S094, sample V-27. **O**–**Q**. *Eisenackitina barrandei* Paris and Křiž, 1984. **O**. ZPAL Ch 8/161S001, sample V-04. **P**. ZPAL Ch 8/152S017, sample V-05. **Q**. ZPAL Ch 8/152S058, sample V-13. **R**. *Ramochitina* ef. *longispina* (Wrona, 1980), ZPAL Ch 8/172S077, sample V-03. **S**. *Ancyrochitina* cf. *aurita* Wrona, 1980, ZPAL Ch 8/159S02, sample V-27. **T**–**V**. *Anthochitina* ex. gr. *superba* Eisenack, 1971. **T**. ZPAL Ch 8/152S056, sample V-25. **U**. ZPAL Ch 8/159S042, sample V-25. **V**. ZPAL Ch 8/159S082, sample V-25. **W**. *Angochitina* sp., ZPAL Ch 8/159S160, sample V-13. **X**. *Eisenackitina* sp., ZPAL Ch 8/159S042, sample V-03. **Y**. *Eisenackitina bohemica* (Eisenack, 1934), ZPAL Ch 8/159S127, sample V-13. **X**. *Eisenackitina* sp., ZPAL Ch 8/159S127, sample V-23. Scale bars 50 µm.

could reflect a cold water environment related to the cooling period preceding the S–D boundary event near the SIDECIE onset (see below, and Figs. 22, 23, and 25), rather than high energy hydrodynamic conditions. The abundant occurrence of the similarly structured vesicles of *Eisenackitina barrandei* and *E. bohemica* higher in the Klonk Event interval could partly have an analogous nature.

In the upper part of the section, in the SIDECIE slice, an abundant and diversified chitinozoan assemblage occurs again, dominated by *Ancyrochitina* and *Angochitina*. These are species of a fundamentally different group of Chitinozoa belonging to Prosomatifera, characterized by thin-walled highly ornamented vesicles, with spines, ramified processes and vellum. This abundance could be the result of progressive warming during the Lochkovian in the area.

In summary, the distribution of chitinozoans at Dnistrove demonstrates specific dynamics across the major S–D isotopic perturbation (see also Paris and Grahn 1996), comparable only in part with the other pelagic and benthic groups analysed above (Fig. 23).

**Other groups**.—The S–D transition in Podolia is characterized overall by a slight diversity drop in the impoverished organic-walled microphytoplankton, by contrast with the di-

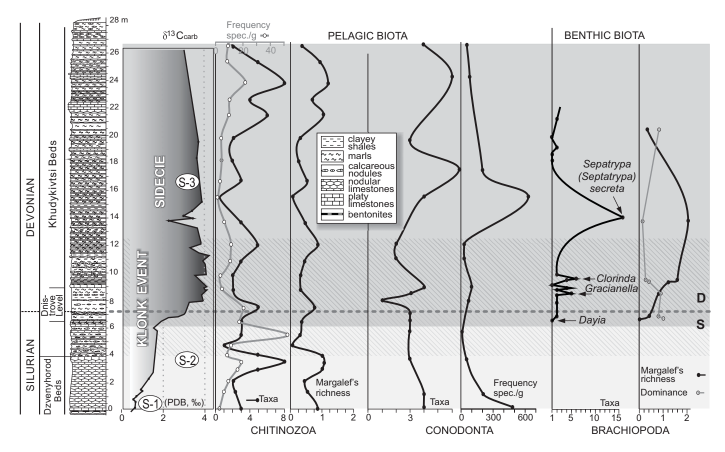


Fig. 23. Faunal dynamics through the Klonk Event and S–D transition carbon isotope excursion (SIDECIE) at Dnistrove. Shades of gray denote two slices of the SIDECIE.

verse older Silurian (Wenlock-Ludlow) acritarch-chlorophyte (prasinophyte) assemblages (see Le Hérissé et al. 2009: fig. 8). In the Dnistrove Level (Fig. 4), palynomorphs in four samples are poorly preserved in dark shaly lithologies due to pervasive oxidation processes, and only the floras from limestone layers are crudely recognizable (Paweł Filipiak, personal communication 2011). Together with infrequent acritarchs, leiospheres and miospores (Apiculiretusispora?), tubular remains with pseudo-cellular cuticles of an enigmatic land plant(?) group Nematophytales (Laevitubuls? and Porcatitubulus) are commonly recovered. The clearly mixed marine-terrestrial character differentiates the Podolian palynofacies from the coeval, more offshore Bohemian assemblage, with acritarchs, leiospheres and mazueloids (Brocke et al. 2006). Large prasinophytes are widespread in both successions (see SOM 6), and probably indicate episodic eutrophic and/or hypoxic conditions, as known e.g., from the Frasnian-Famennian mass extinction boundary beds (Filipiak 2002; see also summary in Le Hérissé et al. 2009).

From a faunal viewpoint, not very frequent trilobites (*Proetus, Acaste*, and *Calymene*), brachiopods (atrypids, *Delthyris*, *Isorthis, Daya*, and *Zygospiraella*), bivalves and rugosan horn corals are quoted from the uppermost Dzvenyhorod Beds (see lists in Nikiforova and Predtechensky 1968: fig. 29; Nikiforova et al. 1972; Nikiforova 1977; Koren et al. 1989). The topmost 1.6 m of the Silurian at Dnistrove are specifically

characterized by occurrences of crinoids, orthocone nautiloids, fistuliporid bryozoans, trilobites (Proetus, Acastella), suspension-feeding bivalves (stationary epifaunal Actinopteria, facultatively mobile infaunal Lunulicardium and Dualina), and ostracods (Libumella, Kuresaaria). In particular, the remnants of the large free-floating crinoid Scyphocrinites elegans, including bulb-shaped loboliths, are very common, forming several lenticular and locally graded bioclastic intercalations, up to 10 cm thick (see Fig. 5G). This unique acme horizon is widely recognizable worldwide (e.g., Zhivkovich and Chekhovich 1986; Jahnke et al. 1989; Havlíček and Štorch 1990; Ferretti et al. 1998; Chlupáč and Hladil 2000; Vacek 2007; Lubeseder 2008; Verniers et al. 2008; Donovan and Lewis 2009; Valenzuela-Ríos and Liao 2012). Within the upper Dnistrove Level, a thin black shale horizon with abundant specimens of the guide graptolite Monograptus uniformis angustidens was found (Fig. 4; see Nikiforova 1977). Conversely, numerous large shells of nautiloid Podolicoceras giganteum are patchy distributed in a ca. 2 m interval thick around the system boundary.

Above the S–D boundary, in addition to common bioturbating infauna (Fig. 5E, F), the brachiopod *Septatrypa*and *Sphaerirhynchia*-dominated faunas are accompanied by rather rare but ubiquitous crinoid ossicles, trilobites (*Acastella*), nautiloids (*Risoceras*), and occasionally also bivalves (*Actinopteria, Panenka, Mytyliarca*), gastropods (*Platyce-* *ras*), corals (*Pachyfavosites*, solitary rugosan *Pseudomicroplasma*), graptolites (*Linograptus*), bryozoans, ostracods (*Richina*), and tentaculitids.

### Discussion

As stressed by many authors, the Silurian global biogeochemical perturbations were regularly associated with stepwise extinction events and restructurings, grouped "at the very beginning or even prior" to the major isotopic event (Munnecke et al. 2010: 301; see also Jeppsson 1997, 1998; Munnecke et al. 2003; Cramer and Saltzman 2007a, b; Loydell 2007; Calner 2008; Lehnert et al. 2010; Melchin et al. 2012: fig. 21.11; Molloy and Simpson 2012; Noble et al. 2012). In particular, Lehnert et al. (2010: 320) stated precisely for the Ireviken Event: "(...) faunal extinctions are connected to time intervals of warming before the establishment of more stable and cooler conditions during the main Sheinwoodian glacial". In fact, since the Jeppsson's (1990, 1997, 1998, 2005) model of oceanic episodes, cooling pulses and, at least in the Early Silurian, expanding ephemeral Gondwanan ice sheets are usually considered responsible for the ecosystem turning points (Lehnert et al. 2007a; Loydell 2007; Calner 2008; Žigaitė et al. 2010; Sadler et al. 2011; Giles 2012; Kozłowski and Sobień 2012; Melchin et al. 2012; Stanley 2012). Conversely, Retallack (2009) interpreted pedostratigraphic spikes (i.e., high-precipitation deep-calcic excursions) as correlative very closely with the Silurian carbon cycle disturbances, and corresponding to greenhouse episodes and sea-level highstands. If so, both the climatic and eustatic driving factors, associated with the biogeochemical signals, were probably quite complex, as shown for the Ireviken Event by Lehnert et al. (2010; see also Stanley 2012). For example, as stressed Kozłowski and Sobień (2012), a combination of glacioeustatic lowstand, gustiness and low-latitude aridity might have been critical for the aeolian Fe-dominated nutrient delivery and cyanobacterial blooming. Also Loydell (2007) regarded that intensified atmospheric circulation during icehouse intervals may have caused increased nutrient input via wind-blown dust, resulting finally in enhanced bioproductivity and benthic oxygen depletion.

Several authors (e.g., Wenzel and Joachimski 1996; Lehnert et al. 2007a; Calner 2008: Fig. 3; Žigaitė et al. 2010) explicitly postulated that the Silurian eustatic cyclicity was glacially driven (see a glacioeustatic scenario of the Lau Event; Fig. 4 in Lehnert et al. 2007a), and this assertion is extended lastly to Early and Middle Devonian (Elrick et al. 2009; see also Miller et al. 2011). However, as noted by Munnecke et al. (2010: 404, 407): "In general there seems to be a clear mismatch between  $\delta^{18}$ O data and sea-level reconstructions based on sequence stratigraphy" and "up until now there is no consensus on Silurian climate development, especially for the post-Llandovery". In addition, Nardin et al. (2011) stressed the climatic impact of changes in plate motion and weathering of fresh volcanic rocks, which resulted in a long-term warming trend from the major late Telychian icehouse (Cancañiri glaciation; Brand et al. 2006) to the Middle Devonian.

On the other hand, the stressed diachroneity of biotic and isotopic signals, influenced by the pattern in the best-known Ireviken Event, is an overestimation as the Silurian paradigma. Sadler (2012: 5) acknowledged clearly that "there is no standard numerical method for picking boundaries and peaks of Silurian isotope excursions". More importantly, as confirmed also by the Podolian record, the extinction steps are largely placed in the rising limb of the  $\delta^{13}$ C curves (see Figs. 24, 25), and thus linked to an initial shift in the carbon cycling, but not with its background (= steady) biogeochemical state. There is little doubt that these ecological turning points were driven by an accelerating disequilibrium of fragile habitat parameters and suddenly increasing environmental stress (see e.g., Urbanek 1993), coarsely only mirrored by the measured <sup>13</sup>C enrichment trends.

Significance of the Podolian succession.-Both global isotopic events are unquestionably paired with  $\delta^{18}O_{carb}$  positive excursions in the Podolian sections (Figs. 5A and 7), confirmed by heavy  $\delta^{18}O_{phosp}$  values in Joachimski et al. (2009) and Lehnert et al. (2010), and therefore to largely declining near-surface temperatures in the marine basin (Figs. 24, 25). Rapid climate perturbations, maybe forced by volcanic activity (Lehnert et al. 2010), are also partly supported by the  $\Delta^{13}$ C secular patterns, suggestive of cooling episodes promoted by falling levels of atmospheric  $pCO_2$ , as recorded in negative  $\Delta^{13}$ C shifts. This chemostratigraphical pattern is certainly traced only across the S-D transition. The lower-resolution data for the ESCIE are rather inconclusive but it seems to deny a dominant warming trend, as postulated from the North American  $\Delta^{13}$ C record by Cramer and Saltzman (2007a). In fact, correlation of the  $\delta^{18}O_{phosp}$  and  $\delta^{18}O_{carb}$  value shifts and negative  $\Delta^{13}C$  signals is to some extent vague (compare Fig. 6A with Bickert et al. 1997: fig. 3; Heath et al. 1998: fig. 2; Brand et al. 2006: fig. 12; Cramer and Saltzman 2007a: fig. 5; Lehnert et al. 2010: figs. 3, 4; see also Noble et al. 2012: fig. 5), and further studies are warranted.

Furthermore, the predicted environmental changes are expected to affect biota mostly in pelagic habitats (especially conodonts, graptolites, and trilobites; e.g., Kaljo et al. 1996; Lehnert et al. 2007b; Loydell 2007; Calner 2008; Sadler et al. 2011; ?also radiolarians, Noble et al. 2012), during times of oligotrophic photic-zone conditions and reduced primary production. Faunal responses in the both Podolian sections follow the above scenario only in part, as manifested especially by the conodont crisis of the Ireviken Event (?also by ostracod turnover; Abushik 1971), prior to the major isotopic/climatic ESCIE disturbance. This large-scale biotic crisis did not affect the relatively deeper-water brachiopods and chitinozoans (in taxonomic richness terms). Nevertheless, the Lilliput effect sensu Urbanek (1993), known among Gotland brachiopods during the Lau Event (Gustavsson et al. 2005), is a recognizable phenomenon also in the Ireviken-re-

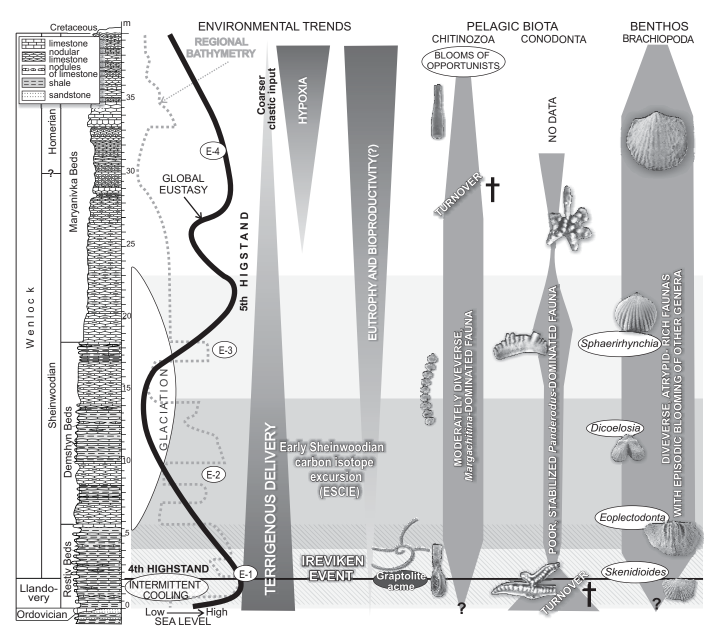


Fig. 24. Scheme of main environmental changes and faunal responses across the Ireviken Event and early Sheinwoodian carbon isotope excursion (ESCIE) in the Kytayhorod succession. Overall eustatic pattern and climatic events compiled from Johnson (2010: fig. 2; see other sea-level curves in Lüning et al. 2000: fig. 9; Cramer and Saltzman 2007: fig. 5; Brett et al. 2009: fig. 20; Ray and Butcher 2010: fig. 1) and Lehnert et al. (2010: figs. 1–3), respectively. Regional bathymetric curve modified from Koren et al. (1989: fig. 1). Note a distinctive discrepancy of the both sea-level patterns in some intervals, probably mostly controlled by regional epeirogenic activity in the Podolian shelf domain. Shades of gray denote three slices of the ESCIE.

lated brachiopod fauna from Kytayhorod. Thus, a regional distinctiveness of the Podolian epeiric domain seems to be manifested, especially in the negligible dynamics of specimen- and species-poor chitinozoan communities across the Ireviken Event and heavy carbon ESCIE interval.

The carbonate ecosystem evolution, from outer shelf toward reefal platform (Koren et al. 1989), was forced mostly by the regional epeirogeny regime paired with a generally regressive eustatic tendency, after the abrupt transgressive onlap inluenced by conspicuous eustatic sea-level rise in late Llandovery (the highstand 4 of Johnson 2006: fig. 1). This synsedimentary tectonism in the shelf basin located near to the mobile Trans-European Suture is revealed by several discrepancies between global and regional sea-level changes, along with rather weak correlation with glacioeustatic effects, implied from the O-isotopic data (Fig. 24; see above). Remarkably, as revealed by the large-scale increase in the <sup>87</sup>Sr/<sup>86</sup>Sr ratio (Brand et al. 2006; Gouldey et al. 2010), the perturbation in long-term carbon cycle and pronounced climatic changes due to increased chemical weathering were likely driven by the initial plate-terrane collisions (Salinic orogeny; see Goodman and Brett 1994; Ettensohn and Brett

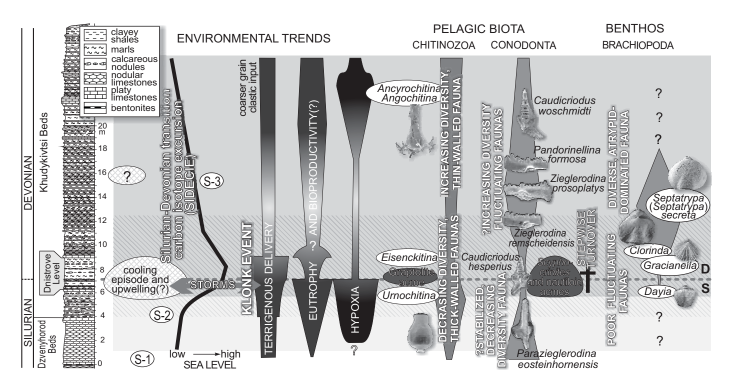


Fig. 25. Scheme of main environmental changes and faunal responses across the Klonk Event and the S–D transition carbon isotope excursion (SIDECIE) in the Dnistrove succession. Sea-level pattern after Walliser (1996: fig. 1; see other sea-level curves in Haq and Schutter 2008: fig. 2; Brett et al. 2009: fig. 20; Johnson 2010: fig. 1; Munnecke et al. 2010: fig. 2 ); for discussion of differently recorded eustasy, as well as climatic setting see Buggisch and Joachimski (2006) and Małkowski and Racki (2009). Shades of gray denote two slices of the SIDECIE.

## 1998; Brand et al. 2006; Cramer and Saltzman 2007a; Kraw-czyk et al. 2008).

The S–D boundary isotopic event appears to be more widely geographically recorded than previously assumed (see Małkowski et al. 2009: fig. 8). The SIDECIE, in various chemostratigraphical variants, in addition to new Laurentian and peri-Gondwanide sites (e.g., Suttner 2007; Kleffner et al. 2009), is recognizable in organic carbon also in the South China, Turkey, and Morocco (amplitude between ca. 3‰ and 8‰; Zhao et al. 2011), as well as in the whole-rock carbonate record in the Canadian Arctic islands (amplitude of ca. 3‰ and 3.5‰ in two sections; Märss et al. 1998), east Baltica (ca. 2‰ only; Kaljo et al. 2012) ; Central Urals (ca. 4‰ in shallow-water facies, but only 2.2‰ in pelagic micrites; Chekhovich et al. 1990), and Subpolar Urals (at least ca. 1.5‰ in shallow-water, coral-rich facies; Yureva et al. 2002).

During this S–D boundary turning point, a moderate faunal changeover, both in benthic and pelagic groups, is recognised in the Podolian epeiric sea exclusively at the very beginning of the prolonged <sup>13</sup>C-enriched SIDECIE timespan, possibly due to progressive dysoxia induced via enhanced bioproductivity by a climatic cooling (e.g., Wenzel and Joachimski 1996; Loydell 2007), an echo of the conspicuous mid-Pridoli glacial episode (Žigaitė et al. 2010: fig. 3). In Podolia, the pelagic graptolite acme and clay-dominated deposition of the Dnistrove Level were clearly linked with a deepening pulse across the S–D boundary (Fig. 25; compare Koren et al. 1989: fig. 105; Kaljo et al. 2012), interpreted as a typical event for the "Bohemian facies" by Walliser (1996; see also Jeppsson 1998; Brett et al. 2009; Vacek et al. 2010, Valenzuela-Ríos and Liao 2012). This sea-level pattern was unexpectedly paired with "regressive" scyphocrinoid tempestite episodes (see Fig. 5G) and nautiloid profusion on the distal hemipelagic shelf that was recurrently punctuated by major storm action (compare Vacek 2007). A similar peculiar facies shift is commonly reported from strata deposited during the Klonk Event (Walliser 1996), e.g., from Central Urals (Zhivkovich and Chekhovich 1986), Sardinia (Ferretti et al. 1998), Bohemia (Hladil and Beroušek 1992; Vacek 2007), China (Jahnke et al. 1989), and Morocco (Lubeseder 2008). As advocated by Vacek et al. (2010: 270): "The facies change close to the S-D boundary and deposition of the Scyphocrinites H[orizon] might predominantly result from a biotic event unrelated to sea-level changes and local subsidence, rather than from sea-level rise/drop". In general terms, comparable biotic dynamics were established in many other localities from different continents (e.g., Havlíček and Štorch 1990; Feist et al. 1997; Kříž 1998; Chlupáč and Hladil 2000; Yureva et al. 2002; Barrick et al. 2005; Suttner 2007; Manda and Frýda 2010; Zhao et al. 2011; Valenzuela-Ríos and Liao 2012). In particular, the transient collapse of the Podolian carbonate factory ("carbonate crisis") corresponded to worldwide reef demise (Copper 2002b; Kiessling 2009), but without coeval loss in graptolite diversity (see Sadler et al. 2011).

On the other hand, the possible influence of eustatic sea-

level rise (?and nutrient-rich upwelling currents; Lubeseder 2008; Manda and Frýda 2010; Histon 2012) remains conjectural in spite of contrary, i.e., regressive bathymetric data from different regions (e.g., House and Ziegler 1997; Carrera et al. 2012), as summarized by Buggisch and Joachimski (2006) and Małkowski and Racki (2009). A tectonic control of the sea-level changes, in the collisional geodynamic setting (Acadian orogeny and a final closure of the Iapetus Ocean; see e.g., Goodman and Brett 1994; Copper 2002b; Krawczyk et al. 2008; Małkowski and Racki 2009; Narkiewicz et al. 2011), is expected also in the Klonk Event timespan. As manifested by thicknesses of the Přidoli-Lochkovian open-shelf successions (above 1 km; Nikiforova et al. 1972; Małkowski et al. 2009), a subsidence rate in the peri-Tornquist foreland basin (?due to dextral transtension) was very high (see Drygant 2003; Sliaupa et al. 2006). During the comparatively very warm timespan (Joachimski et al. 2009; Nardin et al. 2011), this regionally effective factor overrode a regressive, allegedly glacioeustatic tendency (see Žigaitė et al. 2010).

After this overall increased stress episode, the Early Devonian carbonate ecosystem quickly recovered in a shallowing setting. Diverse shelly benthos and pelagic communities temporarily flourished in the Podolian shelf domain (see Nikiforova et al. 1972), still being affected by episodic oxygenation and temperature changes.

## Concluding remarks and implications

The Ireviken and Klonk biotic crises and related global isotope excursions, identified in Podolian sections, reveal opposite environmental and biotic trends, partly influenced by the later Caledonian geodynamic setting of the TESZ domain:

(i) The major Ireviken Event was marked by well-defined temporal changes in detrital input, redox states and, supposedly, bioproductivity, but without correlative relations with the  $\delta^{13}$ C pattern (Fig. 24). Thus, the global biogeochemical disturbance was of minor ecosystem significance in the mostly epeirogeny-controlled sedimentary signature of this epeiric sea, as claimed also for other Laurussian sites in many other studies (see review in Cramer and Saltzman 2007a, b; Calner 2008; Munnecke et al. 2010).

(ii) The depositional and ecological signature of the Klonk Event exhibits some temporal links with the abrupt  $\delta^{13}$ C shift toward the basal Devonian isotopic plateau, although effectively overprinted by the tectonically driven transgressive trend in the crucial interval (Fig. 25). Regardless of specific causes, the isotopic shift is reflected in the carbonate crisis paired with a tendency towards eutrophication, oxygen deficiency and temperature drop at the S–D boundary time. The features collectively point to the uniqueness of the Klonk Event among the Silurian global events, and some similarity already to Devonian transgressive/anoxic episodes (see Walli-

ser 1996; Buggisch and Joachimski 2006; Małkowski and Racki 2009).

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(iii) In the light of the above results, advanced multidisciplinary studies to decipher complex interactions between Silurian ecosystems, forced by climatic, eustatic, tectonic and subsequent biogeochemical (i.e., carbon cycling) factors, and quickly evolving marine and terrestrial biospheres, in terms of effects of the accelerated terrestrialization (see Strother et al. 2010), are necessary in other palaeogeographical domains.

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