Diversity and biostratigraphic utility of Ordovician brachiopods in the East Baltic

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Abstract. The stratigraphy of the Ordovician carbonates of Baltoscandia was initially based, during the 19th century, on the stratigraphical ranges of macrofossils, mainly trilobites, but other fossils (brachiopods, echinoderms and cephalopods) were also used. During the 20th century, their importance in biostratigraphy gradually decreased due to a greater reliance on microfossils, especially conodonts and chitinozoans, which enable accurate correlation of carbonate successions where graptolites are absent or very rare. New methods have further reduced the attraction of macrofossils for biostratigraphy, although they are useful tools in different fields of geology such as palaeobiogeography and palaeoecology. The revised data on species diversity and the stratigraphical distribution of articulated brachiopods with carbonate shells (rhynchonelliformeans) in the East Baltic are used here for the evaluation of their role and potential in the modern stratigraphy of the Ordovician System. The 106 stratigraphical units (mainly formations and members) belonging to 17 Ordovician and the lowermost Silurian regional stages are analysed based on the taxonomic composition of their brachiopod faunas comprising in total more than 400 species. The cluster analysis used in this stratigraphical experiment captures the major dissimilarities between and continuity of the regional subseries, stages and subregional units by the similarity of their brachiopod faunas. Bryozoans, another important group of benthic organisms in shallow-water facies, are analysed for comparison, providing a test for the correlations based on brachiopods.

Key words: brachiopods, multivariate data analysis, biostratigraphy, diversity, East Baltic.

INTRODUCTION

The Ordovician stratigraphical framework of Baltoscandia, including the East Baltic, was established during the 19th and beginning of the 20th centuries (Schmidt 1858, 1881, 1907; Törnquist 1883; Warburg 1910). In Estonia, the stage-level stratigraphy and names of units were defined by Schmidt. The detailed subdivisions of the Lower and lowermost Middle Ordovician in the Klint area of Estonia and NW Russia are those of Lamansky (1905). The stratigraphic ranges of trilobites and a few other fossil groups were the main tools for the correlation of Ordovician strata.

The stratigraphy of the Ordovician in the East Baltic is based traditionally on a combination of lithological characteristics, which define the formations and members (Männil & Meidla 1994), together with biostratigraphic information, including the zonal ranges of fossils. In the lower half of the Ordovician, trilobites are essential for correlation; their zonation was recently revised by Bergström et al. (2013). Nowadays the exact correlation of the carbonate rocks of the Baltic sections is based on a biozonation of the distribution of conodonts and chitinozoans (Nõlvak et al. 2007; Männik & Viira 2012). The use of graptolites in biostratigraphy is limited in these rocks (Paškevičius 1973, 2011; Männil 1976).

The current study deals with the East Baltic region and covers the Baltic States of Estonia, Latvia and Lithuania together with the northwestern part of Belarus. This region represents the main areas of the Estonian and Lithuanian shelves separated by the Livonian Basin (Fig. 1). The easternmost territories with Ordovician strata, including the St Petersburg area, are not considered in this study. Incorporating these data would have required more material from the areas east of the Baltic region, including the Moscow Basin, which is beyond the scope of this study. Notable extensive investigations of the brachiopod faunas, environments and stratigraphy...
of the Floian to lower Darriwilian (Hunneberg to Kunda regional stages) have been carried out in the St Petersburg area (Egerquist 1999; Hansen & Harper 2003; Rasmussen et al. 2007, 2016; Rasmussen & Harper 2008). The bed by bed studies of sections have revealed there a much higher brachiopod diversity than in northern Estonia (Rasmussen et al. 2007). The Lower and lowermost Middle Ordovician have a restricted thickness in northern Estonia (Meidla 1997) and are not as fossiliferous as the sections in the easternmost areas noted above. The facies differentiation and the diversity of benthic faunas in the Baltic Basin are described in several publications (e.g. Paškevičius 1973, 1997, 2000; Jaanusson 1976; Hints & Põlma 1981; Harper 1986; Ropot & Pushkin 1987; Hints 1990a; Laškovas et al. 1993; Hints & Harper 2003; Kaljo et al. 2011). The changes in the composition of the Ordovician biota (e.g. trilobites, brachiopods and bryozoans) along the onshore–offshore profile have been plotted using a dendrogram (Hints 1990a) and illustrated in separate sections (Hints & Põlma 1981; Harper & Hints 2003; Kaljo et al. 2011). The modern geological context of the Baltic region has been discussed in numerous publications commencing with Männil (1966), who described the general features of the basin, its evolution and faunas. Subsequent overviews of Ordovician faunas and facies are reported in Ropot & Pushkin (1987), Paškevičius (1997) and Raukas & Teedumäe (1997). The differences in the development of brachiopod faunas on the Estonian and Lithuanian shelves, and in the Livonian Basin are worthy of note. The benthic biotas of the shelves, predominantly in carbonate depositional regimes, apparently lacked environmental stress apart from the intervals of emergence indicated by the gaps in the deposition of different durations (Laškovas 2000; Meidla et al. 2014). Three different ecological faunas developed in the Livonian Basin, which were influenced by immigrations from the westernmost areas. Rare and very low-diversity faunas in the red-coloured facies during the Early, first half of the Middle, and Upper Ordovician developed in specific environments with strong, multi-directional currents (Kiipli et al. 2008). Low-diversity brachiopod faunas also occur in black shales formed in anoxic conditions (Kiipli & Kiipli 2013) at two stratigraphical levels within the Katian. In the Livonian Basin, most of identified brachiopods belong to the Sandbian–lower Katian and Hirnantian siliciclastic deposits. In the former case these deposits formed in oxygen-rich marine environments during an interval of volcanic activity (Bergström et al. 1995). During the Hirnantian the environments and biota were stressed by the development of the Gondwana icecap (Brenchley et al. 1994). The brachiopods have essential roles in studies of various aspects of palaeobiogeography and palaeoecology. Nevertheless, some rapidly evolving lineages, for example the dalmanelloid and plectambonitoid brachiopods, have been used for local and regional correlation, with limited success, in the Middle (Williams 1976) and Upper Ordovician (Bancroft 1945; Wright 1976) rocks of Avalonia. Brachiopod assemblages have been used to characterize rock units, most notably Cooper’s (1956) benchmark study of the Ordovician brachiopods of the mid-Continent and margins of eastern North America; that pioneer study developed the value of an assemblage-
1. The genus-level taxonomy of the Ordovician brachiopod-dominated benthos. The present study, however, tests the importance of brachiopods in identifying larger, stratigraphic units with precise time constraints and clarifying if, within a regional context, stages, subseries and series can be recognized on the basis of their brachiopod-dominated benthos.

COMMENTS ON BRACHIOPOD TAXONOMY

1. The genus-level taxonomy of the Ordovician brachiopod-dominated benthos follows the volumes of the Treatise on Invertebrate Paleontology (e.g. Harper 2000; Williams & Carlson 2000; Williams & Harper 2000 and Cocks & Rong 2007). These studies synonymized several new names for Estonian strophomenoid genera published by Rõõmusoks (2004).

2. Open nomenclature (species names with cf., aff., sp. nov.) is used in some cases. In other cases those brachiopods are briefly described (Öpik 1930, 1932, 1934; Hints 1975; Tinn 1998; Hints & Harper 2015).

3. The initial data on the stratigraphical distribution of species are complemented or have been revised by other authors. Up to now, the most complete lists of brachiopod and bryozoan species from the East Baltic Ordovician are those of Rõõmusoks (1966, 1970), Ropot & Pushkin (1987) and Paškevičius (1997). The genus-level overviews are available in Hints (1990b) and Hints & Rõõmusoks (1997), and some aspects of the brachiopod diversity and dynamics in Hints & Harper (2003).

4. The brachiopod data from Belarus (Table S3) are based on nine drill core sections (figs 4–12 in Ropot & Pushkin 1987). The stage-level data on brachiopod distribution in the southern East Baltic (text-table 5 in Ropot & Pushkin 1987) are not used here, because we have revised data for the Lithuanian part of the region (Table S2).

5. The brachiopods from the Livonian Basin have been insufficiently studied and are identified at different taxonomic levels (Table S3). An exception is the Hirnantian fauna (Hints & Harper 2015).

COMMENTS ON THE STRATIGRAPHICAL CONTEXT AND CONTROL

1. In the Baltic region, the modern stratigraphic classification of the Ordovician System, chrono-, bio- and lithostratigraphical units and correlation with the global chronostratigraphical units have been summarized in several key publications (Männil 1966; Paškevičius 1973, 1997; Ulst et al. 1982; Ropot & Pushkin 1987; Ebbestad & Högström 2007; Nõlvak et al. 2007; Meidla et al. 2014). The stratigraphical chart in Fig. 2 comprises the main regional stratigraphical units (regional stages) and their correlation with the global stages. The lithostratigraphical units (formations and members) identified in the Lithuanian Shelf (Ropot & Pushkin 1987; Paškevičius 1997) (Fig. 2), are ranked on the chart in stratigraphic order within regional stages without exact correlations with the formations and members in Estonia.

2. Nevertheless there are complications concerning the ranges of species because of different or changing opinions on the identification of sequences (mainly those in the Lasnamägi to Kukruse stages) by different authors (e.g. figs 17 and 26 in Rõõmusoks 1970). According to Rõõmusoks (1970, fig. 26) and Männil (1986), stage C2 (Kukruse) of Öpik (1930, 1934) corresponds to the uppermost Uhaku and lowermost Kukruse stages and Öpik’s stage C3 (Idavere) corresponds to the Kukruse Regional Stage (RS).

3. The brachiopod data from the Lasnamägi RS possibly need future revision because, based on the characteristic graptolites (Männil 1976, fig. 2), in the Lasnamägi section (Tallinn) (Rõõmusoks 1970, fig. 11) part of that stage should be included in the Uhaku RS.

4. The lower part of the Haljala RS (C3), comprising the Tatruse Formation (Fm.) and Vasavere Member (Mb.), corresponds to the Idavere RS and the upper part to the Jõhvi RS in Rõõmusoks (1970).

5. Studies of the early Katian reefs (Vasalemma Fm.) in northern Estonia (Kröger et al. 2014) have suggested a Keila age for the reef complex, which earlier was partly included in the Oandu RS (Männil 1960; Rõõmusoks 1970). The Sakum Mb. of the uppermost Vasalemma Fm. is considered as transitional between the reefs and the carbonate deposits of the Hirnuse Fm. (Männil 1990). According to new data (Kröger et al. 2014, 2017), the Sakum Mb. overlies the reefs and its topmost part could be younger than the Hirnuse Fm. Therefore, the ranges of several brachiopods may require adjustment. Some common species (e.g. Bassettella alata Hints, Neoplatystrophia lutkevichi (Alichova), Vellamo oanduensis Öpik, Ilmarinia dimorpha Öpik) from the Hirnuse Fm. of the Oandu RS and the Vasalemma Fm. presumably appear first during the Keila in reef environments and may thus no longer indicate an Oandu age.

6. The continuous transition from the Ordovician to Silurian is missing in the East Baltic due to gaps of different durations (Paškevičius 1973, 1997; Nestor & Einasto 1997; Abushik et al. 2007; Männik 2014). The most diverse, early Rhuddanian brachiopod fauna
noted in this study has been reported from the Estonian shelf (Varbola Fm.) and basinal deposits (Öhne Fm.) of the Juuru RS in the Central East Baltic (Rubel & Rõõmusoks 1970; Rubel et al. 1984; Rubel 2011). In Belarus, the gap is much more extensive than in other parts of the East Baltic and the oldest Silurian is of Telychian age (Abushik et al. 2007).
MATERIAL AND METHODS

The Ordovician brachiopods of the East Baltic have been described in a number of substantial monographic studies (Öpik 1930, 1932; Alikhova 1951, 1953; Alikhova et al. 1954; Rõõmusoks 1959, 2004; Rubel 1961, 2011; Hints 1975; Cocks & Rong 2000; Rubel & Wright 2000) and in numerous shorter papers (Öpik 1932, 1933a, 1933b; Oraspöld 1956, 1959; Rubel 1963; Rõõmusoks 1964, 1981, 1985; Hints 1973, 1979, 1986, 1993, 2012, 2014; Paškevičius 1973, 2016; Rubel & Popov 1994; Tinn 1998; Zuykov 1999; Paškevičius & Hints 2016). These data are supplemented with those from several stratigraphical reviews (Jaanusson 1956; Männil 1966; Rõõmusoks 1966, 1970). The Ordovician and lowermost Silurian brachiopods on the Estonian Shelf are assigned to about 400 species in addition to numerous taxa under open nomenclature. On the Lithuanian Shelf the number of brachiopod species is about a half of that on the Estonian Shelf (Alikhova et al. 1954; Paškevičius 1997). Most of them are also present on the latter shelf. More than 150 taxa of brachiopods occur in the periphery of the Livonian Shelf in northwestern Belarus (Ropot & Pushkin 1987). Some of these species are unknown or have somewhat different stratigraphic distributions in the rest of the East Baltic.

About 100 rhyynchonelliforme brachiopods are present in the Livonian Basin. However, many of them are identified only to genus or higher taxonomic levels.

The source data on bryozoans (275 species; see Table S6 at http://doi.org/10.15152/GEO.233) used in this paper are based on the faunal lists in Rõõmusoks (1966, 1970) where original data by Ralf Männil (Männil 1966; Rõõmusoks 1966, 1970). The Ordovician and lowermost Silurian bryozoans on the Estonian Shelf are assigned to about 400 species in common to the number of taxa under open nomenclature. On the Lithuanian Shelf the number of bryozoan species is about a half of that on the Estonian Shelf (Alikhova et al. 1954; Paškevičius 1997). Most of them are also present on the latter shelf. More than 150 taxa of bryozoans occur in the periphery of the Livonian Shelf in northwestern Belarus (Ropot & Pushkin 1987). Some of these species are unknown or have somewhat different stratigraphic distributions in the rest of the East Baltic.

The Ordovician brachiopods of the East Baltic have been taken into account. The bryozoans in (1966, 1970) where original data by Ralf Männil (Männil 1966; Rõõmusoks 1966, 1970). The Ordovician and lowermost Silurian brachiopods of the East Baltic using the neighbour-joining algorithm and the Raup–Crick and Simpson similarity coefficients (Hammer & Harper 2006). These analyses enable us to visualize the similarity of stratigraphical units according to the presence and absence of brachiopod species or their generic composition and highlight how these fossils support the accepted stratigraphy and its correlation.

The original material of the rhyynchonelliforme brachiopods used in this study is housed at the Institute of the Geology (Tallinn Technical University), Geological Museum (Tartu), Natural History Museum (Tallinn), Latvian Museum of Natural Sciences (Riga), Institute of Geosciences (Vilnius University) and Belarussian Research Geological-Prospecting Institute (BELNIGRI, Minsk). The large collections, studied by Alikhova, are housed at the Chernyshev Central Scientific Geological and Prospecting Museum (CNIGR museum) in St Petersburg.

THE MULTIVARIATE ANALYSIS

The multivariate technique of cluster analysis (Hammer & Harper 2006) is widely used in palaeobiogeography for the identification of faunal differences between palaeocontinents, regions and faunas of various ages (Tychsen & Harper 2004; Harper et al. 2013), more specifically, in studies of the Ordovician biodiversification (Harper 2006) and faunal turnovers (Zhan et al. 2008). The present use of cluster analysis is somewhat unconventional and differs from those applications, because of its stratigraphically-oriented goals. The neighbour-joining algorithm and the Raup–Crick and Simpson similarity coefficients (see Hammer & Harper 2006) enable visualization of the similarity between different stratigraphical units based on the taxonomic composition of brachiopods. The first coefficient is a probabilistic index based on presence/absence data. The Simpson similarity coefficient characterizes the ratio of the species in common to the number of taxa in the smaller set. The dendrograms based on two coefficients reveal some differences useful in the interpretation of the faunal similarities of separate units.

Cluster analyses of stratigraphical units based on brachiopod composition for the Estonian Shelf

In the East Baltic, the development of brachiopod faunas, their diversity changes and role in the benthic faunas differ essentially between the shelves and the basinal part of the basin (Fig. 3). The brachiopod data from the Estonian Shelf are from 41 stratigraphical units (Fig. 2), arranged into three main clusters using the Raup–Crick
and Simpson similarity coefficients (Fig. 4). One cluster comprises the units from the Hunnebergian RS (B1a, units 2, 3) to the Kunda RS (B3) (units 3–10). The Aseri RS (unit 11) belongs, unusually, to that cluster. On the stratigraphical chart (Meidla et al. 2014), and in the cluster based on the Simpson similarity coefficient, it is tied to the overlying units. The units from the Lasnamägi RS (C1b) to Keila RS (D2) (units 12–22) form the second cluster which together with the first one are separated from all other Ordovician units in the interval from the Oandu RS (D3) to Porkuni RS (F2). The major Ordovician faunal change in the shelf facies at the Keila–Oandu transition is a well-known biotic crisis (Meidla et al. 1999) following the Guttenberg carbon isotopic event (GICE, Ainsaar et al. 2010). Both dendrograms generated using the Raup–Crick and Simpson similarity coefficients (Fig. 4A, A1) indicate very clearly that during the Ordovician, the Estonian Shelf was inhabited by three different sequential faunas and the main faunal change occurred at the Keila–Oandu transition. In a cluster analysis of the global ranges of orthide and strophomenide taxa (Harper et al. 2009), the main partition occurred during the mid-Darriwilian, probably corresponding to the split in clusters between the Ontika and Purise subseries. In the East Baltic, this level (boundary between the Lower and Middle Ordovician by earlier interpretations; see Männil 1966) marks also a notable faunal renovation, however, it has not received enough attention in comparison with that at the Keila–Oandu boundary.
Cluster analyses of stratigraphical units based on brachiopod composition for the Lithuanian Shelf

The dendrogram, based on the brachiopod fauna from the Lithuanian Shelf (Fig. 5), differs from that of the Estonian Shelf due to contrasts in Ordovician brachiopod dynamics (see Fig. 3). The dendrogram generated using the Raup–Crick similarity coefficient (Fig. 5A) comprises three main clusters as in the previous case (Fig. 4). Several units (62–71) in the lowermost Ordovician are difficult to assign to clusters due to insufficient brachiopod data from these units. Only units characterized by the brachiopods of the Kunda RS (B3) (units 65–67) form a separate part of the cluster (Fig. 5).

Units 72–77 (Lasnamägi RS to the upper Haljala RS) form a cluster with variable similarities between units. They have low similarities with the overlying units, captured in another cluster comprising the units from the Keila RS to Rakvere RS. The close similarity of the brachiopod faunas of the Auleliai (78) and Vilučiai Fms (79) supports the Keila age of the former unit (Paškevičius 2016). Earlier it was included within the Haljala RS (Paškevičius 1997). The largest cluster, which comprises units from the Nabala RS (F1a) to Porkuni RS (F2) (units 83–90), is similar to the cluster of units on the Estonia Shelf. The dendrogram generated by the Simpson similarity coefficient (Fig. 5A1) comprises only two main clusters. Both dendrograms (Fig. 5) show that, in contrast with the Estonian Shelf, units of the Keila and Oandu RSs belong to one and the same cluster. The continuity in brachiopod distribution corresponds presumably to the stratigraphically more complete sections in Lithuania than in Estonia (Hints et al. 2016). In Lithuania, the boundary between the Rakvere (E) and Nabala (F1a) RSs appears more marked. The units of these stages belong to different clusters, whereas based on the Raup–Crick coefficient (Fig. 5A), the Keila–Rakvere units are closer to the uppermost strata, but based on the Simpson coefficient (Fig. 5A1), have closer relationship with older strata. The boundary between the Rakvere and Nabala RSs is biostratigraphically easy to recognize by the presence of the chitinozoan zonal species Armoricochitina reticulifera (Grahn) in the lowermost Mõntu or Paekna Fms (Männil & Meidla 1994; Nõlvak et al. 2007) of the Nabala RS.

On the periphery of the Lithuanian Shelf, in the northwestern part of Belarus (units 92–107), the key changes in the brachiopod faunal composition at the Keila–Oandu transition are more similar to those in the Estonian Shelf than the rest of the Lithuanian Shelf (Fig. 6). This is

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**Fig. 5.** Cluster analysis of the Ordovician stratigraphical units of the Lithuanian Shelf (in Lithuania). The dendrogram is compiled based on the Raup–Crick (A) and Simpson (A1) similarity coefficients and a neighbour-joining algorithm (Hammer & Harper 2006) using brachiopod data from Lithuania (Table S2). For indices of the regional stages and numbers of the units see Fig. 2. The regional subseries are given to the right of the dendrogram. P, Puikule Fm., Silurian.

**Fig. 6.** Cluster analysis of the Ordovician stratigraphical units using the data on brachiopods from Belarus (Ropot & Pushkin, 1987) (Table S3). The dendrogram is compiled based on the Raup–Crick (A) and Simpson (A1) similarity coefficients and a neighbour-joining algorithm (Hammer & Harper 2006). For indices of the regional stages and numbers of the units see Fig. 2. The regional subseries are given to the right of the dendrogram. K, Kohila Subseries; O, Ontika Subseries.
clear on both dendrograms generated based on the Raup–Crick and Simpson similarity coefficients (Fig. 6A, A1). The units (101–104) of the Oandu (D3) to Pirgu (F1c) RSs belong to the same large cluster highlighting the main faunal changes at the Keila–Oandu transition. The Livonian Shelf in Lithuania includes at some stratigraphical levels (Oandu–Rakvere) brachiopods, which occur also in the offshore part of the Estonian Shelf (Hints et al. 2016), again indicating somewhat deeper-water environments in Lithuania than in Estonia. The changes in shelf bathymetry are illustrated by the brachiopod communities in Lithuania (Paškevičius 2000), belonging to benthic associations BA1–2 during the Early and Middle Ordovician, to BA4 in the Oandu and down to BA5 in the late Katian.

Cluster analyses of stratigraphical units based on brachiopod composition for the Livonian Basin

The dendrograms displaying the faunal similarity between the lithostratigraphical units of the Livonian Basin are preliminary due to insufficient brachiopod data (Fig. 7). Like in northern Estonia, the Aseri RS (unit 46) belongs to the cluster together with units of older strata (clusters with units 42–46). The Pirgu RS (units 57, 58) and Porkuni RS (units 59, 60) are indicated on the dendrogram compiled by the Raup–Crick similarity coefficient as a separate cluster together with the lowermost Silurian (unit 61). The interval from the Kukruse RS (C2; unit 49) to Vormsi RS (F1a; unit 56) belongs to one cluster with variable levels of similarity between the units. The dendrogram based on data from the Livonian Basin displays unexpected and probably unreliable results due to insufficient baseline data.

COMPARISON OF THE BRACHIOPOD AND BRYOZOAN FAUNAS

Brachiopods are associated with diverse associations of bryozoans in the shelf facies. In Estonia, 275 Ordovician bryozoan species have been identified (Table S6). Their diversity trend through the Ordovician differs, though, from that of brachiopods (Fig. 3). Moreover, the total number of bryozoan species in several stratigraphical units exceeds that of brachiopods (Fig. 3). Many bryozoan species share several (up to 16) succeeding units (Table S5) increasing in diversity with the appearance of new species. The bryozoan diversity increases from the late Uhaku RS (C1c), being the highest in the Sandbian Haljala RS (C3–D1). In contrast to brachiopods, however, the diversity of bryozoans decreases towards the end of the Ordovician (Fig. 3), which follows the global trend in the evolution of the phylum (Taylor & Ernst 2004; Ernst 2018).

The cluster analysis of the stratigraphical units based on the bryozoan species composition (Fig. 8A) shows
The dendrogram generated from bryozoan data differs from that based on brachiopods in the step-by-step structure of the dendrogram. Such a structure is dependent on the large number of bryozoan species sharing many succeeding units (Table S6). The bryozoan data do not show strong faunal differences at the Keila–Oandu transition; a separate cluster comprises the units from the Keila RS to Rakvere RS. The Kukruse bryozoan fauna seems to be rather specific, as the corresponding units [Lower and Upper Viivikonna Fm. (15, 16)] belong to the cluster separated from the others.

The bryozoans from Belarus (Table S7; Fig. 8B), which are analysed at the stage level, are arranged on the dendrogram according to the groups of units corresponding to the regional subseries. They show essential differences between faunas at the Keila–Oandu boundary as revealed by the dendrogram based on the brachiopods from Estonia.

DEVELOPMENT OF THE BRACHIOPOD FAUNA IN THE EAST BALTIC

The brachiopod fauna of Baltica developed during the movement of the palaeocontinent from high to low latitudes, beginning with the early stages of the isolation of the Baltic Province and at the end of the Ordovician with its interface with the Kosov Province (Harper et al. 2013). The dominant brachiopods belong to Orthida and Strophomenida (Harper et al. 2004, 2017; Harper 2006; Curry & Brunton 2007). In the Livonian Basin, the leading factor in modification of the brachiopod faunal succession is the repeated and drastic changes of environments and the immigration of faunas adapted to new environments. This is related, for example, to the immigration of the elements of the deep-water Foliomena Fauna and the latest Ordovician Hirnantia brachiopod fauna in the Livonian Basin (Hints & Harper 2015; Harper & Hints 2016) together with the faunas of black shales and red-coloured deposits at different stratigraphical levels.

The brachiopod data summarized here, in total more than 400 species from the shelfal and basinal parts of the Baltic Basin (Tables S1–S5), display four intervals characterized by high diversities (Figs 3, 9C). These intervals in the Baltic Basin are correlated with global trends in the development of brachiopod faunas using the time slices of the Ordovician Period (Webby 2004) (Fig. 9B).

In Baltoscandia, the oldest rhynchonelliformean brachiopod (Apheoorthina daunus Walcott = Orthis christianiae Kjerulf) have been reported from the Tremadocian Ceratopyge Limestone in Sweden (Walcott 1912; Rubel 1961). In the East Baltic, this species together with the oldest representatives of Plectella, Panderina and Paurorthis (unpublished data of Ralf Männil) occurs in the Hunnebergian RS (B1a) of Lithuania, which corresponds to time slices 1c–2a. In the Estonian Shelf, the brachiopod diversity is high in the Volkov (B2) and Kunda (B3) RSs corresponding to time slices 3 and 4.
diverse biota. That circumstance and the practical richness deposits contributed to the development of a highly region, unusual environments with kukersite kerogen-caused by essentially environmental differences, occurs pod diversity of the Uhaku (C1c) and Kukruse (C2) RSs, in a regional context a marked difference in the brachiopod faunas, especially of plectambonitoids and strophomenoids (Harper et al. 2004, 2013; Rasmussen et al. 2007, 2016). In a regional context a marked difference in the brachiopod diversity of the Uhaku (C1c) and Kukruse (C2) RSs, caused by essentially environmental differences, occurs between the Lithuanian and Estonian shelves. In the latter region, unusual environments with kukersite kerogen-rich deposits contributed to the development of a highly diverse biota. That circumstance and the practical importance of kukersite kerogen as a raw material invigorated the studies of brachiopods (Bekker 1921, 1924; Õpik 1928, 1930, 1934). The contemporaneous pack- and wackestones on the Lithuanian Shelf were not especially fossiliferous and the drill core material was inadequate for detailed research.

In the Estonian Shelf, the third brachiopod diversity high occurs on the Sandbian–Katian transition (Keila RS) at the beginning of time slice 5. In contrast with the Estonian Shelf, a diverse brachiopod fauna on the Lithuanian Shelf is found only in the Sandbian and lower Katian (Haljala to Oandu RSs of time slices 5b, 5c; Figs 3A, 9B). The number of species in separate formations of these stages varies within limits close to those in Estonia (Fig. 3A). However, the species diversity drops beginning with the late Katian Vormsi RS towards the end of the Ordovician. The brachiopod dynamics in the Belarus part of the Estonian Shelf (Fig. 3B) differs from that in Lithuania (Fig. 3B) in having a diversity peak at the Haljala RS (C3-D1), which follows a decreasing trend in biodiversity.

The fourth brachiopod diversity rise on the Estonian Shelf occurs during the late Katian Vormsi (F1b) and Pirgu (F1c) RSs, which correspond to time slices 6a, 6b (Fig. 9B). Some diversity decrease is observed in the brachiopod associations of the Tudulinna (unit 32, Fig. 2) and Halliku (unit 34, Fig. 2) Fms that were deposited in the somewhat deeper-water environments of the outer part of the Estonian Shelf. The brachiopod diversity drop at the end of the Ordovician is noticeable on shelves (Fig. 3) and continuous transition to the Silurian is interrupted by gaps. The earliest Silurian (unit 41) brachiopod fauna has 13 genera and one species common with the Ordovician.

In the Livonian Basin, few plectambonitoid brachiopods have been identified among the trilobite-dominated fauna (Männil 1963) of the red-coloured deposits of the Early and most part of the Middle Ordovician (Hunneberg to Aseri RSs). In the Sandbian and early Katian, the brachiopods were represented mainly by small plectambonitoids and orthides, characteristic of the Livonian Basin (Fig. 3). The black shales in two stratigraphical levels of time slices 5c (Keila? RS) and 5d (Vormsi RS) comprise specific associations similar to the Chonetoidea/Sericoida–lingulid association in Norway (Hansen 2008) and in other regions as the deepest association of sowerbyellid–strophomenid–dalmanellid fauna along the onshore–offshore transect (Jaanusson 1984). The red-coloured Katian lithologies of the Pirgu RS (6a, b) comprise brachiopods in common with those of the Jonstorp Fm. in Sweden [Sowerbyella (Rugosowerbyella) rosettana] (Jaanusson 1982).

**BIOSTRATIGRAPHICAL IMPORTANCE OF BRACHIOPODS IN THE EAST BALTIC**

The stratigraphical units from the Estonian Shelf are divided into three main groups (clusters) based on brachiopod composition (Fig. 9). This suggests the occurrence of three evolutionary faunas in the Baltic Basin. One cluster connects the units of the Onitka Subseries (Hunneberg–Kunda RSs, the Lower Ordovician and lower part of the Middle Ordovician). The second cluster comprises the Purste and Kurna subseries (Lasnamägi–Keila RSs) forming the most part of the Viru Series (Fig. 2). The position of the Aseri RS is somewhat unclear. In some cases it is tied with the older, in some cases with the overlying units. The largest cluster comprises all units from the Oandu RS up to the top of the Ordovician (=Vinni subseries of the uppermost Viru Series, and the Kohila and Atla subseries of the Harju Series). The dendrograms generated from the Estonian and Belarus brachiopod data (Figs 4, 6) indicate that the uppermost units of the Viru Series (Vinni Subseries; Oandu and Rakvere RSs) are aligned biostratigraphically more closely with the units of the Harju Series.

The dendrogram based on the Lithuanian data also delineates three main clusters, but these are slightly different from those generated from the Estonian data. Specifically, the Haljala–Oandu interval is characterized by a high brachiopod diversity on the shelf. The dendrogram based on data from the periphery of the Lithuanian Shelf in Belarus (Fig. 6) corresponds well
to that generated from Estonian data. The units in the interval from the Billingen RS (B1) to the Keila RS (D2) and from the Oandu RS (D3) to the Porkuni RS (F2) belong to different clusters.

The dendrogram compiled using data from the Livonian Basin is preliminary. Insufficient data and stratigraphic problems concerning the correlation of some units (Blidene and Mossen Fms) render a few clusters unreliable. The Aseri RS (C1a) possibly requires additional investigation. In all the dendrograms its corresponding units have more similarity with the older rather than with the younger units. At least part of the Aseri RS shares the *Eoplacognathus pseudoplanus* conodont Zone with the Kunda Stage (Nõlvak et al. 2007).

In spite of the differences in faunal dynamics and the regional stratigraphy of the Ordovician sections in the East Baltic, the dendrograms based on the Raup–Crick and Simpson similarity coefficients contain clusters comprising the units of the subseries (Fig. 9A, A1). An exception is the Aseri RS mentioned above, whose stratigraphic position requires further investigation. The position of the Keila RS in the cluster together with the Oandu and Rakvere RSs (Fig. 5) is presumably driven by the wider stratigraphical distribution of several species in the Keila–Oandu interval in Lithuania or it could indicate problems with regional correlations.

The present study shows that cluster analysis, a robust method for processing multivariate data (Hammer & Harper 2006), gives useful results not only for the global analysis of faunas but also in regional studies of stratigraphy and the vertical distribution of different faunal groups.

The development of the brachiopod fauna of the Baltic Basin indicates that the levels of main changes in its composition do not coincide with the levels of the boundaries of the global chronostratigraphical units of the Early, Middle and Late Ordovician. Nevertheless, the trends in the diversification of the Baltic brachiopods follow the global trends. The oldest brachiopod fauna prior to the Kunda RS (B3) characterizes the interval of replacement of siliciclastic deposits with carbonate lithologies. The upper part (Oandu and Rakvere RSs) of the Viru Regional Series (upper Darrwilian–early Katian) comprises a brachiopod fauna more closely related to succeeding Late Ordovician faunas.

CONCLUSIONS

1. More than 500 taxa of Ordovician rhynchonelliformean brachiopods occur in the 106 environmentally distinct lithostratigraphical units of the East Baltic. Based on the similarity in the composition of brachiopods, the stratigraphical units are partitioned into three clusters: (1) the units of the Ontika Regional Subseries, (2) the units of the Purtse and Kurna subseries and (3) the units of the Vinni, Kohila and Atla subseries. Dendrograms for different parts of the region show some specific deviations.

2. Short stratigraphical ranges of species delineate the stratigraphical units of different categories up to the stages and subseries levels.

3. Bryozoans appear to have less biostratigraphical value than brachiopods, mainly due to the longer stratigraphical ranges of many species.

4. The faunal crisis at the boundary between the Keila and Oandu RSs is well documented on the dendrograms. The shelfal brachiopod faunas of the upper part of the Viru Regional Series (the Vinni Subseries) are much closer to those of the Harju Series than to the rest part of the Viru Series. This crisis is less pronounced in the basinal facies.

5. The faunal changes on the transition between the Ontika and Purtse regional series needs further investigation.

6. Cluster analysis using two similarity coefficients is a useful tool that captures the similarity of different stratigraphical units based on their faunal composition. This analysis validates the stratigraphical charts and highlights the biostratigraphical role of the rhynchonelliformean brachiopods and global trends in their evolution.

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Supplementary online data

Supplementary online data associated with this article can be found at http://doi.org/10.15152/GEO.233. These include the stratigraphical distribution of rhynchonelliformean brachiopods in the Estonian Shelf (Table S1), Lithuanian Shelf (Table S2), Belarus (Table S3), Livonian Basin (Table S4), the stratigraphical distribution of all brachiopods in the East Baltic part of the Baltic Basin (Table S5), and the stratigraphical distribution of the bryozoans in the Estonian Shelf (Table S6) and in Belarus (Table S7).
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University of Kansas, Boulder, Colorado, and Lawrence,
Kansas.
Brahhiopoodide mitmekesisuse muutumine ja biostratigraafiline tähtsus Balti Ordoviitsiumis

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On esitatud andmed ligi 500 brahhiopoodi taksoni levikust 106 Ordoviitsiumi–Siluri läbilõike erinevas stratigraafilises üksuses (lademes, alamlademes, kihistus või kihi stikus), mis on eristatud Baltikumi eri piirkondades (Eestis, Lätis, Leedus ja Valgevenes). On analüüsitud brahhiopoodide taksonoomilise koosseisu muutusi ajas ja võrreldud seda sammalloomade koosseisu muutustega. Töös on kasutatud varem publitseeritud, kuid taksonoomiliselt ja stratigraafiliselt korrigeeritud ning uusi andmeid liikide leviku kohta (on esitatud lisadena seitsmes tabelis). Tänapäeval on makrofossiilide, sealhulgas brahhiopoodide kasutatavus biostratigraafias oluliselt vähenedud, kuivõrd läbilõige korreleerimiseks ja biostratigraafiliste piiride määramiseks on hõlpsam rakendada erinevaid mikrofossiile. Käesoleva töö eesmärgiks on näidata brahhiopoodide kasutatavust biostratigraaffias, tuginedes nende liigilise koosseisu erinevate stratigraafiliste üksustes. Selleks on kasutatud Hammeri ja Harperi (2006) väljatöötatud Past3 programmi kobaranalüüsi. Brahhiopoodide leviku põhjal dendrogrammes esinevad erinevad seoseid seina-sisese, väljaspoolse, seda paljude uusi andmeid ja võrdlemist. Dendrogrammid näitavad erinevate stratigraafiliste üksuste või kivimkehade faunade erinevusi ajas ja erinevate stratigraafiliste üksuste või kivimkehade faunade uurimisel. Võimalus on esitada näitlikud seoseid, mis võimaldavad otsida kasutatava pandeega, mis näitavad kasutatava pandeega. Vaid toovad esile ka võimalikke revisjoni või edastatist uurimist aspekte biostratigraafias.