The Middle Miocene of the Fore-Carpathian Basin
(Poland, Ukraine and Moldova)

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ABSTRACT:


Studies of Miocene sediments in the Fore-Carpathian Basin, conducted by geologists from the University of Warsaw have provided new insights on the distribution of the facies infilling the basin, particularly in the forebulge and back-bulge zones. The origin of the large-scale sand bodies, evaporitic deposits and large-scale organic buildups is discussed, described and verified. These deposits originated in variable, shallow marine settings, differing in their water chemistry and the dynamics of sedimentary processes, and are unique with regard to the fossil assemblages they yield. Many years of taxonomic, biostratigraphic, palaeoecologic and ecotaphonomic investigations have resulted in the identification of the fossil assemblages of these sediments, their age, sedimentary settings and post-mortem conditions. Detailed studies were focused on corals, polychaetes, most classes of molluscs, crustaceans, echinoderms, and fishes.

Key words: Lithofacies; Fossil assemblages; Middle Miocene; Fore-Carpathian Basin; Poland; Ukraine.

INTRODUCTION (A. Radwański, A. Wysocka and M. Górka)

This paper reviews and updates the investigations of researchers from the Faculty of Geology in Warsaw on the mid-Miocene of the Fore-Carpathian Basin in southern Poland, western Ukraine and Moldova. In palaeogeographic terms, the studied area belonged to the northern compartment of the Central Paratethys that extended, in mid-Miocene times, from Moravia, through Poland and Ukraine, to Moldova and Romania (Text-fig. 1A, B).

The investigations into the Miocene of the Fore-Carpathian Basin began at the end of the 18th century with the paper by Jaśkiewicz (1787), who reported molluscan shells from the Korytnica Clays, and concluded on their marine origin. The palaeontological assemblages have been studied gradually, until the 1920’s, when the period of more intense work on faunal assemblages of the Miocene, which lasts till the present, commenced (see chapter ‘Overview of selected taxonomic groups’). The palaeontological groups treated in the literature range from leaf flora and microfossils, through numerous invertebrates, to large marine mammals. Not only the taxonomic composition but also the interspecific relations (e.g., symbiosis, commensalism, parasitism) and the eco-taphonomic history among the described organisms have been reported.

Besides their stratigraphic importance, the numerous taxa and/or organic assemblages appear to be
The Fore-Carpathian Basin is defined herein as a palaeostructural term, a circum Carpathian basin, in which the Miocene marine domain is usually treated as a basin. It has been named and understood, however, either as the Fore-Carpathian Basin, or the Carpathian Foredeep. These two terms, in various papers, are treated either as synonymous or as distinct units. Short comments to both of these terms are given below.

The Miocene Fore-Carpathian marine domain is usually treated as a basin. It has been named and understood, however, either as the Fore-Carpathian Basin, or the Carpathian Foredeep. These two terms, in various papers, are treated either as synonymous or as distinct units. Short comments to both of these terms are given below.

The present account is organized as a series of separate chapters (or subchapters), prepared by different authors, to present selected aspects of the lithofacies, evolution, palaeontology and palaeoecology of the Fore-Carpathian Basin.

**FORE-CARPATHIAN BASIN VS. CARPATHIAN FOREDEEP (A. Radwański and A. Wysocka)**

The Miocene Fore-Carpathian marine domain is usually treated as a basin. It has been named and understood, however, either as the Fore-Carpathian Basin, or the Carpathian Foredeep. These two terms, in various papers, are treated either as synonymous or as distinct units. Short comments to both of these terms are given below.

The Fore-Carpathian Basin is defined herein as a palaeogeographic unit, a Middle Miocene marine domain whose water stretched out from the Carpathian margin. In Poland, it ranged as far as the belt of the present-day Central Polish Uplands. In Ukraine, it reached to the south-eastern margin of the Podolian Platform (a part of the East-European Craton) (Text-fig. 1B). The basin was underlain by topographically diversified land, and characterized by variable depths.

On the contrary, the Carpathian Foredeep is a geotectonic term, a circum Carpathian basin, in which the sediments, marine or non-marine, accumulated due to geotectonic subsidence of its basement (Text-fig. 1B). The sediments may have attained remarkable thicknesses masking completely the topography of the substratum. In the latter case, the basin could have been facies-uniform. So defined, the Carpathian Foredeep corresponds to the Outer Foredeep sensu Oszczypko (2006, see also Oszczypko and Oszczypko-Clowes 2011).

In terms of the nomenclature used for the foreland basin system (after DeCelles and Giles 1996), a foreland basin system is defined as consisting of: (1) a wedge-top, (2) foredeep, (3) forebulge, and (4) backbulge depozones. The type and thickness of sediments occupying these depozones depends on their location at the time of deposition, rather than their geometrical relationship with the thrust belt (DeCelles and Giles 1996). During the latest stages of the Carpathian thrust-and-fold belt development, the Fore-Carpathian foredeep depozone was formed as a result of the immense load of the growing nappes stacking one on the other, moving outwardly from the depocentre. This caused a high subsidence rate just in front of the orogenic front with the progressive movement of the depocentre to the north. The Carpathian forebulge belt was formed as an isostatic response to the subsidence in the foredeep, fringing the foredeep well north of the Carpathian front. In recent topography, the Carpathian forebulge is marked by the Upper Silesian High, Cracow Upland, Miechów Upland, the Holy Cross Mountains (Text-fig. 1B) (e.g., Kutek and Glazek 1972; Radwański 1974; Marcinowski and Radwański 1983), and the Lublin Upland with the Roztoce Hills ranging as far as Lviv in the Ukraine (Text-fig. 1B) (Wysocka 2002, 2006; Jankowski and Margielewski 2015). The situation in the Ukrainian and Moldovian part of the fore-Carpathian region is less clear, as no distinct forebulge zone may be recognised (although it may be supposed that the Medobory Hills could have originated as such). Consequently, the vast area covered by Miocene deposits outside the Fore-Carpathian foredeep and forebulge depozones, should be treated as the Fore-Carpathian back-bulge basin (Text-fig. 1B). According to DeCelles and Giles (1996), this zone should be characterised by shallow water sediments accumulated in a broad zone of flexural subsidence cratonward of the forebulge. The Fore-Carpathian forebulge depozone is expected to be a site of local unconformities, condensations, stratal thinning, fault-controlled depocentres, as well as synsedimentary and postdepositional block faulting, e.g., in the sandy sequences of the Opole Minor in the Ukraine (Pazdro 1953; Wysocka et al. 2012, fig. 2).
For almost two centuries, the geological record of the Fore-Carpathian Basin has attracted geological studies (e.g., Staszic 1815; Alth 1850; Hilber 1882; Lomnicki 1897, 1898; Teisseyre 1900; Nowak 1938; Kudrin 1966; Ney et al. 1974). Numerous new geological and geophysical studies, related to prospecting for hydrocarbon sources, started recently (e.g., Oszczypko 1996, 2006; Krzywiec 1999; Dziadzio 2000; Porębski et al. 2002; Krzywiec et al. 2005, 2008; Mastalerz et al. 2006; Popadyuk et al. 2006; Ślączka et al. 2006; Pietsch et al. 2010; Warchoł 2011; Lis and Wysocka 2012; Gozhyk et al. 2015).

The Miocene succession of the Fore-Carpathian Basin starts with transgressive siliciclastics, occasionally with thin coaly interbeds, clays and marls of Karpian (Garecka and Jugowiec 1999) or Early Badenian age (Text-fig. 2). Typical transgressive lithostratigraphic units are the Baranów (Baraniv in Ukraine) Beds as well as Skawina and Pińczów Formations. These shallow-water, high-energy deposits do not exceed 100 m in thickness (e.g., Wysocka 2002; Mastalerz et al. 2006), are overlain by regionally persistent evaporitic beds (Text-fig. 2) and referred to as the Wieliczka and Krzyżanowice formations (Tyras suite in Ukraine). They correspond to the lower part of nannoplankton zone NN6 (Peryt and Peryt 1994). The relatively low basin diversity marked by the evaporitic beds suggests the intermittent lack of basin diversification into the foredeep, forebulge, and back-bulge depozones. The return to the basin subdivision into the three zones is well revealed by the supra-evaporitic deposits of the Machów Formation (Dashava Fm. in Ukraine), exceeding 2000 m in thickness in the axial zone of the foredeep (over 5000 m in the Ukrainian foredeep) (Text-fig. 3; and Table 1).

The large-scale sandy-bodies, evaporitic beds and large-scale carbonate build-ups of the forebulge and back-bulge depozones of the Fore-Carpathian Basin are discussed in the following chapters.
Large-scale sandy-bodies of the forebulge and back-bulge depozones of the Fore-Carpathian Basin (A. Wysocka and A. Radwański)

The Middle Miocene sandy and organodetrital deposits of the Roztocze, Opole Minor and Ternopil area cover c. 45,000 km². Recently, these deposits have been studied intensively (see e.g., Radwański and Wysocka 2001; Wysocka 2002; Górska et al. 2006; Radwański et al. 2006; Wysocka and Jasionowski 2006; Górska et al. 2012; Peryt et al. 2012a; Radwański et al. 2012; Wysocka et al. 2012; Radwański et al. 2014, and Table 1), and the results are shortly summarized below.

The sandy and organodetrital deposits are easily accessible in huge quarries and sand-pits revealing their 3-D facies geometry (e.g., Radwański and Wysocka 2001; Wysocka et al. 2012; Radwański et al. 2014). However, their facies interpretation and correlation are insufficient because of the lack of new and reliable cartographic and stratigraphic data. Consequently, there is no single general stratigraphic scheme for the entire area of the Fore-Carpathian Basin (Text-figs 2 and 3). Based on published schemes, the investigated lithofacies are correlated with the Lower Badenian Baraniv, Mykolayiv, and Naraviv beds, and with the Upper Badenian Pidhirtsi, Ternopil, and Buhliv beds, as well as with the Lower Sarmatian sands (Text-fig. 3). The large-scale sandy-bodies of the forebulge and backbulge depozones of the Fore-Carpathian Basin are individualized in the Roztocze and Opole Minor (Lviw area) and in the Ternopil area (Text-fig. 3). In these areas, several tens of exposures show a variability of sedimentary structures and a diverse biotic content (Text-fig. 3 and Table 2).

The sandy sequence of the Roztocze and Opole Minor (the so-called Mykolayiv Sands, see Wysocka et al. 2012) extends along the south-western margin of the East European Craton and rests upon a Laramide basement of Upper Cretaceous (Maastrichtian) marls, the topography of which was determined by Palaeogene erosion and denudation. The resultant topography, having been locally exposed today, allows the estimation of the pre-Miocene morphological denivelations (Pazdro 1953; Wysocka et al. 2012). All have been filled more or less completely with a mass of sands, the thickness of which ranges, consequently, from a dozen metres to nearly nil. The petrographic content of these sands is very monotonous, as they are composed of quartz, with a variable but minor admixture of glauconite, clay minerals, as well as calcareous dust and bioclasts. All bioclasts are calcitic, as the aragonitic ones have been completely leached out during diageneis. Typically, the larger bioclasts are heavily armoured with sand grains due to pressure-solution processes (Radwański and Wysocka 2001; Radwański et al. 2012).

A big variety of different sedimentary structures is typical of this sandy-sequence. Within the particular sets, both depositional and deformational structures of variable scale are present (Table 2), and are commonly burrowed (Text-fig. 4) (Wysocka 2002; Wysocka et al. 2012). These sandy bodies are interpreted as migrating megaripples of various kinds, sand bars, slumps, and/or delta-slope bodies. An uneven bottom morphology controlled the current directions (Wysocka 2002) until this

<table>
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<th>Section</th>
<th>Most important bibliography, including older reference data</th>
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<td>Radwański 1969, 1973</td>
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<td>Smerdyna</td>
<td>Rutkowski 1976; Laptas 1992; Roniewicz and Wysocka 2001; Leszczyński and Nemec 2015</td>
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<td>Nawodzice</td>
<td>Baluk and Radwański 1968; Nosowska 1997</td>
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<td>Świniary</td>
<td>Kowalewski 1929; Blaszk 1965; Pawłowski 1965; Radwański 1973; Kenig and Wysocka 1996; Radwański and Wysocka 2004</td>
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<td>Korytnica</td>
<td>Baluk 1974; Baluk and Radwański 1977, 1984a, b</td>
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<td>Opatówka Valley</td>
<td>Czapowski 1976; Czarnecka 2011</td>
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<td>Pińczów-Busko area</td>
<td>Radwański 1969; Studencki 1988a, b</td>
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<td>Chmielnik region</td>
<td>Rutkowski 1976; Czapowski and Studencka 1990; Leszczyński and Nemec 2015</td>
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<td>Gleboviti</td>
<td>Radwański and Wysocka 2001; Wysocka 2002; Radwański and Wysocka 2006</td>
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<td>Strach</td>
<td>Lomnicki 1999; Wysocka 2002; Wysocka and Jasionowski 2006</td>
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<td>Lozyna</td>
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<td>Birky</td>
<td>Wysocka 2002</td>
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<td>Yasnyska</td>
<td>Wysocka 2002</td>
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<td>Sykhiv</td>
<td>Teisseyre 1938; Wysocka 2002</td>
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<tr>
<td>Pidjarkiv</td>
<td>Jahn 1937, Malicki and Jahn 1937; Szórényi 1953</td>
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Table 1. Summary of selected published geological data on the sandy and organodetrinitic lithofacies in the forebulge and backbulge depozones of the Fore-Carpathian Basin
Topography was smoothed by sedimentation to form an extensive offshore zone. As long as the topographic relief remained, slumping and liquefaction, or even seismic shocks, were locally important to produce homogenized sand bodies up to 15 m thick (Wysocka 2002; Wysocka et al. 2012). Later, the sand bodies were influenced by storm agents (with scouring and residual lags), or by waves and currents, to produce sand bars of variable heights (up to 4, rarely to 10–15 m).

A poorly known, large-scale sand-body is exposed in the Ternopil area, in the huge Mohila sand-pit and the small, unexplored Lozova sand-pit. It occurs to the south west of the Medobory reefal-complex and is separated from the back-reef sandy facies (Jasionowski et al. 2006). In the Ternopil area, the sandy-complex is characterised by a progradational set of large-scale tabular cross-stratification. It is covered by a sandy unit with structures indicative of extremely shallow-water environments (Wysocka et al. 2006). Another unusual feature that has attracted attention in the Lozova section is a large-scale landslide of biogenic material, transported from the Medobory reefal-complex (Wysocka et al. 2002).

Text-fig. 3. Position of the studied sections in Ukraine against the regional (as used traditionally) and standard zonations (compiled from Petryczenko et al. 1994, Andreyeva-Gregorovich et al. 2008 and Górka et al. 2012)
Text-fig. 4. Selected sedimentary features of the large-scale sedimentary complexes of the Roztocze, Opole Minor and Podolia; for symbol explanations see Table 2. A – progradational set of large-scale tabular cross-stratification (Smerdyna, PL); B – progradational set of large-scale tabular cross-stratification, downlapping on plane-stratified sands (Sykhiv, UA); C – large-scale channel-like structures (Dch, underline by dashed line) cut in ripple cross-stratified calcarenites (Dr, p) (Józefów, PL); D – large-scale cross-section of a giant scour (underline by dashed line) filled with structureless sands (Hlukhivets, UA); E – cosets of ripple, trough and tabular cross-stratified calcarenites (Pardysówka, PL); F – channel-like infills (base out-lined) cut into cosets of tabular and trough cross-stratified sands (Khorosno, UA); G – cosets of synsedimentary deformed layers (Khorosno, UA); H – strongly bioturbated, ripple cross-stratified sands (Dr) with numerous burrows of callianassid decapods (Bb) (Romanniv, UA)
socka et al. 2006), pointing to a high bottom relief during sedimentation and the isochronity of sandy and reefal facies in the area (Text-fig. 3). To the north east of the Medobory reefal-complex, a sandy sequence is exposed e.g., in the Ohryzkovtsi section, a very remarkable section (Jasionowski et al. 2006) that is terminated by oolitic sands. Undoubtedly, the sandy sequence in the Ternopil area should be intensively investigated in the future.

All sandy bodies discussed above are intriguing because of the huge mass of pure quartz-sands lying on a nonclastic basement and representing a regional accretion prism which, in the Ukrainian part of the Fore-Carpathian Basin, had carpeted the pre-Miocene topography at the margin of the East-European Craton, and having been gradually downthrown south-westerly towards the foredeep depozone. This prism had been dependant on the pre-Miocene topography of the region, the synsedimentary tectonics, as well as the dynamic conditions of the transgressive Middle Miocene (Badenian) overflood. The supply of a huge, terrestrial sand-mass is inferred to have successively been continued from the Palaeogene wastes of the East-European hinterland of Podolia and Volhynia (Wysocka et al. 2011; Radwański et al. 2014). The mode of its delivery remains as yet unknown: neither fluviatile or aeolian transport, nor deltaic spread may be recognised by the sedimentary structure of the sand sequences studied. All display a variety of structures typical of shallow- to very shallow-marine, high-energy bottom conditions.

Summing up, the lateral and vertical facies succession in the vast area of the forebulge and back-bulge depozones of the Fore-Carpathian Basin point to a full transgressive-regressive cycle, controlled by the evolution of the Carpathians during the Badenian-Sarmatian times. The problems of detailed correlation as well as the synchronity/diachronity of the particular facies still remain unsolved. The next fascinating and unsolved problem is the influence of the Middle Miocene Climatic Optimum, followed by the subsequent drastic cooling, on the facies pattern as well as on the biotic diversity (for preliminary discussion see Wysocka et al. 2012 and Radwański et al. 2014).

**Badenian evaporites** [Maciej Bąbel]

The evaporitic event that took place in the Central Paratethys in Badenian time has been referred to as the Badenian salinity crisis by analogy to the famous Messinian crisis in the Mediterranean area (Peryt 2002, 2006a). The widespread deposition of Ca-sulphates, Na-chlorides and, locally, K Mg salts took place at that time in several apparently interconnected basins or areas: the Fore-Carpathian Basin, the Slovak Basin, the Intra-Carpathian Basin and the Transylvanian Basin (Liszkowski 1989; Peryt 2006a) (Text-fig. 1C). Among these basins, the Fore-Carpathian Basin was the largest, and its northern part is the best studied. In this area, the primary evaporite gypsum deposits are excellently exposed, particularly in the Ukraine. The halite facies as well as a large part of the carbonate facies with native sulphur deposits are hidden in the subsurface (Aleksenko 1967; Gąsiewicz 2000b; Garlicki 2008; Bukowski 2011).

This short review attempts to present the main new ideas concerning evaporite sedimentology and stratigraphy developed during the last few decades of studies conducted in the northern part of the Fore-Carpathian Basin, mainly in the gypsum depositional zone. Great progress in the studies of Badenian evaporites was initiated by the publications by Kwiatkowski (1972) and Garlicki (1979). The halite facies are only briefly mentioned, and the extensive geochemical and isotopic studies are not reviewed at all.

**Analyses of the Badenian gypsum facies**

At the end of the last century, an older view that the large gypsum crystals occurring in the Badenian deposits are a primary evaporite sediment (Kreutz 1925; Gawel 1955), was fully established (Bąbel 1986, 1987; Kasprzyk 1991, 1993a,b; Peryt 1996; Petrichenko et al. 1997). At that time it was also proved that the large (several metres in size; Text-fig. 5A) domal structures composed of such gypsum crystals, recorded in the Badenian, are also primary structures that formed on the evaporite basin floor (J. Tokarski in Flis 1954, p. 21; Koltun and Roskosh 1969; Bąbel 1986, 2005b, 2007a; Kasprzyk 1993a, b; Kasprzyk et al. 1999; Pokalyuk et al. 2009).

Deposits composed of large primary gypsum crystals (> 2 mm in size) are called selenites (Warren 1982; Bąbel 2004a) and represent a significant part of the Badenian sulphate evaporites. Application of the facies analysis methodology to these and other primary deposits in the Fore-Carpathian Basin (Polish part) has revealed a novel and more detailed picture of evaporitic environments in the marginal zone of the basin (Kasprzyk 1991, 1993a,b; Bąbel 1999a,b, 2005b). Further sedimentological and facies studies conducted in the Ukraine, Moldova and in the Czech Republic (Peryt 1996, 2001; Pery et al. 1997; Petrichenko et al. 2006).
### Depositional Structures

<table>
<thead>
<tr>
<th>Structures</th>
<th>Characteristics</th>
<th>Interpretation</th>
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<tbody>
<tr>
<td>Giant-scale high-angle foreset stratification $D_{ox}$</td>
<td>Medium- to coarse-grained sands, occasionally gravels; average thickness of single set about 4-10 m</td>
<td>Fault scarp-attached bar formed by the avalanching of littoral sediments</td>
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<tr>
<td>Large-scale tabular cross-stratification, occasionally in form of cliniforms $D_x$</td>
<td>Medium- to coarse-grained sands; average thickness of single set about 2-4 m, maximum up to 10-15 m; commonly of diagonal and tangential type</td>
<td>Migration of straight-crested bars or progradation of a delta slope; high-energy shallow-water environment</td>
</tr>
<tr>
<td>Trough cross-stratification $D_t$</td>
<td>Medium- to coarse-grained sands; set thickness from 10 to 50 cm</td>
<td>Migration of curved-crested ripples; high-energy shallow-water environment</td>
</tr>
<tr>
<td>Ripple cross-lamination $D_r$</td>
<td>Fine-grained sands; heights of individual ripples range between 2 and 10 cm; dominate asymmetrical and climbing (ripple laminae-in-phase and ripple laminae-in-drift) ripples, occasionally wavy and flaser lamination</td>
<td>Migration of wave and wave-formed current ripples; shallow-water environment, (?) up to intertidals</td>
</tr>
<tr>
<td>Plane bedding $D_p$</td>
<td>Various sands; thickness of sets rarely exceeds 10 cm</td>
<td>Lower- or upper-stage plane beds; low- to high-energy shallow-water environment</td>
</tr>
<tr>
<td>Large-scale scours $D_s$</td>
<td>Large-scale elongated scours filled with fine-grained, well-sorted sands; depth up to 10 m, widths range up to 20 m; concave sharp and erosional bottom surface; basal part massive, top parts with traces of lamination and numerous fold and diapir deformations</td>
<td>Grain flow in the form of underwater slump-scours; sediment movement caused by overloading of slope, rapid drawdown and destruction of frontal part of a delta body, and/or by seismic shock</td>
</tr>
<tr>
<td>Channel-like fills $D_{ch}$</td>
<td>Channel-like elongated forms filled with massive fine- to coarse-grained sands and/or organodetrital material; sharp concave lower boundaries, depth up to 3 m, width to 5 m</td>
<td>Storm surge channels, locally filled by lag deposits</td>
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### Biogenic Structures

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<tr>
<th>Structures</th>
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<tr>
<td>Individualised burrows $B_b$</td>
<td>Syn- or post-depositional activity of echinoderms (echinoids and sea-stars), shrimps and crabs, as well as undetermined sea-anemones, bivalves, and holothurians</td>
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<tr>
<td>Maze of ‘burrow-ghosts’ of various size, totally blurring primary depositional structures $B_m$</td>
<td>Total destruction of depositional structures, formed by long-termed activity of over-populated mass-abundant bottom dwellers</td>
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### Deformational Structures

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<th>Structures</th>
<th>Characteristics</th>
<th>Interpretation</th>
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<tr>
<td>Slump (fold and diapir) structures $F_f$</td>
<td>Highest parts of scour in fills, vertical or overturned anticlines and synclines, commonly detached, with strongly deformed bend zones; up to 2 m height</td>
<td>Caused by collapse of large-scale bars or delta slopes and/or by seismic shock</td>
</tr>
<tr>
<td>Convolute bedding $F_c$</td>
<td>Small-scale fold deformations underlain or overlain by non-deformed deposits; deformed layer thickness up to 30 cm; anticlinal parts are steep and chevron-like, synclinal are flat and blunt</td>
<td>Result of reversed density gradient or sediment liquefaction; caused by overburden by overlying deposits, seismic shock and/or movement of liquefied sediment</td>
</tr>
<tr>
<td>Pseudonodule structures $F_p$</td>
<td>Disturbed bedding in form of ovate or spherical masses surrounded by massive sands</td>
<td>Fluid-loss structures, produced by escaping water, breaking upward through a primary lamination; caused by gravity mass movement or deformational shear of currents</td>
</tr>
<tr>
<td>Structureless $F_s$</td>
<td>Sands completely devoid of any sedimentary structures; bodies up to 15 m thick</td>
<td>Homogenisation caused by rapid sedimentation, total bioturbation, redeposition, and/or liquefaction</td>
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1997; Bąbel 2005b, 2007b) have shown that at the stage of Ca-sulphate deposition the evaporite basin was a very shallow, flat and broad platform occupied by a system of more or less (5-20 m) deep saline pans, dominated by the deposition of coarse and giant-crystalline selenites, and semi-emerged evaporite shoals where the selenite deposition was associated with fine-grained gypsum precipitation, as well as with the formation of gypsum microbialite and/or stromatolite deposits.

The Badenian gypsum facies were shown to be very similar to the famous Messinian gypsum facies (e.g., Lugli et al., 2010), with the exception of some facies, which are very specific and typical of the Badenian. They include the giant gypsum intergrowths facies (Text-fig. 5B, C; Bąbel 1987, 1999a, b; Peryt 1996), never recorded outside of the Badenian basins and widespread microbialite and stromatolite facies (Text-fig. 5D, Kwiatkowski 1970, 1972; Kasprzyk 1993c; Peryt 1996; Bąbel 2005b, 2007b; Bąbel et al.

Text-fig. 5. Gypsum facies. A – primary selenite dome (Skoviatyn Quarry, UA); B – giant gypsum intergrowths facies (Zhalybory, UA); C – giant gypsum intergrowths facies (Gacki, PL); D – gypsum microbialite facies passing upward into the grass-like selenite facies (Pisky Quarry, UA); E – sabre gypsum facies with conformably oriented selenite crystals (Ustechko, UA)
Selenite facies – controlling factors

The selenite facies are very specific. They are created by the in situ growth of gypsum crystals on the bottom of the evaporite basin, and hence are classified as autochthonous evaporites (Vai and Ricci-Lucchi 1977). Although the mechanism of sediment accretion is clear and seemingly simple (Bąbel 2007a, b), the selenite deposits display a great variability of facies, sometimes drastically different and incomparable (Ortí 2011). The main visual difference is that within every facies or evaporite bed the gypsum crystals reveal a characteristic crystal morphology, create different types of twins, form specific crystal aggregates or crystalline constructions such as for example domes (Ortí and Shearman 1977; Rodriguez-Aranda et al. 1995; Lugli et al. 2010).

Undoubtedly the differences exhibited by the selenite facies depend on some environmental factors that influence or control the morphology of the growing crystals (such as pH, deviations from stoichiometry, degree of supersaturation, temperature, or foreign admixtures in the solution, and some physical factors such as flow of solution). It seems that the most important factor in the shaping of the morphology of the growing gypsum crystals was the presence of various specific organic substances in the solution (see Cody and Cody 1989, 1991; and reviews in Bąbel 1991; Rodriguez-Aranda et al. 1995; Vogel et al. 2010). It was experimentally proved that even very slight differences in the listed environmental parameters and the presence of only minor amounts of some specific organic compounds in the brine can produce a drastically different gypsum crystal habit observed in various selenite facies. So far, however, these parameters and compounds remain poorly recognized.

Palaeocurrent analysis

In the case of some selenite facies, one factor was recognized as crucial for the creation of the specifically oriented arrangement of crystals or crystal texture, i.e. the flow of brine. As originally suggested by Dronkert (1977, 1978, 1985), in case of the Messinian selenites, the gypsum crystals could grow in an oriented manner, under the influence of a bottom brine current supersaturated with gypsum, with the apices predominantly turned upstream (Pawlikowski 1982; Bąbel 1986; Lugli et al. 2010). This interpretation was fully supported by field studies and statistic measurements of the orientation of crystal apices in the Badenian basin within the laterally continuous sabre gypsum facies beds (Text-fig. 5E). These studies revealed the counter clockwise pattern of brine flow in the basin, interpreted as cyclonic circulation typical of the closed and semiclosed basins of the northern hemisphere (Bąbel 1996, 2002b; Bąbel et al. 1999, 2011, 2015; Bąbel and Becker 2006; Bąbel and Bogucki 2007). The Badenian basin is possibly the only ancient example where the cyclonicbasinal water flow pattern has been detected.

Stratigraphy

The halite Wieliczka Formation is the only formally defined lithostratigraphic unit in the entire Fore-Carpathian Badenian (Garlicki 1994). Wala (1980) attempted to introduce a formal stratigraphic subdivision for the Badenian gypsum deposits of the Nida area and even presented stratotype sections during the conference field trip in 1980. However, his proposal was described only in limited conference materials and remains practically unknown. It was Alexandrowicz et al. (1982) who coined the name Krzyżanowice Formation as the term for the Badenian gypsum evaporite unit in Poland and this name is now widely accepted and used in spite the fact that the formation has not been defined formally. In Ukraine, the coeval evaporite deposits are referred to the Tiras Suite.

Wala (1961, 1963, 1980, and unpublished materials) recognised that the Badenian gypsum deposits of the Nida area in Poland are composed of a set of thin layers showing a nearly perfect correlation, and lettered them alphabetically from a at the base to n at the top. Most of these layers represents selenite beds. Subsequent investigations showed that this sequence of layers is detected in a much larger area of the basin (Kasprzyk 1989, 1991, 1993a), and is also recognizable in Ukraine (Kasprzyk 1995). Kubica (1992, 1994b) working on core material from the northern part of the Fore-Carpathian Basin recognised seven laterally continuing thin lithostratigraphic units within the gypsum sections and marked them with capital letters from A to H. Later this subdivision was recognised in a much larger area of the basin and supplemented by Bąbel (2005a).

Investigations of a number of sections in the entire basin revealed that some of the layers are excellent marker beds. Further sedimentological studies indicated that these beds can be connected with important basin-scale events and interpreted as isochronic or near isochronic. The event-stratigraphy methodology was successfully applied to the Nida Gypsum area (Bąbel 1996, 1999a, b) and to the entire sulphate marginal part.

Models of the evaporitic basin

Before 2000 several more or less justified models of Badenian evaporite basins had been presented. Many authors suggested that the basin was a saline lagoon type (e.g., Kwiatkowski 1972). Garlicki (1979) adopted the so-called “saturated shelf” concept of Richter-Bernburg (1955) and presented the comprehensive model of brine flow in the basin. According to this model, the deposition of evaporite facies was controlled by depth – Ca-sulphates in shallower and Na-chlorides in deeper parts of the basin. This idea was also accepted by Połowicz (1993), Bukowski (2011), and Głuszyński and Aleksandrowski (2016). Similarly Bąbel and Bogucki (2007), who applied limnological terminology to the evaporite basin, suggested a meromictic lake model for the basin, in which halite deposition was restricted to the deepest monimolimnion zone. It was assumed that the selenite deposits were formed only in the well oxygenated and Ca-sulphate saturated mixolimnion zone, on the marginal platform or shelf only. In both models the halite deposition in relatively deep subbasins could be connected with the shallowing and emersion of the surrounding shelves (Bąbel and Bogucki 2007; cf. Bukowski 2011). Liszowski (1989) introduced the model of non-evaporitic deposition for the Ca-sulphates and Na-chlorides assuming that the entire mass of salts derived from the formational brines was expelled to the foreland basin from the tectonically squeezed flysch deposits during the thrust movements of the Carpathian nappes.

Analyses of Badenian sulphate facies integrated with geochemical studies led to the introduction of a new basin model. The marine salina model, formerly mentioned by a few authors, was suggested for the Fore-Carpathian Basin, particularly for its northern Ca-sulphate margin by Peryt et al. (1995), Peryt (2001, 2006a), Bąbel (2004b, 2007b) and Cendón et al. (2004). This model assumes that the basin is a depression separated from the sea by a topographic barrier, and is supplied with seawater seeping through it or overflowing this barrier, occasionally in the form of short-term marine transgressions. The water level in the basin is situated below the seawater level and the basin is in fact a saline lake supplied with marine water.

Gypsum depositional models

The western part of the Ca-sulphate basin (Poland and the Czech Republic) shows different development of the gypsum facies than in the eastern and east-southern part. In the west zone of the basin the sections commonly contain clay intercalations and large pore structures are present between the selenite crystals. Some subfacies, such as skeletal palisade subfacies of the giant gypsum intergrowths, are present only the west (Bąbel 1987, 1999a, b). The giant selenite domes composed of the flat-lying variety of sabre crystals occur only to the east (Text-fig. 5A). It seems that climate and supply of meteoric water to the basin were the main control of these “megafacies” features and Schreiber et al. (2007) suggested the dry shore and wet shore model of evaporite deposition for these two zones.

Carbonate facies

Carbonate facies associated with Badenian gypsum deposits commonly form separate bodies. Among many varieties of limestone two are apparently the most important: the so-called Ratyn Limestone and the sulphur-bearing limestone, both representing a petrological puzzle.

During the earliest studies, the limestones associated with gypsum were called “supra-gypsum limestone” because they were commonly found in the topmost part of the gypsum successions (Alth and Bieniasz 1877), although in many sections they also form a lateral equivalent of gypsum. Lomnicki (1897) introduced the term Ratyń limestone (Ratyń – in Polish, Ratyn – in Russian, Ukrainian and English spelling) for a variety of such a limestone occurring at Ratyn Hill and its environs in Lviv (see Peryt 2006b). Later the term was expanded to embrace other types and varieties of limestones associated with the Badenian gypsum deposits. The Ratyn Limestone facies is difficult to define precisely because the deposits include numerous varieties of limestone, generally chemical in character (massive, dense, and cavernous, commonly without macrofauna, although with outstanding exceptions) and of an unclear origin (see review of the meanings of Ratyn Limestone in Peryt et al. 2012b).

The Ratyn Limestone occurs mostly in Ukraine in a belt of exposures situated north of the Badenian gypsum extent, as well as appearing in the topmost part of the Badenian gypsum commonly covering a karstified surface. The recent petrological and geochemical study of the Ratyn limestone from several representative outcrops in Ukraine by Peryt et al. (2012b) indicates that the limestones are mostly the product of meteoric (vadose-phreatic) diagenesis of a primary evaporitic peloidal carbonate, containing in situ grown lenticular gypsum crystals, deposited in a saline lagoon (or possibly a saline environment; Peryt and Kasprzyk
The sulphur-bearing limestone represents an economically important facies forming large native sulphur deposits both in Poland and Ukraine. There are two opposite views concerning the origin of these sediments (Kwiatkowski 1966), one assuming an entirely post-sedimentary, so-called epigenetic origin via replacement of primary gypsum deposits during burial, and the other suggesting a primary origin at the bottom of the evaporite basin or during very early diagenesis. Most Polish authors accept the epigenetic hypothesis strongly supported by the presence of pseudomorphs after gypsum crystals as well as unchanged gypsum strongly supported by the presence of pseudomorphs. The findings and documentation of the largest natural gypsum intergrowth (Bąbel 1987, 1991), and the recognition of an uncommon type of gypsum intergrowth (Bąbel 1987, 1991), and the findings and documentation of the largest natural crystals so far recorded in Poland (Bąbel 2002a; Bąbel et al. 2012). The sulphur-bearing limestone is thought to have been caused by storm agitation, tidal or pseudotidal waves, or gusts of landborne winds.

Other discoveries

During studies of the Badenian gypsum facies several rare geological phenomena were discovered. They include the finding of gypsum ooids in core material from the environs of Smerdyne in Poland (Text-fig. 1C) (Kasprzyk and Bąbel 1986; Bąbel and Kasprzyk 1990), known only from a few occurrences in the world (Tekin et al. 2008; and references in Bąbel et al. 2012). However, Gąsiewicz (2000a, b, 2010) and Olchowy (2015) were recently able to support the syngenetic theory proving that most of the sulphur-bearing limestones show sedimentary structures different from those observed in the gypsum deposits, which are difficult to explain by the epigenetic theory. According to these authors, the sulphur-bearing limestone represents the sedimentary to early diagenetic environment of a Badenian evaporite shoal.

Large-scale build-ups of the back-bulge depozone of the Fore-Carpathian Basin (A. Radwański, A. Wysocka and M. Görka)

A peculiar coralgal facies is recognised in the area between Ternopil and Kamianets Podilskyi (Ukraine), up to Balatina (Moldova), crossing the Moldova-Romanian border as well. It extends for over more than 300 km, within an up to 40 km wide zone in the centre of the forebulge depozone of the Fore-Carpathian Basin. It is subparallel to the north-eastern margin of the foredeep depozone. The bodies of this facies that are the build-ups (‘bioherms’ or ‘reefs’) once formed slightly elevated monadnocks that protruded above an almost flat-shaped landscape. Their patchy range has long been known as the Medobory Hills (Text-figs 1 and 6) available for studies in plenty of natural and artificial exposures. The dominant components of the Medobory Hills include the Upper Badenian coralline algal reefs associated with bioclastic, marly and rhyodoid facies (Korolyuk 1952; Jasionowski et al. 2005, 2006; Radwański et al. 2006, 2011; Studencka and Jasjionowski 2011; Górka et al. 2012). The coralgal facies from the territory of Moldova was also briefly analysed (Janakevich 1969a,b, 1977).

A typical sequence of the large-scale build-ups is exposed e.g., in the Maksymivka, Nihyn, Humentsi, and Haluschyntsi quarries in the vicinity of Ternopil (Text-fig. 1). These are huge (c. 1 km long and some tens of metres deep) open-pits, located along the topographic crest. The exposed build-ups have been referred to as the Medobory Biothermal Complex (Radwański et al. 2006). The complex is an up to 60 m thick biogenic sequence composed of coralline algal colonies of various shapes and sizes (Text-fig. 6D). Isolated coral colonies, mostly of Tarbellastrea and Porites, occur with usually low frequency in some parts of the build-ups (Górska et al. 2012).

The coralgal build-ups vary in size, from the centimetres of rodolith forms to hummocks several metres high (Text-fig. 6A, C). They are developed either as isolated masses, or overlapping each other and are separated by calcarenites or calcirudites (Text-fig. 6A, C and E). In some cases, the fabric of the biohermal structures was soft during their growth (Radwański et al. 2006). As a result, the build-ups are often densely burrowed by alpheid shrimps. Particular burrows are filled with a clayey material full of bioclasts, especially crab remains and echinoid tests. Some build-ups are riddled by borings of the bivalve Lithophaga, whose internal moulds not only of borings, but also of shells are commonly preserved (Studencka and Jasionowski 2011). This points to an extremely shallow, high-energy environment, in which early diagenetic, beachrock-type cementation occurred. The high-energy dynamics of the Maksymivka environment is thought to have been caused by storm agitation, tidal or pseudotidal waves, or gusts of landborne winds.
In the Polish part of the Fore-Carpathian Basin, the faunal content of an open-marine, normal salinity environment typical of the large-scale build-ups of the back-bulge depozone is constant throughout all regions, upon which the Middle Miocene (Badenian) transgression had progressed. Very diverse faunal assemblages, in facies ranging from clays (Baluk and Radwański 1977), through sands (Radwański 1970; Radwański and Wysocka 2004), to pure limestones of the ‘Leithakalk’ type known as the Pińczów Limestones (Radwański 1965; Studencki 1988a), indicate open-marine conditions, with influx from the Indo-Pacific bioprovince (e.g., Radwański 1974; Hoffman 1977; Förster 1979a, b). Consequently, the studied coral facies and other deposits with an open-marine fauna in Ukraine and Moldova are all correlated with those present in Poland (Roztocze area), and are regarded to be coeval in general terms. However, there are still some open questions on the correlation between the Polish and Ukrainian part of the Fore-Carpathian Basin (compare e.g., with Studencka et al., 1998). Moreover, a temporal shift cannot be excluded, as resulting from the two Badenian transgressions spread propagating from the west and south-west (e.g., Baluk and Radwański 1977; Kovač et al. 2007).

The uppermost parts of the coralgal build-ups are eroded and covered by the Lower Sarmatian serpulid-microbialite reefs (Text-fig. 6C, F). Moreover, these Sarmatian reefs are located also at the south-western foot of the Medobory Hills where they build a few tens of meters high rocky hills called the tounra mounds (Text-fig. 6A, B). They are usually arranged in curvilinear chains oriented approximately perpendicular to the orientation of the Medobory Hills (Text-fig. 6B). Such a geometric pattern is still unexplained and awaiting further sedimentological/structural investigations. Additionally, the change in the composition of the organic community and the type of build-ups is interpreted as related to the restricted conditions in the Paratethys during the Early Sarmatian and resulting palaeoenvironmental changes (Pisera, 1996; Jasionowski et al. 2002, 2003, 2006; Jasionowski 2006; Studencka and Jasionowski 2011).

OVERVIEW OF SELECTED TAXONOMIC GROUPS

Some of the sediments infilling the Fore-Carpathian Basin yield unique and very rich faunal assemblages. Many years of meticulous studies have enabled the analysis of their precise taxonomic and stratigraphic position. The relationships between the particular groups of organisms and their eco-taphonomic significance have also been studied. Many of the analysed groups have turned out to be perfect tools to reconstruct the particular sedimentary settings in the Miocene sea. A brief overview of selected groups of organisms, including corals, polychaetes, mollusces, crustaceans, echinoderms, and fish otoliths is presented below.

All specimens illustrated herein, except Notocholhis tigrina Röding, 1798 of Text-fig. 11Q, which is from the private collection of P. Degórski, are housed in the S.J. Thugutt Geological Museum (prefix MWGUW) of the Faculty of Geology of the University of Warsaw.

Miocene corals from the Fore-Carpathian Basin (M. Görka)

Corals from the Fore-Carpathian Basin were first described by Pusch (1837), Zejszner (1845), and Reuss (1847, 1871). Eichwald (1853) stated the number of the known coral species to be three, but soon after this number quickly increased to over 20 (Siemiradzki 1918; Friedberg 1928; Kowalewski 1930). These early reports were summarized by Dembińska-Rózkowska (1932), who reported 42 species and subspecies. The corals from Korytnica were described by Stolarski (1991) and Roniewicz and Stolarski (1991).

Prior to the paper by Studencki (1979), who reported on a small Early Badenian coral patch-reef in Grobie (discovered by Ryszard Wrona), no coral build-ups were recorded from the Miocene of the Holy Cross Mts. area. The reef was briefly mentioned by Müller (1984), who later described it as the northernmost coral reef in the Neogene and presented its coral assemblage (Müller 1996). Further studies on the coral build-up in Grobie were conducted by Görka (2002), who concluded that its most important coral component was the hermatypic species Tarbellastraea reussiana (Milne-Edwards and Haime, 1850) (Text-fig. 7A) supported by Porites col legniana Michelin, 1842. This particular taxonomic composition of the coral assemblage, dominated by only two species, is a typical phenomenon in the reefal build-ups of the Fore-Carpathian Basin (e.g., Müller 1996; Radwański et al. 2006; Görka et al. 2012). Miocene coral reefs are more common in the Roztocze area (e.g., Pisera 1978, 1985; Müller 1996). The reefs of Roztocze are coeval with a large system of barrier reefs stretching across western Ukraine, north-western Moldova, down to Romania (e.g., Korolyuk 1952; Janakevich 1977; Pisera 1996; Radwański et al. 2006; Studencka and Jasionowski 2011; Görka et al. 2012). This Late Badenian barrier reef is well seen in morphology where it forms the hilly range known as the Medobory Hills. When com-
Text-fig. 6. Spatial relationships of the biohermal facies in the Medobory area, and details of their lithology and main frame-builders. A – Schematic geological cross-section through the Medobory area. Abbreviations: P – Pidhirtsi Beds, Tb – Ternopil Beds (biothermal facies), Td – Ternopil Beds (detrital facies), B – Vyshhorodok Beds and Buhliv Beds, V – Volhyn Beds (biothermal facies in yellow), after Jasionowski et al. 2006; B – Overall view of the vicinity of Nihyn, to show the spatial relationship between the Medobory Hills and the tountry chain; C – Panoramic view of the north-eastern wall of the Humentsi Quarry to show the spatial distribution of the lithofacies; D – massive and laminar thalli of coralline algae (Upper Badenian, Humentsi, UA); E – fine-grained organodetrital limestones from the interbiohermal area (Upper Badenian, Humentsi, UA); F – microbialite limestone with a high content of bivalve shells, serpulid tubes, and numerous growth caverns (Lower Sarmatian, Humentsi, UA)
pared to Roztocze, it becomes obvious that the dimensions of the reefal bodies grow significantly to the south; in Ukraine the thickness of the bioherms reaches almost 100 m. Nevertheless, the taxonomic composition of the coral assemblages is only slightly modified; *T. reussiana* and *Porites* (in this case *P. vindobonarum prima* Kühn, 1925) still predominate (Text-figs 7B, C and 8A-C).

Also noticed are *Siderastrea cf. italica* (Defrance, 1826) (Text-fig. 7D) and *Montastrea* sp. (Text-fig. 8D). The taxonomic composition of the coral assemblage shows a significant similarity with other Miocene coral reefs of the Tethys (Esteban et al. 1996; Hayward et al. 1996) and Paratethys (Friebe 1991, 1993; Müller 1996; Riegl and Piller 2000; Górka 2002).

The shapes of the coral colonies (see Korolyuk 1952; Radwański et al. 2006; Górka et al. 2012) vary from massive, lamellar, through knobby, to branching and/or platy forms (Text-fig. 7B). Large quantities of *Porites* may also indicate a significant influx of suspended material of terrigenous origin (see Müller 1984; McCall et al. 1994; Esteban 1996; Jasionowski et al. 2005, 2006).

The important role played by hermatypic corals in the ecosystem is particularly emphasized by the presence of diversified borings and/or pits formed by numerous groups of organisms inhabiting the cenostome. Aragonitic skeletons are regarded as a hard substrate, ideal for settling. With the exception of organisms that inhabited the coral colonies only due to their hardness but could also settle on inanimate hard substrates (boring bivalves of the genera *Lithophaga*, *Gastrochaena*, and *Jouannetia* (Text-fig. 8C); see Radwański et al. 2006; Studencka and Jasionowski, 2012; Górka et al. 2012; Text-fig. 7C, D), there were also animals closely connected to corals, probably due to some kind of interspecific relations: the cirrped crustaceans of the genus *Creusia* (Baluk and Rad-
wański 1967; Wysocka et al. 2012) and the micropoly-lychaetes *Josephella commensalis* (Bałuk and Radwański 1997).

In a similar manner to the other invertebrate groups of the Fore-Carpathian Basin, corals were ubiquitous during the Early Badenian climatic optimum (see Müller 1996; Górka 2002; Radwański et al. 2006) and disappeared with the advent of the salinity crisis. They re-appeared with the return of open-marine conditions in the latest Badenian. The presence of corals is evidence of at least warm-temperate or even subtropical waters during the Middle Miocene of the Fore-Carpathian Basin (Radwański et al. 2006; Górka et al. 2012).

**Polychaetes from the Korytnica Basin** (U. Radwańska)

The vast majority of polychaete worms are marine, being one of the most common components of present-day shallow-marine communities (Bianchi 1981; Jäger 1983). Based on the mode of life, the Class Polychaeta Grube, 1850 is subdivided into two orders: the Sedentaria Lamarck, 1818, and the Errantia Audouin and Milne-Edwards, 1832. Representatives of sedentary polychaetes live usually permanently in a more or less lithified tube, rarely in burrows or borings, and their fossil potential is relatively high.

Most of the tube-dwelling polychaetes are suspension feeders, and thus they require permanent water mobility to keep the branchial crown free from the sediment. Others have feeding tentacles extending from the tube opening and creeping along the mud, picking up organic particles. Favourable bathymetric conditions for most of the present-day genera are referenced as littoral, usually inter- or shallow subtidal; some, however, are found at bathyal depths (Hayward 1977; Bianchi 1981; Jäger 1983; Radwańska 1994a).

The first tube-dwelling polychaetes from the Ko-
rytnica Clays were reported by Pusch (1837, p. 181). Subsequently, various materials from Poland, Volhynia and Podolia were studied by Dembińska (1924), and more recently the group was monographed by Jakubowski and Musial (1977, 1979a, 1979b); Radwańska (1994a, 1994b); Baluk and Radwański (1997) and Radwański and Wysoka (2004).

Sedentary polychaetes from the Korytnica Clays

The most diversified sedentary polychaete assemblage was recognised from the Korytnica Clays, which comprises 32 species- or genus-level taxa (Radwański 1964, 1969, 1970; Radwańska 1994a, b; Baluk and Radwański 1997). They are represented by rock-boring forms of the family Spionidae and Sabellidae, tube-dwelling forms (Radwańska 1994a), as well as commensal micropolychaetes (Baluk and Radwański 1997) and unique, opercular caps (Radwańska 1994b). Also reported was the ichnofossil Helicotrephichnus commensalis Kern, Grimmer and Lister, 1974, occurring within the columnellae of some gastropods, and attributed to borings of present-day spionid polychaetes that are commensal to the hermit crabs occupying empty gastropod shells (Kern 1979; Baluk and Radwański 1984b; Radwańska 1994a). Another ichnofossil, Tibikoria sanctacrucensis Baluk and Radwański, 1979, being aggregates of faecal pellets, was ascribed to the life activity of the present-day species Heteromastus filiformis (Claparède, 1864).

Tube-dwelling polychaetes: This is most abundant group among sedentary polychaetes (Text-fig. 9). Variable tube morphology, mode of coiling, sculpture, and relation to the substrate are the basis of their species-level taxonomy. Particular species reveal phyloetic relationships either to Late Mesozoic or to modern polychaete faunas.

The studied polychaete assemblage includes also very peculiar material consisting of tiny calcareous tubes, observable under SEM-magnifications (Baluk and Radwański 1997), settled within the calyces of some colonies of the scleractinian coral Tarbelliastraea reussiana. These tubes are ascribed to the life activity of micropolychaetes commensal to this coral and are accommodated into the extant genus Josephella Caulery and Mesnil, 1896 to represent a separate species Josephella commensalis Baluk and Radwański, 1997.

O per cular caps: This is a very unique group among the tube-dwelling polychaetes (Radwańska 1994b; Text-fig. 10), representing isolated parts of the opercular system closing the tubes of some polychaetes. The group was described, in open nomenclature, as opercular caps comparable to those of the present-day species of the genus Vermilopsis Saint-Joseph, 1894. Ten Hove and Nishi (1996) considered these caps to belong to the serpulid genera Spirobanchus Blainville, 1818 and/or Pomatoceros Philippi, 1844. Unfortunately, tubes of these genera have not been found in the Korytnica Clays (Radwańska 1994a), nor were the opercula found in situ.

Rock-boring polychaetes: These polychaetes contain three species: Polydora ciliata (Johnston, 1838), P. hoplura (Claparède, 1869), and Potamilla reniformis (O.F. Müller, 1771). All of the taxa recorded are adapted to live permanently in their characteristically shaped borings. These polychaetes mostly inhabited abrasion surfaces along the rocky seashores of the Korytnica Basin, zones affected by strong waves and currents, and on any skeletal material available within the Korytnica Clays (Radwański 1964, 1969, 1970; Radwańska 1994a).

Distribution of sedentary polychaetes in the Korytnica Basin

Sedentary polychaetes are relatively rare in the Korytnica Basin, both in the rocky-shore facies of the oyster shell bed and within the clay facies. In both facies, polychaete tubes were found either isolated, probably having been detached from any soft substrate, or as epibiontic forms encrusting skeletal material (primarily, coral colonies and mollusc shells). The former ones, except for the free-living species Ditrupa cornea (Linnaeus, 1767) and the commensal micropolychaete Josephella commensalis, were certainly attached to such soft substrates as e.g., sea plants, or to some soft-bodied animals. In the clay facies, the tube-dwelling polychaetes usually occur as gastropod epizoans, and are rarely represented by single specimens of spirobids or serpulids.

Gastropods (M. Złotnik)

Text-fig. 10. Middle Miocene (Badenian) opercular caps of the tube-dwelling polychaetes *Vermiliopsis* Saint-Joseph, 1894 from the Korytnica Clays. A – Cap comparable to those of the present-day species *Vermiliopsis monodiscus* Zibrowius, 1968, outer view, MWGUW PK-096; SEM × 50; B-C – Caps comparable to those of the present-day species *Vermiliopsis infundibulum* (Philippi, 1844), B – outer view of a low-conical specimen, MWGUW PK-097; C – outer view of a high-conical specimen, MWGUW PK-098; SEM × 50; D-F – Caps comparable to those of the present-day species *Vermiliopsis labiata* (O.G. Costa, 1861), D, E – outer view of low-conical specimens, D - MWGUW PK-099, E - MWGUW PK-100; SEM × 30; F – outer view of a high-conical specimen with a small cross, MWGUW PK-101; SEM × 50. Photographs by C. Kulicki
dencka and Popov 1996; Studencak et al. 1998, 2012) they are usually the most abundant macrofossils occurring in the sandy and especially in the clay facies. Gastropod shells, or rarely their internal moulds, were reported from numerous exposures of the entire Fore-Carpathian Basin, e.g., from Niskowa (Bałuk 1970), Andrychów (Krach and Nowak 1956) and Benczyń (Krach and Książkiewicz 1949), Miechów (Krach 1947), Przeciszów (Alexandrowicz and Krach 1963), Gliwice Stare (Krach 1954), to Korytnica (Friedberg 1911-1928; Kowalewski 1930; Bałuk 1975, 1995, 1997, 2003, 2006), numerous localities in Roztocze Hills area (Jaikuowski and Musiał 1977, 1979a,b, Krach 1981, Piwowarski 1977). Calcareous nannoplankton and dinoflagellate cysts locate the Korytnica Clays (Radwański 1969; Hoffman 1977), located in the southern part of the basin and recognised only from a single locality, Eichwald (1853), Hörnes (1856), and Hoernes and Auinger (1879) included the Korytnica gastropods into their classic monographs on the Miocene malaco fauna. The most recent Baluk’s (1975, 1995, 1997, 2003, 2006) monographs include over 550 prosobranch species representing 65 families, with 33 new species recognised (Baluk 1975, 1995, 1997, 2003, 2006; Baluk and Jakubowski 1968). The highest taxonomic diversity is noted among the Turridae (101 species). However, 20 species or more, are noted also among the Muricidae, the Nassariidae, the Mitridae, the Cerithidae, and the Rissoidae. The total number of gastropod species from the Korytnica Clays exceeds 800 species, including more than 250 opisthobranch species (Baluk, in preparation). According to Baluk, the gastropods from the Korytnica Clays reveal the strongest affinity to the gastropods from the Vienna Basin (see Hörnes 1856; Hoernes and Auinger 1879; Harzhäuser and Kowalke 2002).

The relative abundance of gastropods varies in the succession. The relative number of gastropod shells ranges from 10% to 85% of all invertebrate macrofossils. The lowest and highest frequencies were recognised from deposits predominated by corbulid bivalve mollusces [Corbula gibba (Olivii, 1792)] and those predominated by turritellid gastropods (Turritella badensis Sacco, 1895), respectively. The relative biovolume of the turritellids recognised from the latter deposits reaches up to 70% of the total biovolume of all intrafaunal macrofossils (Hoffman 1977).

The gastropod taxonomic composition varies between particular parts of the Korytnica Basin. Its littoral zone is occupied mostly by Nassarius limatus (Chemnitz, 1786), Petaloconchus interruptus (Lamarck, 1818), N. korytnicensis (Baluk, 1997), N. restitutianus (Fontannes, 1879), Alvania spp., Caecum sp. and cerithids (see Baluk and Radwański 1977; Hoffman 1977; Zlotnik 2003). The coastal swamps (Radwański 1969; Hoffman 1977), located in the southern part of the basin and recognised only from a single locality, were dominated, i.a., by Neritina picta, Nassula schoeni (Hoernes and Auinger, 1882), Melanopsis sp., Turrillia erronea Cossmann in Friedberg 1914, Terebralia sp. and Natica pseudoredecepta Friedberg, 1923 mostly of which are rare or absent from other parts of the basin.

The major off-shore part of the Korytnica Basin was dominated by Nassarius korytnicensis, N. restitutianus, N. styriacus (Auinger in Hilber, 1897), Turritella badensis, Turboella spp., cerithids, Odostomia spp., Eulina sp. and Notocochlis tigrina (=Natica tigrina in Baluk 1995). Nassarius korytnicensis, T. badensis and N. styriacus dominate the upper and/or uppermost part of the succession and the latter species

Most gastropods from the Korytnica Clays are characterised by an exceptionally good preservation state. The shell surface is usually shiny, and the details of the sculpture are perfectly preserved. The protoconch is usually present and easily distinguishable from the te-
gratein (Text-fig. 11Q1, respectively). On the other hand, some gastropod shells are corroded and/or bear biocorossional traces, a record of the activity of clini-
did sponges, boring pelycopods and polychaetes. The shell surface may also be altered by encrusting and et-
ing organisms: polychaetes, barnacles and bryozo-
ans (Bałuk and Radwański 1977).

Due to their exceptional preservation state and extremely high taxonomic diversity, the group has been the subject of intensive investigation for over 200 ye-
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Gastropods from the Korytnica Clays – taphonomy, taxonomy, abundance and mortality patterns

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leoconch (Text-fig. 11R, S). In a few cases the colour pattern is also preserved. This is well recognisable in Neritina picta Férrusac, 1825 or Notocochlis tigrina (Röding, 1789) (=Natica tigrina in Baluk 1995) (Text-
fig. 11L and Text-fig. 11Q1, respectively). On the other hand, some gastropod shells are corroded and/or bear biocorossional traces, a record of the activity of clini-
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1928, 1928; Kowalewski 1930), Eichwald (1853), Hörnes (1856), and Hoernes and Auinger (1879) included the Korytnica gastropods into their classic monographs on the Miocene malaco fauna. The most recent Baluk’s (1975, 1995, 1997, 2003, 2006) monographs include over 550 prosobranch species representing 65 families, with 33 new species recognised (Baluk 1975, 1995, 1997, 2003, 2006; Baluk and Jakubowski 1968). The highest taxonomic diversity is noted among the Turridae (101 species). However, 20 species or more, are noted also among the Muricidae, the Nassariidae, the Mitridae, the Cerithidae, and the Rissoidae. The total number of gastropod species from the Korytnica Clays exceeds 800 species, including more than 250 opisthobranch species (Baluk, in preparation). According to Baluk, the gastropods from the Korytnica Clays reveal the strongest affinity to the gastropods from the Vienna Basin (see Hörnes 1856; Hoernes and Auinger 1879; Harzhäuser and Kowalke 2002).

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frequently occurs only in the central part of the basin (see Baluk and Radwański 1977; Hoffman 1977; Zlotnik 2003).

Besides small and medium-sized species, which predominate in both abundance and biovolume in the bulk samples, many large-sized gastropods were collected, although their relative abundance in the bulk samples is usually low. Some of these large gastropods are important components of the trophic web (Hoffman 1977) and diagnostic of the distinguished palaeocommunities (see Baluk and Radwański 1977). Among them, Clavatula polonica (Pusch, 1837), Murex friedbergii Cosman and Peyrot, 1924, Ancila glandiformis (Lamarck, 1810), Tudica rusticula (Basterot, 1825) and Semiacassis miolaevigata Sacco, 1890 are the most abundant (see Text-fig. 11).

The mortality patterns of some Korytnica gastropods were studied by Hoffman (1976) and Kowalewski (1990). The investigations show remarkable differences between the studied species. The mortality rates of Murex friedbergi, Nassius restitutianus, Notocochlis tigrina and Pyrgulina indistincta (Montagu, 1808) are nearly constant for their postlarval life. A very high proportion of individuals dying in the early stages was observed for Turboellula badensis. Raphitoma hispidula (Jan in Bellardi 1847) and Turbonilla scala (Eichwald, 1853) are characterised by a relatively high juvenile mortality, followed by a plateau of adult vigour, when the mortality was very low. In gerontic stage, the probability of death rapidly increases for these species (Hoffman 1976). The analysis of the contribution of predatory boring gastropods to the mortality of the Korytnica gastropods demonstrates that, generally, the mortality patterns do not depend on the predation by muricid and/or naticids, even when the predation pressure was very high (Hoffman 1977; Kowalewski 1990).

### Nassariidae gastropods and local stratigraphy of the Korytnica Clays

The Korytnica Clays lie almost horizontally (also confirmed by geophysical data - see Szymanko and Wójcik 1982). Representatives of the family Nassariidae, the most abundant gastropods in the Korytnica Clays (see Baluk and Radwański 1977; Hoffman 1977; Kowalewski 1990; Zlotnik 2003), are crucial for local stratigraphy (Zlotnik 2003). Based on five nassariid assemblages (Nassarius restitutianus, N. restitutianus-korytnicensis, N. korytnicensis, N. koryntcensis-limatus and N. schoenni), the Korytnica Clays succession was subdivided into five eocratigraphic units: Lower restitutianus, Lower restitutianus-korytnicensis, Upper restitutianus, korytnicensis, and Upper restitutianus-koryntcensis. The entire sequence reflects temporal changes in the relative size of two large gastropod populations: of N. restitutianus and of N. korytnicensis (Zlotnik 2003). This nassariid-based stratigraphy appears to almost the entire Korytnica Basin with the exception of the littoral zone, characterised by the schoenni and koryntcensis-limatus assemblages (Zlotnik 2003, text-fig. 7). The nassariid-based correlation was confirmed by many geological and palaeobiological data.

The appearance of the korytnicensis assemblage in the upper part of the succession seems to be the most significant ecological event hitherto recognised from the entire sequence of the Korytnica Clays. The occurrence of several strictly tropical faunal elements and their broad correlation with the most taxonomically diversified community III of Baluk and Radwański (1977), suggest that the appearance of this assemblage was caused by a warming event. If this preliminary hypothesis (Zlotnik 2003) is correct, then the entire succession of nassariid assemblages could be interpreted as a record of climatic fluctuations that took place in the early Middle Miocene.

### Palaeoecology of the Korytnica gastropods

The gastropods represent almost all members of the trophic web, viz. predators (e.g., Notocochlis, Murex, Clavatula, Raphitoma), scavengers (Nassarius), browsers (e.g., Turboellula, Alvania, cerithids), parasites (e.g., Odostomia, Eulima), and suspension feeders (Turri-
tella, Petaloconchus) (Hoffman 1977). The group is the member of the trophic nucleus within 4 out of 5 macrobenthic assemblages recognised from the Korytnica Basin by Hoffman (1977). Nassarids, turritellids, Turboella, Pyrgulina, Odostomia, Turbonilla, Alvania, and naticids are members of the trophic nucleus in the Turboella-Loripes assemblage; Nassarius and turritellids – in the Corbula-spatangoid assemblage; turritellids – in the turritellid assemblage; and Clavatula – in the Pteromeris-coral assemblage.

Noteworthy, the dominant gastropod taxa recognised by Bałuk and Radwański (1977) from the clay succession differ from those recognised by Hoffman (1977) or Złotnik (2003). These differences reflect different sampling and sieving methods discussed earlier in this paper.

Both the overall character of the gastropod fauna and the occurrence of gastropods typical of warm waters indicate tropical/subtropical conditions during the sedimentation of the Korytnica Clays. According to Bałuk and Radwański (1977), the occurrence of such genera as Terebra, Rostellaria, Architectonica, and large-sized species of the genera Cypraea, Conus, Strombus, Triton, and Galeodes indicate tropical conditions. The same is true for the genus Parastrophapia represented in the Korytnica Clays by Parastrophapia radwanskii (cf. Bałuk 1975). The subtropical/subtropical conditions of the clay sedimentation are also confirmed by the occurrence of the bivalved gastropod Berthelinia (see Bałuk and Jakubowski 1968; Bałuk and Radwański 1977). Finally, the extremely high taxonomic diversity of the Korytnica gastropods also indicates warm climatic conditions, as in recent communities biodiversity is positively correlated with temperature (Weiner 1999). Warm climatic conditions (tropical and/or subtropical) are also suggested by the non-gastropod animal groups recognised from the Korytnica Clays (for review see Bahuk and Radwański 1977 and references therein; see also Schultz 1977; Radwańska 1984, 1992; Rögl and Brandstätter 1993). Warm water, subtropical conditions in the Central Parathetys realm in the Badenian are also confirmed by the results of Studencka et al. 1998, Studencki 1999 and Harzauser et al. 2003. According to Sitár and Kováčová-Slamková 1999, Böhme 2003 and Kvaček et al. 2006, the Badenian climate of the Central Parathetys realm represents a part of the Miocene Climatic Optimum.

The Korytnica Clays sequence and the overlying members were interpreted by Bałuk and Radwański (1977) as formed in a progressively shallowing basin with depths changing from c. 60 m at the beginning of sedimentation to almost 0 m at its end. Normal marine salinity is confirmed by the occurrence of corals (Dembinska-Różkowska 1932; Bałuk and Radwański 1984b; Stolarski 1991), echinoids (Mączyńska 1977, 1987), asteroids (Kaczmarska 1987), crinoids (Radwańska 1987), and some benthic foraminifera (Walckiewicz 1975, 1977b; Rögl and Brandstätter 1993).

Drilling gastropods from the Korytnica Basin

Three major groups of gastropod drillers were recognised from the Korytnica Basin, viz. naticid, muricid and cassid drillers. Drill holes in the mollusce shells attributed to the predatory activity of naticid and/or muricid gastropods were reported from the Korytnica Clays by Hoffman et al. (1974), Kowalewski (1990), and Złotnik (2001). Drill holes on echinoid tests attributed to cassids were recognised from both the Korytnica Clays (Ceranka and Złotnik 2003) and from the overlying Heterostegina Sands (Ceranka and Złotnik 2003; Złotnik and Ceranka 2005), the deposits of which filled the basin in the terminal stage of its development (see Bałuk and Radwański 1977; Gutoński 1984). The fossil remains of the drillers and traces of their predatory activity usually co-occur; however, in some cases the number and/or size of the shells of the drillers remain in evident contrast with the number and/or size of the drill holes (see Ceranka and Złotnik 2003). Examples of the drilling activity of predatory gastropods from the Korytnica Basin are presented in Text-fig. 12A–C.

The detailed quantitative analysis of the drill holes has provided many important data on the palaeobiology of the drillers. Among 122 species of prey attacked by naticids and/or muricids, the highest drilling frequencies were recognised for the gastropods Pyrgulina interstincta (Montagu, 1808) (41%) and Tornatina truncatula (Bruguère, 1792) (40%), and for the bivalves Meretrix sp. div. (43%) and Glycymeris deshayesi (Mayer, 1868) (40%). On the other hand, some gastropods [e.g., Ficus sp. div., Ranella marginata (Martini, 1777)] have been attacked neither by naticid nor muricid drillers (Hoffman et al. 1974). According to the latter authors, the naticids had more frequently attacked bivalve molluscs whereas the muricids preyed mostly upon gastropod prey. This result is, however, disputable, because the use of drill hole morphology as a criterion for distinguishing between naticid and muricid drillers (applied by Hoffman et al. 1974) has been undermined by Złotnik (2001). The correlation between the size of the predator and the size of the prey predicted by Kitchel’s et al. (1981) model of predator – prey interaction was not a rule in the case of naticid drillers from the Korytnica Basin. A strong correlation between the size of the prey and the size of the preda-
tor was recognised for naticid drillers preying upon the gastropod *Nassarius restitutianus*. The lack of any correlation between the discussed variables was noted for naticids preying on the bivalves *Corbula gibba* (Olivi, 1792) and *Venus nux* Gmelin, 1791 (= *Venus multilamella*, Lamarck, 1818) (Kowalewski 1990).

Kowalewski (1990) indicated that naticids were at least in some cases size-selective drillers. The central region of the shell of the bivalve *Corbula gibba* and the two last whorls of *Nassarius restitutianus* (= *Nassa hoernesi* in Kowalewski 1990) were drilled by naticids preferentially. Złotnik (2001) showed that the predatory behaviour of naticid drillers can change during the driller ontogeny. Larger naticids chose a more attractive (gastropod) prey, and when preying upon bivalves (*Corbula gibba*), they also displayed a stronger site selectivity.

Both naticid and murid attacks were very effective, with more than 95% of all drill holes complete (sensu Kitchel et al. 1981), enabling prey consumption (Hoffman et al. 1974; Kowalewski 1990; Złotnik 2001).

Patterns of drilling predation of cassid gastropods on echinoids from the Korytnica Basin were recognised by Złotnik and Ceranka (2005). Echinoid prey, represented by three species of *Echinocyamus* (*E. linearis* Capeder, 1906, *E. pusillus* Müller, 1776 and *E. pseudopusillus* Cotteau, 1895) were drilled presumably by one cassid species, *Semicassis miolevigata* (Ceranka and Złotnik 2003; Złotnik and Ceranka 2005). The drilling predation intensities varied among the prey species and ranged from 3.4% for *E. linearis* up to 36% for *E. pusillus*. An intermediate value of drilling predation (15%) was recognised for *E. pseudopusillus*. The intensities of drilling predation recognised for some of the prey species (*E. pusillus*) varied also between (but never within) the sublithofacies of the investigated deposits (*Heterostegina* Sands). Drilling predation of cassids was both size- and site-selective. Larger individuals of *E. linearis* and *E. pusillus* were attacked more frequently and the aboral side of the test of all *Echinocyamus* species was drilled preferentially. An extremely high concentration of drill holes was observed in the apical disc and petals. Results obtained for the most abundant prey (*E. linearis*) indicate that the predatory behaviour of large cassids was somewhat different from those typical of small cassids. Large cassids drilled and consumed their prey...
almost always individually, whereas small cassids sometimes preyed upon the urchins in a group. Large cassids displayed also a higher site-selectivity. Cassids from the *Heterostegina* Sands were extremely efficient drillers, as all drill holes are complete and unrepaired. This means that all attacks on the echinoids were successful (Złotnik and Ceranka 2005).

Cassid drill holes in *Echinocyamus* from the *Heterostegina* Sands significantly complement the information on the size and structure of cassid populations recorded by body fossils. The high number of small drill holes recognised from the sands indicates the occurrence of a large number of small individuals in the cassid population, although neither small shells nor small internal moulds of cassids were preserved in the fossil record of the Korytnica Basin (Ceranka and Złotnik 2003).

**Crustacean predation upon gastropods**

Many gastropod shells bear traces of crustacean predation. The predatory activity of hermit crabs has been reported by Radwański (1969) and Baluk and Radwański (1977). Traces of stomatopod predation upon the Korytnica gastropods have been described and analysed by Baluk and Radwański (1996). According to Baluk and Radwański (1977), “particular species of Korytnica gastropods were attacked by hermit crabs to a very variable extent. Of the long siphonated forms, the most common two *Clavatula* species, *C. laevigata* (Eichwald, 1853) and *C. asperula* (Pusch, 1837), became the prey in very low percentage, whereas very uncommon *Tudicla rusticula* was damaged in most specimens collected. A low percentage of damage is also noted in a common species *Ancilla glandiformis* whose shell may be however nipped completely”. These preliminary results may indicate that hermit crabs were very selective when they chose their gastropod prey, however, it should be emphasized that the suggested species-selectivity, although very probable, should be treated as only a preliminary hypothesis. This is because the investigation of Baluk and Radwański (1977) focused only on a few, selected gastropod species and no numerical data have been presented. My field observations evidently indicate that many other gastropods (mainly nassarids) were also frequently attacked by crabs (Text-fig. 12F–H).

Stomatopod predation upon gastropods was the subject of more detailed studies. The percentage of the attacked individuals varied distinctly between 31 investigated species of prey and ranges from 0.6% for *Notocochlis tigrina* to 54.4% for *Nassarius dujardini* (Deshayes, 1844) (=*Sphaeronassa dujardini* in Baluk and Radwański 1996). Stomatopod predation upon gastropods was site-selective. Based on the different morphology of the traces they were attributed to two different stomatopod families: Gonodactylidae and Squillidae (Baluk and Radwański 1996). Some of the shells bearing traces of crab or stomatopod predation are evidently repaired (see Baluk and Radwański 1977 and 1996 for crabs and stomatopods, respectively). This indicates that gastropods could have sometimes survived attacks of crustacean predators, both crabs and stomatopods.

Traces of stomatopod predation observed on the gastropod shells (circular to sub-circular holes and elongate scars – see Text-fig. 12D, E) are, in fact, the only record of the occurrence of stomatopods in the Korytnica Basin, as no remains of these low mineralized crustaceans were preserved in the fossil record of the discussed deposits.

**Scaphopods, chitons and cephalopods** (M. Złotnik)

Fossil scaphopods are common in the Middle Miocene deposits of the Fore-Carpathian Basin. Although the number of their species does not exceed 20, their shells were reported from numerous exposures located in a broad area extending from the Silesia Upland to the Roztocze Hills, e.g., from Andrychów (Krach and Nowak 1956) and Benczyn (Krach and Książkiewicz 1949) near Wadowice, Gliwice Stare (Friedberg 1911–1928; Krach 1954), Brzeżnica near Bochnia (Krach 1960), Zglobe (Friedberg 1938), Wieliczka (Reuss 1867), Małoszów (Krach 1947), Niskowa (Skoczylasówna 1930; Baluk 1965, 1970), and Korytnica near Kielce (Friedberg 1911–1928; Kowalewski 1930; Baluk 1972, 1984). They were also reported from the Ukrainian part of the Carpathian Foredeep Basin (Friedberg 1928, 1938).

The most diversified scaphopods were recognised from the Lower/Middle Badenian Korytnica Clays, a unique facies deposited in a shallow bay during the transgression of the Parathethys sea onto the southern slopes of the Holy Cross Mts. in Central Poland (see Radwański 1969; Baluk and Radwański 1977 for geological setting; Martini 1977; Dudziak and Łuczewska 1991; Rögl and Brandstätter 1993, Gedl 1996; for stratigraphic position). Among 13 scaphopod species (including 4 new ones) recognised in the clays, *Dentalium fossile* Schroeter, 1783 and *Fustiaria mio- caenica* (Boettger, 1901) are the most abundant taxa (Baluk 1972). In some parts of the clays, the relative biovolume of the latter species reaches up to 10% of the total biovolume of all invertebrate fossils (Hoffman 1977). The pattern of geographic distribution of particular scaphopod species recognised by Baluk (1972)
shows that Dentalium fossile, D. badense Partsch in Hönes, 1856, and Entalina tetragona (Brocchi, 1814) were the most widespread scaphopods across the Polish part of the Parathetys Sea.

Chitons were recognised from Niskowa (Bałuk 1965), Korytnica (Baluk 1971, 1984), Gierszowice Rybnica1, Rybnica2 and Nawodzice (Studencka and Studencki 1988b), Monastyrz and Trzęsiny (Jakubowski and Musiał 1977, 1979a) and from numerous localities in Ukraine (Studencka and Dulai 2010). The most diversified assemblage was recognised from the Korytnica Clays. Among 17 species of chitons (including 5 new ones) recognised from the Korytnica Clays, Chiton denudatus Reuss, 1860 and Lepidopleurus decoratus (Reuss, 1860) are the most abundant species (Baluk 1971). The latter species occurs also in the much less diversified chiton assemblage from Niskowa (Baluk 1965). The results of Studencka and Studencki (1988b) show that taxonomic diversity of chitons declines during Badenian.

Cephalopods are represented by a single, endemic cuttlefish species, Sepia sanctacrucensis Baluk 1977, reported only from the Korytnica Clays. Fragmented parts of the sepions (“os sepiae”) are the only remains of this species preserved in the fossil record (Baluk 1977, 1984). Both chitons and Sepia occur in Miocene deposits very rarely (Baluk 1971, 1977). Selected representatives of all discussed molluscs are presented in Text-fig. 13.

Scaphopods and chitons from the Korytnica Clays are very similar to those recognised from the Miocene deposits of the Vienna Basin (Baluk 1971, 1972). The occurrence of the genus Sepia in the Parathetys has been discussed broadly by Baluk (1977), who concluded that Sepia, due to its relatively low fossilization potential, is very rarely present in the fossil record.

The occurrence of scaphopods, chitons and cephalopods indicates normal marine conditions (salinity) during the sedimentation of the Korytnica Clays. Chitons, reported exclusively from the upper and uppermost parts of the clays, indicate very shallow depths of the sea during the late stage of the development of the Korytnica Basin (Baluk 1971; Baluk and Radwański 1977). Finally, the occurrence of representatives of the polyplacophoran genera Crasedochiton and Cryptoplax recognised from the clays by Baluk (1971) is considered as indicative of tropical conditions by Baluk (1971) and Baluk and Radwański (1977). All these palaeoecological interpretations are in agreement with the results obtained from the investigations of many other animal groups, which inhabited the Korytnica Basin in the Middle Miocene (for review see Baluk and Radwański 1977 and chapter “Gastropods” in this paper).

Crustaceans of the Fore-Carpathian Basin (M. Górka)

The first decapod remains were found in Wieliczka (Reuss 1867) and it was concluded at that time that crustaceans are present in very diverse lithofacies, from coarse-grained cliff conglomerates, through sands, clays, to carbonates, of Early and Late Badenian age.

Cirripeds: Badenian crustaceans were first reported by Baluk and Radwański (1967), who described a new cirriped species, Creusia sanctacrucensis Baluk and Radwański, 1967, that inhabited coral colonies (see also Baluk and Radwański 1984a). Subsequently, they described another species of a cirriped crustacean, Trypetesa polonica Baluk and Radwański, 1991, considered as an obligate commensal to the hermit crabs (Baluk and Radwański 1991). Diverse reworked cirriped species from the Lower Badenian Mykolaiv sands of Ukraine were described by Wysocka et al. (2012), who reported abundant barnacles Balanus sp., goose barnacles Scalpellum sp., and unique specimens of Creusia sp. (Text-fig. 14A).
Decapods: The studies on decapods intensified in the 1970’s with the continuous works of Wacław Bahuk and Andrzej Radwański (Radwański 1967; Bahuk and Radwański 1968; Bahuk 1970) in the south-eastern parts of the Holy Cross Mts. Radwański (1973) reported the remains of a Callianassa shrimp from the Lower Badenian sands of the Holy Cross Mts. area and attributed numerous Ophiomorpha nodosa Lungdren, 1891, burrows to it. Later, he published on decapod-attributable burrows from various Miocene sites from western Ukraine: burrows attributed to the shrimp Alphaeus sp. from the reeval deposits of Upper Badenian limestones of the Ternopil Beds (Radwański et al. 2006); the burrows O. nodosa attributed to the Callianassa shrimp and Psilonichnus ichsp., attributed to the ghost crab Ocypode sp., from the Lower Badenian Mykolaiv Beds (Radwański et al. 2012; Wysocka et al. 2012).

The first monographic descriptions of the Polish Badenian decapods appeared in 1979, with Förster’s (1979a) publication on Miocene crabs from southern Poland. It was followed by a paper devoted exclusively to the decapod assemblage of the Lower Badenian deposits of the Korytnica Basin (Förster 1979b), in which 23 forms representing at least 10 genera were recorded. Förster’s designations were reviewed and revised by Müller (1984, 1996). The latter of these papers (Müller 1996) must be treated as the most extensive and up-to-date for the recognition of decapods from the Polish part of the Fore-Carpathian Basin. It presents 36 species from different parts of the basin – not only from the biohermal (reefal) deposits on the southern slopes of the Holy Cross Mts. (see Görka 2002) and the Roztocze area, but also from other lithofacies, varying from clayey, through muddy, to sandy deposits. It also states that decapod assemblages were present in reeval facies in two chronostratigraphic levels – in the Lower Badenian (predominantly the Holy Cross Mts. area and the uppermost Badenian (Roztocze). The taxonomic composition of these assemblages is almost identical to the assemblages recorded in other reefal deposits of Miocene of the Polish part of the Fore-Carpathian Basin. After their re-appearance on the Korytnica Clays (e.g., Mączyńska 1977, 1987; Radwańska and de Angeli 2010; Ósso and Stalennuy 2011; Górka 2006; Radwański et al. 2012), they finally disappeared due to the decisive environmental changes at the Badenian/Sarmatian boundary (Studencka and Jasionowski 2011; Görka et al. 2012).

Miocene Echinodermata of Poland (U. Radwańska)

The Echinodermata are relatively common in the Middle Miocene of the Polish part of the Fore-Carpathian Basin and are represented by five classes: Holothuroidea, Crinoidea, Asteroidea, Echinoidea, and Ophiuroidea (not described yet).

The most diversified echinoderms are known from the Korytnica Clays (e.g., Maczyńska 1977, 1987; Walkiewicz 1977a; Kaczmarska 1987; Radwańska 1987; Ceranka 2007a), of the Korytnica Basin, and from the sands of Nawodzice (Bahuk and Radwański 1984, 1993; Radwański et al. 2006; Gatt et al. 2012). The most diversified echinoderms are known from the Korytnica Clays (e.g., Maczyńska 1977, 1987; Walkiewicz 1977a; Kaczmarska 1987; Radwańska 1987; Ceranka 2007a), of the Korytnica Basin, and from the sands of Nawodzice (Bahuk and Radwański 1984, 1993; Radwański et al. 2006; Gatt et al. 2012).
1968; Nosowska 1997) and Świniary of the Raków-Klimontów area (Radwański 1973; Maćzyńska 1988; Radwański and Wysocka 2004; Ceranka 2007a) on the southern slopes of the Holy Cross Mts.

**Holothurioidea**: Holothurian sclerites have only been described from the Korytnica Clays (Walkiewicz 1977a). The assemblage comprises 32 forms belonging to the Stichopitidae, Calcilamnidae, Theeliidae, and Synaptitidae. The indirect proof of the presence of free moving holothurians in the Korytnica Clays are the otoliths of fishes of the family Carapidae (Radwańska 1992), which, in the majority, are commensals to these holothurians. Sclerites with a circular outline (Theeliidae, Synaptitidae) recognised in the Korytnica Clays should be ascribed to burrowing holothurians, whereas sclerites developed as perforated plates (Calcilamnidae) and elongated perforated rods (Stichopitidae) presumably belong to free moving species (Walkiewicz 1977a).

**Free-living crinoids**: The free-living crinoids have been described from Pińczów (Radwański 1977), Łagiewniki near Busko (Tasakowski 2013), and Korytnica (Radwańska 1987, 2003). In the “Leithakalke” type limestones at Pińczów only one centrodorsal with a radial ring of *Discometra rhodanica* (Fontannes, 1879) has hitherto been reported by Radwański (1977). The relatively rich assemblage of free living crinoids (Text-fig. 15) of the order Comatulida A.H. Clark, 1908, was described from the littoral facies of the Korytnica Basin (Radwańska 1987). This assemblage comprises *Sievertsella polonica* (Radwańska, 1987) of the family Comasteridae A.H. Clark, 1908 (Text-Fig. 1/1a–1c), and members of the family Himerometridae A.H. Clark, 1908; *S. polonica* was also described from Łagiewniki (Tasakowski 2013).

All recent genera of the subfamily Comasterinae and the family Himerometridae, are regarded as subtropical and tropical (Clark 1931), with both recent and
fossil representatives of these groups (Clark, 1931; de Castro Manso and Souza-Lima, 2017) confined to the Indo-Pacific realm. This general statement matches well the hitherto recognised tropical and/or subtropical conditions prevailing during the Middle Miocene (Badenian) development of the Korytnica Basin (e.g., Baluk and Radwański, 1977, 1979; Radwańska and Radwański, 1984).

**Asteroida:** Asteroids are relatively common in the Middle Miocene of the Fore-Carpathian Basin. They were reported from Nawodzice (Baluk and Radwański, 1968; Nosowska, 1997), Świniary (Radwański, 1973; Radwański and Wysocka, 2004), Korytnica (e.g., Kowalewski, 1973; Baluk and Radwański, 1977; Kaczmarska, 1987), and from Niskowa near Nowy Sącz (Baluk, 1970). Most of the asteroids (Text-fig. 16) are represented by isolated ossicles (see Kaczmarska, 1987; Nosowska, 1997), but entirely preserved skeletons (Astropecten navodicensis Nosowska, 1997 and Ceraster polonicus Nosowska, 1997) have been reported from Nawodzice (see Baluk and Radwański, 1968; Radwański, 1973; Nosowska, 1997), and interpreted as being buried in life position (Baluk and Radwański, 1968).

The assemblages (Kaczmarska, 1987; Nosowska, 1997) described from the Nawodzice Sands and the Korytnica Clay Clays compare well with the environmental conditions recognised in the Middle Miocene (Badenian) area, typified by shallow, open-marine, and/or subtropical waters, and show certain affinities to the recent asteroid fauna of the Indo-Pacific area.

**Echinoida:** Echinoids (Text-fig. 17) represent the most diversified and numerous group of Echinodermata recognised from the Middle Miocene of the Fore-Carpathian Basin. They were first reported by Roemer (1870), and subsequently studied by Kowalewski (1930), Radwański (1973), Baluk (1975), Maćzyńska (1977, 1979, 1987, 1988, 1993, 1996), and Ceranka (2007a).

In the rich material, consisting of over 37,000 complete tests and almost 10,000 fragments, Ceranka (2007a) recognized 20 species of 16 genera, 11 families and 7 orders. The assemblage is dominated by irregular forms of the genera Clypeaster Lamarck, 1801 and Echinocyamus van Pelssum, 1774, of the order Clypeasteroida. Fairly common are representatives of the order Psammechinoida, such as Schizaster L. Agassiz, 1835 and Spatangus Gray, 1825. Among regular forms, the commonest are the Cidaroida, of the genus Eucriaris Pomel, 1883, and Echinoida, of the genera Tripneustes L. Agassiz, 1841, Schizechinus L. Agassiz, 1841, and Psammechinus L. Agassiz et Desor, 1846.

Among all echinoid localities, noteworthy is the Świniary locality (see Radwański and Wysocka, 2004), with its echinoid species Psammechinus dubius (L. Agassiz, 1840) occurring gregariously (Text-fig. 17E) preserving spine canopies and Aristotle’s lanterns in position. All echinoids were jumbled together in haphazard positions, some of them inverted on the sole, showing that some were dead, but others still alive, when the layer was deposited (see Radwański and Wysocka, 2004, text-fig. 5). Such taphonomic premises indicate that the echinoids were transported in a huge mass of sand, from which live specimens could escape only occasionally (Radwański and Wysocka, 2004).

The Middle Miocene echinoids of the Fore-Carpathian Basin comprises widely distributed forms. Their ecological requirements (particularly the occurrence of the genera Clypeaster and Echinolampas) indicate warmer seas, including the tropical zone, and well-marked affinities with the Indo-Pacific bioprovince.

**Miocene Echinodermata of Ukraine (M. Górka)**

The first notes on Ukrainian echinoderms were by Niedźwiedzki (1879), Hilber (1882) and Lomnicki (1897, 1898). More serious studies started later, in the 1950s, on the echinoids by Szörényi (1953) and the asteroids by Kudrin (1957). More recently, Ukrainian echinoderms were studied by Radwański and Wysocka (2001), Kroh (2005), Radwański et al. (2006), Górka et al. (2012), and Wysocka et al. (2012), and a monographic description of the group was published by Radwański et al. (2014).

The Lower Badenian sandy deposits of the Mykolaiv Beds have yielded the most ubiquitous assemblage of echinoderms from western Ukraine. This facies is distributed in the vicinity of Lviv (see Wysocka, 2002) and stretches to the south as far as the Dniester valley and to the east where it reaches the town of Perekop.

The Upper Badenian Ternopil Beds are the second echinoid-bearing lithofacies of western Ukraine. Four echinoid species were described from the biohermal (reefal) limestones of this unit, and two species from the biodetrital unit. The same two species are present also in the topmost part of the coeval Pidhirtsi Beds (see Radwański et al. 2014).

**Starfish:** Only one species of starfish, *Astropecten forbesi* Heller, 1858, has been recorded from the Mykolaiv Beds. Although a few relatively well preserved specimens have been collected (Text-fig. 18A), larger remains of starfish, excepting isolated ossicles, are relatively rare (Radwański et al. 2014).

**Echinoids:** The taxonomic composition of the echinoid assemblage is closely related to the facies and age of the investigated strata. The assemblage from the Mykolaiv Beds is dominated by “irregular” echinoids (Text-fig. 18), with 10 species recognised: *Parascutella cf. paulensis* (L. Agassiz, 1841), *Parmulechinus* sp., *Echinocyamus* sp., *Clypeaster* sp., *Echinolampas* sp., *Conolampas* sp., *Spatangus cf. austriacus* Laube, 1871, *Hemipatagus ocellatus* (Defrance, 1827), *Echinocardium leopolitanum* Radwański et al. 2014 co-occur with “regular” sea urchins: *Stylocidaris polyacantha* (Reuss, 1860) and *Psammechinus dubius* (L. Agassiz, 1840).

On the contrary, the ratio between irregular and regular echinoids is more balanced in the Upper Badenian biohermal Ternopil Beds (Text-fig. 19), in which *Brissus unicolor* (Leske, 1778) and *Rhabdobrissus tarnopolensis* Radwański et al. 2014 co-occur with “regular” *Eucidaris desmoulinii* (Sismonda, 1842) and *Arbacina catenata* (Desor in L. Agassiz and Desor, 1846). Only two species have been recorded in the organodetrital Ternopil Beds and in the Pidhirtsi Beds: *Parascutella gibbercula* (De Serres, 1829) and *Spatangus cf. austriacus* Laube, 1871.

**Fish otoliths** [U. Radwańska]

Fish otoliths from the Middle Miocene of southern Poland were reported by Friedberg (1924), then by Chaine and Duvergier (1928), and Kowalewski (1930). Taxonomic studies were later performed by Šmigielska.
All fish otoliths (Text-fig. 20) from the Middle Miocene of southern Poland belong to the teleosts. Of the three types of paired (left and right) otoliths present in teleost fishes, all but one specimen of the investigated material represent the saccular otoliths, i.e., the sagittae. Radwańska (1992) monographed 147 species representing 50 families. Among the recognised families, the Myctophidae and Sparidae are characterised by high taxonomic diversity (14 species). The Congridae, Gadidae, Capridae, Triglidae, and Gobiidae are represented by at least 5 species. The total number of otolith specimens exceeds 22,600.

Taxonomically the richest otolith assemblage comes from the Korytnica Basin (Text-fig. 20), where it is represented by 105 species of 47 families (Radwańska 1992), with the majority of species confined to the littoral zone. Noteworthy in the assemblage are ubiquitous fish otoliths of the family Gobiidae, 1832. They were found together with tiny claws, which have recently been reported by Karasawa et al. (2014) as those representing the shrimp family Alpheidae Raubesque, 1815. This co-occurrence suggests a commensal relationship of gobies with snapping shrimps.

The second important group in the Korytnica Basin is the family Pomadasysidae, and primarily the species Brachydeuterus latior (Schubert, 1906), which is the most numerous (over 3,300 specimens) taxon recognised in the assemblage. This species is an important environmental indicator suggesting tropical and subtropical environments. Noteworthy in the assemblage are ubiquitous fish otoliths of the family Gobiidae, 1832. The Korytnica assemblage comprises also other tropical elements, such as those representing the shrimp family Alpheidae Raubesque, 1815. This co-occurrence suggests a commensal relationship of gobies with snapping shrimps.

The otolith assemblage from Nawodzice is relatively poor, both regarding its abundance and taxonomic diversity (10 species from 7 families). Nawodzice represents an extremely shallow-marine basin, evidently too shallow to be a permanent habitat of any fishes.

Relatively rich is the otolith assemblage from Rybnica (4,106 specimens), although it is taxonomically poorly diversified (33 species of 18 families). The most abundant is the family Gadidae. The ecological requirements of the assemblages from Nawodzice and Rybnica indicate the dominance of otoliths, which belonged to fishes that inhabited littoral waters with a normal salinity, but all the taxa recognised are much tolerant of thermal conditions, and thus cannot give precise information about the climate.

The otolith assemblage representing the deepest setting among the localities studied is the one from Bęczyn. It is a rather monotonous assemblage dominated by otoliths of the fishes of the family Myctophidae, which live currently mainly in warm, open oceanic waters.

Otolith assemblages have been described also from the Łychów area and from Niskowa, representing, however, rather poorly diversified faunas.

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