Middle Miocene serial killers: drilled gastropods from the south-western margin of the Central Paratethys, Croatia

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1. INTRODUCTION

Drilling predation is one of the most studied biotic interactions in the fossil record, and papers dealing with this topic have been extensively published since the 1960s (see KLOMPMAKER et al., 2019 and references therein). Information retrieved from the fossil traces, in this case preserved as drill holes, provide insight into the predator-prey relationship and other aspects of palaeoecology. According to KLOMPMAKER et al. (2019, Fig. 2), the majority of the research papers on drilling predation refer to molluscs; the majority of predation evidence occurs as a drill hole, with predominantly molluscs being both prey and predator. Fossil and extant gastropod families recorded as predators include the Capulidae (parastic), Cassidae, Eulimidae (parastic), Marginellidae, Muricidae, Nassariidae and Naticidae (e.g., CARRIKER, 1981; KABAT, 1990; ZLOTNIK, 2001; KLOMPMAKER et al., 2019 and references therein). The most studied shell drilling predators among them are the naticids and muricids.

While there are many papers dealing with predatory drillings on recent mollusc (bivalves and gastropods) shells, only a few of them document such behaviour in the middle Miocene (Badenian) of the Central Paratethys (e.g., HOFFMANN et al., 1974; GÖRÖG & SOMODY, 1988; KOWALEWSKI, 1990; HOFFMEISTER & KOWALEWSKI, 2001; ZLOTNIK, 2001; SAWYER & ZUSCHIN, 2011; GANIĆ et al., 2016). In the aforementioned predation studies on the middle Miocene molluscs from Central Paratethys, there are no data from Croatia. Here, we report for the first time drilling analysis on the middle Miocene gastropods from Croatia, on the south-western margin of the Central Paratethys.

The Croatian Natural History Museum (CNHM) in Zagreb hosts a large collection of the middle Miocene (Badenian) marine gastropods from the Zaprešić Brijeg locality near Samobor, 30 km west of Zagreb, Croatia (Fig. 1), where many of the shells have predation marks in the form of drill holes. The Zaprešić Brijeg fossils (gastropods and bivalves) were originally collected at the beginning of the 20th century from the highly fossiliferous „light yellow sandstones” outcropping in local vineyards of Zaprešić Brijeg (also known as Zaprešić Breg, Zaprešički Vinograd or Vrhovčak). Bulk sampling was organized by the curator of the Samobor Museum, Stepan Orešković, and performed by the local residents (e.g., PAVLOVSKY, 1957). The collection was later housed in the CNHM.

Our objective was to count drilling frequencies in the Zaprešić Brijeg material, and to compare them to the available data from the middle Miocene (Badenian) of the Central Paratethys (e.g., GÖRÖG & SOMODY, 1988; HOFFMEISTER & KOWALEWSKI, 2001; SAWYER & ZUSCHIN, 2011; GANIĆ et al., 2016) (Fig. 2). In this study the focus is on: (1) the frequencies of the drill holes present on the gastropods; and (2) their probable predator(s).

2. GEOLOGICAL SETTING

The Zaprešić Brijeg locality lies west of the centre of Samobor, and 2 km south-east of the village of Dubrava, towards the village of Vrhovčak, around 30 km from Zagreb (Fig. 1).

This area is part of the rift-type North Croatian Basin (NCB), located in the south-western part of the Pannonian Basin System (PBS) surrounded by the Alps, Carpathians and Dinarides (e.g., PAVELIĆ & KOVAČIĆ, 2018) (Fig. 2). Palaeogeographically it
belonged to the south-western margin of the Central Paratethys (e.g., RÖGL, 1998; PAVELIĆ, 2001; HARZHAUSER & PILLER, 2007; KOVÁČ et al., 2007, 2017, 2018; PILLER et al., 2007; PAVELIĆ & KOVAČIĆ, 2018). Miocene marine transgressions did not cover the entire Croatian part of the Pannonian Basin System simultaneously, and the basement is therefore unconformably overlain by Neogene deposits of various ages (e.g., PAVELIĆ & KOVAČIĆ 2018 and references therein).

The presented Zaprešić Brijeg locality is part of the Samoborsko gorje Mt. (Fig. 1). The Neogene basement of the Samoborsko gorje Mt. is diverse and most commonly composed of Triassic dolomites (GORJANOVIĆ-KRAMBERGER, 1894; ŠUKLJE, 1929), with, at some localities a volcanogenic-sedimentary complex of Cretaceous age, and less often a Palaeogene clastic-carbonate complex (ŠIKIĆ K. et al., 1978, 1979; VRSALJKO et al., 2005 and references therein). The oldest Miocene sediments are Badenian (Langhian) deposits overlying Triassic dolomites. VRSALJKO et al. (2005) describe algal shallow-water limestones as the most common lithotype at the beginning of the transgression. They are overlain by silty marls. BAKRAČ et al. (2010) recognized two Badenian marine transgressive-regressive cycles (TB 2.4 and TB 2.5) in deposits of the neighbouring Vrhovčak section. The Late Badenian marine transgressive-regressive cycle TB 2.5 could be a consequence of a short, but intense marine transgression due to eustatic sea level rise with the opening of the connection between the Central Paratethys and the Indo-Pacific Ocean (VRSALJKO et al., 2005; KOVÁČ et al., 2007; BARTOL et al., 2014).

History of the Zaprešić Brijeg locality investigations

The geology of the Zaprešić Brijeg locality was first described by GORJANOVIĆ-KRAMBERGER (1894). The basement of the Neogene sediments is composed of Triassic dolomites, and the roof sediments are defined as the “Leithakalk”. Below the “Leithakalk”, white and light yellow sandstones occur containing a large amount of mollusc shells and other fossils. GORJANOVIĆ-KRAMBERGER (1894, 1896) collected and determined the fossil fauna (foraminifera, molluscs, balanids, fish and mammals), and assumed that these Neogene sediments belonged to
Table 1. Analyzed gastropod families from the CNHM collection with drilling predation data. Abbreviations: n = number of specimens; DHS = successfully drilled shells; UDHS = unsuccessfully drilled shells; DF (%) = drilling frequency; PE (%) = prey effectiveness.

| FAMILY       | n  | DHS | UDHS | DF (%) | PE (%) |
|--------------|----|-----|------|--------|--------|
| Trochidae    | 1063| 633 | 113  | 5.72   | 15.15  |
| Pisanidae    | 1   | 0   | 0    | 0      | 0      |
| Neritidae    | 32  | 0   | 0    | 0      | 0      |
| Melanopsidae | 2   | 0   | 0    | 0      | 0      |
| Pachychilidae| 11  | 0   | 0    | 0      | 0      |
| Cerithiidae  | 508 | 65  | 15   | 12.8   | 18.75  |
| Littorinidae | 1   | 0   | 0    | 0      | 0      |
| Scaliidae    | 1   | 0   | 0    | 0      | 0      |
| Potamididae  | 4459| 201 | 36   | 4.51   | 15.19  |
| Turritellidae| 1253| 181 | 28   | 14.45  | 13.4   |
| Rissoidae    | 3   | 0   | 0    | 0      | 0      |
| Seguenziidae | 2   | 0   | 0    | 0      | 0      |
| Zebroidae    | 1   | 0   | 0    | 0      | 0      |
| Strombidae   | 8   | 0   | 0    | 0      | 0      |
| Aporrhaidae  | 13  | 1   | 0    | 7.69   | 50     |
| Calyptraeidae| 14  | 0   | 0    | 0      | 0      |
| Naticidae    | 90  | 5   | 4    | 5.56   | 44.44  |
| Triviidae    | 1   | 0   | 0    | 0      | 0      |
| Eratoidea    | 6   | 0   | 0    | 0      | 0      |
| Cassidae     | 2   | 0   | 0    | 0      | 0      |
| Charoniidae  | 6   | 0   | 0    | 0      | 0      |
| Epitonidae   | 1   | 0   | 0    | 0      | 0      |
| Eulimidæ    | 1   | 0   | 0    | 0      | 0      |
| Muricidae    | 150 | 10  | 2    | 6.67   | 16.67  |
| Tucidae      | 17  | 1   | 0    | 5.88   | 0      |
| Babyloniidae| 1   | 0   | 0    | 0      | 0      |
| Buccinidae   | 2   | 1   | 0    | 50     | 0      |
| Nassariidae  | 2428| 34  | 4    | 1.4    | 10.53  |
| Columbellidae| 11  | 0   | 0    | 0      | 0      |
| Melongenidae| 11  | 1   | 0    | 9.09   | 0      |
| Fasciolariidae| 5  | 0   | 0    | 11.11  | 0      |
| Voluitidae   | 18  | 0   | 1    | 0      | 100    |
| Mitridae     | 3   | 0   | 0    | 0      | 0      |
| Costellariida| 8   | 0   | 0    | 0      | 0      |
| Olivida      | 1   | 0   | 0    | 0      | 0      |
| Ancillaridae | 70  | 2   | 2    | 2.86   | 50     |
| Cancellariida| 14  | 2   | 0    | 14.29  | 0      |
| Conidae      | 12  | 0   | 0    | 0      | 0      |
| Borsoniidae  | 18  | 1   | 0    | 5.56   | 0      |
| Drillidae    | 13  | 3   | 1    | 23.08  | 25     |
| Clavatulidae | 1813| 122 | 17   | 6.73   | 12.23  |
| Turridae     | 10  | 2   | 1    | 20     | 33.33  |
| Terebridae   | 4   | 1   | 1    | 25     | 50     |
| Pyramidellida| 1   | 0   | 0    | 0      | 0      |
| Acteonidae   | 15  | 0   | 0    | 0      | 0      |
| Ringiculidae | 15  | 0   | 0    | 0      | 0      |
| Cylichnidae  | 5   | 0   | 0    | 0      | 0      |
| Retusidae    | 1   | 0   | 0    | 0      | 0      |

the „Mediterranean age“ (= Badenian), and compared them with the Grund Formation near Vienna, which he considered to be contemporaneous with the Zaprešić Brijeg sediments based on the fossil fauna and distribution of the gastropod Pereirea gervaisi (VEZIAN, 1856) (previously Pereiraia gervaisi) in Croatia, Slovenia and Hungary. The Grund Formation belongs to the middle Miocene – Langhian, that is, in the Central Paratethys chronostratigraphy, the Badenian (e.g., HARZHAUSER et al., 2003, CORIC et al., 2004). The collected fauna also contains a freshwater species, Vitta picta (FERUSSAC, 1823) (old name Neritina picta FERUSSAC, 1823), which was probably brought into the marine sediments by river flows. ŠUKLJE (1929) pointed out that the most abundant fossils at Zaprešić Brijeg are gastropods, with the genera Pleurotomaria and Cerithium as the most numerous, and confirmed the conclusions of GORJANOVIĆ-KRAMBERGER (1894, 1896). PAVLOVSKY (1957, 1960) continued the determinations of the Zaprešić Brijeg gastropods, and assigned the stratigraphic age of the locality to the middle Miocene – Langhian / lower Badenian (in original: „Lower Tortonian“), comparing it to the “Grund” layers. BAJRAKTAREVIĆ (1978), based on the Zaprešić Brijeg foraminifera, concluded that the age of the sediment is middle Miocene – Serravalian / upper Badenian (in the original: „Upper Tortonian“ (Ammonia (= Rotalia) beccarii zone)). Based on the presence of the genera Cerithium, Pirenella and Nassa in the Zaprešić Brijeg assemblage, VRSALJKO (1990) interpreted the palaeoenvironment as shallow marine with sporadic brackish and freshwater areas. MADERIC (2017) described naticid and muricid drillings from Zaprešić Brijeg within his BSc Thesis, based on gastropods housed at the Faculty of Science, University of Zagreb.

The age of the Zaprešić Brijeg horizons is still not precisely determined. We visited the area several times, in the hope of finding the fossiliferous horizon and conducting further research, but these deposits are no longer available at the surface due to the expansion of settlements, plantations and vineyards. However, due to the presence of the marine gastropod species Pereirea gervaisi (VEZIAN, 1856) in the assemblage, the age could be further restricted to the upper part of the middle Badenian or to the upper Badenian (Upper Langhian – Serravalian), close to the N5 - NN6 nannofossil zone boundary (BARTOL et al., 2014; MIKUŽ & ŠOSTER, 2014).

3. MATERIALS AND METHODS

Analyses of the drilled gastropod shells from Zaprešić Brijeg were made on 11063 specimens from the CNHM collections, inventory numbers 1419-1544, and 10305-10414 (Table 1, Supplement 1). The analyzed fauna was taxonomically categorized into families according to the World Register of Marine Species (WoRMS). Revision of the gastropod fauna in the collection is beyond the scope of this work. This material comprises representatives of marine (44), marine and brackish (2), and brackish (2) families. Four families live as infauna (Naticidae, Turritellidae, Aporrhaidae and Nassariidae), while the other families live as epifauna. Looking at the feeding type of the gastropods, they are interpreted as carnivores, suspension feeders, detritivores and grazers. Two families from freshwater and terrestrial environments were excluded from the analyses due to their non-marine origin.

The first step was to distinguish the drilled from non-drilled gastropod shells. We counted the shells with one and more drillings visible, and based on that, the shells were divided into two groups, drilled and non-drilled gastropod shells. Drilled gastro-
pod shells were further categorized as (a) successfully drilled shells (DHS), (b) unsuccessfully drilled shells (UDHS), and (c) multiply drilled shells (MULT-DHS) (Supplement 1). A successfully drilled gastropod shell has one complete drill hole with one entrance in the shell (Figs. 3A, 3B, 3C, 3D), and an unsuccessfully drilled shell (Figs. 3E, 3F) has one incomplete drill hole with only the outer layer of the shell drilled, and no entrance hole. Multiply drilled shells have more than one successful and/or unsuccessful drill holes (Figs. 3G, 3H, 3I, 3J), showing the traces of predator drillings and parasitic holes on the shell. In this study we focused on the successfully drilled shells (DHS) and unsuccessfully drilled shells (UDHS) showing the predator drillings made by gastropods, not taking into consideration multiply drilled shells and their agents.

Analysis of predation intensity was determined as drilling frequency (DF) (Lower Taxon Frequency (LTF) according to KOWALEWSKI, 2002), which aims to estimate the frequency of interactions for a given taxon of prey (here family) measuring the rate of prey mortality due to the drilling predation (e.g., KELLEY & HANSEN, 1993; KOWALEWSKI, 2002; SAWYER & ZUSCHIN, 2011; HUTCHINGS & HERBERT, 2013; GANIĆ et al., 2016; PAHARI et al., 2016; SARKAR et al., 2016):

$$\text{DF (LTF)} = \frac{D_K}{N_K},$$

where K is a lower taxon target (here family), $D_K$ is the number of specimens of that taxon that contain one successful predation trace (drill hole) and $N_K$ is the total number of specimens of that taxon in the sample (after KOWALEWSKI, 2002). In the text we use the term drilling frequency (DF) for the LTF (Lower Taxon Frequency).

The estimate that provides some measure of the predator’s failure is called the escalation parameter (KOWALEWSKI, 2002). The relative frequency of failed attacks is often referred to as „prey effectiveness“, which documents traces of unsuccessful attacks (e.g., repair scar or healed drill hole) as described in KOWALEWSKI (2002):

$$P_E = \frac{T_F}{T_T},$$

where $P_E$ is prey effectiveness computed as $T_F$, the number of a particular trace fossil of predatory origin that records failed attacks (e.g., the number of incomplete and healed drill holes) divided by $T_T$, the total number of these trace fossils (e.g., the total number of drill holes) (e.g., KOWALEWSKI, 2002; SAWYER & ZUSCHIN, 2011; HUTCHINGS & HERBERT, 2013; GANIĆ et al., 2016; PAHARI et al., 2016; SARKAR et al., 2016).

To indicate a probable predator, we categorized drill holes as made by naticid, muricid or an undeterminable agent, following the descriptions in e.g., HOFFMANN et al., 1974; CARRIKER, 1981; KABAT, 1990; KOWALEWSKI, 2002, 2004. Naticid drillings are parabolic (straight or oblique) and the drill hole is countersunk (outer drill hole diameter is larger than the inner diameter). Muricid drillings are cylindrical, with nearly straight edges.

Abbreviations used in the figures and tables in the following sections are as follows (alphabetically):

- DF (%) = drilling frequency; DH = drill hole; DHS = successfully drilled shells; MULT-DH = multiple drill holes; MULT-DHS = multiply drilled shells; n = number of specimens in the sample; PE (%) = prey effectiveness; UDHS = unsuccessfully drilled shells.

4. RESULTS

Among the 11063 analysed middle Miocene (Badenian) gastropod shells from the Zaprešić Brijeg locality from the CNHM, 1024 shells were drilled (DHS, UDHS and MULT-DHS), among which 746 shells belong to DHS and UDHS, 6.7% of the studied sample (Supplement 1). The distribution of these categories is shown in Fig. 4. Within the drilled sample (9.3% of the total), 61.8% exhibit one complete drill hole.

The most numerous families exhibiting drillings in the sample are the Potamididae (4459 specimens), Nassariidae (2428 specimens), Clavatulidae (1813 specimens), Turritellidae (1253 specimens), Cerithiidae (508 specimens), Muricidae (150 specimens).
mens), and Naticidae (90 specimens) (Fig. 5, Supplement 2). The Potamididae have the highest number of the successfully drilled shells (201), followed by the Turritellidae (181), Clavatulidae (122), Cerithiidae (65), and other families (Fig. 5A, Supplement 2). Potamididae live in the marine and brackish environments as epifauna. Others (Clavatulidae and Cerithiidae) live in marine environments as epifauna, and the Turritellidae are a marine semi-infauna (Supplement 1).

Figure 5B. shows the distribution of the unsuccessfully drilled shells, again with the Potamididae family showing the highest number of unsuccessfully drilled shells (36) as well as the highest number of specimens. The number of unsuccessfully drilled shells is relatively small compared to the successfully drilled shells (Fig. 5A). Other families are represented with less than 30 undrilled shells (Supplements 1 and 2).

The average rate of prey mortality due to the drilling predation (DF) for the whole analyzed gastropod sample is 5.72% (Table 1, Supplement 1). When we look at the drilling frequencies of the most numerous families in the sample (Supplement 2), DF ranges from 1.4% (Nassariidae) to 14.45% (Turritellidae).
Relative frequency of failed attacks (PE) for the analyzed sample is 15.15% (Table 1, Supplement 1). Looking at the numerously represented families in the sample (Supplement 2), PE ranges from 10.53% (Nassariidae) to 44.44% (Naticidae).

Drilling differences between infauna and epifauna in the analyzed sample are shown in Supplements 3 and 4. The average DF of the infauna (5.84%) is slightly higher than the DF of the epifauna (5.66%), while the average PE of the infauna (14.34%) is lower than the PE of the epifauna (15.57%) indicating that the infauna is less successful at resisting the drilling attack. This result may be affected by the biased preservation of epifaunal gastropod shells, while epifaunal shells are less resistant to various taphonomic processes.

Categorization of the gastropods based on their feeding type and drilling occurrence is shown in Supplement 5. Most gastropods represented in the sample are grazers (5029 specimens). The highest DF is observed in the suspension feeders (14.29%), represented mostly by turritellids, and the lowest among the carnivores (3.89%). The highest PE occurs within the detritivores (50%), and the lowest within suspension feeders (13.4%).

To analyze which prey predators prefer, we looked at the shape of the drill holes and categorized them as naticid drill holes, muricid drill holes and undeterminable, where the attacker is uncertain (Fig. 6, Supplements 6 and 7). Fig. 6A shows the distribution of the categorized drill holes in the sample within the most represented families with successfully drilled gastropod shells. As shown in Fig. 6A, naticid drillings prevail, especially on the shells of the Turritellidae. Examination of the unsuccessfully drilled shells (Fig. 6B), shows that the attacks are mostly undeterminable, and, if they are determinable, they are categorized as naticid drillings, with the majority of the prey belonging to the Turritellidae.

5. DISCUSSION

Analyses of drilling predation can offer insight into the palaeoecology of fossil molluscs (e.g., KOWALEWSKI, 2002). Available published papers on drilling predation describing the Central Paratethys Miocene molluscs as prey and predators, geographically cover the present day area of Poland, Bulgaria, Hungary, Austria, Slovakia and Serbia (Table 2 and references therein).
Comparison between the Miocene marine deposits of Europe (Boreal and Paratethys provinces – from the present areas of Austria, Slovakia and Hungary) was analyzed by Hoffmeister & Kowalewski (2001), giving the spatial patterns on predation on the Burgudian and Langhian molluscs. The authors conclude that numerous studies on predator-prey relationship are based on local palaeoecological interpretations and/or long-term temporal trends. Since predation data can significantly vary from one locality or province to another due to the spatial variation in predation, Hoffmeister & Kowalewski (2001) suggest that analyses should concentrate more toward documenting spatial variation in predation analyzing multiple sites (e.g., Sawyer & Zuschin, 2011). In this presented predation analysis, it was not possible to include more sites into the palaeoecological study (as well as in e.g., Ganić et al., 2016). In the published studies on the Central Paratethys Badenian gastropod predation so far, data from Croatia are missing. In the following section we describe and compare drilling predation on gastropods from the Zaprešić Brije locality with the available data obtained from the other trace fossil studies on the Badenian gastropods from the Central Paratethys (Görög & Somody, 1988; Sawyer & Zuschin, 2011, and Ganić et al., 2016).

5.1. Gastropod Drilling Predation in the Central Paratethys during the Badenian

Drilling predation studies on the middle Miocene (Badenian) gastropods from the Central Paratethys have been made on the collections from Hungary (Görög & Somody, 1988) and bulk samples from the Vienna Basin (Austria and Slovakia, in Sawyer & Zuschin, 2011) and Serbia (Ganić et al., 2016). Görög & Somody (1988) describe drilling and settlement traces on Badenian gastropods from the sand pit at Várpalota in Hungary. The Várpalota fauna (further in the text we refer to it as V-H), with 6620 specimens (after Table 1 in Görög & Somody (1988)) is dominated by the herbivorous gastropod families Potamididae and Turritellidae. The main predators are the naticids and muricids. Sawyer & Zuschin (2011) analyze drilling predation on molluscan assemblages from the lower and middle Miocene (upper Burgudian/Karpathian, Langhian-lower Serravalian/Badenian) localities in Austria and Slovakia (Vienna Basin; further in the text we refer to it as VB). The bulk sample consists of 39234 mollusc shells (bivalves, gastropods, scaphopods and polyplacophorans), and the most numerous are gastropods (22292 specimens, of which 9101 are the Badenian gastropods). The authors consider the muricids and naticids as the most likely predators. Mollusc predation from the southern part of the Central Paratethys is described in Ganić et al. (2016), on the middle Miocene (Badenian) bulk sample from the Rakovica stream sands near Belgrade (further in the text we refer to it as RSB-S). Authors analyze 2301 shells of molluscs (gastropods and bivalves; 1617 gastropod specimens) drillings, and conclude that carnivorous gastropods are dominant in the sample and the main predators were muricids and naticids.

In Supplement 8, we show the abundance comparison between Badenian gastropods from the Zaprešić Brije locality (further in the text is referred to as ZBL-C), with part of the recorded Badenian gastropods from V-H and RSB-S, based on both published and data presented herein for gastropods present in all three localities. The abundance from VB is not shown because that study includes both Karpathian and Badenian localities, with given abundance for only part of the gastropods represented in the Badenian deposits. The most numerous gastropods in ZBL-C and V-H belong to the Potamididae, with the most abundant number of specimens belonging to the genera Pirenella and Terebratalia (Supplement 8), while in RSB-S the potamids are not recorded. The Nassariidae and Turritellidae families are also abundant at the ZBL-C and V-H localities, and are present in the RSB-S. Among the turritellids the most numerous is the genus Turritella, and from the Nassariidae, the genus Nassarius is the most abundant at the ZBL-C, Dorsanum at V-H, and Nassia at RSB-S. When we compare the distribution of the gastropod predators, naticids and muricids, naticids prevail at V-H and RSB-S (genus Natica), while in ZBL-C prevail muricids (genus Ocinebrina). The most likely predators at all the described localities, including the VB, are the Naticidae and Muricidae. It should be noted that when we compare the present fossil fauna, differences could be possible due to the gastropods classification. For example, in some cases one genus belongs to one family in one register, and in another register it is assigned to the other family. The numerical ratio of complete analyzed gastropod samples from the above described Central Paratethys localities is shown in Figure 7.

As presented in Sawyer & Zuschin (2011, Table 3), mollusc drilling frequency (DF) for the Karpathian (upper Burgudian) and Badenian (Langhian and lower Serravalian) of the Central Paratethys is 7.5%, which is lower in comparison with the neighbouring Miocene areas of the Boreal province (24.5%) and the entire Paratethys (14%) (Hoffmeister & Kowalewski, 2001). The Badenian gastropods of VB display a DF of 9.2%. The DF of the RSB-S gastropods is a similar value, 9.65%. In contrast, DF in the ZBL-C gastropods is significantly lower: 5.72% (Table 3). There are no data for the DF and PE for the V-H fauna, only the list and number of damaged specimens (see Table 1 and 2 in Görög & Somody, 1988).

Results for the PE again differ between the compared areas. The PE of the VB Badenian gastropods at 2.2% (Sawyer & Zuschin, 2011, Fig. 6B) is lower than the PE of ZBL-C which equals 15.15%, and the PE of 7.18% of the RSB-S gastropods (Table 3).

Similarities between the DF and PE of the most drilled gastropod shells between ZBL-C, VB and RSB-S are shown in Table 3. Comparing the drillings of the most represented gastropods from the ZBL-C, with the neighbouring Badenian areas in the Paratethys, the highest DF in the marginal area of the Central Paratethys is in the Turritellidae (14.45%, present paper, and 17.33% in RSB-S). Cerithiidae also have a higher DF value in the marginal areas of the Central Paratethys (12.8% in present paper, and 14.41% in RSB-S). The predator gastropods, Naticidae and

![Figure 7. Ratio of gastropod numbers between the compared localities.](Image 308x76 to 550x215)
Muricidae, have lower DF values in the marginal Central Paratethys areas, than the average of the Central Paratethys (Table 3). The families with the higher DF values in the marginal area have lower values than the Central Paratethys values based on the results from the localities in Austria and Slovakia, and vice versa.

Comparison of the PE shown in Table 3, indicates that the Badenian gastropods from the Central Paratethys marginal areas have higher PE than the average of Central Paratethys. The highest PE in the marginal area have Naticidae (44.44% in present study, and 25% in RSB-S).

As shown in SAWYER & ZUSCHIN (2011), DFs could fluctuate within similar environments at a single locality. It should be mentioned that comparison on drilling predation between the ZBL-C (this paper), VB and RSB-S is based on uneven gastropod abundances (see Figure 7). The number of specimens within families greatly differ at the mentioned study areas. Also, SAWYER & ZUSCHIN (2011) made an analyses on multiple sites, and GANIĆ et al. (2016) and this paper only on one locality. Differences in values between naticids and muricids (predator gastropods) could also be the consequence of a smaller amount of predator naticids in marginal areas (90 specimens ZBL-C; 49 specimens in RSB-S) than in the Central Paratethys average (403 specimens in VB).

### 5.2. Probable Predators of Zaprešić Brijeg Gastropods

Muricids and naticids are the most likely gastropod predators in the Zaprešić Brijeg. Both groups use a highly specialized accessory boring organ (ABO) which aids the drilling process chemically by secreting an acid which affects prey shell layers by dissolving them and excavates a hole through which the prey tissue is extracted (e.g., CARRIKER, 1981; KABAT, 1990). However, they can also attack on the surface (e.g., HOFFMAN et al., 1974; PAHARI et al., 2016). Naticids do not attack in groups. As described in KABAT (1990), the sequence of prey capture events by extant naticid includes detection of the prey, evaluation, seizure, covering and immobilization of the prey, wrapping in the dilated foot of the naticid, dragging for some distance and covering into sand for commencement of borning. As naticids enfold the prey with their foot and look for the boring place, multiple naticid drillings on one shell could be the consequence of the prolonged time of boring or the fact that naticids do not recognize if the prey is alive or dead and they drill shells which are already drilled (e.g., HOFFMANN et al., 1974). It is possible that naticids just produce multiple drill holes during a single attack (e.g., KABAT, 1990), or as described in GÖRÖG & SOMODY (1988) the double borings may have been produced by two predators or by one which needed more space to extract the food.

Our assumption here is that the predator drilling was fatal to the prey, and shells having one successful drill hole can possibly support this, although a possibility remains that the prey could escape the predator (KOWALEWSKI, 1990, 2004). Successfully drilled shells with regenerated drillings (repair scar on the shell) and/or incomplete drillings can be a sign of an unfinished attack from a predator (e.g., CHATTOPADHYAY & BAUMILLER, 2007, 2010; HUTCHINGS & HERBERT 2013).

Other potential drillers, except Muricidae and Naticidae, could be the gastropods Marginellidae, Capulidae, Nassariidae, and Cassidae, as well as cephalopods, who are all capable of drilling holes in the shells (e.g., KABAT, 1990; ZLOTNIK, 2001; KLOMPMAKER et al., 2019). Marginellidae are not present in the analyzed sample, so we can discard them as other potential drillers at the studied site. Capulidae are ectoparasitic symbionts of molluscs and echinoderms, with cylindrical drill holes similar to muricid drilling and leave an attachment scar on the host shell (KABAT, 1990). Capulidae are not present in the analyzed sample either, and an attachment scar in the surrounding area of the drill hole is not recorded, so we can also discard them. Nassariidae (2428 specimens) are mobile scavengers and predators, but in the recent nassariids the drill hole diameter is not larger than 0.25 mm, and drillers are mostly juvenile nassariids (ZLOTNIK, 2001). However, we can exclude nassariids due to the fact that they are mostly herbivores or scavengers, in addition to the fact that the studied drillings do not correspond to nassariid drum hole measurements (e.g., KABAT, 1990; KOWALEWSKI, 1990; ZLOTNIK, 2001). Cassidae (2 specimens) mostly drill echinoid tests (e.g., KABAT, 1990), and rarely bivalve shells (KABAT, 1990; ZLOTNIK, 2001), so they are not the potential drillers here. Cephalopods (octopuses) are shell boring predators of molluscs, with drillings of an irregular or oval outline and extremely small inner drill hole diameter (e.g., KABAT, 1990). We have recorded smaller and larger drill hole diameters in the analyzed sample, but the drillings were circular so we dismiss cephalopods as possible predators here.

The highest DF from the Zaprešić Brijeg occurs in the family Turritellidae, with naticid drill holes as the most abundant drillings (Fig. 6A, 6B, Supplements 6 and 7). In the successfully drilled shells there are drill holes that look like muricid drillings, as well as undeterminable drill holes (Fig. 6A). In the unsuccessfully drilled shells, the drillings are recognized as naticid or undeterminable (Fig. 6B). Looking at the analyses of the presented gastropod sample, the other naticid choices of prey were mostly potamidids, clavatulids and cerithiids (Fig. 6A), although these
three families live as epifauna, and the naticids are characterized as predators of infaunal organisms. However, it was noted that muricids also attacked shallow-burrowing bivalves in more offshore habitats, explained by the hypothesis of atypical feeding behaviour, residual nature of the fossil assemblage, smaller fossilization potential of the epifauna, and misidentification of naticid drillings for muricid ones (KOWALEWSKI, 1990). Most of the drill holes in the unsuccessfully drilled shells belong again to three epifaunal families (potamidids, clavatulids and cerithiids), beside turritellids, and most of the drillings are undeterminable in origin, except the majority of drillings on turritellids (Fig. 6B). As shown in Fig. 6, there are more naticid drillings in the sample than the muricid ones, although the number of naticids in the sample (90) is smaller than the muricids (150). Although muricids attack more epifaunal organisms and gastropods, this sample is characterized by most of the drillings being either of naticid or undeterminable origin. Possible reasons for the prevalence of naticids over muricid drillings are: (1) originally there were more naticids in the sample – due to the taphonomic processes they were not present at the locality because of their fragile shells that are more easily carried away in suspension than the muricid shells; (2) a possible rapid burial event – living as infauna, naticids find more easily their way to the surface than the muricids who are epifauna; and (3) mistaking muricid drillings for the naticids, and vice versa, and not recognizing the undeterminable drillings. Other organisms that can produce naticid- and muricid-like drill holes include nematodes, flatworms, octopods and other gastropods (KOWALEWSKI, 2004). At the neighbouring V-H locality GÖRÖG & SOMODY (1988) give an analyses of the naticid and muricid borings. Although the Muricidae are less abundant in the sample, their drillings prevail, with most drillings on Potamididae (see Table 2 in GÖRÖG & SOMODY, 1988). Naticid drillings prevail on the Turritellidae and Nassariidae, which is in accordance with the predators mode of life. Distinguishing naticid from muricid drilling solely on morphological criteria can be difficult because the drilling mechanisms are similar (ZLOTNIK, 2001). Transitions from naticid- to muricid-type of drilling have been noticed (HOFFMANN et al., 1974), and also, the same species can drill holes in different shapes (KOWALEWSKI, 2004). Therefore the determination of the drill hole shape in our sample should be considered as only a probable indication of predator identity, especially given the great number of drill holes categorized as undeterminable (Fig. 6).

5.3. Comparison with the Recent Assemblages
Most of the recent research papers dealing with the origin of drillings in gastropod shells are based on laboratory experiments, and focused mostly on predator behaviour (e.g., CHATTOPADHYAY & BAUMILLER, 2007; CASEY & CHATTOPADHYAY, 2008, CHATTOPADHYAY et al., 2014). ZLOTNIK (2001 and references therein) mentioned that the laboratory induced behaviour (e.g., by starvation) differs from nassariid drilling behaviour in nature. In the samples collected from the Northern Adriatic Sea, turritellids were the most commonly drilled (SAWYER & ZUSCHIN, 2010). They were mostly drilled by naticids, but since they live buried directly under the sediment surface, they are more susceptible to both muricid and naticid drillers (SAWYER & ZUSCHIN, 2010 and references therein), as it is also noted in our sample (Fig. 6, Supplements 6 and 7), and e.g., TULL & BÖHNING-GAENSE (1993). ALLMON (2011) described the ecology of the Turritellidae, pointing out that recent turritellids are mainly sessile semi-infaunal suspension feeders living in shallow waters of full-marine salinity and temperatures below 20°C, but also occur as an active group which can crawl on the surface and thrive at a variety of depths, salinities and temperatures, and with probably changed environmental tolerances through history. As pointed out by ALLMON (2011) turritellid drilled shells indicate that their most frequent drilling predators are naticids, which is in concordance with the fossil data presented here and compared to the other studied samples from the Central Paratethys area.

6. CONCLUSIONS
Information retrieved from the drilled gastropod shells provides a closer insight into the biotic interaction in this part of the Panonian Basin System during the Badenian. The Zaprešić Brijeg collections offer the possibility of understanding such biotic interaction due to the abundance of marine gastropod species and the abundance of the drilled shells.

There are several gastropod families recognized as predators, which attack other molluscs, and the results of the presented data analysis point to the Naticidae and Muricidae as the most probable predators. In the studied sample, epifaunal Potamididae have the highest number of the drilled shells, with the majority of them being successful. Comparing the DF of infauna versus epifauna, it is slightly higher in the case of infauna (5.84%, and 5.66%, respectively). DF comparison between the feeding type is the highest for the suspension feeders (14.29%), represented mostly by the turritellids, and is lowest for carnivores (3.89%). The average rate of the gastropod DF at Zaprešić Brijeg is 5.72%, which is lower than the Badenian gastropods DF average of 9.2% for Central Paratethys as given in SAWYER & ZUSCHIN (2011, Fig. 6A), and gastropods DF of 9.65% from the south-eastern margin of the Central Paratethys (GANIČ et al., 2016). Average gastropods PE of the presented assemblage is 15.15%, which is significantly higher than the PE of 2.2% in the Central Paratethys (SAWYER & ZUSCHIN, 2011, Fig. 6B), and PE at the south-eastern margin of the Central Paratethys of 7.18% by GANIČ et al. (2016).

Comparison of the presented data with previously published data from the neighbouring Paratethys areas shows that the Badenian gastropod families with higher DF in the marginal area of the Central Paratethys have lower DF than the same families in the Central Paratethys. In addition, the Badenian gastropods from marginal areas have higher PE than the average of the Central Paratethys. The highest DF at the Zaprešić Brijeg is recorded in the infaunal suspension feeding Turritellidae, with the most abundant naticid drillings. This is in concordance with their life habitat and known predators from both the fossil and recent data.

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