The oldest post-Paleozoic (Ladinian, Triassic) brachiopods from the Betic Range, SE Spain

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Triassic brachiopods from the Betic Range were unknown hitherto. Herein we describe the first brachiopod occurrences in the early Ladinian of this domain referable to a new genus and species *Misunithyris goyi* derived from three localities of the south-Iberian Triassic outcrops. The analysis of internal and external characters of this brachiopod allowed to characterize systematically and biogeographically this fauna in a chronostratigraphic interval when the paucity of brachiopod records is attributable to the entire peri-Iberian epicontinental platform system established in the westernmost Tethyan margin. The new record is endemic to the Betic Range and represents a new faunal constituent of the multicostate zeillerids stock. This fauna inhabited the epicontinental seas of the Sephardic bioprovince since a closer affinity with the low-latitude Tethyan assemblages is revealed. The possible linkage of the Triassic stock with the Early–Middle Jurassic multicostate zeillerid representatives suggests feasible phylogenetic relationships between both groups.

Key words: Brachiopoda, Zeillerioidea, palaeobiogeography, Triassic, Ladinian, Spain.

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Introduction

The Triassic Period represented one of the most significant episodes in the evolutionary history of brachiopods, inasmuch as severe biotic turnovers took place between the Permian–Triassic extinction event and the end-Triassic extinction boundary, two of the big five extinctions during Phanerozoic (Raup and Sepkoski 1982). Most of the diverse and abundant Paleozoic brachiopod faunas became extinct in the end-Permian extinction (Gould and Calloway 1980; Sepkoski 1996). After this biotic crisis, marine benthic biota underwent a new radiation in the late Early–Middle Triassic (cf. Hallam and Wignall 1997; Payne et al. 2004), contributing to the definitive replacement by the new evolutionary Mesozoic fauna. Brachiopods recovery started in the Olenekian–early Anisian (Chen et al. 2005, 2015; Shen et al. 2006; Zakharov and Popov 2014; Ke et al. 2016; Gaetani 2016), and experienced a new peak at community-level for the brachiopod clades during the Mesozoic (cf. Ager 1965; Sandy 1995; Dulai 2001; Tomašových 2006, and references therein).

In this timespan, brachiopod communities proliferated in the shallow epicontinental seas in several paleogeographical domains of the Paleo- and Neo-Tethys Ocean (e.g., Bittner 1890; Siblík 1972, 1998; Dagens 1974; Vörös and Pálfy 1989; Pálfy 1991, 2003; Pálfy and Török 1992; Senkowiczowa and Popiel-Barczyk 1993; Kaim 1997; Torti and Angiolini 1997; Hagdorn and Sandy 1998; Benatov 2001; Feldman 2005, 2013), but in the westernmost Tethyan basins the record is very scarce. Prior to the widely accredited Early Jurassic bloom in brachiopod diversity (e.g., Jiménez de Cisneros 1923; Baeza-Carratalá 2011, 2013; Baeza-Carratalá and García Joral 2012), no other Mesozoic brachiopods have been reported from elsewhere in the Betic Range hitherto. The only pre-Jurassic brachiopod evidence in this area is restricted to the indefinite exceptional record of a few cross-sections in the Early Devonian of the Nevada-Filábride complex (Laborda-López et al. 2015).

In the South-Iberian Paleomargin, collecting effort in numerous Triassic localities over the last decades resulted in extensive collections of Ladinian nautiloids and ammonoids (Pérez-Valera 2015; Pérez-Valera et al. 2017). In contrast,
only three of these localities yielded brachiopod specimens suitable for taxonomy and having appropriate stratigraphic framework based on ammonoids (Pérez-Valera 2015; Pérez-Valera et al. 2017). It has also substantiated the record of brachiopods in the Triassic from the Betic Range for the first time, resulting in a new genus and species, allowing detailed descriptions of its external and internal structure. This record in the Betic Range is also remarkable because it makes available new data among the sparse brachiopod occurrences in this chronostratigraphic interval. In fact, this paucity of brachiopod records is not confined to the Betic Range, but is attributable to the entire peri-Iberian epicontinental platform system established in the westernmost Neo-Tethys margin. Brachiopod occurrences into these Triassic epicontinental seas around Iberia have only been reported in the Iberian Range through the monospecific Lingularia bursts (Márquez-Aliaga et al. 1999, 2007; Escudero-Mozo et al. 2015; Giannetti et al. 2017), and four specimens attributed to Coenothyris sp. (Márquez-Aliaga et al. 1994; Escudero-Mozo et al. 2015) and C. vulgaris (Schmidt 1935), all of them mainly Ladinian in age, plus the sporadic Anisian Mentzelia (M. mentzeli, M. sp.), and Koeveskalliina koeveskalyensis, mainly from Menorca and the Catalonian Coastal Range (Schmidt 1935; Escudero-Mozo et al. 2015).

Finally, this study enables the establishment of palaeobiogeographical affinities, by analyzing the similarity of the recorded fauna with those assemblages from several well-documented basins from the Germanic, Tethyan, and Sephardic bioprovinces.

**Institutional abbreviations.**—DCTMA, Earth and Environmental Science Department collections, University of Alicante, Spain.

**Other abbreviations.**—L, length; W, width.

### Material and methods

Amidst the profuse Triassic outcrops prospected in the Betic Range by the authors, only a total number of six brachiopods were collected and studied from three localities (Fig. 1). Taxonomic supra-generic assignments follow the published determinations: mainly by KAESLER and Selden (1997–2007). Recent supplementary data (MANCEÑIDO 1993a; BAEZA-CARRALÁ and GARCÍA JORAL 2014) are considered for the systematic arrangement into the zeilleriids. The main biometric parameters (in mm) and indices were measured for the taxonomic analysis when it was possible. Internal structure was studied using the conventional method of taking acetate peels after making transverse serial sections. The distance between serial sections was 0.1 mm. High resolution microphotographs of acetate peels were taken using an optical microscope (Nikon CF60 E600POL). All specimens were...
coated with magnesium oxide prior to photographing. All specimens figured and acetate peels are deposited in the collections of Earth and Environmental Sciences Department (University of Alicante, Spain).

The ammonoid zonal scheme used is based on the standard zones proposed by Balini et al. (2010) modified by recent chronosтратigraphic data from Pérez-Valera (2015) for the Betic domain. Paleobiogeographical analysis performed is plotted on a paleomap slightly modified after Pérez-López and Pérez-Valera (2007). It is worth noting that the paleobiogeographic relationships are inferred on the basis of the unique record in the Triassic Betic Domain hitherto; consequently the accuracy of the analysis would be improved if further brachiopod faunas were to be found in the future.

Geographical and geological setting

The Betic-Rif orogen is an extremely arcuate orogen that comprises the south of Spain and the north of Morocco, both connected by the so-called Gibraltar Arc, a structure formed in the Miocene (Platt et al. 1995; Balanyá et al. 2007). The north branch of this orogen is the Betic Range (Fig. 1), traditionally divided in two different domains: the Internal and External zones. Three tectonic complexes constitute the Internal Zones: Nevado-Filábride Complex, Alpujárride Complex, and Maláguide Complex, structurally stacked from bottom to top. These complexes are limited by the so-called “South-Iberian Triassic” (Fig. 1; Pérez-López 1998). All Triassic units have been included in facies and epicontinental carbonates of Muschelkalk facies, with pelagic fossils (e.g., ammonoids) containing brachiopods, can represent the deeper part of the next parasequence, and therefore, the alternation of marly limestone beds and marls to together with bivalves, ammonoids, and nautiloids in a cm-thick alternation of marls and thin-bedded marly limestone levels (Fig. 2). In Talave and Calasparra sections, this part is located over a regional unconformity that represents the maximum flooding surface, with presence of abundant cephalopods (ammonoids and nautiloids) and other representative fauna (Pérez-López et al. 2005). This member is constituted by a succession of marly limestone beds, lutes, thin bedded limestone and bioclastic limestone beds (tempesites) showing a progradational pattern in a high-stand stage (Pérez-Valera and Pérez-López 2008).

The stratigraphic arrangement and facies present in the Siles and Cehegin formations show the evolution of an epicontinental platform that was emplaced over the siliciclastic deposit of Buntsandstein facies (Arroyo Hurtado sections, Fig. 2). Brachiopods have been found occasionally together with bivalves, ammonoids, and nautiloids in a cm-thick alternation of marls and thin-bedded marly limestone levels (Fig. 2). In Talave and Calasparra sections, this part of the succession is situated over a bioclastic key-level that probably mark the end of a regressive parasequence and therefore, the alternation of marly limestone beds and marls with pelagic fossils (e.g., ammonoids) containing brachiopods, can represent the deeper part of the next parasequence, as proposed by Pérez-Valera and Pérez-López (2007). In the Arroyo Hurtado section, bioclastic levels with borings at the top mark the end of the regressive parasequences, and brachiopods have been found in levels together with ammonoids, developed over the bioclastic beds.

The recent finding of a large number of ammonoids has allowed a new biostratigraphical framework in the Muschelkalk carbonates of the Betic Range in which three ammonite biochronozones have been distinguished (Fig. 2), all belonging to the Ladinian stage (Pérez-Valera J.A. 2005; Pérez-Valera et al. 2011; Pérez-Valera 2015): (i) Eoprotrachyceras curionii Zone (lower part of Fassanian, lower Ladinian), which is subdivided into the Eoprotrachyceras curionii and Gevanites awadi subchronozones; (ii) Gevanites epigonus Zone (upper part of Fassanian, lower Ladinian); and (iii) Protrachyceras hispanicum Zone (lower part of Longobardian, upper Ladi-
Fig. 2. Ladinian lithostratigraphical sections showing the occurrences of *Misunithyris goyi* in Talave, Calasparra, and Arroyo Hurtado sections. Ammonite zones and subzones after Pérez-Valera (2015).
nian). This is consistent with the previous data provided by bivalves, conodonts, and nautiloids of Ladinian age (Márquez-Aliaga et al. 2001; Plascencia et al. 2007; Pérez-Valera et al. 2017). All the levels containing brachiopods in the three studied outcrops are situated in the Gevanites epigonus Zone (Fig. 2) and they can be therefore attributed to the uppermost Fassanian (lower Ladinian).

**Systematic palaeontology**

Phylum Brachiopoda Duméril, 1805  
Subphylum Rhychonelliformea Williams, Carlson, and Brunton, 1996  
Class Rhychonellata Williams, Carlson, and Brunton, 1996  
Order Terebratulida Waagen, 1883  
Suborder Terebratellidina Muir-Wood, 1955  
Superfamily Zeillerioidea Allan, 1940  
?Family Zeilleriidae Schuchert in Schuchert and Brunton, 1996  
Genus Misunithyris nov.

*Etymology:* From Mišňař, ancient name of the current Mundo River; in the Mundo River valley, the most significant outcrops of specimens from which this genus is erected are found.

*Type species:* Misunithyris goyi sp. nov. (by monotypy); see below.

*Diagnosis.*—As for the type species by monotypy.

*Remarks.*—The supra-generic systematic arrangement of *Misunithyris* is debatable, depending on the diagnostic criteria selected. Exceptional concurrence of several internal and external features can make attributable this genus to different groups within Terebratulida. On the one hand, the new genus herein erected shows common features with the superfamily Dielasmatoidea Schuchert, 1913, which includes some genera with enveloped dental plates and cardinal process. However, the assignment to Dielasmatoidea can be problematic because *Misunithyris* does not evidence a dielasmoid-type brachidium (sensu Dagys 1974 or Smirnova 2008). In this sense, supplementary elements such as median ridges or vertical plates (even not forming part in the development of brachidium, as stated by Dagys 1974 and Smirnova 2008) have not been observed in the internal structure of *Misunithyris.* Furthermore, some Dielasmatoidea representatives show septum-supported architectures and often short-looped developments (e.g., Adygella Dagys, 1959; Dielasmina Waagen, 1882; Tunethyris Calzada, Peybernes, Kamoun, and Youssef, 1994). In addition, crural bases are given off dorsally instead the distinctive crural progress revealed in *Misunithyris.*

Within Dielasmatoidea, higher similarity was expected with the anteriorly multicostate stock attributed to the Permian *Dielasmina* Waagen, 1882 and *Hemiptychina* Waagen, 1882 and the Permian–Triassic *Costoconcha* Jin, Sun, and Ye, 1979, since besides the anterior ribbed pattern, all genera share quite a few beak features and the presence of cardinal process, but the rest of the internal structure is totally different, mainly referred to the dental plates and the crural development, which is clearly a brachidium-supported structure in the first stages.

Another dielasmatoid morphotype, widely distributed and to some extent contemporary with *Misunithyris* is represented by *Coenothyris* Douvillé, 1879. The species of this Triassic genus display comparably dorsal plates, undeveloped or fused with the thickened shell wall, evident cardinal process, and long loop. Conversely, it evidences an initial septum-supported structure, notable septalium and, especially, the external features (such as smooth shell, often with strong uniplication) are entirely different (e.g., Popiel-Barczyk and Senkowiczowa 1989; Török 1993; Senkowiczowa and Popiel-Barczyk 1996; Kaim 1997; Pálfy 2003; Feldman 2005).

Arrangement within Zeillerioidea Allan, 1940 is the most plausible option, mainly because of the presence of dental lamellae, the well-developed and large dorsal median septum, and a clear zeilleroid-type brachidium (sensu Smirnova 2008) with a long-looped development, not connected to the median septum. However, some characters do not fully agree with the various families up to now determined in Zeillerioidea.

*Misunithyris* shares with the family Eudessidae Muir-Wood, 1965 the envelopment of dental plates, the presence of a cardinal process and most of the beak features. The most remarkable difference to Eudessidae are in a short dorsal median septum and the crural bases given off dorsally, as well as the usual growth of a median cardinal plate of the later group and not perceived in *Misunithyris.* Some external diagnostic criteria are very different as well, since even showing Eudessidae multicostate shells, *Misunithyris* shows a marginal ribbing pattern instead the entire multicostate shell-length of Eudessidae. Folding pattern is also unrelated as the conspicuous dorsal sulcus developed by *Misunithyris* is not shared with any Eudessidae representatives.

The affinities with the family Zeilleriidae are found in several subfamilies. Some representatives of the subfamily Vectellinae Baker, 2006 exhibit a long-looped development and cardinal process as a knob or poorly developed callus. However, Middle–Late Triassic representatives of this subfamily are very different in both external and internal features to the new established genus. It is the case of *Fletcherithyroides* Dagys, 1977, *Aulacothyroides* Dagys, 1965, and *Parantiptychia* Xu and Liu, 1983, consisting on smooth morphotypes, also showing a long stage of crura supported by septal pillars. Probably the clearest affinity in this subfamily is found in the Upper Triassic—lowermost Jurassic *Tauromenia* Seguenza, 1885, due to comparable anterior ribbing pattern and the beak features, as well as the well-developed dorsal median septum (Alméras et al. 2007; Baeza-Carratalá and García Joral 2012) and long-looped brachidium (Alméras et al. 2007).
Finally, there are several arguments for and against assigning *Misunithyris* to the subfamily Zeilleriinae Schuchert, 1929 as was defined in Kaesler and Selden (1997–2007). The septum is clearly well-developed and the brachidium evidences a long-looped progression. Conversely, the presence of cardinal process is atypical in this subfamily, although some genera show primitive lobes as massive callus or small knobs (e.g., *Antipychina* Zittel, 1880; *Kolymithyris* Dagys, 1965). On the other hand, some of them also reveal envelo ped dental lamellae instead the typical strong and unen veled dental plates characteristic of this subfamily. As for *Tauromenia*, the closer external affinities are found in the anteriorly ribbed representatives of this subfamily, i.e., *Calpella* Owen and Rose, 1997 and *Parathyridina* Schuchert and Le Vene, 1929, the internal structure of which remains poorly known except for the presence of a prominent medio septum (e.g., Cooper 1983; Baker 2006; Alméras et al. 2010b). In this sense, very close internal and external affinities have been recognized with the recently erected multicostate genus *Menathyris* Feldman, 2013, except for the cardinal process and the anterior folding pattern.

It must be kept in mind that the subfamily Zeilleriinae Schuchert, 1929 was recently split into several subfamilies by Baeza-Carratalá and García Joral (2014) on the basis of the hinge plates-crural bases relationship, thus resulting three subfamilies (Aulacothyrinae Babanova, 1964, Zeilleriinae Schuchert, 1929, and Securininae Baeza-Carratalá and García Joral, 2014). Attending to this criterion, *Misunithyris* clearly shows a *Bakonyithyris*-type pattern and might be arranged into Aulacothyrinae. However, the representatives of Aulacothyrinae (sensu *Menathyris* Feldman, 2013) have not evidenced cardinal process so far, and the progress of dental plates is rather different.

Summarizing, the arrangement of *Misunithyris* into family Zeilleridae is the best plausible determination, but does not fully agree with the currently defined subfamilies. Combining external and internal features, it seems to be closer to the multico state zeilleriid stock, such as *Menathyris* Feldman, 2013, *Calpella* Owen and Rose, 1997, or *Tauromenia* Segu enza, 1885. A new subfamily might be erected, either with *Misunithyris* as monotypic taxon, or together with the aforementioned genera, but further studies of internal structures in their representatives will be required to establish the validity of this approach, emphasizing examination of cardinal process and brachidium architecture in addition to the already known shared features.

**Stratigraphic and geographic range.**—*Misunithyris* is so far a monospecific genus recorded in the *Gevanites epigonus* Chronozone of the Fassanian (lower Ladinian, Middle Triassic) from the Betic Range (Fig. 2).

*Misunithyris goyi* sp. nov.

Fig. 3.

**Etymology.** In a tribute to Antonio Goy (Complutense University of Madrid), prominent Triassic and Jurassic palaeontologist, to whom the authors are very indebted for long term teaching and collaboration.

**Type material.** Holotype: DCTMA-BQ-TA1.2 (Fig. 3F; Table 1) from the Talave section. Paratype: DCTMA-BQ-AH2.2 (Figs. 3B, 4, 5). Intraspecific variability is shown through the paratypes (Fig. 3) and measured in Table 1.

<table>
<thead>
<tr>
<th>Specimen</th>
<th>L</th>
<th>W</th>
<th>T</th>
<th>W/L</th>
<th>T/L</th>
<th>T/W</th>
<th>R</th>
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<td>17.56</td>
<td>12.78</td>
<td>10.83</td>
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<td>0.62</td>
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<td>13.00</td>
<td>12.44</td>
<td>8.20</td>
<td>0.96</td>
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<td>0.66</td>
<td>9</td>
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<tr>
<td>BQ-TA1.1</td>
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<td>16.60</td>
<td>~8.44</td>
<td>0.95</td>
<td>0.48</td>
<td>0.51</td>
<td>9</td>
<td>9p</td>
</tr>
<tr>
<td>BQ-TA1.2</td>
<td>28.34</td>
<td>24.43</td>
<td>17.01</td>
<td>0.86</td>
<td>0.60</td>
<td>0.70</td>
<td>13</td>
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<td>21.74</td>
<td>15.77</td>
<td>15.57</td>
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<td>0.71</td>
<td>0.98</td>
<td>11</td>
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<td>16.64</td>
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<td>0.81</td>
<td>0.65</td>
<td>0.80</td>
<td>11</td>
<td>9p2s</td>
</tr>
</tbody>
</table>

**Type locality.** Talave section (External Betic Zone, SE Spain).

**Type horizon.** Upper member of Siles Formation, Fassanian (early Ladinian); *Gevanites epigonus* Biochronozone.

**Material.**—Six specimens (BQ-CL1.1, BQ-CL1.2, BQ-TA1.1, BQ-TA1.2, BQ-AH2.1, and BQ-AH2.2), including type material, with diverse state of preservation. All individuals show articulated but generally fragmented and encrusted valves. Internal recrystallization is visible in two specimens.

**Diagnosis.**—Medium-sized and ventribiconvex zeilleriid, pyriform in outline with strong sub-labiate beak and epityrid/permesothyrid foramen. The lateral commissure is straight and the anterior one is clearly sulcate. Sharp ribs (8–13) appearing only near the anterior margin (3–5 in the median sulcus), often bifurcate and occasionally intercalate. Deltidial plates present and well-developed striate cardinal process. Crural bases are given off dorsally and hinge plates initially fused and subparallel. Dorsal septum long, exceeding the length of the brachidium. Long descending branches, distally with ventral progression.

**Description.**—**External features** (Fig. 3): Medium-sized ventribiconvex shells (Table 1), pyriform to subbovidial in dorsal outline. W/L ratio is rather variable, but all specimens show L > W. Thickness is about 3/5 of the length. Maximum convexity lies in the posterior third while maximum width lies in the anterior third of the shell. Maximum thickness is observed near the mid-length. The beak is massive, sub-labiate, and strong, erect to slightly incurved, with a medium-sized epityrid to permesothyrid foramen; the beak ridges are well-perceived and blunt, developing short and narrow interareas. The lateral commissure is straight and the anterior one is clearly sulcate, with a wide and arcuate dorsal median sinus. Sulcation is generally shallow, being more pronounced in the larger specimens.

Ribbing pattern is distinctive of this species. Ribs are absent in the posterior third of the shell; costation consists of 8 to 13 ribs on each valve, 3–5 of which can be present in the median sulcus. Ribs are wide, strong, and square in cross-section, becoming stronger and sharper near the anterior margin. They are often bifurcate; the thicker specimens
possess a set of weaker, intercalated ribs. Costation is more evident in the ventral valve and on the flanks of the dorsal one. In the median dorsal sulcus, ribs are only noticed from the mid-length onwards. Concentric, dense and strong growth lines are present on the entire surface giving a reticulate aspect to the shell.

**Internal structure** (Figs. 4, 5): This species shows a sub-octagonal to ovoidal delthyrial cavity in cross-section where the remains of a short pedicle collar and well-developed deltoidal plates are visible. Dental plates are difficult to distinguish as they are enveloped in a thickened-shell wall (Fig. 5). They are short and subparallel. Hinge teeth are massive with crenulations, inserted in broad, shallow, and also crenulated sockets; small denticula are also discernible. It has a noticeable cardinal process, striated and raised by a high cardinal platform where a central cavity is located under the myophore (Fig. 5). Hinge plates are initially horizontal, fused and subparallel. The dorsal median septum is long and well-developed up to 2/3 of the shell. A short elliptical septalium is discernible. Crural bases are located in the ventral area of the hinge plates but an incipient dorsal thickening emerges clearly showing an early dorsal development (Fig. 4), suggesting a *Bakonyithyris*-type hinge plate/crural base inter-relation (sensu Baeza-Carratalá and García Joral 2014). The descending branches of the brachidium show subparallel and vertical plates and they are developed in the commissural plane in the posterior third of the shell, acquiring a slight ventral development anteriorly. The transverse band is not visible.

**Fig. 3.** Intraspécific variability of zeillerid brachiopod *Misunithyris goyi* gen. et sp. nov. of the lower Ladinian from the South-Iberian Triassic platform, the Arroyo Hurtado (A, B), Calasparra (C, E), Talave (D, F) sections; in dorsal (A1–F1), anterior (A2–F2), and lateral (A3–F3) views. A. BQ-AH2.1. B. BQ-AH2.2, sectioned in the present work. C. BQ-CL1.1. D. BQ-TA1.1. E. BQ-CL1.2. F. BQ-TA1.2, holotype. All specimens were coated with magnesium oxide.
Fig. 4. Internal structure of zeillerid brachiopod *Misunithyris goyi* gen. et sp. nov. (BQ-AH2.2) from the lower Ladinian of the South-Iberian Triassic platform, the Arroyo Hurtado section. **A.** Transverse serial sections through the specimen (distance from the beak in mm). **B.** Microphotographs of acetate peels showing the hinge plates-crural bases relationship: crural bases are located in the ventral part of the hinge plates but a dorsal thickening emerges towards an early dorsal development. Section at 4.10 mm (*B₁*) and 4.30 mm (*B₂*) from the apex. Abbreviations: cb, crural bases; dp, deltidial plates; ld, dental lamellae; ms, median septum; pc, cardinal process; sp, septalium.
Remarks.—In addition to the main biometric ratios (Table 1), intraspecific variability mainly lies in the number of ribs present in each valve, depending on the bifurcation or intercalation. The smooth posterior stage is also variable between ½ and ¼ of the shell. A single specimen shows an exceptionally wide sulcus giving to the anterior commissure an almost rectimarginate aspect, but analyzing in detail this individual the sinus can be noticed from the early stages in the posterior areas of the shell. Thus, even width of sulcus is rather stable, it ranges from mid-width in the BQ-TA1.2 specimen up to wider developments (BQ-AH2.1 and BQ-CL1.1) occupying nearly the entire anterior margin. On the other hand, maximum width can be shifted toward the anterior third, resulting in a more trigonal dorsal outline.

As it is discussed above in the supraspecific assignment, this species shows high singularity in the external and internal traits, probably because it is an endemic taxon. The closer affinity is shown with the Lower Jurassic (mostly anteriorly) multicostulate zeilleriids *Tauromenia polymorpha*, *T. brevicostata*, and *Fimbriothyris* spp. (Seguenza 1885; Dubar 1942; Elmi et al. 2003; Alméras et al. 2007; Baeza-Carratalá and García Joral 2012), mainly recorded in the southern part of Western Tethys and even with the Middle Jurassic Eudesiidae representatives recorded in both Tethyan margins and the Mid-East platforms (e.g., Alméras 1987; Cooper 1989; Mukherjee et al. 2000; Alméras et al. 2010b), having in common with these last taxa several internal features (cardinal process present in several taxa, long median septum) but not sharing the brachidium architecture and the ribs covering the whole surface of the shell in the Middle Jurassic taxa.

Among the Triassic representatives, the closer affinities are found in *Menathyris wilsoni* Feldman, 2013 from Israel, virtually contemporary with the Betic material, sharing the sulcate shape of the shell, the marginal ribbing pattern and comparable internal structure, except for the cardinal process, absent in *M. wilsoni*; however, sulcus is shallower and
narrower, anterior commissure is generally rectimarginate and ribs are simple without bifurcation or intercalation in such species. Curious similarity concerning the internal structure is observed in one specimen of *Coenothyris* with a central umbonal chamber in the cardinal process, similarly to the Betic material, and several *Aulacothyridae* species developing wide cardinal process, all of them depicted by Senkowiczowa and Popiel-Barczyk (1996) in the Anisian–Ladinian from Poland.

Stratigraphic and geographic range.—In the Betic Range, this new species has been recorded together with biostatigraphical marker of the Gevanites epigonus Biochronzone (Pérez-Valera F. 2005; Pérez-Valera et al. 2011; Pérez-Valera and Popiel-Barczyk 1996) in the Ladinian of the Fassanian (early Ladinian).

Discussion

Paleobiogeographical implications.—During the Triassic Period, brachiopod bioprovinces are mainly defined for the Late Triassic, where the taxonomic diversity was higher, being therefore the biochoremas more definite, with up to five high-rank biochoremas subdivided in several bioprovinces (Dagys 1993; Manceñido 2002). In the Ladinian, brachiopods were less diverse and their record is very rare in several regions and therefore their biogeographical differentiation pattern is unclear (Dagys 1993). In fact, in several Tethyan basins such as the Caucasian, brachiopods are totally absent during the Ladinian (Ruban 2006a, b) leading to postulate a severe crisis on brachiopod fauna likely as a results of an unsteady recovery after the end-Permian event (Ruban 2017).

The low-latitude Tethyan brachiopod faunas or Southern Tethyan Sub-Realm (cf. Dagys 1993; Manceñido 2002) comprise several minor-rank biochoremas usually referred not only for establishing the brachiopod fauna affinities but also for cephalopods, conodonts, bivalves and other groups (Hirsch 1987; Parnc et al. 1985; Pérez-López et al. 2003; Pérez-Valera et al. 2015). Thus, the affinity of the faunal components from these Southern Tethyan basins can be discriminated, at least, into the Germanic, Tethyan, and Sephardic bioprovinces (Fig. 6).

The attribution of the studied material to a new genus and species seems to point out to the presence of endemic brachiopod fauna in the Betic Range. On the other hand, independently from the argued supra-generic assignment of *Misunithyris*, this genus undoubtedly corresponds to an unusual and scarce morphotype within the multicostrate zellerids. The unique morphology of this group makes it a suitable tool for paleobiogeographic correlations and biostatigraphical calibrations (e.g., Ager and Walley 1977; Almèras 1987; Baeza-Carratalá and García Joral 2012). In this sense, *Misunithyris* finds its closer counterpart within the multicostrate zellerids in the coeval *Menathyris* Feldman, 2013, if we compare the internal and external diagnostic features of both genera. *Menathyris* was recorded in the early Ladinian from Negev (southern Israel), integrated into the Sephardic bioprovince (Feldman 2013).

Thus, a Sephardic affinity can be suggested for the new brachiopod fauna recorded in the Betic Range, since in addition to the possible link to the Israeli zelleriids (Fig. 6), it has to be taken into account that, in the Ladinian, this bioprovince is typified by the low faunal diversity and high degree of endemicity developed in epicontinental seas (Page 1996), just as in the Betic case. This endemicity is not only applicable for the brachiopods but also for the nektoplanktonic biota affinities (e.g., Tozer 1981; Pérez-Valera 2015). The geographical/reproductive isolation assumed for the nektoplanktonic fauna would become even more evident in groups with short larval stages such as brachiopods.

Conversely to the Sephardic province, the Germanic and Tethyan bioprovinces are more prolific in brachiopod diversity in the entire Triassic (Dagys 1993), recording a variety of rynchonellid and spiriferinid taxa and, in the Ladinian, even emerging new taxonomic groups (Dagys 1993) such as Pennospiriferiidae or Angustothyrididae. Significant blooms of Koninckinacea, Dielasmatoidea, Neorettiziinae, Aulacothyropsidae, and Vectellinae also occurred in this timespan in both bioprovinces (e.g., Bittner 1890, 1892; Siblik 1972, 1998; Dagys 1974; Popiel-Barczyk and Senkowiczowa 1989; Vörös and Pálfy 1989; Pálfy 1991, 2003; Pálfy and Tórók 1992; Senkowiczowa and Popiel-Barczyk 1993; Kaim 1997; Torti and Angiolini 1997; Hagdorn and Sandy 1998; Benatov 2001; Klug et al. 2005; Ruban 2010).

As with ammonites, the biogeographic boundaries of Sephardic bioprovince in the Ladinian are unclear toward the North of the Betic Range (Pérez-Valera 2016). In the closest peri-Iberian epicontinental platforms, Middle Triassic brachiopods are only represented in the Levantine Sector of the Iberian Range through the monospecific pervasive *Lingulario* busts (Mármol-Aliaga et al. 1999, 2007; Escudero-Mozo et al. 2015; Giannetti et al. 2017), and very few specimens attributed to *Coenothyris* (Schmidt 1935; Márquez-Aliaga et al. 1994; Escudero-Mozo et al. 2015) all of them mainly Ladinian in age, plus the Anesian *Mentzelia* from Menorca and the Catalonian Coastal Range (Schmidt 1935; Escudero-Mozo et al. 2015).

Consequently, it can be inferred that influences from the epicontinental seas of the Germanic or Tethyan provinces could occasionally have taken place in the Northern peri-Iberian platforms by means the immigrants *Coenothyris* and *Mentzelia* (Fig. 6). Nevertheless, both *Coenothyris* and *Mentzelia* can be regarded as ubiquitous taxa since, even they are more common in the Alpine Muschelkalk facies, they are world-wide distributed in basins such as Northern Caucasus (Ruban 2010), Hungary (Tórók 1993; Pálfy 2003); Southern Alps of NW-Italy (Torti and Angiolini 1997); Northern Alps and Slovakia (Siblik 1972, 1988, 2001), Poland (Popiel-Barczyk and Senkowiczowa 1989; Senkowiczowa and Popiel-Barczyk 1993; Kaim 1997; Szulc 2000), and even in the Sephardic bioprovince from Israel (Feldman 2005).

In this sense, it has to be taken into account that Feldman...
(2005) separated the endemic *Coenothyris oweni* from those of the Germanic bioprovince. For this reason, the occurrence of both genera (*Coenothyris* and *Mentzelia*) in the peri-Iberian platform system is not fully indicative to establish the boundary of the Sephardic vs. Tethyan/Germanic influences (Fig. 6).

Finally, it is substantiated that the multicostate zeilleriids counterparts of *Misunithyris* exceeded the end-Triassic extinction event. In the Early Jurassic the genera *Tauromenia* and *Calpella* continued inhabiting the wide and shallow platforms developed in low-latitude seas of the Western Tethys as deduced from their record in the High/Middle Atlas (Dubar 1942), Western Algeria (Elmi et al. 2003; Alméras et al. 2007), Eastern Sicily and Calabria (Di Stefano 1887; Taddei-Ruggiero and Vörös 1987), Apennines (Pozza 1989), Gibraltar (Owen and Rose 1997), Cephalonia (Manceñido 1993a) and Eastern Subbetic (Baeza-Carratalá and García Joral 2012). These occurrences are mostly uppermost Sinemurian–lowermost Pliensbachian in age. The exceptional record of *Tauromenia* from the South Armorican Massif in the *Uptonia jamesoni* Zone is slightly younger (Alméras et al. 2010a). On the other hand, the genus *Fimbriothyris*, besides its Mediterranean distribution, shows a more profuse presence in the NW-European platforms from the late Pliensbachian onwards (Deslongchamps 1856; Alméras 1987).

In fact, this restricted geographical distribution makes this group a useful paleobiogeographical tool. In the Early Jurassic, multicostate zeilleriids distribution reveals an E-W dispersion pattern from the intra-Tethyan basins to the Atlantic areas (Ager and Walley 1977; Baeza-Carratalá and García Joral 2012). Thus, it can be emphasized that multicostate zeilleriids were not recorded in the NW-European
basins during Triassic and earliest Jurassic times (except for the youngest South Armoricain Massif record, Alméras et al. 2010a, which fit well in the E-W route). They disappeared from the Mediterranean Betic Ranges when the epicontinental/epi-oceanic duality became more evident from the early Pliensbachian onwards. The ulterior Middle Jurassic diversification of these forms and their migration to the NW-European platforms typified by the fully costate genera \textit{Menathyris} and \textit{Fauré 2008). An entirely ribbed and bifurcate extreme form of the authors (except for Delance 1974) seem to agree in \textit{Fimbriothyris} as possible ancestor or, at least, as the possible Lower Jurassic counterpart of this clade (Ager and Walley 1977; Alméras 1987; Mukherjee et al. 2000; Alméras and Fauré 2008). An entirely ribbed and bifurcate extreme form was depicted by Deslongchamps (1863) within the intra-specific variability of the genus \textit{Fimbriothyris}, which was postulated as possible predecessor of this group (Alméras 1987; Alméras and Fauré 2008).

Regardless their mono-versus polyphyletic origin, most of the authors (except for Delance 1974) seem to agree in \textit{Fimbriothyris} as possible ancestor or, at least, as the possible Lower Jurassic counterpart of this clade (Ager and Walley 1977; Alméras 1987; Mukherjee et al. 2000; Alméras and Fauré 2008). An entirely ribbed and bifurcate extreme form was depicted by Deslongchamps (1863) within the intra-specific variability of the genus \textit{Fimbriothyris}, which was postulated as possible predecessor of this group (Alméras 1987; Alméras and Fauré 2008).

The new genus herein erected shares significant internal and external features such as the ribbing pattern, cardinal process, or dental plates structure with this possible evolutionary lineage (Table 2). Taking into account the above paleobiogeographical analysis (Fig. 6), from an anagenetic perspective it can be suggested that the Triassic stock made up by \textit{Misunithyris} gen. nov. (probably together with \textit{Menathyris} Feldman, 2013) might represent the precursor linkage taxon of this lineage which had its initial homeland in the low-latitude Tethyan basins during Triassic, such as the Sephardic bioprovince.

**Conclusions**

We report first record of brachiopod fauna in the Triassic from the Betic Range, as well as the oldest occurrence amidst Mesozoic brachiopod assemblages in this domain.
The new finding is assigned to a new genus and species (*Misunithyris goyi*), which reveals a very characteristic external and internal structure, emphasizing the distinctive dental lamellas, a prominent cardinal process and its ribbing pattern. These features together with the septal development and the brachidium architecture make feasible its arrangement within the superfamily Zeillerioidea, evidencing high similarity with the multicostate zeilleriids stock. The most plausible supra-generic position is found within family Zeilleriidae, although several features are also shared with several zeilleriid families, not discarding the erection of a new subfamily in a future including other taxa with similar structure.

The occurrence of *Misunithyris goyi* in the Betic Range has been accurately established in the *Gevanites epigonus* Biochronzone of the Fassanian (lower Ladinian) and it can be used therefore as a valuable correlation and calibration tool.

A Sephardic paleobiogeographic affinity is suggested, since this fauna is so far endemic to the Betic Range and finds its possible counterparts within the multicostate zeilleriids integrated into the epicontinental platforms of the Sephardic bioprovince.

From an anagenetic perspective, it is suggested that this new genus might represent the precursor stock of the Early–Middle Jurassic multicostate zeilleriids, probably originated in the low-latitude Tethyan basins (such as the Sephardic bioprovince) during the Triassic.

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