Delamaran trilobites from the La Laja Formation, San Juan, Argentina

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Abstract. The La Laja Formation lies exposed almost continuously along the western margin of two hill ranges, i.e., the Sierra Chica de Zonda and Sierra de Villicum. This limestone unit bears a remarkable fauna of Cambrian trilobites. In this work we analyse the Delamaran trilobite fauna from the upper part of the El Estero Member and the lower third of the Soldano Member of the La Laja Formation. Taxa described are Glossopleura walcotti Poulsen, Glossopleura lodensis (Clark), Fieldaspis sp., and Prozacanthoides zondaensis n. sp. The Poliella denticulata, Mexicella mexicana?, and Glossopleura walcotti Zones are documented; these had been previously included in a hiatus. Trilobites-based correlations are established between the La Laja Formation and olistoliths of the Western Precordillera. Additional correlations are also suggested between the Precordillera and different localities in Laurentia. Finally, we discuss the biogeographic implications of the described material. Delamaran taxa reported from the La Laja Formation are characteristic and so far restricted to Laurentia. Analysis of the trilobite fauna from this unit supports the allochthonous origin of the Precordillera.

Resumen. Trilobites delamarianos de la Formación La Laja, San Juan, Argentina. La Formación La Laja aflora de forma prácticamente ininterrumpida a lo largo del margen occidental de las sierras Chica de Zonda y Villicum, San Juan, Argentina. Sus calizas son portadoras de una importante fauna de trilobites cámbricos. En este trabajo se describe y analiza la trilobitofauna del Delamariano de la Formación La Laja, ubicada entre los últimos metros del Miembro El Estero y el tercio inferior del Miembro Soldano. Se determinaron las biozonas de Poliella denticulata, Mexicella mexicana? y Glossopleura walcotti, las que en trabajos recientes han sido incluidas en un hiato faunístico. Los taxones determinados en este trabajo son: Glossopleura walcotti Poulsen, Glossopleura lodensis (Clark), Fieldaspis sp. y Prozacanthoides zondaensis n. sp. Se realizaron correlaciones entre los niveles de la Formación La Laja y olistolitos de Precordillera Occidental. También se correlacionaron los niveles de Precordillera con otros en diferentes localidades de Laurentia. Los géneros documentados en este trabajo para el Delamariano de la Formación La Laja son característicos y restringidos a Laurentia. El análisis de toda la fauna de la Formación La Laja, apoya el modelo alóctono para el origen de Precordillera.

Key words. Cambrian. Delamaran. Trilobites. Argentina. Precordillera.

Palabras clave. Cámbrico. Delamariano. Trilobites. Argentina. Precordillera.

Introduction

Trilobites from the La Laja Formation are fundamental for understanding the early history of the Precordillera terrane. Different authors have studied the faunas contained in this unit, dealing mainly with its biostratigraphic aspects. However, systematic studies of the trilobites are scarce and this study on the Delamaran trilobites from the La Laja Formation is a contribution towards broadening knowledge on this important group of fossils.

Kayser (1876) and Stelzner (1923-1924) described the first Cambrian trilobites from the Argentine Precordillera (San Juan Province). These trilobites were later revised by Harrington and Leanza (1943), who identified the existence of the Middle Cambrian in the Precordillera. Poulsen (1958) acknowledged the presence of the Glossopleura Zone in the Cambrian of Argentina based on specimens of Glossopleura Poulsen, 1927, from San Isidro (Mendoza Province). This established the first zonation for intercontinental correlation of the Cambrian in the Precordillera. In addition, Poulsen (1958) inferred the Glossopleura Zone in the La Laja Formation (Quebrada La Laja and Quebrada Juan Pobre in Sierra Chica de Zonda), based on species it has in common with the same Zone in exposures at San Isidro, such as Mendospidella digesta (Leanza, 1947) and Kistocare mendozanum (Rusconi, 1945).

Bordonaro (1986) recognized the Bonnia-Olenellus Zone in the El Estero Member of the La Laja Formation based on the occurrence of Olenellus zondaen-
sis Bordonaro, 1986, *Bonna villicumica* Bordonaro, 1986, *Antagmus argentinus* Bordonaro, 1986, and *Bristolia* sp. This fossil assemblage was recently revised by Foglia and Vaccari (2006a) and referred to the uppermost part of the upper *Bonna-Olenellus* Zone. The *Bonna-Olenellus* Zone was also documented within the Cerro Totora Formation at Quebrada La Angostura (La Rioja Province), based on the presence of *Arcuolenellus megafrontalis* (Vaccari) (see Vaccari, 1988; Astini and Vaccari, 1996). Vaccari and Bordonaro (1993) recognized the *Bonna-Olenellus* Zone in olistolith C1 at Los Túneles, on the basis of the record of *Olenellus?* sp., *Prozancanoides* sp. (see discussion of the genus below), and *Sombrerella argentina* Vaccari and Bordonaro, 1993.

Bordonaro (2003a, 2003b) mentioned a hiatus spanning the *Plagiura-Poliella* and *Albertella* zones and also possibly the *Glossopleura* Zone in the Eastern Precordillera. However, Foglia and Vaccari (2006b) recognized the presence of the *Glossopleura* Zone and also possibly the *Albertella* Zone in the Soldano Member of the La Laja Formation. Records of the *Glossopleura* Zone in the Western Precordillera are limited to reports of its occurrence in the olistoliths of Ojos de Agua and Los Ratones (Bordonaro and Banchig, 1990, 1995, 1996). Poorly preserved material at these localities was identified as *Glossopleura* aff. *leona* Lochman and *Glossopleura* sp. More recently, Bordonaro et al. (2008) published a biostratigraphic model for the early Marjumian in the Precordillera and correlated different sections in the Eastern and Western Precordillera. Finally, Bordonaro and Pratt (2008a, 2008b) recognized the *Eokochaspis nodosa* Zone in the lowermost part of the Soldano Member, and the *Glossopleura* and *Elmussia* zones in the middle and upper parts of this unit. It is important to point out that none of these contributions illustrated specimens of these taxa from the La Laja Formation and references are restricted to the text only.

In this work, two species of *Glossopleura* are described and illustrated, documenting precisely for the first time the stratigraphic range of the *Glossopleura wallcotti* Zone in Eastern Precordillera. Material collected from beds immediately below the base of the *Glossopleura wallcotti* Zone is also illustrated and discussed from a biostratigraphic point of view. This material comes from the *Poliella denticulata* Zone at the Cerro Molles section and possibly from the *Mexicella mexicana* Zone at the Quebrada La Laja section. Localities mentioned in the text are shown in figure 1.

Cambrian trilobites from the Precordillera in general (e.g., Poulsen, 1958; Borrello, 1963; Vaccari, 1994) and those from the La Laja Formation in particular, show close affinities with Laurentian faunas. In this sense, data drawn from the material described herein may help resolving the recent controversy on the Gondwanan (e.g., Finney et al., 2003, 2005; Finney, 2007; Aceñolaza and Toselli, 2007) or Laurentian (e.g., Vaccari, 1994; Astini et al., 1995, 1996; Thomas and Astini, 1999; Benedetto, 2004) origin of the Precordillera.

### Localities and stratigraphy

The La Laja Formation is exposed with significant lateral continuity along the western slope of the Sierra de Villicum and the Sierra Chica de Zonda. Originally described by Borrello (1962), the unit is mainly composed of carbonate deposits with mixed carbonate-siliciclastic intervals, especially in the lower part of the unit (e.g., Bordonaro, 1980; Keller, 1999; Gómez and Astini, 2006). Baldis and Bordonaro (1981) defined four members in the La Laja Formation, namely from bottom to top El Estero, Soldano, Rivadavia, and Juan Pobre members. Based on sedimentologic and stratigraphic differences, Gómez and Astini (2006) added the Las Torres Member between the Juan Pobre Member and the overlying Zonda Formation. According to Gómez and Astini (2006), the facies associations of the La Laja Formation are interpreted as representing shallow marine environments, ranging from relatively
Delamaran trilobites from the La Laja Formation

Deep subtidal to very shallow subtidal-intertidal settings. The material illustrated herein comes from the lower third of the Soldano Member and the upper part of the El Estero Member (see figure 2).
All material is housed in the Centro de Investigaciones Paleobiológicas (CIPAL), Universidad Nacional de Córdoba, Córdoba, Argentina (prefixed CEGH-UNC).

Systematic paleontology

Order Corynexochida Kobayashi, 1935
Family Dolichometopidae Walcott, 1916

Genus Glossopleura Poulsen, 1927

Type Species. Dolichometopus boccar Walcott, 1916.

Remarks. Sundberg (2005) recently revised the type species G. boccar (Walcott) based on numerous collections; he highlighted its intraspecific variability and synonymized at least six species defined after introduction of G. boccar. Some of the characters mentioned by Sundberg as variable are pygidium convexity, degree of expansion of the frontal lobe, development of cranidial furrows, and depth of pleural furrows of the pygidium. The author suggested that some variable features such as the presence of a prominent S1 in the specimens preserved in shales are preservational differences and do not represent diagnostic specific characters. Glossopleura species are in need of revision, as many of them were based on single specimens and their variability ranges remain unknown. Such a study lies beyond the scope of the present contribution.

Glossopleura walcotti Poulsen, 1927

Figure 4.10-23

1927. Glossopleura walcotti Poulsen; Poulsen, p. 268, pl. 16, figs. 20-30.
1964. Glossopleura walcotti Poulsen; Poulsen, p. 25, pl. 1, figs. 2-4.
1979. Glossopleura walcotti Poulsen; Palmer in Palmer and Halley (part), p. 79, figs. 8-6, 11-13, 16-19, non figs. 14-15.
1995. Glossopleura aff. Iconi Lochman; Bordonaro and Banchig, pl. 2, figs. 13-14.
2009. Glossopleura walcotti Poulsen; Benedetto et al., fig. 2g.

Material examined. One articulated specimen, 44 cranidia, 12 librigenae and 30 pygidia; CEGH-UNC 23686/CEGH-UNC 23778.

Occurrence. Beds LLSL01, LLSL02, and LLSL03, Soldano Member, La Laja Formation, La Laja section; beds CMSL02 and CMSL03, Soldano Member, La Laja Formation, Cerro Molles section; beds EZSL04 and EZSL05, Soldano Member, La Laja Formation, Esteros de Zonda section.

Description. Cranidium outline trapezoidal, width/length ratio varying between 1.05 and 1.20. Glabella clearly defined by abaxially concave axial furrows; furrows usually ending in very distinct pit located in front of anterior end of palpebral lobe. Glabella transversally and longitudinally convex; frontal lobe clearly expanded laterally. Ratio between maximum and minimum width of glabella ranging between 1.5 and 1.7. Glabella with four pairs of very shallow and poorly defined glabellar furrows. Pair S1 strongly sloping inwards and backward, describing an angle varying between 45º and 60º respect of sagittal line. Pair S2 oriented almost transversally; S3 and S4 clearly oriented anteriorly. Palpebral lobes well defined by abaxially convex furrows; intraocular areas convex, width (trans.) about 0.30-0.35 of cranidium maximum width. Occipital furrow well defined, relatively wide. Occipital ring conspicuous, uniformly wide transversally. Occipital ring showing slight posterior node in some specimens. Anterior branch of facial suture starting at anterior end of palpebral lobe and then running outwards and anteriorly; posterior branch developing from posterior end of palpebral lobe and running outwards and slightly backwards to meet posterior furrow of cranidium, then turning rearwards and slightly outwards. Cranidium ornamented with fine lines in sinuous terraces on anterior part of frontal lobe of glabella. Weak ornamentation resembling fingerprints around occipital node.

Librigena with convex circumocular suture; posterior furrow well defined and with clear margin. Margin is flat and widening posteriorly. Librigena ornamented with lines in sinuous terraces, beginning perpendicular to lateral edge, then sloping posteriorly and becoming almost parallel to outline. These lines continue onto the genal spine and perpendicular to it.

Seven thoracic segments; pleurae ending in dorsally flattened spines; spines slightly backwards. Thoracic segments with middle node in axial ring. Although not all pleurae are the same size, a macropleura was not observed. Ornamentation of thorax characterized by fine impressions similar to fingerprints covering axial nodes, and by fine sinuous lines in terraces located at ends of pleurae.

Pygidium semicircular, sometimes elliptical in larger specimens; width/length ratio in the latter varying between 1.6 and 1.7. Axis prominent and strongly convex, relatively short, cylindrical or slightly conical; five axial rings and a terminal piece; inter-ring furrows moderately marked. Pleural furrows narrow and well-marked, continuing along pleural field to join border furrow. Border concave and well defined by border furrow. Pygidium ornamented with nodes on the axis, around which develops a pattern similar to the one described for pleural segments. Lines in sinuous terraces beginning perpendicular to edge of furrow, then turning anteriorly and running parallel to outline of pygidium.

Discussion. The specimens from the Precordillera
are indistinguishable in their cephalic and pygidial characters from the type material of *Glossopleura walcotti* Poulsen from the Cape Wood Formation in northwestern Greenland (Poulsen, 1927, p. 268, pl. 16, figs. 20-30). They are also identical to part of the material referred to this species in the Carrara Formation of the Great Basin (Palmer in Palmer and Halley, 1979, pl. 16, figs. 6-8, 11-13, 16-19). However, two pygidia referred to *G. walcotti* by Palmer (Palmer and Halley, 1979, pl. 16, figs. 14-15) show granules on the pleural ribs of the pygidium. This feature is not present in the type material from Greenland or in any of the specimens from the Precordillera, suggesting that the material from the Carrara Formation may probably belong to a different species.

*Glossopleura lodensis* (Clark, 1921)

Figure 4.1-9

1921. *Bathyuriscus howelli lodensis* Clark.
1928. *Dolichometopus lodensis* (Clark); Resser, p. 10, pl. 3, fig. 9.
1935. *Glossopleura lodensis* (Clark); Resser, p. 34.
1979. ? *Glossopleura lodensis* (Clark); Palmer in Palmer and Halley (part), p. 78, pl. 16, figs. 1-3, 10 non figs. 4-5, 9

**Material examined.** Twelve cranidia, four librigenae and 17 pygidia; CEGH-UNC 23779/CEGH-UNC 23810.

**Occurrence.** Beds LLSL04, LLSL05, Soldano Member, La Laja Formation, La Laja section.

**Description.** Cranidium nearly trapezoidal; width/length ratio varying between 0.95 and 1.10. Glabella clearly defined by axial furrows, straight and slightly curved outwards at anterior end. Glabella transversally and longitudinally convex; frontal lobe expanded anteriorly and slightly laterally. Maximum/minimum ratio of glabella width 1.1-1.3. Four pairs of superficial and indistinct glabellar furrows. Pair S1 (only observed in some specimens) sloping inwards and backwards, meeting sagittal line at an angle of 40°-50°. Pair S2 oriented almost horizontally; pairs S3 and S4 clearly oriented inwards and forwards. Palpebral lobes well defined by abaxially convex furrows. Intraocular areas convex and reaching 0.25-0.30 of cranidium maximum width. Occipital furrow clearly defining constantly wide (transversally) occipital ring. Posterior branch of facial suture starting strongly divergent outwards and slightly backwards respectively of posterior furrow of cranidium, then turning backwards to meet posterior border of cranidium. Anterior part of frontal lobe of glabella ornamented with fine sinuous terraced lines.

Librigena with convex and relatively narrow genal field and very well defined posterior and border furrows. Border flat, its width increasing towards posterior region. Genal spine beginning at genal angle and running posteriorly and outwardly. Ornamentation of librigena consisting of lines in sinuous terraces, beginning perpendicularly to middle sector of border furrow, then becoming parallel to outline; lines continuing into genal spine, almost perpendicular to axis of spine.

Pygidium outline semicircular; width/length ratio 1.7-1.9. Pygaxis prominent, strongly convex, cylindrical, including five to six axial rings and ending in a terminal piece. Pleural furrows very distinct and strongly marked right down to edge of pygidium. Lateral and posterior borders concave, only defined by change in slope. Pygidium ornamented with discrete lines in sinuous terraces near border, less noticeable than in other species of the genus.

**Discussion.** *Glossopleura lodensis* (Clark) was defined on the basis of an articulated specimen from the Mohave Desert and was later illustrated again by Resser (1928, pl. 3, fig. 9). The most distinctive characteristic of this species is a long glabella, with relatively short palpebral lobes, a long spine in the fifth thoracic segment, and a pygidium with pleural furrows that cross the border. Although articulated specimens are not available in our collection, cranidia and pygia with the same characteristics as the holotype of *G. lodensis* occur in the same levels.

Palmer referred some specimens from the Carrara Formation to this species (Palmer and Halley, 1979, pl. 16, figs. 1-5, 9-10). These specimens show a great variability, and probably more than a single species is represented. Only the specimen depicted by Palmer (1979, pl. 16 fig. 10) exhibits pleural furrows that cross the wide border and thus could be tentatively assigned to *G. lodensis*.

*G. lodensis* as revised here differs from *G. walcotti* by the shorter palpebral lobes, the longer glabella, and by the pleural furrows that distinctly cross the wide border of the pygidium.

Family ZANANTHOIDIDAE Swinnerton, 1915

**Genus** Prozacanthoides Resser, 1937

**Type species.** *Olenoides stissingensis* Dwight, 1889, p. 147, pl. 6, figs. 9-5.

**Comments.** Palmer (in Palmer and Halley, 1979, p. 97) indicated that the diagnosis of *Prozacanthoides* given by Resser (1937) was not based on the features of the type species, i.e., *Olenoides stissingensis* Dwight, 1889. Likewise, based on a revision of the type material, Palmer pointed out the characters that differentiate *Prozacanthoides* from the rest of the genera included in the family, such as the relatively short palpebral lobes, the subparallel anterior branch of the facial suture, the subtriangular posterolateral projection, and the three strong pairs of spines on the
lateral margins of the pygidium, of which the posterior pair is the longest and most separate (Palmer and Halley, 1979, p. 97). Considering these characters, Palmer restricted the genus Prozacanthoides to the type species. We fully concur with this author and following his concept we define here a new species from the section at Quebrada La Laja.

Prozacanthoides zondaensis n. sp.

Material examined. Seventeen cranidia, one librigena, four hypostomes and twenty seven pygidia; CEGH-UNC 23637/CEGH-UNC 23685.

Occurrence. Bed LLEE04. El Estero Member, La Laja Formation, La Laja section.

Etymology. After Sierra Chica de Zonda.

Holotype. CEGH-UNC 23640. Pygidium.

Diagnosis. Prozacanthoides with three pairs of tubular marginal spines; posterior pair very long and stout.

Description. Cranidium trapezoidal; glabella elongate, rectangular, clearly convex transversally, defined by parallel or slightly inwardly divergent axial furrows. Four pairs of glabellar furrows; pair S1 deepest of the four and oriented backwards; pair S2 almost transversal to axis of glabella; S3 and S4 (when observable) distinctly oriented anteriorly. Some specimens showing well-defined pit at intersection of S4 and axial furrow. Anterior border of cranidium well defined, concave, external curvature radius always larger than curvature radius of glabella frontal lobe. Preglabellar furrow always conspicuous, its width increasing abaxially. Intraocular areas convex, always lower than glabella. Palpebral lobe long, convex, and clearly defined by furrow. Anterior branch of suture beginning at anterior end of palpebral lobe, then running outwards and anteriorly, obliquely crossing pre-glabellar furrow. Posterior branch of suture beginning at posterior end of palpebral lobe, then running clearly outwards and slightly backwards to contact posterior furrow of cranidium. Occipital furrow always very well defined, although narrow and shallow. Occipital ring triangular and transversally convex. No specimens with nodules or spines in occipital ring. Ornamentation of cranidium consisting of only fine terraced lines on anterior margin, running almost parallel to head outline.

Librigena convex, triangular, with well defined — albeit narrow and shallow marginal furrow. Margin convex, its width increasing backwards to gradually form genal spine. Librigena ornamented with fine lines in sinuous terraces running parallel to margin outline; lines becoming clearer and denser on genal spine.

Pygidium trapezoidal (excluding spines); axis prominent with conical outline defined by strong axial furrows, comprising four rings and a terminal piece. First two rings very strong, defined by clear and not very deep furrows; third ring less noticeable (although not always present), with posterior furrow very shallow and only seen in some specimens. Furrows of fourth ring are moderately developed. Terminal piece conical, end rounded. Pleural areas wide and convex, with three conspicuous pleural furrows, apparently coinciding with axis rings and delimiting pygidium spines. Only best preserved specimens showing interpleural furrow coinciding with posterior furrow of first axial ring; small pit visible at its intersection with the axial furrow. Border of pygidium defined by change in slope. Three pairs of robust tubular spines along lateral borders; spines slightly flattened transversally, their length increasing backwards. Posterior pair the longest and the most separated from the rest. Posterior margin of the border showing highest character variation within species; usually straight, but can also be faintly convex, with two pairs of spines or protuberances, or completely smooth. Ornamentation of pygidium consisting of small lines in sinuous terraces on the posterior border and on marginal spines.

Discussion. The main problem arising in any discussion of this genus is that Prozacanthoides stissingensis (Dwight) is the only species of the genus and the illustrations provided by Dwight (1889) are inadequate for any meaningful comparison. In spite of these limitations, strong similarities are observed between our material and P. stissingensis, enough to accommodate our material in Prozacanthoides. Nonetheless, Prozacanthoides zondaensis n. sp. differs from P. stissingensis in the tubular marginal spines of the pygidium (in contrast to the flat ones in P. stissingensis), and in the longer and more robust posterior pair of spines.

Aguaraya acutispina Rusconi, 1955, type species of Aguaraya Rusconi, 1955, and based on a single pygidium from San Isidro (Mendoza), bears a strong resemblance with the pygidia of the new species. The specimen illustrated by Rusconi (1955) has a pygidium with a similar outline, a prominent axis and three pairs of strong marginal spines, of which the posterior one is the longest (Rusconi, 1955, pl.1, fig. 2). However, it is different from P. zondaensis n. sp. in having a relatively shorter axis, a longer posterior edge and two pairs of small spines along the posterior margin. Some of the specimens in our collection bear two protuberances at the posterior margin that could be homologous to the marginal spines of A. acutispina (i.e., pl. 2, fig. 9, this contribution). If this were so, both species could be included in the same genus. It is also important to point out that according to Palmer (in litt., 1995), some
paratypes of *P. stissingensis* show a quadrangular posterior margin with short spinules. A revision of the species of this genus would clarify the relationships among them and identify a possible synonymy.

The material illustrated by Vaccari and Bordano 1993 (pl. 1, figs. 10, 12, 13) as *Prozacanthoides* ? sp. and *Prozacanthoides* n. sp. was also analyzed and compared. Direct observation of the material indicates that the specimen illustrated in figure 10 (CORD-PZ 8729) is congeneric with the material from the La Laja Formation, while the specimens in figures 12 and 13 do not belong to *Prozacanthoides*, due to the transversal width of the posterior border of the pygidium, and the shape and size of the spines. These specimens are perhaps more closely related to the genus *Mexicaspis* Lochman, 1948.

Genus *Fieldaspis* Rasetti, 1951

**Type species.** *Fieldaspis furcata* Rasetti, 1951.

*Fieldaspis* sp.

1968. ? *Fieldaspis* cf. *F. superba* Rasetti; Norford, p. 35, pl. 4, fig. 9, 11; pl.5, figs 1-3.

**Material examined.** Two pygidia, partially preserved. CEGH-UNC 23811/CEGH-UNC 23812.

**Occurrence.** Bed CMEE08, El Estero Member, La Laja Formation, Cerro Molles section.

**Description.** Pygidium outline trapezoidal (excluding spines). Axis convex, relatively conical, with four distinct rings and terminal piece. Axial furrows sinuous and shallow, deeper at intersection with rings. Pleural fields moderately convex, with three pairs of deep pleural furrows. First pair oriented almost transversally in its proximal part, strongly curving backwards distally. Interpleural furrows shallow. Lateral border furrow wide, essentially delimited by distal deepening of pleural furrows. Border well defined at antero-lateral corner, slightly widened and more concave at postero-lateral corner, reaching backwards into marginal spines. Posterior margin curved and marginal spines strongly oriented backwards, slightly outwards and upwards. External surface covered by small granules on axis and pleural fields, and by anastomosed lines on border.

**Discussion.** These pygidia can be clearly referred to *Fieldaspis* Rasetti, 1951, and they are close to *F. superba* Rasetti (Rasetti, 1951, p. 162, pl. 16, figs. 10-18; Sundberg and McCollum, 2003, p. 343, figs. 11.7,11.11) in their general shape, and the course and depth of the pleural furrows. Nonetheless, they differ in the outline of the edge, which is oriented backwards and slightly inwards in the Argentine species, while it is oriented backwards and outwards and with the marginal spines more outwardly directed in *F. superba*.

Pygidium of *Fieldaspis* sp. resembles those of *Fieldaspis* cf. *F. superba* Rasetti (Norford, 1968, p. 35, pl. 5, figs. 1, 3) in the general outline as well as in the shape of the axis. The only difference is that the first pair of pleural furrows is more transverse in *Fieldaspis* sp., while in the Canadian specimens it is curved, directed backwards and the axis is narrower posteriorly. It is possible that these materials could be conspecific.

**Biostratigraphy**

Cambrian biostratigraphy is based essentially on the distribution of benthic faunas of trilobites. Because of their strong endemism, correlation at an intercontinental scale is difficult and most of the stages within the series have not been established yet. This is why local biostratigraphic frames are widely used in continental correlations. Because of the characteristics of the fauna described here, it will be referred to the schemes established for the Cambrian of Laurentia (see figure 3). The middle Cambrian biostratigraphic schemes for Laurentia were tradi-
Figure 4. 1-9, *Glossopleura lodensis* (Clark, 1921), all specimens from La Laja section, Soldano Member / todos los ejemplares de la sección La Laja, Miembro Soldano. 1, pygidium, dorsal view / pigidio, vista dorsal, CEGH-UNC 23790 level LLSL04. 2, 3, pygidium, dorsal and posterior views / pigidio, vistas dorsal y posterior, CEGH-UNC 23805, level LLSL04. 4, 5, pygidium, dorsal and posterior views / pigidio, vistas dorsal y posterior, CEGH-UNC 23793, level LLSL04. 6, cranidium, dorsal view / cranidio, vista dorsal, CEGH-UNC 23800, level LLSL04. 7, cranidium, dorsal view / cranidio, vista dorsal, CEGH-UNC 23796, level LLSL04. 8, librigena, dorsal view / mejilla libre, vista dorsal,
tionally defined based on the occurrence of a single genus or genera association (Rasetti, 1951; Lochman and Wilson, 1958). During the last decades, biostratigraphic research of great detail was undertaken mainly by Sundberg (1994), Eddy and McCollum (1998), Sundberg and McCollum (2000, 2003), McCollum and Sundberg (2007) among others. These contributions represent a source of abundant and detailed data on the different successions and faunal associations of Laurentian localities. Moreover, these authors proposed more precise biostratigraphic schemes based on species or species associations.

Trilobites of the La Laja Formation allow recognition of the Delamaran Stage. This stage was postulated by Palmer (1998) for the interval of the Corynexochid biomere and represents the first stage of the Lincolnian Series of the Cambrian of Laurentia. The base of this stage, in its type section -Delamar Mountains, eastern Nevada (Pioche Shale)- is characterized by the presence of abundant silicified specimens of *Eokochaspis nodosa* (Sundberg and McCollum, 2000). The upper boundary of the stage is defined by the occurrence of *Proehmaniella basilica* (Resser, 1945) at the base of the *Proehmaniella* Subzone of the *Ehmaniella* Zone that defines the base of the next stage, i.e., the Topazan (Sundberg, 2005). Trilobites so far recorded in the Delamaran Stage are ptychoparids related to *Kochaspis* Resser, 1935, and corynexochids of the families Zacanthoididae and Dolichometopidae (Palmer, 1998). The zones established for this stage are: *Eokochaspis nodosa* Zone, *Amecephalus arrojosensis* Zone, *Poliella denticulata* Zone, *Mexicella mexicana* Zone and *Glossopleura walcotti* Zone. The zones of *Poliella denticulata*, *Mexicella mexicana* (possibly) and *Glossopleura walcotti* are recognized in the La Laja Formation. A scheme of the zones recognized in the La Laja Formation is shown in figure 5.

**Poliella denticulata Zone**

Recently, and after a series of papers focused mainly on the lower part of the Middle Cambrian, Sundberg and McCollum (1997, 2000, 2002, 2003) defined - in the Eastern region of Nevada - three zones located between the last record of oleneellids and the base of the *Mexicella mexicana* Zone. These are, in ascending order, *Eokochaspis nodosa* Zone, *Amecephalus arrojosensis* Zone and *Poliella denticulata* Zone. The authors also subdivided the youngest of these zones into five different faunal assemblages, i.e., the *Poliella denticulata*, *Fieldaspis celer*, *Syspacephalus longus*, *Poliellates gloriosa*, and *Kochiella augusta* assemblages.

It is important to point out that species of *Fieldaspis* (see Rasetti, 1951) occur in three of the five assemblages mentioned and that these species are practically restricted to the *P. denticulata* Zone. In this contribution we document the record of *Fieldaspis* sp. at Cerro Molles. Some specimens collected twenty meters below the first record of *Propazanchothodes* at Quebrada La Laja are very close to *Fieldaspis bilobata* Rasetti, 1951. Yet, because of the scarce and relatively poorly preserved material (one pygidium and one cranidium), its assignment remains uncertain. *Fieldaspis* sp., from the top of El Estero Member (Cerro Molles section), closely resembles *Fieldaspis* cf. *superba* Norford, 1968, and they are probably conspecific. *Fieldaspis superba* Rasetti, 1951, was recorded in the Pioche Shale, in the *Kochiella augusta* association, according to the scheme of Sundberg and McCollum (2003). According to the authors this association would be the equivalent to *Kochaspis* Zone defined by Palmer and Halley (1979) in the Carrara Formation.

In the La Laja Formation, *Fieldaspis* sp. constitutes -along with an association dominated by ptychoparids- the earliest Middle Cambrian fauna recorded to date in Eastern Precordillera. The occurrence of *Fieldaspis* sp. in Cerro Molles allows us to infer the presence of the *Poliella denticulata* Zone at the top of El Estero Member. However, we can not determine with certainty to which zonule or assemblages our material belongs. It is important to mention that Rasetti (1957) suggested a lineage including *F. bilobata* Rasetti, 1951, *F. furcata* Rasetti, 1951, *F. celer* (Walcott, 1917) and *F. superba* Rasetti, 1951. Likewise, Sundberg and McCollum (2003) found a morphologic gradient along the three species of *Fieldaspis* in the Pioche Shale (*F. bilobata*, *F. celer* and *F. superba*) that further supported the idea of Rasetti. The morphologic gradient suggested by both authors...
Figure 5. 1-17, 22-26, Prozacanthoides zondaensis n. sp., all specimens from La Laja section and level LLEE04. 1, pygidia, top: dorsal view of CEGH-UNC 23641; bottom: dorsal view of CEGH-UNC 23640. 2-4, pygidium, dorsal, lateral and posterior views. 5, cranidium, dorsal view. 6-8, cranidium, dorsal, lateral and anterior views. 9, pygidium, dorsal view.
consists of a progressive transformation occurring in the posterior margin of the pygidia, from a clearly bilobed margin in *F. bilobata* at one end, to the strong pair of spines in *F. superba* at the other. The intermediate stages could be represented by *F. furcata* and *F. celer*. Each species of *Fieldaspis* identified in the Pioche Shale belongs to a different assemblage, and the morphologic succession coincides with younger assemblages. Since the material from Precordillera shows a strong resemblance to *F. superba*, it is possible to suggest that the studied assemblages could be correlated with the *Kochiella augusta* association. Study of the ptychopariid associated with *Fieldaspis* at Cerro Molles should help to confirm or reject this tentative correlation.

According to data available up to now, *Fieldaspis* sp. is the only clear representative of the *Poliella denticulata* Zone in Precordillera, and its record is restricted to the upper levels of the El Estero Member at Cerro Molles. Although this restriction prevents further correlations with other localities from Precordillera, it becomes a very valuable tool when correlating with Laurentian localities. In this sense we can affirm that the top levels of the El Estero Member (Cerro Molles section) bearing *Fieldaspis* sp. (*Poliella denticulata Zone*), can be correlated with those that yield *Fieldaspis cf. F. superba* in the Mackenzie district, northeastern Canada (Norford, 1968); and with the ones that contain *Fieldaspis superba* (*Fieldaspis superba* faunule of the Plagiura-Kochaspis Zone) in the Mount Whyte Formation in Alberta, Canada (Rasetti, 1951). Likewise, the levels with *Fieldaspis* sp. of the La Laja Formation can be correlated with the blocks assigned to the top of the Cap Kent Formation in northeastern Greenland (Poulsen, 1964); and with the levels that contain *Fieldaspis superba* at the top of the *Poliella denticulata Zone* in the Pioche Shale Formation, Nevada (Sundberg and McCollum, 2003).

**Mexcilla mexicana Zone**

The *Mexcilla mexicana* Zone (McCollum and Sundberg, 2007) was recently defined in replacement of the *Albertella* Zone (Deiss, 1939), a criterion that is shared here. In the Grassy Springs Member of the Pioche Shale, Nevada, Eddy and McCollum (1998) recognized three distinctive associations, *i.e.*, *Albertellina espinosa* Subzone, a middle unnamed subzone, and *Albertella highlandensis* Subzone. This subdivision corresponds to three different assemblages that do not reflect a temporal succession, but a particular spatial distribution in the basin. Palmer (in Palmer and Halley, 1979) proposed a similar scheme for the same interval in the Carrara Formation where the *Albertella Zone* was subdivided in three faunules, *i.e.*, *Ogygopsis* faunule, *Zacanthoidid* faunule, and *Albertella-Mexcilla* faunule. However, McCollum and Sundberg (2007) disagreed with the stratigraphic relationships suggested for these associations, and placed the *Ogygopsis* Zonule and the temporally equivalent *Zacanthoidid* Zone between the uppermost *Mexcilla mexicana* Zone and the lowermost *Glossopleura walcotti* Zone.

The *M. mexicana* Zone cannot be recognized with certainty in the La Laja Formation. In the upper part of the El Estero Member exposed at Quebrada La Laja a trilobite association dominated by *Prozacanthoides* is recorded. This genus does not have a distinctive position in the biostratigraphic schemes of Laurentia. This is because at the single locality where it is found it is the only representative of the Middle Cambrian, rendering impossible to determine its relative position with respect to the other zones of the same age (Palmer and James, 1979). Furthermore, no diagnostic element of this trilobite zone has been recognized in the preliminary study of the ptychopariids, dorypygids and zacanthoidids associated with *P. zondaensis* at Quebrada La Laja. In any event, *Prozacanthoides zondaensis* n. sp. is found in the La Laja Formation below the first record of the *Glossopleura walcotti* Zone and 20 m above the record of *Fieldaspis* sp. Therefore this association could yet correspond to the *Poliella denticulata Zone* or else represent the *M. mexicana* Zone. If the latter alternative were correct, then the restricted development of the *Mexcilla mexicana* Zone in the La Laja Formation (as well as the one in the *Poliella denticulata Zone*) significantly contrasts, for instance, with what was documented for the Pioche Shale in Nevada. Possibly, this particular record in the Precordillera is a result of the presence of a condensed interval, a hiatus in the record, or a combination of both factors.

The beds with *Prozacanthoides zondaensis* n. sp.,...
from the upper part of the El Estero Member at Quebrada La Laja, are correlated with those that bear Prozancanthatoides? documented by Vaccari and Bordonaro (1993) in the block within the olistostrom of the Los Sombreros Formation (at Río Jáchal). The levels with Prozancanthatoides in the La Laja Formation are correlated with the levels that yield Prozancanthatoides stissingensis (Dwight) Dutchess County, New York. It is important to note that the succession exposed at the latter locality has been referred to the Middle Cambrian with no further precisions (Landing, 2007). Accordingly, taking into account the correlation herein proposed, Dwight’s locality could be placed below the Glossopleura Zone - between the uppermost part of the P. denticulada and possibly the M. mexicana Zones.

Glossopleura walcotti Zone

Rasetti (1951) analyzed the faunas of the Middle Cambrian of British Columbia and identified three different faunules in the Glossopleura Zone, i.e., Glossopleura merlinensis faunule, Glossopleura boccar faunule, and Polypleuraspis insignis faunule. Some of the species in these faunules were later considered synonyms and others are just found at their type localities; therefore use of this scheme in other regions is uncertain. The Glossopleura Zone in northeastern Greenland was subdivided by Poulsen (1964) into two different faunules, i.e., Glossopleura faunule and Clavaspidella faunule. Within the Glossopleura faunule are found the three faunules identified by Rasetti (1951) for Alberta. We do not agree with the criteria used by Poulsen (1964) when he included the Clavaspidella faunule within the Glossopleura Zone. Instead it may be more closely related to the Ehmaniella Zone. Sundberg (2005) documented the Glossopleura Zone in the Chisholm Formation, Utah, based on a record of Glossopleura boccar. Finally, McCollum and Sundberg (2007) revised the zonation of the Delamaran of the Great Basin and defined the Glossopleura walcotti Zone, by the FAD of Glossopleura walcotti Poulsen (1927).

Glossopleura walcotti is one of the best represented zones in the Precordilleran La Laja Formation. It has been recorded at the Quebrada La Laja section (five levels), the Esteros de Zonda section (two levels), and the Cerro Molles section (two levels), becoming the main element for correlation among the different sections of the lower Middle Cambrian of the La Laja Formation. These records are associated -at the three sections- to a noticeable lithofacial change. This change takes place a few meters above a surface interpreted as transgressive by Keller (1999). In general, it coincides with the base of the Soldano Member in the studied localities.

The beds carrying the fauna assigned to the Glossopleura walcotti Zone in the La Laja Formation are correlated with part of the blocks of Ojos de Agua and Los Ratones in Western Precordillera, which bear Glossopleura sp. and Glossopleura aff. leona (Bordonaro and Banchig, 1990, 1996). In Laurentia the Glossopleura walcotti Zone is widely distributed. The levels with Glossopleura walcotti documented in the Soldano Member of the La Laja Formation are correlated with those bearing the same species in the Cap Wood Formation in northeastern Greenland (Poulsen, 1964). They are also correlated with the ones that contain G. walcotti in the Carrara Formation (Palmer and Halley, 1979).

Biogeographic considerations

The Precordillera has faunal and tectosedimentary features that are notably different from other Paleozoic successions along the western margin of South America. These peculiar characteristics led different authors to formulate hypotheses explaining the presence of this terrane in western Argentina. Nowadays, and on the basis of stratigraphic (e.g., Bond et al., 1984; Astini et al., 1995, 1996; Thomas and Astini, 1999), paleomagnetic (Rapalini and Astini, 1998), and faunal evidence (Vaccari, 1994; Benedetto, 2004) there is widespread consensus on the Laurentian origin of Precordillera. Nevertheless, other authors are inclined towards a parautochthonus origin respect of the margin of Gondwana (e.g., Baldis et al., 1989; Loske, 1993; Aceñolaza et al., 2002; Finney, 2007).

Benedetto et al. (2009) analysed the biogeographic affinities of the early Paleozoic faunas of the Precordillera. Particularly, they suggest that trilobite evidence throughout the entire Cambrian succession of Precordillera points towards a strong faunal identity with Laurentia, and at the same time towards the absence of faunal exchange with coetaneous areas of East Gondwana. Besides, Precordilleran trilobites are mostly recorded in shallow environmental settings. It is widely accepted that “inshore biofacies” are mostly confined to a single palaeocontinent (Fortey and Owens, 1978, 1987) and, hence, these associations are among the best biogeographic markers. In the parautochthonous model proposed by Aceñolaza et al. (2002) and recently reviewed by Finney (2007), during the Cambrian the Precordillera is located adjacent to the margin of Gondwana, close to Antarctica and Australia. According to this model Antarctica, Australia, and Precordillera were in a similar latitudinal position, with carbonate platforms developed over all three areas, and no oceanic barriers separating them. As is well known, Cambrian faunas from Antarctica and Australia are closely re-
lated and belong to the Redlichiid Realm, whereas no faunal elements of this realm are shared with Precordillera. If the Precordillera and Antarctica-Australia platforms were connected as depicted in the parautochthonous reconstruction by Finney (2007, fig. 5), a fluid faunal exchange among these blocks should be expected in both directions. Finney (2007) argues that the westward direction of equatorial oceanic currents is responsible for the lack of tropical Gondwanan elements in the Precordillera. However, during Furongian-Tremadocian times active larval dispersal into the Andean platforms from Australia, New Zealand and South China is well documented (Vaccari et al., 2006; Vaccari and Waisfeld, 2008). If Cuyania was adjacent to Gondwana, as Aceñolaza et al. (2002) and Finney (2007) proposed, a question remains that needs to be explained using biogeographic arguments: why are faunal affinities of Precordillera exclusively Laurentian until the Tremadocian? On the other hand, if Laurentian faunas were carried into the Precordillera by west-flowing equatorial currents (Finney, 2007, fig. 10) it is hard to explain why they did not reach the East-Gondwanan carbonate platforms.

One of the arguments used to sustain the parautochthonous model and refute the allochthonous one is based on the comparison of faunal diversity between the Middle Cambrian of Laurentia and Precordillera (Finney et al., 2005; Finney, 2007). From our perspective, this is an inappropriate comparison not only because of the difference in scale (the whole oriental margin of Laurentia against Precordillera), but also because of the great variety of environments present in the different Laurentian localities which have not been recorded in Precordillera. Accordingly, the supposed difference in diversity between the faunas of Precordillera and Laurentia is largely a sampling artifact. One hundred years of intense research in Laurentia against a stage of incipient study of the Cambrian faunas of Precordillera additionally explain this sampling bias. Recent studies (Foglia and Vaccari, 2006a, 2006b; Bordonaro et al., 2008; Benedetto et al., 2009) and the present contribution provide new elements to assess the different contrasting arguments. The genera documented in this work for the Delamaran of the La Laja Formation (Fieldaspis, Prozacanthoides and Glossopleura), as well as those documented by Bordonaro (Bordonaro et al., 2008; Bordonaro and Pratt, 2008a) for other intervals in the La Laja Formation are characteristic of -and restricted to- Laurentia. It is important to point out that neither the allochthonous model nor the parautochthonous one question the affinities of the faunas of Precordillera with those of Laurentia. The differences between these models surface when it comes to explaining the presence of this fauna in Precordillera. Clearly, the most parsimonious interpretation is to place the Precordillera terrane close to Laurentia.

Conclusions

The trilobite faunas collected in the limestones of the La Laja Formation allow referring the lower third of the Soldano Member and the upper part of the El Estero Member to the Delamaran Stage. The Poliella denticulata, Mexicella mexicana? and Glossopleura walcotti zones are present in this interval. The taxa that enabled identification of these zones are: Fieldaspis sp., G. walcotti and G. lodensis.

The restricted development of the P. denticulata Zone and the uncertainty about the record of the M. mexicana Zone in the El Estero Member suggest either the record of a condensed section or a hiatus at the contact with the Soldano Member.

Correlations were established between the La Laja Formation and different Laurentian lithostratigraphic units, as well as with blocks of Western Precordillera. Because of their age, these blocks could represent lateral equivalents of the La Laja Formation. According to Gómez and Astini (2006), some of them constitute the only record of relatively deep facies and can be used to reconstruct at least partially the Precordillera platform during the Middle Cambrian.

The fauna illustrated in this work has undeniable Laurentian affinities. Further and detailed studies of the entire fauna from the La Laja Formation will be a key to establish the location of Precordillera with greater precision in paleogeographic reconstructions, and also in determining the timing of the inferred break-away from Laurentia.

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