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# STRATIGRAPHY, PALEONTOLOGY AND DEPOSITIONAL ENVIRONMENTS OF THE LOWER PERMIAN ROBLEDO MOUNTAINS FORMATION OF THE HUECO GROUP, ROBLEDO MOUNTAINS, NEW MEXICO

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# Abstract

Early Permian fossil localities, including numerous tracksites, in the southern Robledo Mountains of Dona Ana County, New Mexico, cover an area of approximately 20 km2 . Lower Permian strata exposed here belong to four formations of the Hueco Group (ascend ing order): Shalem Colony, Community Pit, Robledo Mountains and Apache Da m Formations. With the exception of the Robledo Mountains Formation, the Hueco Group is dominated by shallow water ma rine facies. The Robledo Mountains Formation is as much as 125 m of ma rine carbona tes and shale, intercalated with siliciclastic red-beds that comprise about one-third of the uni t's thickness. At more than 30 localities, the red beds in the study area contain extensive invertebra te a nd vertebra te (tetrapod-footprint) trace fossils and a megafossil plant assemblage composed mainly of Wnlchin. Marine facies of the upper part of the Robledo Mountains Forma tion contain an extensive late Wolfcampian assemblage of megafossil invertebrates, dominated by brachiopods and bryozoans, with considerable numbers of molluscs (bivalves, gastropods, a few specimens of ammonites), and numerous indeterminate crinoids. on-fusulinid foraminifera ns and ostracods dominate the microfossil assemblages. Conodonts from the lower part of the Robledo Mountains Forma tion, found in stra ta that bracket most of the tracksites, indicate a late Wolfcampian n (= la te Artinskia n) age.

Carbonates of the Robledo Mow1tains Formation were deposi ted in relatively quiet shallow- water shelf environments below active wavebase. They show a trend from restricted circulation (brackish?) wa ters in the lower part of the forma tion to more open normal marine waters in the middle a nd u pper parts of the formation. Most of the 34 red-bed tracksites in the Robledo Mountai ns Formation occur a t one stratigraphic level and th us represent a mega tracksite tha t encompassed a t least 20 km2. Tracksites were formed on siliciclastic tidal flats during early stages of rising base level (transgression).

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# STRATIGRAPHY, PALEONTOLOGY AND DEPOSITIONAL ENVIRONMENTS OF THE LOWER PERMIAN ROBLEDO MOUNTAINS FORMATION OF THE HUECO GROUP, ROBLEDO MOUNTAINS, NEW MEXICO

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ABSTRACT: Early Permian fossil localities, including numerous tracksites, in the southern Robledo Mountains of Dona Ana County, New Mexico, cover an area of approximately 20 km<sup>2</sup>. Lower Permian strata exposed here belong to four formations of the Hueco Group (ascending order): Shalem Colony, Community Pit, Robledo Mountains and Apache Dam Formations. With the exception of the Robledo Mountains Formation, the Hueco Group is dominated by shallow water marine facies. The Robledo Mountains Formation is as much as 125 m of marine carbona tes and shale, intercalated with siliciclastic red-beds that comprise about one-third of the unit's thickness. At more than 30 localities, the red beds in the study area contain extensive invertebra te and vertebra te (tetrapod-footprint) trace fossils and a megafossil plant assemblage composed mainly of Wnlchin. Marine facies of the upper part of the Robledo Mountains Formation contain an extensive late Wolfcampian assemblage of megafossil invertebrates, dominated by brachiopods and bryozoans, with considerable numbers of molluscs (bivalves, gastropods, a few specimens of ammonites), and numerous indeterminate crinoids. on-fusulinid foraminifera ns and ostracods dominate the microfossil assemblages. Conodonts from the lower part of the Robledo Mountains Formation, found in strata that bracket most of the tracksites, indicate a late Wolfcampian (= late Artinskian) age.

Carbonates of the Robledo Mow Itains Formation were deposited in relatively quiet shallowwater shelf environments below active wavebase. They show a trend from restricted circulation (brackish?) waters in the lower part of the formation to more open normal marine waters in the middle and upper parts of the formation. Most of the 34 red-bed tracksites in the Robledo Mountains Formation occur at one stratigraphic level and thus represent a mega tracksite that encompassed at least 20 km<sup>2</sup>. Tracksites were formed on siliciclastic tidal flats during early stages of rising base level (transgression).

#### INTRODUCTION

The Robledo Mountains (Fig. 1) are a wedge-shaped horst of Paleozoic and Cenozoic rocks tilted southward  $10^{\circ}$  to  $14^{\circ}$  (Hawley et a l., 1975). This horst 1 ies along the western ma rgin of the southern Rio Grande rift and exposes a 500+ m thick, carbona te-dominated section of Paleozoic strata overlain locally by eogene siliciclastics and cut locally by Cenozoic intrusives (Seager et al., 1987).

In the southern portion of the Robledo Mountains, numerous fossil-footprint localities (tracksites) are known from the southwestern quarter of T22S, RIE and the northeastern quarter of T22S RIW, Dona Ana County. Discovered and developed by Jerry P. MacDonald, these tracksites represent the most scientifically significant record of Permian tetra pod footprints in the world (Lucas et al., 1994b, 1995; Haubold et al., 1995a; Hun t et al., 1995b). The h-ack-bearing shata are intercalated with ma rine sed iments that contain an extensive invertebrate biota (e.g., Kietzke and Lucas, 1995; Kozur and LeMone, 1995; Kues, 1995). Our purpose here is to review the stra tigra phic and depositional context of these tracksites. In this article, MM H refers to the New Mexico Museum of Natural History and Science, Albuquerque.

#### LOCATION AND METHODS

An a rea of about 20 km<sup>2</sup> (Fig. 2) is delimited by 43 known fossil localities in secs. 19-20, 29-30, T22S, RIE and secs 23-26, T22S, RIW (Lucas et al., 1995, table 1, figs. 2-3). Seager et al. (1987) and Lucas et al. (1995) recently mapped the geology of this area at scales of 1:250,000 and 1:24,000, respectively (Fig. 2).

Our stratigraphic conclusions are based on seven stratigraphic sections of the Robledo Mountains Formation (Fig. 3) described in detail and published by Lucas et al. (1995), as well as three other sections of the strata bracketing this formation published by Lucas et al. (1998).

#### PREVIOUS STUDIES

## Regional Geology

Regional geologic maps have encompassed the Robledo Mou nta i ns (Kottlowski, 1960; Seager et al., 1987) as have broad regional studies of Permian stratigra phy in southern New Mexico (e.g., Kottlowski, 1963; Jordan, 1971, 1975). Sedimentologica 1 studies of the intertongued Abo-Hueco stra ta of the Robledo and Dona Ana Mow1tains were published by Mack and Ja mes (1986) and Mack et al. (1988, 1991). Seager et al. (1976) presented a stratigraphic section of Wolfcampian rocks in the Robledo Mountains. LeMone et al. (1967, 1971a, b, 1975) published brief paleontological and micro-facies analyses of the Robledo Mow1tains and Apache Dam Formations of the Hueco Group in the study area. Lucas et al. (1995) integrated much of this information into a detailed report on the stratigraphy and depositional history of the track-bearing interval.

Mack and associates studied the sedimentology of the trackbearing strata in general (Mack and James, 1986; Mack et al., 1988, 1991), and Lucas (1993) studied the sedimentology of NMMNH locality 846 in particular. Lucas (1993), Hunt et al. (1993, 1994a, b) and Lucas et al. (1994a,b) reported some initial results of scientific study of the tracksites, followed by the more detailed reports in Lucas and Heckert (1995).

## Invertebrate Paleontology

Until 1995, there had been little previous study of the Wolfcampian marine faunas of the Hueco Group in the Robledo Moun tains, in spite of the highly fossiliferous nature of these sh ata. Shumard (1859) noted the "Upper Carboniferous" stratified na ture of Robledo Mountain, but reported no fossils from that range, although he did note late Paleozoic taxa from several parts of the nearby San Andres, Caballo and Fra Cristobal Mountains. Shumard (1886, p. 106) reported "Productus costatus, Athyris s 11btilita and Plwrotomaria, Chemnitizia, and S traparollus of undescribed species" from the Robledo Mountains. Although Shumard (1886) identified the fossiliferous stra ta as Carboniferous, these taxa were almost certainly collected from the Permian Hueco Group. Not only is the Hueco most of the accessible marine stra ta in the Robledos, but taxa reported by Shumard (1886), including Omphalotrochus ("Ple11rotomaria"), Wilkingia ("Cllemnitzia") and Euomphalus (Straparol/11s), are abundant and conspicuous in the Hueco (Kues, 1995).

Intertonguing of uncommon nonma rine red-bed "Abo" and more typical "Hueco" facies in the Robledos has been known for decades (e.g., Dunham, 1935; Thompson, 1942, 1954), but detailed studies of this section are considerably more recent (e.g., Jorda n, 1971; LeMone et al., 1975; Mack and James, 1986; Lucas et al., 1995). In spite of the extensive studies of Hueco faunas from cor rela tive sha ta to the east (e.g., Orogrande area, Hueco Mountains, Sierra Diablo area), the invertebra te fa una of the Hueco Group in the Robledos Mountains remained rela tively unstudied.

Thom pson (1954) reported a few Wolfca mpian fusulinids from the lower Hueco in the Robledo Mountains, and commented that most of the prolific fusulinid fa u na remained to be studied. LeMone et al. (1971 a,b) provided an abbrevia ted list of taxa and a lso published a brief study of the stra tigraphy and fa una! assemblages Loft the Robledo Mountains Apached Dam Macqueres in

## This area(LeMone et al., 1975) Kozur and LeMone (1995)

presented additional information on the conodonts of the Robledo Mountains Formation, and Kues (1995) was the first to document characteristic marine invertebrate taxa from the Robledo Mountains Formation.

# Mega-tracksite

The history of vertebra te fossil collecting in the Robledo Mountains is limited to the history of tracksite collecting, and was documented in detail by MacDonald (1994, 1995). Although a few track specimens had been collected from the Robledo Mountains over many years, collecting increased with the opening of a public quarry (the Community Pit) in red beds of the Hueco Group. However, there was only one popular account of early collecting (Ratkevich, 1980), and only one specimen (of *Dimetropus*) ended up in a museum (Hunt et al., 1993). In 1987, MacDonald made a major find (NMMNH locality 846), and the next year he informed the BLM, who administer the area. MacDonald made extensive collections, and several popular articles were written about the Robledo tracks (e. g., Bowlds, 1989a, b; Garretson, 1989; MacDonald, 1990, 1992; Stewart, 1992). In 1990, the U.S. Senate passed a bill funding the study of these tracksites. Hun t et al. (1993) and Lucas et al. (1994b) published preliminary studies of the tracksites, and Lockley and Hunt (1995) illustrated several specimens.

Schult (1994) wrote his doctoral disser ta tion on the Robledo tracksites and authored three papers on this topic (Schult and Fa rlow, 1992; Schult, 1995a, b). Schult (1994, 1995b) listed the occmrence of 23 teh-apod ichnogenera in the Robledo Mountains Forma tion, but this purported ichnofa una includes some ichnotaxa known only from pre-Permian sh-ata (e.g., *A11thracop11s*) as well as forms only reported from eolia n dune facies (e.g., *Laoporus*), so we believe that this ichnotaxonomic evaluation is largely incorrect.

Ichnofossils have long been known from Lower Permian red beds of Europe, and their extensive history of study is welldocumented (e.g., Haubold, 1971, 1984). Vertebrate and



FIGURE I. Generalized geological map of the Robledo Mountains showing location of the study area in Figure 2 (based on Seager et al., 1987).

invertebra te hackways are also known from many localities in Lower Permian red beds of the America n Southwest (e.g., Gilmore, 1926; Hunt et al., 1990; Lockley and Madsen, 1993; Hunt et al., 1995b). However, most localities yield only a few taxa, and few have been studied in detail. The Lower Permian localities in the Hueco Group of the sou thern Robledo Mountains surpass all others i n quantity, quality, and diversity of ichnotaxa. f ndeed, they represent the most scientifically important t Ea rly Permia n terrestrial ichnofa una known (Lucas et al., 1994a).

#### STRATIGRAPHY

Three formations of the Hueco Group were mapped in the stud y area (in ascending order): Communi ty Pi t Formation, Robledo Mountains Formation, and Apache Da m Formation (Fig. 2). The lowest formation of the Hueco Group in the Robledo Mountains, termed the Shalem Colony Formation by Lucas et al. (1998), crops out well to the north of the mapped area. For purposes of this study we will concentrate on the Robledo Mountains Formation, with only minor attention to the bounding Community Pit (below) and Apache Da m (above) Formations.

# Community Pit Formation

The Community Pit Forma tion of the Hueco Group crops out in the northeastern and north-central parts of the study area (Fig. 2). Brownish-gra y and grayish-orange packstone and micritic limestone and shale/siltstone domina te the Community Pi t Forma tion. No red-bed siliciclastics are present in the Community Pi t Forma tion; the base of the overlying Robledo Mountains Forma tion is mapped at the base of the stratigraphically lowest red bed s. The Community Pit Forma tion of the Hueco Group in the map area is about 85 m thick. Well-preserved permineralized logs of gymnospermou s wood are present in gray calca reous shale about 3 m below the top of the Community Pit Forma tion at locality 3016 (Tidwell and Munzing, 1995). These logs clearly floa ted into and were buried in a shallow marine environment as drif twood.

#### Robledo Mountains Formation

Strata previously referred to as the Abo Tongue, Abo Formation, or Abo-Hueco Member in the Robledo and Dona Ana



FIGURE 2. Geological map of the study area and structural cross sections in the southern Robledo Mountains (after Lucas et al., 1995). Map units are: Plun = Community Pit Formation; Phr = Robledo Mountains Formation; Phu = Apache Dam Formation; Qa = Quaternary alluvium; QTsf = Santa FeGroup; Ti = intrusive.

Mountains (Seager et al., 1976, 1987; Mack and James, 1986; Mack et al., 1988, 1991) were named the Robledo Mountains Member of the Hueco Forma tion by Lucas et al. (1995) and are raised in rank to the Robledo Mountains Forma tion of the Hueco Group by Lucas et al. (1998). The type section of the Robledo Mountains Forma tion is our section G (Fig. 3; Lucas et al., 1995, figs. 4,7), which was described previously by Jordan (1971) and Lucas et al. (1995). At its type section, the Robledo Mountains Forma tion is 125.4 m thick. Most of the section is interbedded marine shale and nod ular limestone (34%) and nonmari ne red-bed sandstone (33%). Ledgy marine limestones (12%) and shale (13%) ma ke up most of the rest of the section; red-bed siltstones and mudstones are a very minor component.

Jordan (1971) and Krainer and Lucas (1995) provided detailed descriptions of the lithology of the Robledo Mountains Formation. Most Robledo Mountains Formation limestones are micri tic and rela tively unfossiliferous. Fossil iferous limestones are mostly bioclastic wackestones and packstones, some of which are domina ted by shell ma terial of tubular foraminiferans (especially *Tolypammina, Hypemmmina, Ammovertella, Globivalvulina, Hemigordius* and *Tuberitina*) (Krainer and Lucas, 1995) and ostracods (Kietzke and Lucas, 1995). Less common lithologies are bioclastic and foraminiferal grainstones. Calca reous sha les, typically yellowish gray in color, a re of ten interbedded with Robledo Mountains Forma tion limestones, and these strata yield most of the invertebrate macrofauna described by Kues (1995).

Red-bed strata of the Robledo Mountains Forma tion are domina ted by grayish red to pale red, fine-grained, micaceous, li tha renitic sandstone. Typical sedimenta ry structures include laminae and/or ripple laminae. A few sandstones are troughcrossbedded, hummocky bedded or have herringbone crossbeds. Raind rop impressions, mudcracks, leaf impressions and tetrapod footprints are common on bedding planes.

These characteristics suggest that most of the Robledo Mountains Formation is of marine origin and thus consists of characteristic Hueco Group li thologies-fossiliferous carbonates and calcareous shales. About one-third of the unit is red-bed siliciclastics that represent the intertonguing of facies typically associated with the Abo Formation to the north with facies of the marine Hueco Group. This is the basis of previous references to this i nterval as Abo Tongue, Abo Formation or Abo-Hueco Member, even though the bulk of the unit consists of typical Hueco Group marine facies. For this reason, we follow Lucas et al. (1995, 1998) and assign the Robledo Mountains Formation to the Hueco.

Lucas et a l. (1995) reported seven measmed stra tigraphic sections that encompassed all or part of the Robledo Mountains Forma tion in the study area (Fig. 3). These sections demonstra te that virtually all the red-bed tracksi tes in the Robledo Mountain s are at the same stratigraphic level, just above a highly distinctive limestone bed. Correla tion of the sections is based not just on this bed, but on an extremely fossiliferous marine ca lca reous shale/nodular limestone interval in the upper part of the Robledo Mountains Forma tion (Kues, 1995; Lucas et al., 1995) and on the base of the Apache Dam Forma tion of the Hueco Group (Lucas et al., 1995, 1998). Most of the h-acksi tes in the Robledo Mountains Forma tion thus constitute a rnega tracksite that covered at least 20 km<sup>2</sup>.

Incomplete sections (top missing) of the Robledo Moun tains Formation crop out in the Dof\a Ana Mountains northeast of the Robledo Mountains in T21S, RIE. Seager et al. (1976, p. 10-12, fig. 6, sheet 1) mapped the distribution and described a measured section of these rocks, which they referred to both as Abo Formation and as Abo Tongue. The preserved Robledo Mountains Formation in the Dof\a Ana Mountains is 81 m thick and consists of interbedded marine limestone/shales and red-bed siliciclastics similar to the strata exposed in the Robledo Mow1tains. We do not extend recognition of the Robledo Mountains Formation further to the east, into the San And res Mountains, to encompass homotaxial rocks-upper Abo Tongue of Bachman and Myers (1969)-because these strata are wholly red beds and best referred to as Abo Formation.

#### **Apache Dam Formation**

Lucas et al. (1998) na med the Apache Da m Forma tion of the Hueco Group for strata formerly termed the upper member of the Hueco Forma tion in the Robledo Mountains. This is the youngest Permia n stratigraphic unit exposed in the study area. It is ex tensively faulted and in truded i n the north ern portion of the study area and caps esca rpmen ts to the south . Jorda n (1971), LeMone et al. (1971a, 1975), Simpson (1976) and Lucas et a l. (1998) have studied the Apache Da m Forma tion in detail. These stra ta are mostly dark gray and brownish-gray algal-pla te limestones, thin biostromes and interbedded siltstones. They con tain a fossil biota domina ted by phylloid algae, cora ls and gashopods (LeMone et a l., 1971a, 1975). Total thickness of the Apache Da m Forma tion is about 122 m Gordan, 1971), though only about 62 m are exposed in the study area.

#### MICROFOSSILS

The Robledo Mountains Forma tion of the Hueco Group in the southern Robledo Mountains prod uces diverse and prolific microfossiJ assemblages domina ted by non-fusulinid forarniniferans and ostracods (Kietzke and Lucas, 1995; Kozu r and LeMone, 1995). Most of these microfossi ls are from a yellowish gray calca reous shale at the base of unit 30 of measured section C of Lucas et al. (1995; Kietzke and Lucas, 1995; Kues, 1995) in the upper part of the Robledo Mountains Forma tion (Fig. 3).

Kohn and Dewey (1990) described some ostracods from the A pache Darn Forma tion of the Hueco Group in the Robledo Mou nta ins. They concluded that these ostracods, domina ted by ba irdeaceans, indica te sha llow nea rshore marine conditions of normal salinity. We believe a similar environment is indica ted by the rnicrofossil assemblage described by Kietzke and Lucas (1995), which is slightly lower stra tigraphically than the assemblage described by Kohn and Dewey (1990).

Most of the non-ostracods in the upper Robledo Moun ta ins Forma tion assemblage suggest shallow marine condi tions. For example, holothmioids and arnmodiscid foraminifera ns indica te sha llow ma rine wa ters of normal salinity (e.g., La ne, 1964). Spirorbids suggest very shallow wa ters, whereas tetra taxid for ams indicate shallow to sublittoral waters (Lane, 1964; Stevens, 1966). Some of the ostracods in this assemblage are eurytopic, such as Rectobairdia, Acmtia and Hollinella (Melnyk and Maddocks, 1988a). Other species a re characteristic of muddy, nearshore waters: *Healdia silliplex*, *Mollocentina lewisi* and *Bairdia beedei* (Melnyk and Maddocks, 1988a). Sansabella is also characteristic of nea rshore marine wa ters (Kaesler and Denver, 1988; Kaesler et al.,1990), but Cavellilla edlllistonae is more typical of offshore wa ters, though it too can be found in neashore deposits (Melnyk and Maddocks, 1988a). Thus, most of the ostracods and other microfossils indica te a shallow, nea rshore marine environment of

normal salinity (Kietzke and Lucas, 1995).

Most of the Robledo Mountains Forma tion ostracods are longranging taxa found in Pennsylva nian and Lower Permia n strata. The exception is Cnvellinn edmistonne, which first appears at or close to the base of the Kindle/In aff. K. ftssi /obn interval zone of la test Wolfcampian-Leona rdian age in Texas (Melnyk and Maddocks, 1988b). On face value, this suggests a latest Wolfca mpian-Leonard ian age for the uppermost part of the Robledo Mountains Forma tion of the Hueco Group, an age assignmen t consistent with the la test Wolfcampian age determined by conodonts for strata lower in the Robledo Mountains Forma tion (Kietzke and Lucas, 1995; Kozur and LeMone, 1995; Lucas et a l., 1995). Indeed, the Wolfcampian-Leonard ian boundary may be close to the boundary between the Robled o Mountains and Apache Dam Forma tions of the Hueco Group in the Robledo Mountains.

#### **MACROINVERTEBRATES**

LeMone et al. (1971a,b, 1975) and Kues (1995) presented the most recent studies of the invertebra te macrofa una in the Hueco Group of the Robledo Mountains. The collections made by LeMone et al. (1971a,b, 1975) were from a variety of localities, but the more than 70 ma rine invertebrate taxa reported by Kues (1995) were collected from near the top of the Robledo Mountains Formation, in an approximately 10-m-thick interval of gray to tan shale and limestone, just below the highest red sandstone bed (Kues, 1995, fig. 3). In general, this assemblage is domina ted by brachiopo ds, bivalves, and gastropods, as well as ra rer representatives of other stenoha line groups, such as bryozoans, echinoids, crinoids, cora ls, sponges, na utiloids, and sharks (Kues, 1995). Here, we summarize the analysis of Kues (1995) and its bearing on the age of the Robledo Mountains Formation.

In the Robledo Mountains, fusulinids have been documented only from the Shalem Colony Formation of the Hueco (Thompson, 1954), and only two, long-ranging species found there are a lso present in the type Hueco, preventing detailed correla tion based on fusulinids (Kues, 1995). Furthermore, fusulinids are exceptionally rare in the Robledo Mountains and Apache Dam Forma tions of the Hueco Group (LeMone et a l., 1975). LeMone et al. (1975) argued for a late Wolfcamp ian age for these units based on a suite of invertebra te taxa said to be indica tive of that age, although many taxa they listed have long temporal ranges (Kues, 1995).

Among the macrofossil inver tebra tes, amrnonoids and brachiopods provide the best indicators of the age of the Robledo Mountains Formation (Kues, 1995). Kues (1995) reported two amrnonoids from the upper Robledo Moun tains Forma tion of the Hueco Group: Properrinites bosei (Plummer and Scott) and Metnlegocerns bnuloronx (White). Of these, Properrinites was originally described from the late Wolfca mpian Admira l Formation of Texas, and Metnlegocerns was first reported from the Leonardian Clyde Formation, which also yielded a more advanced species of Properrinites, P. rnnminsi (Kues, 1995). However, as Kues (1995) noted, Miller and Pa rizek (1948) reported specimens nearly identical to the Robledo Mountain Forma tion amrnonoids from the midd le Hueco, of probable l a te Wolfcampia n age, nea r Orogrande, New Mexico. Thus, the available amrnonoid evidence suggests that the Robledo Moun tains Forma tion is of late Wolfcampia n age, as Kues (1995) concluded.

The brachiopod fauna of the upper Robledo Mountains

Forma tion consists of a t least 16 genera and 19 species and includes numerous Wolfcampian species, and some species previously reported from strata no older than Leonardian. Kues (1995, p.64-65), however, noted that "the age significance of [Leonardian) species in the upper Hueco of the Robledo Mountains is somewhat equivocal, as Cooper and Grant (1972-1977) studied only a rela tively small number of Hueco species and the possibility of longer stratigraphic ranges (into the Hueco) for some of their Leonardian species exists." This is perhaps especially true of the most abundant brachiopod in the Robledo Mounta ins Formation, which is closely related to or conspecific with *Squamnria moorei* Muir-Wood and Cooper, and was first described from the earliest Leonardian Clyde Formation. Otherwise, many of the taxa reported by Kues support a latest Wolfcampian age for the Hueco Formation.

#### TRACKSITES

There are 33 localities that yield tehapod hacks in the Robledo Mountain s (NMMNH localities 846, 2811-2839, 2849-2852). The loca li ties are scattered over an area of about 20 km<sup>2</sup> in a structurally complex area, but they have been correlated by careful mapping and stratigraphic analysis (Lucas et al.,1995). By far the most important of these is NMMNH locality 846, which has been extensively quarried by J. P. MacDonald and has yielded most of the recovered specimens. This locality produced hacks from multiple stratigraphic levels (25) and preserved multiple layers of undertracks of some layers.

Popular accounts of the Robledo Mountains ichnofa una have suggested that it contains an unprecedented level of diversity (e.g., MacDona ld, 1994). This is also suggested by Schult (1994, 1995a, b). However, we agree with Ha ubold et al. (1995a) and bel ieve that the ichnod iversity has been greatly overestimated. This is in part due to the extraord inary wide range of gait- and substratum-influenced va riations of track morphology exhibited by this ichnofa una (Ha ubold et a l., 1995a), termed extramophological variation by Peabody (1948). This is partly the result of varying substra tum conditions (e.g., moisture). In addition, there are large numbers of undertracks (which can often be associated with original tracks). Fmthermore, iclrnotaxonomic inflation of the Robledo ichnofa unas was caused in part by a confusing ichnotaxonomic litera ture and a lack of intercontinental studies of Permian tracks, but the latter is beginning to change (e.g., Ha ubold et a l., 1995a, b; McKeever and Ha ubold, 1996; Haubold, 1996). The ichnotaxonomy u tilized here follows Haubold et a l. (1995a), who described and i llustra ted many Robledo specimens.

Bntrnchichnu s and Limnopu s are tracks of small and large temnospondyl amphibians, respectively (Haubold, 1971). The most common track is Dromopu s (Fig. 4), the track of an araeoscelid. Hyloidichnu s is the track of a ?diadectid, Dimetropus is the track of a large sphenacodontid pelycosa ur, and Gilmoreichnus probably represents a smaller pelycosa ur (Haubold, 1971). Bones of these trackmakers are generally not known from the Abo Forma tion in southern New Mexico (Vaughn, 1969; Berma n, 1993; Lucas et al., 1995).

The Robledo tracksites provide abund ant information for paleoecological studies. Schult (1994, 1995a) compared ichnotaxa between different layers at tracksites and related them to substra tum differences. However, as noted above, his iclrnotaxonomy, which forms the basis of his paleoecology, is at odds with ours, so we cannot easily evaluate his results.



It is clear that there are d ifferences in ichnotaxonomic composition between layers at NMMNH locality 846. MacDona ld 's layer 16 consists almost entirely of tracks of *Dromopus agilis*, whereas tracks on layers 4, 5 and 6 are almost exclusively *Batrachichnu s delicatulus*. Many of the *Batraclzichnus* specimens on layers 4-6 exhibit t a tridactyl morphology. Other layers (e.g., layer 10) have n umerous specimens of both *Batra chichnus* and *Dromopus* s. Several layers (e.g., layers 10, 21) conta in abundant trackways of *Dimetropus* nicolasi. There seems to be some evidence for a separa tion of *Dromopus* and *Batrachichnus* on different surfaces, which might have ecologica l significance.

The Robledo h-ackways have important potential for the study of locomotor evolution in the Paleozoic. The trackwa ys of *Dimetropus* are particularly interesting in that they suggest that traditional skeletal mounts of sphenacodontids need to be modified. These restorations suggest a wide gait, and that trackwa ys should preserve a tail drag. Robledo samples of *Dimetropus* indicate a relatively narrow gait and no tail drag (Hunt et al., 1993; MacDonald, 1994). In terestingly, Small (1993) found, in mowlting a *Dimetrodon* skeleton for the Denver Museum of Na tural History, that he could not articulate the limbs to achieve the traditional sprawled posture of this animal.

Schult (1994) and MacDonald (1994) considered the abundance of amphibian trackways in the Robledo tracksites to be a problem because they believed that these ichnofa unas formed on a saline tidal flat. Schult (1994) went to great lengths to discuss the very limited salinity tolerances of modern amphibians. Lucas et al. (1995) noted that: (1) Robledo tracksites were made on tidal flats, but were not right at the shoreface and were not necessa rily subject to saline wa ters when the tracks were made: (2) the salinity tolerances of modern amphibians are i relevant as lissamphibians are very distantly related to Early Permian amphibians; and (3) tracks of spiders and other terreshial invertebra tes not tolerant of high salinity are common at the Robledo tracksi tes (Brad dy, 1995). We add other features that are relevant to this discussion: (1) the primi tive (and xerophytic?) conifer Wa/chia is common at tracksites (Hunt, 1983; Hunt et al., 1993); (2) despite claims to the contrary (e. g., Hunt, 1993), the only a mphibia n family ever to have a docu mented high-sal inity tolerance is the Triassic Trema tosa uridae; (3) ver tebra te tracks are rare on saline tida l flats, whereas infa una! invertebrate traces are abund ant (Frey and Pember ton, 1986, 1987)-the latter are absent at Robledo tracksites; and (4) there are no differences between the ichnotaxonomy of the Robledo sites and those of other redbed sites in u nequivocally freshwa ter settings (Haubold et al., 1995a; Hunt et al., 1995a, b, c). Therefore, we conclude that there is no evidence that the Robledo tracks were formed on a saline tidal flat.

Severa l ver tebra te tracksi tes conta in an abundance of inver tebrate trails (notably NMMNH localities 846, 2851) or pla nt fossils (notably NMMNH loca lity 2828) (Braddy, 1995). The invertebra te trails are mostly of arachnids and arthropods, a lthough eurypterid and limulid tracks also are present (Bradd y, 1995). The plaeoflora is almost monospecific and consists principally of *Walchia piniformi s*. This primi tive conifer is the dominant plant in Ea rly Permia n floras in New Mexico (Hunt, 1983). La rge fronds are preserved a t severa l localities, notably NMMNH 2828. The abundance of this plant may reflect i ts true abundance or it may be a taphonomic artifact. *Walchia* is usually considered to be xerophytic beca use of its need le-like leaves and is taken to indicate at least a seasonally dry clima te, although the

fossil logs reported by Tidwell and Munzing (1995) from the underlying Community Pit Formation lack well-defined growth rings, indicating a less seasonal climate. Studies of paleosols and oxygen isotopes confirm a seasonally dry climate in southern New Mexico during the Early Permian (Mack et al., 1991).

Lucas et al. (1995) demonstra ted that the majori ty of the Robledo tracksites occur at one stratigra phic level over an area of 20 km<sup>2</sup> and thus constitute a megatracksite (*sensu* Lockley, 1991). The Robledo Mountains mega tracksite is unique for a number of reasons: (1) it is the only pre-Middle Jurassic mega tracksite; (2) it is the only mega h-acksi te to include abundant invertebra te trails; (3) it has a much more diverse tetrapod ichnofauna than any other mega tracksite; (4) it is the only mega tracksite to be dominated by small (< 20 cm pes impression length) tetrapod tracks; (5) it is the only mega tracksite to occur in red beds; and (6) it is the only mega tracksite not to include dinosa ur footprints.

The Robledo Moun ta ins tracksi tes are the only Permian tracksites that can be correla ted without question to the global standa rd ma rine biochronology. Most of the Robledo ichnota xa are identical with, or have close relatives in, Eu ropea n teh-apod tracks (Haubold et al., 1995a), which have been used as the basis of local biostra tigraphies (e.g., Boy and Fichter, 1988). The presence of *Batrachichnu s* (similar to *Anthichnium*), *Gilmoreichnus*, *Hyloidichnus* and *Dimetropu s* /*eisnerianus* i n the Robledo ichnofa una of Europe (e.g., Gand and Ha ubold, 1988, figs. 1-2). La te Autwuan is consistent wi th a la te Artinskian age for the Robledo ichnofa una.

The Robledo Mou n tains tetra pod ichnofa una is broadly similar to those found in other Early Permian hacksites in western Nor th America (Hun t et al., 1995a, b, c). It differs from fluvialfacies ichnofa unas from the Abo Forma tion in centra l New Mexico (e.g., Hunt et al., 1995c) by including greater numbers of *Batrachichnus* and *Dimetropus* and in having fewer specimens of *Limnopus*. The Robledos ichnofa una differs from that of the Hermit Shale of Arizona and Sangre de Cristo Forma tion of northeastern New Mexico in lacking *Parabaropus* and *lchniotherium*, which seem only to have been present in more inland ichnofa unas (Hunt et al., 1995c).

Wolfcampian ichnofaunas of the American Southwest are taxonomically very similar to those in the Rotliegend of Europe, and many ichnospecies from both regions appear to be conspecific (Haubold, 1996). One iclmotaxon wluch is conspicuously absent from the Robledo ichnofa una, and from other U.S. tracksites, is *Amphisauropus*. The apparent absence of this iclmotaxon in Nor th Amer ica may be due to biogeographic or paleoecological (intermonta ne vs. lowland deposition) factors.

# DEPOSITIONAL ENVIRONMENT AND CYCLIPTY

Jorda n (1971, 1975), Mack and James (1986) and Mack et al. (1988, 1991) have carried out sedimentolog ical studies of the Robledo Mountains Forma tion and adjacent Lower Permian strata. More recently, Krainer and Lucas (1995) evalua ted the microfacies of Robledo Mountain Forma tion limestones and Lucas et al. (1995) described the stratigra phy and sedimentology of the Robledo Mountains Forma tion . These workers concluded that the Community Pit Forma tion of the Hueco represent shallow, marine shelf environments, whereas the Apache Dam Forma tion of the Hueco consists of shallow marine shelf limestone and sil tstone, including a diverse biota that allowed Jorda n (1971, 1975) and LeMone et al. (1971a, b, 1975) to recognize a variety of

biofacies. The Robledo Mountains Forma tion represents a complex intercala tion of siliciclastic tidal flat (red beds) and shallow marine shelf (limestones and calcareous shales) deposits. All of the Robledo tracksites are in the tidal flat deposits.

Our observations support in a general way the conclusions of Mack and associates regarding the depositional environments of the Robledo Mountains Formation. The dominance of micritic limestones in the marine facies of the Robledo Mountains Formation indicate deposition in a quiet environment on a shallow shelf. Some limestones, dominated by small foraminiferans and ostracods, suggest restricted (brackish?) depositional environments, whereas bioclastic wackestones and packstones with diverse, brachiopod- and bryozoan-dominated megafaunas suggest normal marine conditions (Krainer and Lucas, 1995).

The distribution of limestone fades in the Robledo Mounta ins Formation indicates a "deepening" upward or transgressive u pward trend within the member (Lucas et al., 1995). Thus, ostracod- and foramini feran-rich limestones are most abundant in the lower part of the Robledo Moun tains Forma tion, whereas megafa una-rich wackestones and packstones domina te limestones of the upper part of the member. We interpret this as a trend from resh-icted circula tion marine environments low in the Roblec!o Mounta ins Forma tion to more open shelf marine environments in the midd le to upper part of the unit. The transition occurs above the mega tracksite level.

The predomina nce of micri tic facies types within the fossi li ferous limestone horizons of the Robledo Mountains Forma tion indica tes deposition in a quiet wa ter environment of a shallow shelf, below active wave base (also see Mack and James 1986; Krainer and Lucas, 1995). LeMone et al. (1971a, b) suggested wa ter depths of less than 10 m.

Limestones containing a restricted fauna composed mostly of ostracods and /or small fora minifera ns point to a restricted (?brackish) depositional environment (Krainer and Lucas, 1995). The bioclastic wackestones/ packstones with a d iverse fa una indicate normal marine conditions and deposition in a quiet water depositional environment below the active wave base. Limestone



FIGURE 4. Tracks of the ichnogenus Drolllopl1s from NMMNH locality 846. Scale in cm.

horizons are frequently under- and overlain by ostracod-rich shales of a brackish environment.

Within the Robledo Mountains Forma tion, ostracod- and foraminiferan-rich limestones a re more abundant in the lower part, whereas bioclastic wackestones domina te in the upper part. Thicker limestone horizons are composed of ostracod mudstones at the base, grading upward into ostracod wackestones, which in turn grade into foraminiferal wackestones and grainstones and f inally into bioclastic wackestones (Lucas et al., 1995). The bioclastic wackestones frequently are overlain by ostracod- and foraminifera n-rich wackestones and mudstones. This reflects grada tion from a restricted environment at the base to an open shelf environment t in the central part or on top ("deepening upward" or transgressive trend), and grada tion to a restricted environment ("shallowing upward" or regressive trend) to the top of thicker limestone horizons.

Mack and Ja mes (1986) interpreted red-bed silicicla stics of the Robledo Mou n ta ins Forma tion as representing three tidal-fla t facies: (1) ripple-la mina ted sandstones deposi ted on i n tertida l sandfla ts nea r mean low tide; (2) "mixed sandstone-shale" deposited landwa rd of the ripple-la mina ted sandstones, on an in ter tidal fla t; and (3) nod ula r (pedogenic calcrete) shale deposited in a supra tidal setting. We agree with the interpreta tion of Mack and Ja mes (1986) that the red-bed siliciclastics of the Robledo Mountains Forma tion represent tidal-fla t facies, but d i ffer in our interpreta tion of specific facies. This d ifference reflects our view of deposi tional cyclici ty (transgression-regression) i n the Robledo Mountains, which is essentially diametrically opposed to that of Mack and associa tes (Fig. 5). Our in terpreta tion, however, is restricted to the mega tracksite level, which we have studied in great detail.

Mack and associa tes viewed li mestones immedia tely below and above packages of red beds in the Robledo Mou n ta ins Forma tion as maximum points of transgression. Overlying red beds were interpreted as largely regressive, with the next transgression beginning in the middle (symmetrical cycle) or upper (asymmetrical cycle) portion of the red-bed package (Fig. 5). Like Mack and associa tes, we agree that limestones bounding



FIGURE 5. Depositional cycles of the Robledo Mountains Formation as interpreted by Mack and associates compared with our interpretation.

red-bed packages in the Robledo Mountains Forma tion represent maximum hansgression, or, more accurately stated, local sea-level highstand (Lucas et al., 1995). However, we view the subsequent regression as an event that did not lead to accumulation of sed imen t. Instead, the lowering of local base level produced by the regression (lowstand) resulted in the development of an unconformity surface on top of the high-stand limestone. During the subsequent transgression, sedimen t bega n to accumula te as base level began to rise. In the case of the mega tracksite level, the patchy distribution of thin shoreface sandstone (Fig. 3, section A, unit 3), thick shoreface sandstone (Fig. 3, section G, unit 16), tidal flat sandstone and siltstone (Fig. 3, section C, unit 4) and very localized delta foresets (at locali ty 2851) provide strong evidence of the infilling of an irregula r, incised landsca pe (paleotopography) developed on top of a highstand marine limestone (Luca s et al., 1995). Vuggy recrysta lliza tion of the top of the h-ansgressive limestone underlying the mega tracksite level a lso suggests subaeria l exposure (Lucas et al., 1995).

Using the megatracksi te as the best studied example (it is an asymmetrica l cycle in the terminology of Mack and associa tes), continued base-level rise formed extensive tidal flat environments leading to deposits covered with a wide range of invertebra te and vertebra te tracks. Continued rising base level caused paleosols to form on top of the tidal flats until they were flooded over by marine wa ters that deposited the next h ighstand carbona te. The existence and in terpreta tion of symmetrical cycles identified by Mack and associa tes is problema tic; none are present in our detailed measured sections of the Robledo Mountains Forma tion (Fig. 3).

The differences between our interpretations of depositional cyclicity in the Robledo Mountains Formation and those of Mack and others are both observational and conceptual. The principal difference between our observations and those of Mack and associates is that we did not observe calcareous marine shales overlying transgressive limestones. Instead, relatively coarse-grained sediments directly overlie the limestones and fine upward into siltstones and shales. If, as we argue, the top of the marine limestone is an unconformity and/or lowstand, then the elastic sediments above that unconformity and the next marine limestone above the elastics form a fining upward sequence (Fig. 5). This fining upward sequence cannot readily be interpreted as regressive, because regression usually produces a coarsening-upward sequence (Dalrymple, 1992).

This highlights the conceptual differences between our interpretation and th t of Mack and associates. As Dalrymple (1992, p. 212) observed "no modern examples of regressive, prograding tidal systems are sufficiently well documented to serve as a model" and "there are also surprisingly few ancient examples [of regressive prograding tidal systems]." Regression in these environments is characterized by erosion and sediment bypassing as base level falls. Although regressive prograda tion may backfill some estuaries (e.g., Dalrymple et al., 1990), it seems unlikely that much sediment accumulates or is preserved in a tidal flat system during regression. For this reason, it makes much more sense to interpret sed iments in the Robledo Moun tains Formation as largely those that accumula ted during hansgression (Fig. 5). We thus *view* the hacksites as having formed on intertidal flats during transgression (Fig. 6).

Schult (1994) concluded that beca use of the tidal flat origin of the Robledo Mou ntains hacksi tes, the a mphibia ns who made many of the tracks were tolerant of high salinities. To support this conclusion, he reviewed the literature on salinity-tolerance in living amphibians, pointing out that a few salamander and frog taxa can tolerate a salinity of 40% seawater for extended periods of time.

The following evidence, however, runs contrary to Schult's (1994) conclusion that salinity-tolerant amphibians made many of the tracks at the Robledo Mountain sites:

1. Although the Robledo tracksi tes were made on tidal flats they were not righ t a t the shoreface and therefore not permanently subjected to saline wa ters (Fig. 6). Par ticula rly significant is the lack of deposit-feeder bioturba tion-indica tive of the shoreface-a t any Robledo tracksite. Instead, the Robledo deposits appear to have been in the intertidal zone and thus subaerial during low tides when the tracks were impressed (Fig. 6).

2. A few living lissa mphibian s capable of tolerating high salini ty is irrelevant to the salinity tolerances of Paleozoic temnospondyls. Lissa mphibians are d ista nt rela tives of temnospondyls; they are distinct subclasses of the class Amphibia. Furthermore, a few salinity-tolerant lissa mphibians are hardly representative of the Lissamphibia, almost all of which can only tolerate freshwater. There is essentially no direct evidence of salinity tolerance by temnospondyls, except for the Triassic trematosaurs.

3. Trackways of spiders (*Octopod ichnus*) and other invertebra tes that are not salinity tolerant are common at most of the Robledo hacksites.

4. The conifer *Wnlchin*, commonly preserved as complete lea f impressions at the Robledo h-acksites, must have lived very close to the tid al flats and was probably not salinity tolerant.

We therefore conclude that the Robledo Moun tain tracksites formed on tidal flats du ring rising base level due to transgression. The flats were in the intertidal zone and subjected to frequent subaeria 1 exposu re. Small temnospond y 1 amphibia ns and araeoscelid reptiles were the domina nt tetrapod trackma kers. Scorpions and spiders were the most common invertebra te track ma kers. An extensive forest domina ted by the conifer *Wnlchin* shroud ed the landscape.



FIGURE 6. Block diagram showing inferred depositional environment of Robledo Mow1tains megatracksite (based, in part, on a diagram in Dalrymple, 1992).

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