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Siluro-Devonian trace fossils from the Mereenie Sandstone, Kings Canyon, Watarrka National Park, Amadeus Basin, Northern Territory, Australia

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Nine trackways referable to the ichnogenus Diplichnites are preserved in the upper Silurian to Lower Devonian Mereenie Sandstone at Kings Canyon, Watarrka National Park, Northern Territory, Australia. Eight trackways are consistent with earlier descriptions of D. gouldi, and one trackway could not be assigned to an ichnospecies. The trackways are co-preserved with a range of sub-horizontal burrows referable to Beaconsites and Taenidium, and several vertical burrows, surficial circular traces and a horizontal trail of uncertain identities. The ichnofossil assemblage highlights the diversity of animals present in the late Silurian to Lower Devonian paralic to fluvial environments of central Australia at the time of early colonization of the land's surface. The assemblage is similar to ichnofaunas from coeval strata elsewhere in Australia and throughout Gondwana, and it highlights the potential of this region for further ichnological studies to elucidate the early stages of terrestrialization in the palaeoequatorial belt.

Keywords: ichnofossils, arthropod trackways, eolian and fluvial environments, terrestrialization, late Silurian, Early Devonian.

INVERTEBRATE trace fossils have been found in Australian continental settings from the Early Cambrian (e.g., Walter et al. 1989) and have provided important information on the ecological structure of early terrestrial communities. By the late Silurian and Early Devonian in Australia, a diverse terrestrial invertebrate fauna existed based on the presence of trace fossils described from the Tumblagooda Sandstone in Western Australia (Trewin & McNamara 1994), Major Mitchell Sandstone in western Victoria (Gouramanis et al. 2003), and Mt. Daubeney Formation (Neef 2004a) and Ravendale Formation (Neef 2004b) in New South Wales. Encroachment of animals onto the land’s surface globally at this time was probably facilitated by the contemporaneous diversification of bryophytes and early vascular plants as evidenced by both macrofossil and palynological records (Gray et al. 1974, Hagström & Mehlqvist 2012, Mehlqvist et al. 2015, Edwards & Kenrick 2015).

The upper Silurian–Lower Devonian mixed eolian and fluvial strata forming the upper Mereenie Sandstone in the western section of the George Gill Range (Wells et al. 1970, Bagas 1988) were deposited in sedimentary environments similar to those of the Tumblagooda Sandstone and Grampians Group. During the Silurian and Early Devonian, central Australia was located within 15° of the equator (Pickett et al. 2000, Talent et al. 2000). Thus, the Mereenie Sandstone and its stratigraphic equivalents in western and southeastern Australia preserve the sedimentological and ichnological expression of tropical lowland environments that were being colonized by early terrestrial plant and invertebrate communities.

The Mereenie Sandstone crops out prominently at the iconic Kings Canyon within the Watarrka National Park, approximately 240 km west-southwest of Alice Springs within the Amadeus Basin, Northern Territory, Australia (Wells et al. 1970, Bagas 1988; Fig. 1). The invertebrate trace fossil assemblages of the Mereenie Sandstone have not been described previously, although Wells et al. (1970) recognized Skolithos pipercork from the lower Mereenie Sandstone and Bagas (1988, p. 14) noted “indeterminate trace fossils on bedding planes” near the ‘Garden of Eden’ staircase in the Watarrka National Park that resemble large interstitial burrows. Several other trace fossils occur within the Mereenie Sandstone at Kings Canyon, and some of the major forms are identified and described here for the first time.

Materials and methods

Several ichnogenera are preserved on bedding planes of the Mereenie Sandstone located on or near the Kings...
Canyon tourist walk, Watarrka National Park (Fig. 1). Examples of these traces were photographed during two field campaigns in July 2012 and April 2015. In particular, nine large trackways (T1–T9) were recorded, with four located on the thin, horizontal to shallow-dipping beds forming the natural steps on the eroded ‘beehive’ structures at Cotterills Lookout (24°15′4″S, 131°34′57″E; Fig. 2A, B: T1–T3; Fig. 2F, G: T4), four from an extensive pavement near Kings Canyon Rim Walk navy blue marker 46 (KCRW46; 24°15′8.27″S, 131°34′58.7″E; Fig. 2C–E: T5–T8), and a single specimen from a cemented block forming part of the stairs near KCRW80 (24°15′20.35″S, 131°34′36.1″E; Fig. 2H: T9).

Other traces described herein include a small trail near the top of the northern ascent to the canyon rim walk (24°14′59.96″S, 131°34′25.35″E; Fig. 3); small circular traces exposed in the eastern cliff of the ‘Garden of Eden’ walk (24°14′56.46″S, 131°34′59.71″E; Fig. 4); small backfilled burrows in blocks forming part of the stairs near 24°15′23.11″S, 131°34′33.43″E; Fig. 5); large backfilled burrows on the canyon rim walk (24°14′56.02″S, 131°34′54.11″E; Fig. 6); and a radial burrow-mouth trace near the eastern rim of the canyon (24°15′00.08″S, 131°35′01.08″E; Fig. 7).

Ichnology

*Diplichnites gouldi* Bradshaw, 1981 (Fig. 2A–G)

*Description and interpretation.* Eight trackways of biserial epichnial imprints attributable to *Diplichnites gouldi* were identified (Table 1). They have preserved lengths up to 0.61 m, internal widths of 10–117 mm, and
Fig. 2. *Diplichnites gouldi* trackways with arrows depicting the direction of movement. A, Photo, and B, interpretive diagram of trackways T1–T3 from Cotterills Lookout. C–E, Photos and interpretive diagram of trackways T5–T8 from KCRW46. F, Photo, and G, interpretive diagram of the trackway T4 from Cotterills Lookout. H, *Diplichnites* isp. (trackway T9) found on a boulder as float in cemented (arrows) stairs near KCRW80. Note sunglasses are 145 mm wide in each image and scale bars = 100 mm.
external widths of 36–161 mm. Individual appendage imprints in the smallest trackway are ca 1–7 mm in diameter; those in the largest trackway are ca 19–24 mm in diameter.

Trackways T1–T3 (Fig. 2A, B), T5–T8 (Fig. 2C–E) and T4 (Fig. 2F, G) occur as biserial parallel sequences of circular to ovoid epichnial depressions reflecting individual or compound imprints (Table 1). Epichnial mounds behind individual or grouped imprints are common and reflect sediment pushed backwards during forward propulsion, indicating the direction of movement. Trackways T6 and T7 transition from well-preserved epichnial depressions to parallel, continuous epichnial ridges, where overlying sediment has not been eroded (Fig. 2C–E). The transition from depressions to continuous ridges in T6 and T7 (Fig. 2C–E) is more likely due to preservation, i.e., where overburden has not been removed completely, rather than environmental conditions at the time of the trackway formation (cf. the animals walking from relatively dry to wet sediments as documented by Trewin & McNamara 1994).

The flanks of each trackway comprise sets of two to three closely spaced imprints that are either discrete or formed as superimposed clusters. Where the individual imprints are distinct, the orientation of the medial to lateral imprints are orientated forward (up to 20°), perpendicular or backwards (up to 26°) from the direction of movement. Typically, the distal and central imprints are larger and deeper and most likely to be preserved or identified, whereas the medial imprints are smaller and less commonly preserved. No medial ridges or grooves are present in any of these trackways.

Trackways T1 and T3 are straight and parallel; the animal was travelling eastwards. These trackways cross T2 but it is unclear as to which trackways were formed first (Fig. 2A, B). The T2 animal was travelling south-southeast before turning (29°) to the south. The animal that formed the short T4 was also heading east (Table 1, Fig. 2F, G), and part of the northern section of the trackway is obscured by a *Beaconites antarcticus* burrow (Fig. 2F). It is unclear from the preservation of the bedding plane how many tracks in a series are present (Fig. 2A, B).

The longer trackways (T5–T8, Fig. 2C–E) are all straight with T6 to T8 parallel, and T5 crossing T6 and T7 at angles of 20° and 29°, respectively. The push-back mounds observed behind the individual imprints of T5 indicate that the animal moved eastwards, whereas the animals producing T6 to T8 were moving in an east-northeasterly direction. It is assumed that T5 also crosses T8, but this is obscured by overburden. It is probable that T5 crossed T6 and T7 because T5 has distinct impressions between both T6 and T7, whereas the T6 and T7 imprints become obscure within T5. It is unclear whether T6 to T8 were formed at the same time, but the unidirectional movement and parallel nature of the trackways suggest that this is a possibility.

**Remarks.** There is great morphological and preservation variability both between and within individual *Diplichnites gouldi* trackways (e.g., Bradshaw 1981, 2010, Trewin & McNamara 1994). The forms preserved in the Mereenie Sandstone most closely resemble *D. gouldi* Form B from the Tumblagooda Sandstone, Western Australia (Trewin & McNamara 1994). Similar *D. gouldi* Form B trackways have been described from a range of sedimentary settings including coastal environments (Bradshaw 1981, 2010, Draganits et al. 2001), dry and damp aeolian surfaces (Trewin & McNamara 1994, Retallack 2009), subaqueous fluvial and damp subaerial surfaces (Woolfe 1993, Smith et al. 2003, Neef 2004a) and an emergent sheet-flooded fluvial environment (Neef 2004b). *Diplichnites gouldi* has been ascribed to myriapods (Briggs et al. 1979, Bradshaw 1981, Johnson et al. 1994, Trewin & McNamara 1994, Draganits et al. 2001, Smith et al. 2003, Neef 2004a), trilobites (Bradshaw 2010), eucarcinoids (Trewin & McNamara 1994, Retallack 2009) and xiphosurans (Trewin & McNamara 1994). It is likely that a similar non-marine aqueous to terrestrial trackway-producing arthropod existed in the Amadeus Basin as the Mereenie Sandstone was deposited.

**Diplichnites** isp. (Fig. 2H)

**Description.** This trackway (T9) occurs on a bedding plane within a boulder preserved as cemented float; the original source of the block could not be identified. The trackway is preserved as nine large distinct depressions with no obvious push-back mounds (cf. *D. gouldi*; Table 1, Fig. 2H). The direction of travel is impossible to determine, and the trackway may be an undertrack in which smaller appendage impressions are not preserved (see Goldring & Seilacher 1971).

**Remarks.** This trackway has dimensions that are slightly larger than those described above for *Diplichnites gouldi* (Table 1), and because it lacks smaller appendage imprints it is provisionally retained in a separate ichnological species. However, owing to its different style of preservation (as a probable undertrack), the apparent differences may be taphonomic. An arthropod trackmaker is proposed, but a more precise biological affiliation is not possible based on the available evidence.

**Small indeterminate trail (Fig. 3)**

**Description.** This incomplete linear trail is 242 mm long, 23 mm in maximum width, is truncated by a rock fracture at one end and dissipates via weathering at the other (Fig. 3A). The ichnofossil appears to be preserved as an undertrack, with all components of the trace preserved in positive relief (Fig. 3B). The trail is complex, consisting of weakly repeating sets of individual elements whose centres are ca 30–38 mm apart (Fig. 3C). The centre of the trail is defined by discontinuous sinuous traces (mostly lenticular mounds) that are 2 mm
wide and ca 9–11 mm long. This central trace is flanked closely on either side by one or two paired arcuate or reniform elements up to 2 mm wide and 10 mm long. Around 8 mm beyond the margins of these elements are sets of roughly circular mounds ca 2 mm in diameter and 11–17 mm apart longitudinally. These outermost elements are weakly preserved or absent on one side of the trail.

**Remarks.** This trail is notably smaller than the examples of Diplichnites described above. It also differs in the arrangement and shape of the individual trace components. The direction of movement of the trail producer is difficult to ascertain as there are no clear signs of ‘push-up’ mounds. However, in a few cases, individual imprints appear to be deeper on the right side (Fig. 3B, C) suggesting (if this is an undertrack) that movement was to the left of the illustrated block.

The relatively small size and multiple discrete elements making up the trail suggest an arthropod producer. However, the sinuous form of the central trace is atypical of arthropods. This small trail also lacks the clusters of multiple sharply discrete lateral appendage imprints of early insects (Minter & Braddy 2006) and most other arthropods. Nevertheless, the Kings Canyon trace are broadly similar in size and complexity to some specimens of Stiaria intermedia Smith, 1909 from the Mississippian of USA that consist of repeating sets of central linear, inner flanking lobate and marginal circular elements that have been tentatively attributed to aptyrygote insects, perhaps walking within a moving fluid (Fillmore et al. 2012).

We can not exclude a small fish as a producer of this ichnofossil, in which case the weakly sinuous to reniform character of the central and lateral traces might reflect fin-tip furrows (cf. Wisshak et al. 2004). Among roughly coeval ichnofossil assemblages from Australia, Siskemia sp. of Trewin & McNamara (1994) has a straighter and more continuous central ridge and lacks the inner zone of arcuate structures, and the outer zone has smaller and more densely spaced appendage impressions. There are also some similarities in terms of repeated lobate elements with putative fish traces from the Lower Triassic of Australia (Chen et al. 2012).

*Laevicyclus* isp. Weber & Braddy, 2003 (Fig. 4)

**Description.** These traces occur on the bedding plane at the foot of the metallic staircase leading up from the eastern cliff of the ‘Garden of Eden’ walk (24°14’56.46”S, 131°34’59.71”E) and were originally identified but not described by Bagas (1988). These epichnial traces range from circular to D-shaped with diameters between 68 and 189 mm (Fig. 4). The traces consist of a peripheral or coiled burrow 10–30 mm wide, surrounding a central bowl-shaped depression.
Remarks. The coiled burrow may represent a horizontal feeding trace and the depression may have formed by sediment collapse into a central cavity. Until vertical profiles of these burrows are examined, the threedimensional structure, mode of formation and producer remain speculative. Weber & Braddy (2003) identified similar but much smaller traces ascribed to *Laevicyclus* isp. from Lower Ordovician shallow marine strata of Antarctica.

**Beaconites antarcticus** (Vialov) emend. Keighley & Pickerill, 1994 (Fig. 5)

**Description.** Horizontal to sub-horizontal, straight or gently curved, parallel-sided, cylindrical burrows reaching >150 mm long and 5–13 mm in diameter (Fig. 5A, C). The burrows occur in fine- to medium-grained sandstone and have an infilling dominated by grains similar in size to the surrounding matrix but divided into weak compartments of unequal size by silty meniscate laminae (Fig. 5B, D). The burrows have a distinct lining, <1 mm thick, defined by a finer grain size (silt) and/or different colour to the remainder of the fill (Fig. 5B, D). The burrows are variably orientated within the same exposure (Fig. 5A, C).

**Remarks.** Small, roughly horizontal, lined burrows of this type are relatively common in the Mereenie Sandstone and are attributed to *Beaconites antarcticus* (Vialov 1962) emend. Keighley & Pickerill, 1994. This ichnotaxon was originally described from Devonian fluvial strata of Antarctica (Vialov 1962) but, subsequently, has been reported from a broad range of ages and depositional environments (Keighley & Pickerill 1994 and references therein).

Burrows similar or equivalent to *B. antarcticus* are common in eolian sandsheet and pond facies of the Tumblagooda Sandstone, Carnarvon Basin, Western Australia (Trewin & McNamara 1994) and this taxon is a key element of Bradshaw’s (1981) *Beaconites* Ichnocoenosis’ recognized in Devonian fluvial to paralic deposits of Victoria Land, Antarctica. *Beaconites antarcticus* and *Taenidium barretti* are commonly associated in mid-Palaeozoic fluvial deposits globally (Bradshaw 1981, Allen & Williams 1981, O’Sullivan *et al.* 1986, Marriott *et al.* 2009), and both are represented in the Mereenie Sandstone. The identity of the *B. antarcticus* burrow maker is uncertain, but such traces have generally been attributed to small arthropods or, less commonly, to annelids and even holothurians (Bradshaw 1981, Weber & Braddy 2003).

**Taenidium barretti** (Bradshaw) emend. Keighley & Pickerill, 1994 (Fig. 6)

**Description.** Large, more or less straight, horizontal to oblique burrows filled by muddy sand within beds of fine-grained sandstone. The burrows are up to 470 mm long and 27–84 mm wide, irregularly orientated (Fig. 6A) and variably isolated (Fig. 6B) or clustered (Fig. 6C). The burrows are weakly septate, incorporating poorly developed and somewhat irregular meniscate
backfill compartments that are ca 20–25 mm wide (Fig. 6D, E, H). Regular complex arcuate or *en echelon* markings are developed along the margins of several burrows repeating at intervals of 10–25 mm depending on the size of the burrow (Fig. 6G, I). A few burrows are devoid of surface markings (Fig. 6B), and one bears fine transverse striations (Fig. 6F). The burrows are preserved in the same beds as small vertical burrows (*?*Skolithos sp.; Fig. 6C) and are locally associated with extensive asymmetrical ripple fields (Fig. 6A).

**Remarks.** Originally placed in *Beaconites* Vialov, 1962 by Bradshaw (1981), this ichnospecies was transferred to *Taenidium* by Keighley & Pickerill (1994) on the basis that, although menisci merge at the burrow margins, no distinctive burrow lining is developed. Those burrows lacking obvious surface markings in the Mereenie Sandstone may have been subject to pedestrian attrition. The fine transverse striations on one specimen (Fig. 6F) are interpreted to represent current lineations on a partially exhumed burrow. Most specimens are marked by somewhat irregular meniscate features that are linked to repetitive arcuate marginal traces interpreted to represent sediment push-back traces produced by arthropod appendages. The *Cruziana*-like surface striations developed in a few cases (Fig. 6C) also favour burrow production by a large arthropod (cf. examples illustrated by Gibb et al. 2009). Indeed, the segmentation evident in some burrows (Fig. 6H) is reminiscent of the dorsal tergite impressions of a large arthropod (cf. Fayers et al. 2010, pl. 1).

The burrow illustrated by Bagas (1988, pl. 12) from the Mereenie Sandstone appears to be referable to the same ichnotaxon as the examples described here. Similar burrow forms probably produced by large arthropods are relatively common in Siluro-Devonian strata of eastern Gondwana (Webby 1968, Bradshaw 1981, Trewin & McNamara 1994, Draganits et al. 2001, Gouramanis et al. 2003). As noted by Hocking (1991) and Trewin & McNamara (1994), morphologically similar traces described from diverse localities and ages may have been produced by a range of burrowing organisms. However, the similarity in size to *Diplichnites gouldi* and preservation in the same successions...
Fig. 6. Large, horizontal to oblique *Taenidium barretti* (Bradshaw) emend. Keighley & Pickerill, 1994 burrows. A, Several irregularly orientated burrows associated with a ripple-marked bedding plane. B, Low-angle view of two burrows illustrated in A. C, Cluster of intersecting burrows, some with *Cruziana*-like surface striations. Circled area hosts numerous ?vertical *Skolithos*-like burrow openings. D, E, Horizontal burrows bearing complex surface striations. F, Horizontal burrow bearing fine transverse striations interpreted to represent current lineations on the partially exhumed burrow fill. G, I, Horizontal burrows bearing arcuate or en echelon markings along the margins (arrowed). H, Segmented burrow reminiscent of dorsal tergite impressions of a large arthropod. Scale bars = 10 mm except in A (=100 mm); coin in B and C = 25 mm diameter.
but within separate facies may reflect different behavioural aspects of the same large arthropod (Fayers et al. 2010).

Small burrow openings (Fig. 6C)

Remarks. Small openings to probable vertical burrows are present on the bedding surfaces where the examples of *Taenidium barretti* (e.g., Fig. 6C) are preserved. These traces occur in clusters of several closely spaced openings and commonly in pairs. Unfortunately, vertical profiles of these burrows were not observed, thus making ichnotaxonomic classification impossible, although ichnospecies of *Arenicolites*, *Monocraterion* or *Skolithos* cannot be excluded.

Burrow entrance with radial feeding traces (Fig. 7)

Description. Circular trace fossil (in plan view) exposed on the upper surface of a sandstone bed (Fig. 7A). Trace having a total diameter of 37 mm, incorporating a central featureless zone (15 mm in diameter); outer zone ca 10–12 mm consisting of more or less regular and radially arranged alternating troughs and ridges with rounded crests (ca 20 ridges; Fig. 7B). Troughs broaden and deepen slightly with distance from the centre of the trace. The outer margin of the trace is entire or slightly lobate. The trace is preserved entirely in fine-grained sandstone with no obvious lithological difference between the central zone, ridges or troughs.

Remarks. This circular trace has a superficial resemblance to a medusoid cnidarian. However, most medusoid impression fossils are characterized by concentric contractional muscle or desiccation features. There are also superficial similarities to the putative Silurian ventobiont *Rutgersella* (Retallack 2015) in having a central disorganized zone surrounded by a broad rim of radial grooves and ridges. However, *Rutgersella* differs in being typically elliptical with finer radial components that bifurcate sporadically. The Kings Canyon trace best matches the general characters of the un-named stellate trace fossils illustrated by Collette et al. (2011), although the specific dimensions of the new trace do not match previous examples. The Kings Canyon trace is also reminiscent of an ichnofossil illustrated by Shi et al. (2009) and assigned to *?Lorenzinia* isp., which consists of radially arranged epichnial ridges surrounding a featureless central zone. However, the latter has only five radial ridges and occurs in bathyal turbiditic deposits. There are also similarities with *Dactyloidites* Hall, 1886 in its radial form. However, *Dactyloidites* is generally interpreted to represent a vertical or oblique burrow with a subterranean (i.e., basal) rosette of feeding tunnels (Wilmsen & Niebuhr 2014). By contrast, the Kings Canyon trace is interpreted to represent a central vertical or oblique burrow with a rosette of radial feeding traces on the bedding surface around the burrow mouth. This is consistent with the interpretation by Collette et al. (2011) of Early Jurassic stellate ichnofossils from ephemeral lacustrine deposits that are similar to the modern feeding traces of infaunal bivalves, such as *Macoma tenta*. Without additional specimens and vertical sections, we cannot determine if there are multiple levels of radial features in this trace as represented in some examples of *Dactyloidites* (Fürsich & Bromley 1985).

Discussion and conclusions

The age of the Mereenie Sandstone is poorly constrained owing to a dearth of body fossils and the presence of unconformities at the base and top of the unit. The Mereenie Sandstone overlies the Carmichael Sandstone dated as Late Ordovician (Wells et al. 1970), and underlies the Park Siltstone of Middle to Late Devonian age (Young 1985). The similarities between the ichnofaunas of the Mereenie Sandstone and those of the Tumblagooda Sandstone (Trewin & McNamara 1994),
Major Mitchell Sandstone (Gouramanis et al. 2003), Mt Daubeney Formation (Neef 2004a) and Ravendale Formation (Neef 2004b) elsewhere in Australia favour a roughly contemporaneous (late Silurian–early Devonian) age of deposition.

The Mereenie Sandstone ichnofauna occurs in a succession that hosts extensive shallow-water sedimentary structures (trough, tabular and low-angle cross-bedding, and wave- and current-ripple cross-lamination; Bagas 1988). Some trackways appear to be developed on emergent surfaces with thin silty coatings. Other traces include vertical or sub-horizontal burrows that were clearly produced by subsurface activities in unconsolidated sediment. A few traces appear to be influenced by active water currents during or after their formation. The co-occurrence of Diplichnites, Beaconites, Taenidium and Skolithos-like traces appears to represent a typical ichnofossil association of mid-Palaeozoic paralic to fluvial successions (Bradshaw 1981, Trewin & McNamara 1994, Gouramanis et al. 2003, Marriott et al. 2009), although some similar assemblages may be found in strata as old as the Early Ordovician (Weber & Braddy 2003).

The ichnofaunal assemblage of the Mereenie Sandstone represents an expression of the development of complex ecosystems at the interface of terrestrial and aquatic environments during the mid-Palaeozoic. The assemblage highlights the preliminary steps leading to the more extensive terrestrialization of animals and plants that developed later in the Devonian. Our reconnaissance study of the Mereenie Sandstone ichnofauna highlights the probable presence of several clades of organisms invading paralic to fluvial environments and engaging in a diverse array of behaviours. We contend that future systematic surveys of the extensive exposures of this rock unit will probably reveal more diverse ichnofossil suites that, in turn, will provide a clearer perspective of the early steps towards biotic occupation of freshwater ecosystems and the land surface in the mid-Palaeozoic. In particular, this formation has strong potential for elucidating the initial enchooachment of arthropods (MacNaughton et al. 2002) and even early tetrapodomorphs (e.g., Niedźwiedzki et al. 2010) into shallow river systems and onto land in the Siluro-Devonian palaeoequatorial realm.

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