Controls on the Formation of Microbially Induced Sedimentary Structures and Biotic Recovery in the Lower Triassic of Arctic Canada

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Controls on the formation of microbially induced sedimentary structures and biotic recovery in the Lower Triassic of Arctic Canada

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ABSTRACT

Microbially induced sedimentary structures (MISS) are reportedly widespread in the Early Triassic and their occurrence is attributed to either the extinction of marine grazers (allowing mat preservation) during the Permo-Triassic mass extinction or the suppression of grazing due to harsh, oxygen-poor conditions in its aftermath. Here we report on the abundant occurrence of MISS in the Lower Triassic Blind Fiord Formation of the Sverdrup Basin, Arctic Canada. Sedimentological analysis shows that mid-shelf settings were dominated by deposition from cohesive sand-mud flows that produced heterolithic, rippled sandstone facies that pass down dip into laminated siltstones and ultimately basin mudrocks. The absence of storm beds and any other "event beds" points to an unusual climatic regime of humid, quiet conditions characterized by near continuous run off. Geochemical proxies for oxygenation conditions characterized by near continuous run off. Geochemical proxies for oxygenation (Mo/Al, Th/U, and pyrite framboid analysis) indicate that lower dysoxic conditions prevailed in the basin for much of the Early Triassic. The resultant lack of bioturbation allowed the development and preservation of MISS, including wrinkle structures and bubble textures. The microbial mats responsible for these structures are envisaged to have thrived, on sandy substrates, within the photic zone, in oxygen-poor conditions. The dysoxic history was punctuated by better-oxygenated phases, which coincide with the loss of MISS. Thus, Permo-Triassic boundary and Griesbachian mudrocks from the deepest-water settings have common benthos and a well-developed, tiered burrow profile dominated by Phycosiphon. The presence of the intense burrowing in the earliest Triassic contradicts the notion that bioturbation was severely suppressed at this time due to extinction losses at the end of the Permian. The notion that Early Triassic MISS preservation was caused by the extinction of mat grazers is not tenable.

INTRODUCTION

The post-mass extinction world of the Early Triassic is often considered to represent an unusual time in Earth history when marine recovery from the Permo-Triassic mass extinction (PTME) was delayed by the prevalence of marine anoxia (Hallam, 1991; Twitchett and Wignall, 1996; Woods et al., 2007; Wignall et al., 2010, 2016; Grasby et al., 2013, 2016; Pietsch et al., 2014), and anachronistic facies better known from Cambrian and earlier times were well developed; especially carbonate microbialites (e.g., Bagherpour et al., 2017) and microbially induced sedimentary structures (MISS) in sandstones (Wignall and Twitchett, 1999; Pruss et al., 2004, 2005; Baud et al., 2007; Noffke, 2010). In recent years a broad range of sandstone bedding features have been related to the presence of microbial mats at the time of deposition. These comprise textured surfaces, including small-scale wrinkles and more regular, parallel ridges, that reflect the original microtopography of the mat. They also include impressions of bubbles trapped beneath the mats, all of these features are generally grouped together as MISS (Noffke et al., 2001; Davies et al., 2016).

Only the shallowest waters provided better oxygenated settings and a refuge from the widespread, inimical conditions of the Early Triassic (Wignall et al., 1998; Beatty et al., 2008; Knaust, 2010; Chen et al., 2011; Song et al., 2013; Proemse et al., 2013). MISSs are rarely reported from such environments indicating their poor preservation potential, because the mats were either grazed, or they could not develop in the first place because of the unstable, burrowed sediment surfaces. In contrast, others have argued that Early Triassic open marine conditions were not unusually stressful (Hofmann et al., 2013; Vennin et al., 2015). Instead, the loss of bioturbators during the PTME is said to have been responsible for the preservation of fine lamination (Hofmann et al., 2015), and a reduction in the depth of the sediment mixed layer (Buatois and Mángano, 2011). In this scenario, geochemical evidence for sediment anoxia is attributed to a lack of irrigation by bioturbation (Hofmann et al., 2015). Davies et al. (2016) has also challenged the uniqueness of supposedly anachronistic “Precambrian” MISS and argued that they are in fact commonplace in shallow marine Phanerozoic settings; however most of their post-Cambrian examples are from peritidal or fluvial settings.

This study aims to provide the first sedimentological study of the type locations of all four Early Triassic substages that are found in the Sverdrup Basin of Arctic Canada (Ellesmere and Axel Heiberg islands). The ammonoid-defined age of the strata is independently assessed by constructing a chemostratigraphic record. Having established an age model, sedimentological context and redox variations, we then record and discuss the origin of the abundant MISS that occur in the basin. Finally, we evaluate the broader significance of these occurrences in the debates on the significance of Early Triassic facies and the nature of recovery from the PTME.

REGIONAL GEOLOGY OF STUDY AREA

Present-day outcrops of Sverdrup Basin strata lie in the northern-most part of Nunavut, Canadian High Arctic. The basin extends ~1000 km
Figure 1. Regional paleogeography of the Sverdrup Basin, Arctic Canada, in the Early Triassic (from Embry and Beauchamp, 2008). Lower Triassic regional stratigraphy is given and thicknesses are those measured at Spath Creek/Cape St. Andrews. Red stars mark the field locations. Fm.—Formation.

The four substages of the Early Triassic were established within the Blind Fiord Formation by Tozer (1965, 1967) and defined by their ammonoid content. His four type localities (Griesbach Creek, Diener Creek, Smith Creek, and Spath Creek continuing to Cape St. Andrews; Figs. 1 and 2) were the subject of our study during an expedition in July 2015. The basal 50 m of Confederation Point Member strata, with the exception of the unexposed basal-most meter, were logged in the western bank of Griesbach Creek on Axel Heiberg Island (Fig. 2). The other three sections lie on the Svartfjeld Peninsula, south of Otto Fiord in northwestern Ellesmere Island. The steep, eastern slopes of Spath Creek provide a series of near-continuous outcrops of the lower part of the Blind Fiord Formation, except for the basal 80 m which are covered. Our measured section began at N80° 54.464′ W89° 11.571′ (NAD83) in the upper part of the Confederation Point Member and continued to a level near the top of the Smith Creek Member where a major dolerite sill forms a prominent ridge. Logging was resumed to the northeast, at the same stratigraphic level (with some overlap), on the nearby slopes of Cape St. Andrews (N80° 55.072′ W89° 14.491′) and continued up to the lower beds of the Svartfjeld Member (Fig. 3). In addition to these two locations the same levels were also examined (and sampled) at Smith Creek and at Diener Creek (Fig. 2).

METHODS

Facies and Fossils

Sedimentary logging was undertaken through the basal 640 m of the Blind Fiord Formation at Griesbach Creek and Spath Creek to Cape St. Andrews, and facies were also examined and sampled at Diener and Smith Creeks (Fig. 3). Trace and body fossils were identified in the field and the intensity of bioturbation assessed using the semiquantitative ichnofabric index (II) scale of Droser and Bottjer (1986) which ranges from 1 (no burrows) to 5 (intense burrowing, no primary sedimentary structures preserved). The field observations were supplemented with analysis of 24 thin sections and polished slabs.
Pyrite framboid size analysis is a useful tool for reconstructing ancient redox conditions. In modern environments, pyrite framboids form in the narrow iron-reduction zone developed at the redox boundary, but they cease growing in the more intensely anoxic conditions of the underlying sulfate-reduction zone (Wilkin et al., 1996; Wilkin and Barnes, 1997). If bottom waters become euxinic (i.e., free H₂S occurs within the water column), then framboids develop in the water column but are unable to achieve diameters much larger than 5–6 µm before they sink below the iron reduction zone and cease to grow (Wilkin et al., 1996). Euxinic conditions are therefore characterized by populations of tiny framboids with a narrow size range whereas dysoxic/weakly oxygenated seafloors contain framboid populations that are larger and more variable in size (Bond and Wignall, 2010). Twenty-five samples from Griesbach Creek, Diener Creek, and Spath Creek-Cape St. Andrews were examined using carbon-coated polished chips viewed in backscatter mode at magnification ×2500 under FEI Quanta 650 (at University of Leeds, Leeds, UK) and Zeiss EVO-60 (at University of Hull, Hull, UK) scanning electron microscopes to determine pyrite content, and where present, the size distribution of pyrite framboids. Where possible, at least 100 framboids were measured from each sample (levels bearing statistically robust numbers of pyrite framboids are shown on Fig. 3). Framboid size distributions are plotted with mean diameter versus standard deviation within each sample, which allows comparison of framboid populations in ancient sediments with modern euxinic, anoxic, and dysoxic populations (Bond and Wignall, 2010).

RESULTS AND INTERPRETATIONS

Stratigraphy

The basal part of the Confederation Point Member at Griesbach Creek begins with shale and coarsens upwards to siltstone (Fig. 3). For the most part these strata are blocky although platy siltstone is developed for 3.5 m beginning 6.0 m above the base of the section. In the uppermost part of the measured section, siltstone and tabular sandstone beds are interbedded and, above the level studied here, a further 20 m of sandstone and shelly limestone beds completes the coarsening-upward unit. The basal 20 m of the Confederation Point Member was also examined at Diener Creek where it comprised shale in the lower half that coarsens up to fissile siltstone.

A 180-m-thick section of the upper part of the Confederation Point Member was examined at Spath Creek. Sandstone dominates the succession although there are many siltstone levels as well as several unexposed stretches that are probably dominated by finer-grained strata based on the presence of shale and siltstone chips seen in the ex-situ scree material. A major (30 m-thick) sandstone forms the topmost part of the Member.

The transition between the Confederation Point and overlying Smith Creek members
Figure 3. Composite stratigraphic log with sedimentary fabrics and ichnofabric index (II, after Droser and Bottjer, 1986) values through the lower part of the Blind Fiord Formation seen at Griesbach Creek, Spath Creek, and Cape St. Andrews (Arctic Canada). The position of samples analyzed for their pyrite framboiid size distributions are marked and numbered 1–16 (see Fig. 9). Mb.—Member.
occurs within a 70-m-thick unexposed interval at Spath Creek. This “gap” was examined at Smith Creek where thinly bedded sandstone strata were seen to dominate. The remainder of the Smith Creek Member is well exposed at Spath Creek and consists of gradual grain size fluctuations, on a scale of tens of meters, from siltstone to sandstone (Fig. 3). Superimposed on these changes are several levels where alternations of silty sandstone and sandy siltstone occur on a decimeter scale. Despite these subtle variations, weathering produces distinctive “banded” outcrops that are particularly well developed in the cliff sections on the western slopes of Diener Creek (Fig. 5).

The overlying, 320-m-thick Svartfjeld Member, seen immediately to the northeast of Spath Creek in the continuous section at Cape St. Andrews, is dominated by laminated shales. Prominent, yellow-weathering dolomite concretions and beds, and organic-rich shales are characteristic of the upper part of the member but not in the lower part considered here.

The ammonoid stratigraphy of the Blind Fiord Formation has primarily been based on collections from the same locations we have studied here (Tozer, 1965, 1967). The Dienerian/Smithian substage boundary was placed around the Confederation Point/Smith Creek contact and the Smithian/Spathian substage boundary around the contact between the Smith Creek and Svartfjeld members (Tozer, 1965, 1967). Our $\delta^{13}C_{\text{org}}$ data (Fig. 4) generally confirm these age assignments. Thus, in the upper Confederation Point Member there is a major decline of $\delta^{13}C_{\text{org}}$ values before they stabilize at $-31\%$ in the basal Smith Creek Member. A similar prolonged negative shift has also been reported at Smith Creek (Grasby et al., 2013) and in $\delta^{13}C_{\text{carb}}$ records from Tethys (e.g., Horacek et al., 2007; Korte and Kozur, 2010) where the onset of the negative excursion is placed in the early Smithian. Using this age assignment, the Dienerian/Smithian boundary probably lies in the upper part of the Confederation Point Member, a little lower than suggested by the ammonite records of Spath Creek, and Cape St. Andrews (Arctic Canada). The Th/U ratio (right hand panel) was measured in the field using a Radiation Solutions portable gamma-ray spectrometer (circles), while the triangles record values obtained on the same samples using inductively coupled plasma–mass spectrometry/ emission spectroscopy. There is remarkable correspondence between the two curves, corroborating the application of field portable gamma-ray spectrometry.

![Figure 4. Carbon isotope values ($\delta^{13}C_{\text{org}}$) and trace metal redox proxies measured from the lower part of the Blind Fiord Formation seen at Griesbach Creek, Spath Creek, and Cape St. Andrews (Arctic Canada). The Th/U ratio (right hand panel) was measured in the field using a Radiation Solutions portable gamma-ray spectrometer (circles), while the triangles record values obtained on the same samples using inductively coupled plasma–mass spectrometry/emission spectroscopy. There is remarkable correspondence between the two curves, corroborating the application of field portable gamma-ray spectrometry. Confed. Pt. Mb. — Confederation Point Member; c — clay; s — sand; fs — fine sand.](https://pubs.geoscienceworld.org/gsa/gsabulletin/article-pdf/132/5-6/918/4987619/918.pdf)
reported from the same level elsewhere in the Sverdrup Basin (e.g., Grasby and Beauchamp, 2008). This facies also forms a substantial thickness of strata in the lower part of the Smith Creek Member in all sections. The laminae are defined by fine silt-rich layers interbedded with clay and occur on a millimeter scale.

This low energy facies is interpreted to be the result of suspension-settling of clay minerals with regular, weak traction currents transporting silt grains to the depositional site. The absence of bioturbation suggests anoxic conditions.

**Burrowed Mudstone**

Most of the fine-grained strata of the Confederation Point Member at Griesbach Creek are bioturbated by millimeter diameter burrows of *Phycosiphon* that can form short vertical spirals, oblique to bedding. Dark, clay-rich material is concentrated in the burrow center surrounded by thicker haloes of very fine silt (Fig. 6A). Ichnofabric index varies from 3–4 in the burrowed mudstone and the most intensely bioturbated examples show a tiered profile of some complexity with *Phycosiphon* cross-cutting *Planolites* and in turn being cut by oblique burrows of *Catenichnus* (Fig. 6A). This facies type also has a considerable fossil content that includes common ammonoids, large bivalves (*Claraia, Modiolus, Unionites*, aviculopectinoids), bellerophontids, and bryozoans. The valves of the bivalves are often intensely bored.

We interpret this facies to record a low energy, well oxygenated depositional environment that allowed a diverse benthic fauna to thrive. By the standards of early Griesbachian mudrock environments (e.g., Dai et al., 2018), this is a rich fossil assemblage.

**Laminated, Heterolithic Siltstone**

This facies type is common in the upper part of the Confederation Point Member and throughout the Smith Creek Member where it dominates the 355-m-thick unit. It is mostly silt but the grain size ranges from clay through silt to very fine sand and is composed of both quartz and pyrite grains. Sedimentary structures include millimeter- to centimeter-scale planar laminae with most a few millimeters thick, but they can reach 1 cm (Fig. 6B). Both fining and coarsening upward trends are common within the laminae and thin beds. In the latter case the coarsest quartz sand and pyrite grains are concentrated in the uppermost part of the laminae and the top surfaces can be weakly eroded (Fig. 6B). Fossils are generally rare but include ammonoids and large *Claraia* up to 4 cm in height. Occasionally, *Claraia* populations, dominated by larval shells, can be prolific and cover bedding planes (Fig. 7A).

The repeated normal and reverse grading trends suggest waxing and waning of seafloor currents, an attribute often seen during deposition from hyperpycnal flows generated in shelf settings from river outflows that regularly surge and wane (Mulder et al. 2003; Wilson and Schieber, 2014). Deposition from hyperpycnal flows typically evolves from tractional to suspensional particularly as lofting occurs (Zavala et al., 2011; Steel et al., 2016), and the thicker mud laminae may record this process.

**Heterolithic, Rippled Sandstone**

This facies type shows thin bedding (1–2 cm thick), with fine scale alternation of sand, silt, and mud laminae, although sand-rich examples have few mud laminae. Ripple cross-lamination is abundant (abundantly so in sand-rich developments of this facies: Fig. 8A) and flaser and lenticular bedded strata dominate when mud laminae are more common (Figs. 6C and 6D). Ripples generally show sinuous crests of the linguoid variety and can record migration over tens of centimeters, especially in sand-rich levels (Figs. 8A and 8B). Erosion surfaces/scours with a relief ranging from a millimeter up to (locally) two centimeters are common and overlying laminae either drape broad deflation surfaces or, in the case of the rippled strata, infill the scours (<2 cm deep) (Fig. 8A). Flow directions vary considerably, with the ripples in scours often migrating at a high angle to other ripples (Fig. 6C). Soft sediment deformation, in the form of minor loading, and centimeter-scale growth faults are also present (Figs. 6D and 8B).

The heterolithic, rippled sandstone beds have some attributes that are typically attributed to tidal settings: lenticular and flaser bedding with high-angle ripple flow directions (e.g., Reineck and Wunderlich, 1968; McCave, 1970; Allen, 1984). However, the paleogeographic occurrence of the Blind Fiord Formation in a semi-isolated basin (Fig. 1) is an unlikely setting for tidal activity: a microtidal regime would be anticipated. Also, contemporaneous nearshore/coastal facies show little evidence for tidal activity (Midwinter et al., 2017). The depositional processes recorded by this facies type (fine-scale alternations of ripples formed by traction currents, local scouring, and mud deposition) bear close comparison with experimental work on rapid deposition of mixtures of sand and mud (Baas et al., 2011, 2016). The presence of even small quantities of mud in a sand flow imparts some degree of cohesion to flow with the result that ripples form in a turbulent layer at the bed boundary while an overlying plug flow de-
velops that ultimately deposits as a clay drape (Baas et al., 2016). Deposition thereby occurs as alternate laminae of cohesive and non-cohesive sediment with flow evolution often reaching the point where it can scour its own bed (Baas et al., 2011, 2016). The infill of scours often occurs with upstream facing laminae. All these features are seen in the heterolithic, rippled sandstones suggesting they were formed by decelerating mixed mud-sand flows with intrinsic depositional processes causing grain size segregation. Only recognized recently, such depositional conditions are probably common in the sedimentary record although bioturbation is likely to mask the depositional processes and produce homogenized, muddy sandstone in most cases. The Blind Fiord Formation examples are only weakly bioturbated to ichnofabric index 2 levels (Fig. 3) and burrows mostly consist of Planolites and rarer Catenichnus (Figs. 7C and 7D), probably due to prevailing dysoxic conditions on the seafloor (see below).

**Skolithos Sandstone**

The 30-m-thick, fine- to medium-grained sandstone that forms the top of the Confederation Point Member at Spath Creek shows bioturbation that ranges from intense (ichnofabric index 5) to weakly bioturbated, with a well-bedded level in its center (between 10 and 20 m in Fig. 3). Original sedimentary structures, now partly overprinted by burrows, were planar lamination and occasional ripples. The Skolithos burrows have an unusually narrow diameter for this ichnogenus (~2 mm) but can be up 20 cm in depth (Fig. 7B). They dominate the ichnofabric, and they cross-cut the horizontal burrows of Planolites and occasionally Thalassinoides. This facies also appears to have been important on the basin margin, in the Bjorne Formation, where Midwinter et al. (2017) record it as their facies association three (FA3), although the reported bioturbation intensity is less than seen in the Confederation Point Member and primary current lineation is well developed.

The abundant trace fossils suggest a fully marine, well-oxygenated environment with the dominance of vertical traces typical of nearshore deposition. This is the most proximal facies development in the Blind Fiord Formation. Midwinter et al. (2017) interpreted their FA3 as a braided stream deposit, albeit with a “weak marine influence” due to the trace fossil content. We suggest that the Skolithos sandstone in the Blind Fiord Formation formed in a marine-influenced, outer mouth bar setting where high current velocities produced primary current lineations.

**Redox Conditions**

**Pyrite Framboid Analysis**

Pyrite framboid size-frequency variations show a close correspondence with the facies of...
Microbial structures Early Triassic Arctic Canada

the Blind Fiord Formation. The lower Confederation Point Member at Griesbach Creek shows considerable variation of ichnofabric index (Fig. 3) and the lowest value (II2) occurs at a level that yields abundant pyrite frambooids that plot in the anoxic field (sample 3, Figs. 3 and 9; Table 1). Most Griesbach Creek strata are moderately burrowed (II3–II4) and the associated frambooids plot in the dysoxic field (samples 1, 2, 4, 5, and 6; Fig. 9; Table 1) indicating anoxia occurred within the sediment, while bottom waters were likely weakly oxygenated. The uppermost 15 m of the Griesbach Creek section consists of fully burrowed strata that lacks pyrite indicating well oxygenated conditions. In contrast, samples from the middle part of the Confederation Point Member, which occur in heterolithic siltstones and sandstones seen at the base of Spath Creek section (Fig. 3), are all pyrite rich and contain frambooids that plot in the anoxic to dysoxic (mostly the latter) fields (Fig. 9; Table 1). The overlying Skolithos sandstone lacks pyrite indicating oxygenated conditions once again, which is perhaps not surprising given the intensity of bioturbation. A laminated (unbioturbated) interval within the Skolithos sandstone (sample 8) contains abundant frambooids with a size-frequency distribution that plots in the dysoxic field despite the inferred shallow depositional depth (Fig. 9). The higher levels in the Blind Fiord Formation consist of heterolithic sandstones and siltstones with very little bioturbation (ichnofabric index 2) and abundant grains of pyrite including frambooids that plot in the dysoxic field (Fig. 9).

In summary, the Blind Fiord Formation was, for the most part, deposited under oxygen-poor (dysoxic) conditions although prolonged intervals of much better ventilation are recorded in the Confederation Point Member, especially in the lower Griesachian strata. At times anoxia was even developed in nearshore conditions such as the laminated sandstones of the Skolithos facies. Interestingly, the oxygenated intervals do not link closely with facies or water depth because the best ventilation is recorded in both offshore, burrowed mudstone and the nearshore, deltaic Skolithos sandstone facies.

Trace Metal Concentrations

Concentrations of Mo and U, both absolute and normalized to Al and Th, respectively, provide a useful measure of redox conditions in ancient sediments (Tribovillard et al., 2006). Mo/U variations can also be informative because Mo enrichment occurs in euxinic conditions, when H2S is present, and U uptake occurs under less intense redox conditions, at the Fe (II)–Fe (III) boundary (Wignall, 1994; Algeo and Tribovillard, 2009).

Molybdenum concentrations and Mo/Al ratios are low in the Confederation Point Member (<1 ppm and <0.4, respectively) and increase slightly in the Smith Creek and lower Svarfeld members before a major increase around 30 m above the base of the Svarfeld Member to 10 ppm and 1.8 (Fig. 4). Uranium concentrations show a similar trend to Mo whereas the Th/U ratio shows a gradual and persistent decline from ~5 at the base to <3 at the top of the Spath Creek section. The Th/U values from spectrometer field measurements and mass spectrometer assay give similar values (Fig. 4).

These results generally agree with the redox trend shown by the trace fossil and pyrite frambooids studies presented above: overall better oxygenation in the Confederation Point Member succeeded by prevailing dysoxia at higher levels with anoxia/euxinia seen in the highest studied levels of the lower Svarfeld Member.

Overview of Sverdrup Basin Environments

Depositional conditions within the Sverdrup Basin during the Early Triassic show many attributes typical of epicontinental basin fills: a “bulls-eye” style of facies belts with basal mudrocks surrounded by silt and sand-dominated facies in shallower waters passing into a sandstone-dominated fluvial hinterland (Fig. 1; Embry and Beauchamp, 2008; Midwinter et al. 2017). Variations in the lateral extent of the facies is attributed to eustatic sea-level changes (Embry, 1988). Oxygenation levels were generally very low in the basin and dysoxic conditions occasionally expanded into shallow-water, nearshore environments. Unusually, the redox conditions do not link with these changes of base level. The best oxygenated intervals are seen in the basal facies of the earliest Grieschian,
a time of rapid transgression, and the nearshore facies of the mid Dienerian. A similar style of redox fluctuations is seen in the Boreal shelf setting of Spitsbergen, Norway, where anoxic conditions repeatedly developed in very shallow waters regardless of base-level fluctuations (Wignall et al., 2016).

There is remarkably little evidence for basinal processes in the Blind Fiord Formation. Wave ripples are essentially unknown, even in the nearshore strata recorded here and by Midwinter et al. (2017). Evidence for storm deposition is also lacking; the sedimentary record consists of continuous deposition of thin laminae with no thicker “event beds.” This is despite the considerable fetch of the basin which may have reached 1000 km in extent (Fig. 1). The lack of storm facies is likely to record an unusual Early Triassic climatic regime. It is notable that the Lower Triassic strata of Spitsbergen were deposited at a similar latitude (~45°N) on an ocean-facing shelf and record only minor, rare storm events (Wignall et al., 2016).

The sediment discharge into the Sverdrup Basin was both intense and remarkably continuous. The Early Triassic succession at Spath Creek generally records accumulation in an outer shelf location and reaches nearly 950 m thickness. Taking an estimated 5 million year duration of deposition, gives an impressive sedimentation rate of 190 m/million years and indicates considerable run-off. At no point do individual bed thicknesses exceed a few centimeters indicating repeated minor discharges in a humid regime rather than a more flashy, semiarid style of run-off. The common decimeter scale alternations of siltstone-dominated and sandstone-dominated deposition (Fig. 5) also indicate longer term variations of climate. These were likely on shorter timescales than Milankovitch cycles. Given the thickness noted above, these are likely to reflect run-off variations of ~1000 years frequency.

Having established the depositional regime within the Sverdrup Basin we now present and discuss the abundant MISS occurrences found in the Blind Fiord Formation. These mostly occur on bedding surfaces of the heterolithic, rippled sandstone facies with occasional examples in the laminated, heterolithic siltstones.

Microbially Induced Sedimentary Structures

Wrinkle Structures

Bedding surfaces of the heterolithic, rippled sandstone facies can be smooth, planar, or rippled but the majority show dense, irregular, linear, and gently curving wrinkle structures up to a centimeter in width with an amplitude of a few millimeters (Figs. 10B and 10C). The wrinkle structures sometimes show a consistent trend including sets with a crescentic pattern reminiscent of backfill in trace fossils like Zoophycos (Fig. 10D); however, they lack the outer ventilation shaft, characteristic of such fodinichnial burrows. These wrinkles are not directly comparable with “elephant skin” textures, that are typically cuspatate in cross section with sharp ridges separating the cusps (Bottjer and Haggart, 2007; Gehling and Droser, 2009). The Blind Fiord Formation wrinkles show gently rounded cross sections that resemble the “mat deformation structures” illustrated by Porada and Bouougri (2007), the “large-scale irregular wrinkles” of Noffke (2000) and the “weave fabric” of Gehling and Droser (2009), a variety of their textured organic surface. The wrinkled surfaces are associated with patches of a finely dimpled texture, called “pucker” in Gehling and Droser (2009). Trace fossils are also occasionally seen: these have a much sharper definition than the wrinkles, and consist of short horizontal burrows of Planolites that cross-cut the wrinkles (Fig. 7C).
Kinneyia

*Kinneyia* is a form of MISS consisting of reticulate patterns with sub-parallel, flat-topped ridges (Hagadorn and Bottjer, 1999; Pflüger, 1999). These are occasionally found in the heterolithic, rippled sandstones of the Confederation Point Formation (Hagadorn and Bottjer, 1999; Pflüger, 1999). They occur both in high density, where bubbles are regularly in contact, or as isolated bubbles. Occasional examples appear to have collapsed/burst and show either a central dimple, producing a concentric pattern, or crater. All these features are seen in both ancient and modern microbial mats where they are caused by trapping of gas bubbles that occasionally escape and “burst” through mats (Davies et al., 2016).

**Controls of MISS Occurrences in the Early Triassic**

The abundance of MISS, associated with low intensity burrowing, in the Blind Fiord Formation is a feature more typical of the Ediacaran Period (e.g., Gehling and Droser, 2009). Phanerozoic occurrences, including microbial structures (Noffke et al., 2001), occur over a limited depth range in the offshore transition zone, and require low oxygen conditions to suppress bioturbation and mat grazing (Pflüger, 1999; Mata and Bottjer, 2009). The Sverdrup surfaces are similarly associated with intermediate water depths, where low oxygen conditions prevailed. They are not found in the intensely bioturbated, *Skolithos* sandstone shoreface facies nor in the deeper water mudstone facies, which can be burrowed or laminated. The MISS disappear at the base of the Spathian, which saw transgression and intensification of oxygen restriction, likely due to deepening below the photic zone at this level.

Thus, it appears that both redox conditions and a water depth within the photic zone control MISS occurrences in the Early Triassic. A similar discrete, depth occurrence is seen for microbial mat occurrences in the Neoproterozoic which was favored at shallow water depths (i.e., in the photic zone) in siliciclastic settings where episodic influx of sand and finer sediment favored growth and then burial of mats (Noffke et al., 2002).

Davies et al. (2016) have argued that the reported prevalence of microbial surfaces in the Early Triassic (e.g., Pruss et al., 2004; Chu et al., 2002) is an artifact of over-sampling from this interval, and that they are probably a mundane facet of Phanerozoic conditions. They supported their contention with a survey that highlighted many MISS occurrences. However, most of the post-Cambrian occurrences described in Davies et al. (2016) are from lacustrine or peritidal environments. The examples from the Early Triassic of the Sverdrup Basin are clearly atypical Phanerozoic facies because they are found in an open marine setting. The poorly oxygenated seafloor conditions suppressed burrowing organisms and ensured the preservation of the microbial mats. The mats presumably developed quickly because sedimentation (from flows of cohesive sand-mud mixtures) was frequent. Dysoxic, shelf facies are not uncommon in the Phanerozoic, but these are usually developed in mudstone facies whereas the MISS occurrences of the Blind Fiord Formation are in sandstones.
This suggests that, for MISS to develop in the Phanerozoic, an ideal taphonomic window (cf. Noffke et al., 2002) requiring both dysoxia and sandy substrates was necessary.

Recovery in the Early Triassic

The bioturbation seen in the Blind Fiord Formation has a bearing on models for the recovery of marine communities in the aftermath of the PTME. It has been argued that the Early Triassic recovery of bioturbators was slow (spanning several million years), and stepwise with a progressive increase in burrow size, depth, and tiering complexity (Twitchett, 2006; Chen et al., 2011). However, there are several examples that contradict this notion. Complex, tiered trace fossil communities were developed immediately after the PTME in shallow-water Griesbachian settings of western Canada (Beatty et al., 2008), northern Italy (Hofmann et al., 2011) and in the shelfal and offshore, basinal settings of Ellesmere and Axel Heiberg islands recorded here. While severe, the Permo-Triassic mass extinction clearly did not eliminate mobile infauna and plenty remained to burrow to a range of depths. In the Sverdrup Basin, offshore mudstone contains tiered profiles that record burrow depths of at least ~3 cm while shoreface Skolithos burrows achieved depths of 20 cm. Thus, trace fossil tiering complexity does not provide a good measure of the recovery rates. Local benthic oxygenation levels are the main control on bioturbation and the resulting ichnofabrics rather than the time elapsed since the extinction.

The link between oxygenation and bioturbation is reversed in the model of Hofmann et al. (2015). They argued that the extinction of most bioturbators at the end of the Permian resulted in weakly bioturbated sediments in the Early Triassic that consequently saw anoxic conditions develop close to the sediment surface. Thus, “phenomena that are commonly interpreted as evidence for Early Triassic seawater anoxia may have been caused by the extinction of burrowers” (Hofmann et al., 2015, p. 10). Hofmann et al. (2015) recognized that bioturbated strata are in fact present after the mass extinction, indeed they have recorded examples of burrows themselves (Hofmann et al., 2011), but this evidence is dismissed because the burrowers supposedly only belong to shallow, infaunal guilds. However, our examples show that earliest Griesbachian offshore mudstone horizons have a mixed layer with well-developed tiering. Such ichnofabrics are lost at younger levels of the Blind Fiord Formation where more intense oxygen restriction was prevalent. Thus, dysoxia, not an absence of bioturbators, inhibited bioturbation in the Early Triassic of the Sverdrup Basin.

CONCLUSION

The Early Triassic history of the Sverdrup Basin was highly unusual. The absence of storm beds indicates generally low energy conditions. Sedimentation was dominated by a near-continuous run-off of cohesive flows of sand-mud mixtures that produced distinctive heterolithic strata in mid-shelf settings that passed offshore to siltstones and ultimately mudstones. Dysoxic seafloor conditions predominated, and often extended into shallow water, but oxygenation levels varied considerably. During the earliest Griesbachian conditions were only weakly dysoxic and a tiered burrow profile developed; an unusual occurrence in the immediate aftermath of the Permo-Triassic mass extinction that contradicts claims that the extinction of burrowing organisms caused weak bioturbation at this time. The dysoxic, mid-water depths of the Sverdrup Basin saw the prolonged accumulation of heterolithic, rippled sandstone and siltstone with bedding surfaces that show abundant evidence...
for microbial mats (wrinkle structures, Kinneya, and gas bubbles). The key requirements for this MISS development were a specific set of conditions: fine sand substrates, dyssoxic bottom waters and moderate water depths within the photic zone. Such conditions were unusual in the Phanerozoic, but their widespread presence in Ellesmere Island, and more widely in the Early Triassic, is a reflection of the unusual environmental conditions of this post-extinction world and not, as has been claimed by some, due to the magnitude of the preceding extinction losses.

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