**Palaeoenvironment of Eocene prodelta in Spitsbergen recorded by the trace fossil Phycosiphon incertum**

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**Abstract**

Ichnological, sedimentological and geochemical analyses were conducted on the Eocene Frysjaodden Formation in order to interpret palaeoenvironment prodelta sediments in the Central Basin of Spitsbergen. *Phycosiphon incertum* is the exclusive ichnotaxon showing differences in size, distribution, abundance and density, and relation to laminated/bioturbated intervals. Large *P. incertum* mainly occur dispersed, isolated and randomly distributed throughout the weakly laminated/non-laminated intervals. Small *P. incertum* occur occasionally in patches of several burrows within laminated intervals or as densely packed burrows in thin horizons in laminated intervals or constituting fully bioturbated intervals that are several centimetres thick. Ichnological changes are mainly controlled by oxygenation, although the availability of benthic food cannot be discarded. Changes in oxygenation and rate of sedimentation can be correlated with the registered variations in the Bouma sequence of the distal turbiditic beds within prodeltaal shelf sediments.

The upper part of the Palaeogene in the Central Basin of Spitsbergen (Svalbard region) comprises a progradation of an extensive coalesced delta system (Fig. 1). The basal part, the Frysjaodden Formation, is essentially a prodelta succession which is overlapped by the Battfjellet Formation, representing a prograding delta front locally with coast barrier bars. Above the Battfjellet Formation, the Aspelintoppen Formation is dominated by a coastal plain to delta plain system (Nagy 2005). In this general setting, a detailed characterization of the factors conditioning the environmental context is necessary.

Ichnological analysis can be used to interpret sedimentary environments, including ecological and depositional conditions (Rodríguez-Tovar & Uchman 2004a, b, 2006, 2010; Rodríguez-Tovar, Uchman & Martín-Algarra 2009; Rodríguez-Tovar, Uchman, Martin-Algarra et al. 2009; Rodríguez-Tovar, Uchman, Alegría et al. 2011; Rodríguez-Tovar, Uchman, Orue-Etxebarria et al. 2011; Knaust & Bromley 2012; Monaco et al. 2012; Rodríguez-Tovar, Uchman, Alegret et al. 2011; Rodríguez-Tovar, Uchman, Orue-Etxebarria et al. 2011). In deep-marine siliciclastic systems, which may include slopes or deep-sea fans, physico-chemical factors such as current energy, sedimentation rate, slope instability, substrate character, oxygen and food availability influence ichnological features such as the distribution, diversity, size and abundance of trace fossils. Therefore, ichnological analysis can be used to assess these parameters (Rodríguez-Tovar et al. 2010; Hubbard et al. 2012; Uchman & Wetzel 2012; Wetzel & Uchman 2012).

Another valuable approach to decipher environmental factors, including palaeoredox conditions and palaeoproductivity, is the use of geochemical proxies. Redox conditions in the water column and within the seafloor can be interpreted using redox-sensitive trace elements like Co, Ni and Mo, while the most extensively used proxies for palaeoproductivity reconstructions are Ba/Al, Sr/Al, Ca/Al and P/Ti ratios (see Reolid et al. 2012 and references therein).

The integration of both ichnological and geochemical data, together with sedimentological information, is a very informative strategy to elucidate interpretation of ecological and depositional conditions, which is applied in this paper to a prodelta succession of the Frysjaodden Formation in the Central Basin of Spitsbergen. The investigation is focused on trace fossil characterization, together with
geochemical analyses (palaeoproductivity and redox proxies) and the analysis of sedimentary features, in order to interpret ecological and depositional conditions. Special attention is placed on distal turbiditic sequences, which are more difficult to interpret than proximal ones since some of the diagnostic features (grain size, sedimentary structures, mineralogical components) do not vary significantly.

**Geological setting and stratigraphy**

The Palaeogene of Spitsbergen’s Central Basin (Fig. 1) is a succession 2300 in thickness of mainly siliciclastic mudstones and sandstones deposited in fluvial, deltaic to delta-influenced marine shelf environments. They crop out mostly south of Isfjorden (Fig. 1). Their lithostratigraphy and depositional conditions have been discussed by, for example, Harland (1997), Dallmann et al. (1999) and Nagy (2005).

The Eocene deposits exhibit a coarsening-upward trend ranging from shales to sandstones, evidencing a transgression at the beginning of the Eocene and a subsequent regression. Three formations make up the Eocene succession: Frysjaodden, Battfjellet and Aspelintoppen (Fig. 1). They form the so-called Gilsonryggen Sequence, which commences with shoreline deposits, culminates in prodelta shelf mudstones, continues in delta front sandstones and terminates in the delta plain to alluvial deposition.

![Geological setting of the core samples from the borehole of the Store Norske Spitsbergen Grubekompani BH9/05.](image)

(a) General location of Spitsbergen. (b) Geological map of the south of Spitsbergen with drill site of core BH9/05. (c) Simplified succession of the Eocene of Spitsbergen with Frysjaodden Formation (Fm.), Battfjellet Formation and Aspelintoppen Formation and the interpreted palaeoenvironments (after Nagy et al. 2013).
plain sandstones and mudstones of the Aspelintoppen Formation.

The studied borehole is located on the eastern flank of the Palaeogene Central Basin of Spitsbergen, south of Nordenskjöld Land, at Reindalen (ca. 77° 50’ N, 16° 30’ E; Fig. 1). The core, about 552 m long, includes the Gilsonryggen Member of the Frysjaodden Formation (from 552 to 112 m; Fig. 2), consisting of prodeltal shelf sediments with distal turbidites. Four parts can be differentiated in the Gilsonryggen Member (Figs. 1, 2).

Transitional beds occur at 552–533 m. A conglomerate bed at 552 m marks the lower boundary of the Gilsonryggen Member (Frysjaodden Formation). The basal 3 m consists of siltstone and fine-grained sandstone beds alternating with dark shales. This interval shows a fining-upward trend.

Lower shales occur at 533–344 m and consist of dark grey non-bioturbated shales commonly with fine lamination. Thin layers of bentonites occur at 517 and 511 m. From 500 m upwards, chert pebbles appear occasionally and are interpreted as dropstones transported on the thalli of floating seaweed (Nagy et al. 2013), but other interpretations as being ice-rafted deposits have been also proposed (Dalland 1977; Spielhagen & Tripati 2009). Siderite-cemented beds up to 1 cm thick appear at 500 m, increasing in frequency and thickness upwards. Decimetre-thick silty beds are typical of the upper part of the interval.

Middle shales occur at 344–166 m and consist of shales and subordinate siltstones and sandstones, principally concentrated in the upper part. From 242 m, a very weak upward-coarsening is observed, as seen in the increasing number and thickness of silt layers that usually contain dense but small trace fossils (see below). The upper part of this interval is characterized by large-scale upward-coarsening and upward-finishing parasequences from silt to fine sand. Load structures are frequent, locally in combination with convolute lamination, suggesting high sedimentation rates (Dypvik et al. 2011). Ripple lamination is common and hummocky cross-stratification has been locally observed.

Uppermost sandy beds occur at 166–112 m and comprise sandstones and shales disposed in upward-coarsening sequences. Hummocky cross-bedding, wave-generated ripple lamination and load structures are common. Large trace fossils are abundant, principally in the mixed sand/shale interface. The top of the interval represents the boundary to the sandstones of the overlying Battfjellet Formation, with upward-coarsening lithology leading to cross-beded sandstones in the topmost part. This influx of coarse-grained material reflects the progradation of the Battfjellet–Aspelintoppen delta system (Dypvik et al. 2011).

Turbiditic sequences are well represented in the middle shales. Hence, the selected study core, corresponding from 226.74 to 202.30 m, pertains to this facies.

**Materials and methods**

This research is based on core samples from borehole BH9/05 of the mining company Store Norske Spitsbergen Grubekompani.
Ichnological analysis focused on trace fossil characterization and ichnofabric analysis, with special attention to relative abundance, degree of bioturbation, distribution of burrows and relationships with lithology and sedimentary structures. Detailed observations of trace fossils were made on half-cut sections of selected intervals of the core. High-resolution images were obtained and several digital image techniques were applied to enhance trace fossil visibility (Dorador et al. 2014a, b, in press; Rodríguez-Tovar & Dorador 2014). Locally, the rock surface was wetted with water and/or light-weight oil—a “modified Bushinsky oil technique” (Bromley 1981).

Whole-rock analyses of major elements were carried out on 25 sampling levels using X-ray fluorescence in a PW 1040/10 spectrometer (Philips, Eindhoven, The Netherlands). The content of trace elements was determined using a Scix-Elan 5000 inductively coupled plasma-mass spectrometer (Perkin Elmer, Waltham, MA, USA) at the Centro de Instrumentación Científica at the University of Granada. The instrumental error was ±2% and ±5% for elemental concentrations of 50 and 5 ppm, respectively.

Analysis of redox conditions in the seafloor is approached by means of redox-sensitive trace elements (such as Co, Cu, Mo and Ni) which tend to be less soluble under reducing conditions and become enriched under oxygen-depleted conditions (Wignall & Myers 1988; Calvert & Pedersen 1993; Jones & Manning 1994; Powell et al. 2003; Gallego-Torres et al. 2007; Gallego-Torres et al. 2010; Reolid et al. 2012). To compare trace element proportions in samples, it is usual to normalize element concentrations to aluminium content (Calvert & Pedersen 1993) to minimize lithological effects, assuming that Al content in sediments is contributed by aluminosilicates (e.g., Calvert 1990). The authigenic values for Mo were calculated according to Zhou et al. (2012) as $\text{Mo}_{\text{auth}} = \frac{\left[\text{Mo}\right]_{\text{sample}} - \left[\text{Mo}\right]_{\text{PAAS}}}{\left[\text{Al}\right]_{\text{PAAS}}}$.$\times\left[\text{Al}\right]_{\text{sample}}$. In addition, geochemical proxies are applied to interpret palaeo-productivity, the most extensively used being Ba/Al, Sr/Al and P/Ti ratios (e.g., Turgeon & Brumsack 2006; Gallego-Torres et al. 2007; Robertson & Filippelli 2008; Sun et al. 2008; Reolid & Martínez-Ruiz 2012; Reolid et al. 2012). Uranium and organic matter in the sediment are related in some sedimentary contexts as uranium may form a complex with dissolved fulvic acid in hemipelagic sediments (Nagao & Nakashima 1992). In this sense, the U/Al ratio can serve as a proxy for palaeo-productivity (Reolid et al. 2012).

Thin section analysis of intervals with trace fossils and intervals with parallel lamination was done using scanning electron microscopy with back-scattered electron images to determine the presence and size of pyrite frambooids. Analyses were carried out with a Carl Zeiss Merlin microscope (Standort Göttingen, Vertrieb, Germany) at the Centro de Instrumentación Científico-Técnica at the University of Jaén, Spain. Framboid size distributions have been successfully applied as anoxia and euxinia indicators in fossil marine sediments (Wignall et al. 2005; Shen et al. 2007; Bond & Wignall 2010; Liao et al. 2010).

The record of Phycosiphon incertum

Ichnological analysis in the studied section reveals the exclusive presence of Phycosiphon incertum. Trace fossils are characterized by small size, more or less cylindrical cores, and in some cases flattened, U-shaped lobes, as well as “fish-hook” shapes and pairs of black spots. Dark tubes consisting of fine-grained material are surrounded by a lighter mantle. Spreite structures were not observed, but this could be an effect of cutting. According to size, two groups of traces can be distinguished: (a) the small ones have observed tubes of a maximum length of 2–3 mm, 0.4–0.5 mm width, and marginal tubes approximately 0.2 mm wide; (b) the large forms have tubes 5–10 mm long, 0.8–1 mm wide, and marginal tubes 0.4 mm wide (Fig. 3). Small forms are dominant in the studied sections. The presented features allow to assign all the discussed trace fossils to P. incertum Fischer-Ooster.

According to the emended diagnosis by Wetzel & Bromley (1994), P. incertum refers to an extensive, small-scale spreite trace fossil consisting of repeated narrow, U-shaped lobes enclosing millimetre- to centimetre-scale spreite, branching regularly or irregularly from an axial spreite of similar width. Lobes are protrusive and mainly parallel to bedding, but the plane enclosing their width may lie horizontally, obliquely or even vertically with respect to the bedding plane. This is the widely accepted ichnotaxonomy by synonymization of Anconichnus horizontalis with P. incertum, emending the original identification of P. incertum by including non-bedding-parallel specimens, and showing Anconichnus as a junior synonym of Phycosiphon. A clear description of P. incertum is presented in Bromley (1996: 264–266). The author refers to a complexity-lobed, small-scale spreite structure, surrounded by a thin mantle of pale sediment; the core consists of backfilled dark material, and the spreite is made of the same pale material as the mantle. When mantle
and spreite are difficult to detect, as in a pale matrix, the paired sections of fill where lobes are cut transversely, or “fish-hooks” where they are cut longitudinally, permit conclusive identification. Illustrative reconstructions of the multiple phases of foraging can be found, including three-dimensional models of “phycosiphoniform” burrows (i.e., figure 11.11 in Bromley 1996; Bednarz & McIlroy 2009). See Uchman (1995, 1998, 1999) for discussion of the *Phycosiphon* group, and ichnotaxonomy at the ichnospecies level.

Detailed analysis of *P. incertum* throughout the studied section reveals several patterns according to distribution, abundance and density, usually related to the differentiated size-groups: (a) sparsely and randomly distributed larger forms consisting of disperse, isolated *P. incertum* in a weakly to non-laminated intervals, with variable orientation in respect to the bedding plane, (b) sporadic records of small forms within a laminated interval, occurring as isolated specimens or as patches of several burrows with variable orientation, (c) densely packed small *P. incertum*, mainly parallel to the bedding plane, being located as horizons in laminated intervals and (d) densely distributed small forms (mainly horizontally) occupying a fully bioturbated interval several centimetres in thickness.

**Laminated/bioturbated intervals in the studied section**

Variations in *P. incertum* features within the laminated to bioturbated intervals include: (a) fully laminated intervals, consisting of millimetre-scale laminae with scarce or even absent *P. incertum*, (b) highly bioturbated intervals, either sparsely bioturbated with larger *P. incertum*, or intensely bioturbated intervals with small-sized forms and (c) bioturbated to laminated intervals, appearing as weakly laminated intervals with densely packed horizons or as patches of small *P. incertum*.

An idealized stratigraphic pattern of the differentiated cases is as follows: (a) at the base, a laminated interval with absent or only very scarce *P. incertum*; (b) above, a coarse-grained interval with cross-bedding lamination, showing a slightly undulated (erosional?) at the base and a gradual transition with a bioturbated interval at the top; and (c) finally, a gradual transition to a laminated interval with absent or very few *P. incertum*. This idealized pattern, however, may be present in a number of variations related to the presence of bioturbated horizons in the laminated/non-bioturbated interval (see below).

**Trace fossil record and turbiditic sequences**

The Gilsonryggen Member of the Frysjaodden Formation consists of prodelta sediments with distal turbidites. In the studied section, several cases were distinguished (Figs. 4, 5, 8).

Case A is present in the transition between successive turbiditic sequences: the transition shows a small erosional undulate-shape body composed of the massive—centimetre-thick—graded Ta interval of the Bouma sequence. It cuts into a laminated mud interpreted as the Te interval of the Bouma sequence below. This massive graded interval shows comparatively coarse grains, is lighter in
Fig. 4 (a) Transition between successive turbidic Bouma sequences, showing the thick massive (centimetre-thick) graded interval (Ta) of the Bouma sequence above into the upper laminated mud interval (Te) of the Bouma sequence below, as an erosional undulate-shaped body (Case A in Fig. 8). Note scarce, larger Phycosiphon incertum in Ta (interval 3; 226.72 cm). (b), (c) Centimetre-thick (Case B in Fig. 8) and (d) millimetre-thick (Case C in Fig. 8) cross-laminated interval (Tc) of the Bouma sequence registered above the lower laminated interval (Tb), with a slightly erosional undulated surface. Note scarce record of P. incertum (yellow arrows) at interval 2; 226.32 cm, 226.25 cm and 226.14 cm, for (b), (c) and (d), respectively.
colour, consisting of discrete, scarce, larger \textit{P. incertum} (Fig. 4a). The laminated interval is characterized by the absence or scarce record of both, trace fossils and foraminifera.

Case B is observed in the middle part of a turbiditic sequence. A centimetre-thick cross-laminated interval Tc of the Bouma sequence is registered above the lower laminated interval Tb, with a slightly erosional undulated surface in between. The cross-laminated interval Tc is characterized by the absence/scare record of \textit{P. incertum} and foraminifera, as occurs in the former lower laminated interval (Fig. 4b, c).

Case C (middle part of a turbiditic sequence) is characterized by a millimetre-thick cross-laminated interval (Tc) above the lower laminated interval (Tb), with a slightly erosional undulating surface in between. This millimetre-thick cross-laminated interval also shows the absence or only scarce occurrence of \textit{P. incertum}. This case is similar to case B, but associated to a thinner cross-laminated interval. In some cases, a cross-laminated interval is not observed; the bioturbated interval (Td) therefore overlies the laminated interval (Tb).

Case D refers to the middle–upper parts of the turbiditic sequence. A gradual transition from bioturbated to

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**Fig. 5** Evolution in turbiditic Bouma sequence (Td–Te). (a) Case D in Fig. 8, showing gradual transition from a bioturbated interval with disperse, abundant, large \textit{Phycosiphon incertum} to small, concentrated \textit{Phycosiphon}, and then a laminated interval (interval 1; 222.52 cm). (b) Case E in Fig. 8, showing a laminated interval with horizons of small \textit{Phycosiphon} and finally a laminated interval (interval 1; 222.66). (c) Case F in Fig. 8, showing the transition from a bioturbated interval with disperse, abundant, large \textit{P. incertum} to a laminated interval with patches of small \textit{Phycosiphon} and finely a laminated interval (interval 2; 226.26 cm).
laminated interval (Td–Te), showing progressively decreasing size and abundance of *P. incertum* and disappearance of the (scarce) foraminifera (Fig. 5a) is observed here.

Case E (middle–upper parts of the turbiditic sequence) shows a gradual transition from a bioturbated interval with disperse, abundant, large *P. incertum*, to a thick laminated interval showing alternating horizons with small *Phycosiphon* and thin laminated horizons (Td–Te) without bioturbational structures (Fig. 5b).

Case F (middle–upper parts of the turbiditic sequence) displays a gradual transition from a bioturbated interval with disperse, abundant, large *P. incertum*, to a thick laminated interval showing patches of small *Phycosiphon* and ending in a laminated horizon without bioturbational structures (Td–Te; Fig. 5c).

**Geochemical proxies**

Geochemical analyses of the studied section show a generalized pattern for all the studied proxies—the presence of long-term trends and the absence of short-term fluctuations, together with the presence of scattered, significant peaks of increasing geochemical values. The record of *Phycosiphon* is not directly related to increases in palaeoproductivity proxies, yet the peaks of redox proxies always occur in laminated horizons.

From the studied section, interval 1 (222.66–222.30 m; 34 cm thick) was selected to disclose the observed pattern (Fig. 6). All palaeoproductivity proxies (P/Ti, Ba/Al, Sr/Al and U/Al) show maximum values and a peak coinciding with horizons poorly-bioturbated by *Phycosiphon*. All redox proxies (Co/Al, Cu/Al, Ni/Al, Mo/Al and Moaut) show no variations along the studied interval, except for an important peak, with maximum values, just above the peak of the palaeoproductivity proxies, related to the well-laminated interval directly above the poorly-bioturbated horizons.

The analyses with back-scattered electron imagery reveal further differences between the well-laminated intervals and bioturbated intervals. The well-laminated intervals are richer in clay minerals, pyrite frambooids and phytodetritus than the bioturbated intervals (Fig. 7). The size of the pyrite frambooids is lower in well-laminated intervals (3.6–5.5 μm) than in bioturbated intervals (6.4–11.5 μm).

**Interpretations and discussion**

**Ichnological and geochemical data: incidence of oxygenation and/or food availability**

*P. incertum* traces are interpreted as having been produced by deposit feeders, usually opportunistic, at different tiers in mud and fine sand at variable depths up to 15 cm.
in sediments deposited in water depths below lower shoreface (Fu 1991; Goldring et al. 1991; Wetzel & Bromley 1994; Uchman 1995, 1999; Wetzel & Uchman 2001; Hovikoski et al. 2008; Wetzel 2010). Phycosiphon is considered a pascichnial/fodinichnial structure (Ekdale & Masson 1988; Bromley 1996); the absence of spreite was interpreted as reflecting a pascichnia (grazing) rather than a fodinichnia (mining) behaviour (Ekdale & Masson 1988). Phycosiphon traces have been observed in recent muddy mass-flow deposits in deep-marine settings, but the producer has not yet been recognized (Wetzel 2008). According to Bednarz & McIlroy (2009), producers of phycosiphoniform burrows were small, probably vermi-form organisms.

Phycosiphon is frequently associated with food availability and oxygen conditions. The Phycosiphon producer reworked a relatively small volume of the sediment containing abundant nutritious material. Depth–size grading is typical of colonization; after initial colonization, the trace-maker grew in size and burrowed deeper into the sediment (Wetzel & Uchman 2001; Hovikoski et al. 2008). Orientation of Phycosiphon spreiten varies from inclined in homogeneous muds to horizontal in laminated deposits, probably reflecting the homogeneous distribution of food material (Wetzel & Uchman 2001; Wetzel 2010). Phycosiphon is common in poorly oxygenated sediments, but the absence of a connection to the seafloor indicates that immediate colonization took place in an oxygenated upper layer habitat (Ekdale & Masson 1988; Wetzel & Uchman 2001).

Integrated lithological and foraminiferal analyses of the core shed light on the general palaeoenvironmental conditions (Nagy et al. 2013). The upper part of the Gilsonryggen Member (Frysjaodden Formation) is related to the initial phases of delta progradation concomitant with the development of a highstand system tract. Salinity is a main restricting factor, though it decreases upward (Nagy et al. 2013), but short-term fluctuations may also be considered. The dominance of opportunistic organisms, such as that producing P. incertum, must be interpreted in the context of a stressed habitat resulting from salinity changes and fluctuations. However, the observed ichnological variations suggest the influence of additional environmental factors, including oxygenation and benthic food availability. The scarceness of foraminifera and the restricted occurrence of oxygen-tolerant forms
(Thurammina papillata, Psammomphaera fusca and Tre- chanmina inornata [Nagy et al. 2013]) would confirm the low oxygen availability.

The exclusiveness of P. incertum, the well-developed laminated intervals and the overall dark colour of the deposits all point to anoxic or highly dysoxic sediments. A minor increase in oxygen content to lower dysoxic conditions allowed for a localized colonization by the producer of Phycosiphon. The offset between the peak of palaeoproductivity proxies and the redox proxies may also be related to a consumption of oxygen resulting from increased productivity. Anoxic conditions are corroborated by the small size of pyrite framboinds. Pyrite framboinds only form at the redox boundary where oxygen-bearing and hydrogen sulphide-bearing waters are in contact (Wilkin et al. 1996; Reolid 2014). In oxygenated bottom water settings, pyrite framboind growth occurs in the upper sediment column, just below the sediment—water interface. However, in euxinic settings the redox boundary is within the water column. Framboinds growing in the water column have a limited diameter range, as once they have grown to a certain size, they will sink out of the narrow redox interface zone and into oxygen-free bottom water and sediment where further growth is not possible (Wilkin et al. 1996; Dustira et al. 2013). The smallest mean diameters (3–5 μm) with a very limited size range are indicative of euxinic conditions (Bond & Wignall 2010). Framboinds forming within the sediment under oxic bottom water conditions attain greater diameters on average (7.7 μm) as they reside and grow at the redox interface zone for a longer time (Wilkin et al. 1996). According to Wilkin et al. (1996) and Dustira et al. (2013), mean framboind size <7.7 μm indicates dysoxic conditions, between 5 and 7.7 μm corresponds to dysoxic—anoxic conditions and <5 μm indicates euxinic conditions. Taking into account these ranges of pyrite framboind size, the laminated intervals of study may have developed under euxinic conditions because the mean size observed was 3.6 μm. Since the mean size in the bioturbated intervals was 8.3 μm, framboinds most likely developed in the sediment pore-water there.

Turbiditic sediments and ichnological record

Within deltaic systems, Phycosiphon has been observed mainly in sediments of the delta front, the transition zone between delta front and prodelta, and in the prodelta (i.e., Bann et al. 2008; Buatois et al. 2008; Carmona et al. 2009; Buatois et al. 2011; Weiguo et al. 2011; Buatois et al. 2012), Several papers addressed the immediate colonization of event beds by Phycosiphon trace-makers (Stow & Wetzel 1990; Goldring et al. 1991; Wetzel & Balson 1992; de Gilbert & Martinell 1998). In muddy turbidites Phycosiphon producers are interpreted to be the first organisms colonizing the upper part of a turbidite bed, probably very rapidly, only a short time after deposition (Wetzel & Uchman 2001).

During delta progradation at the studied sections, turbidite deposition was extended, showing a succession of Bouma sequences. These Bouma sequences graded from massive sand deposits at the base to laminated silt and mud, with hemipelagic and pelagic material on top. Models of tiering, trace fossil distribution and ichnofabrics for turbidite/couplets have been proposed (Uchman 1999; Uchman & Wetzel 2011, 2012). Often, a sequential colonization of event layers can be found (Wetzel & Uchman 2001). Newly deposited turbiditic sediments contain relatively well-oxygenated pore water and a relatively high amount of benthic food, allowing initial colonization by small opportunistic deposit feeders, such as mining Phycosiphon producers. Later, when oxygen and food decrease at the level where mining as feeding strategy becomes inefficient, the sediment is colonized by stationary chemosymbionts, such as Chondrites producers. Finally, during the long period before deposition of a new turbidite, the slowly deposited background sediment is colonized by graphoglyptids in well-oxygenated settings (Uchman & Wetzel 2011, 2012). The studied case reveals clear differences with the model proposed by Uchman & Wetzel (2011, 2012), probably related to less favourable conditions, impeding a more diverse and better developed trace fossil assemblage.

When interpreting the turbiditic sediments in the studied section, the particular environmental conditions have to be taken into account, especially a long-term trend of decreasing salinity with short-term fluctuations, an oxygen content fluctuating between anaerobic and dysaerobic conditions, a short-term local input of benthic food and episodic sedimentation affecting all these factors. Variations in the ecological and sedimentological parameters would explain the variable record of the encountered Bouma sequences (Fig. 8) (e.g., Mulder 2011). Their variable development is interpreted according to the observed sedimentological and ichnological features associated with the transition between successive turbiditic sequences or within a single turbidite sequence (Fig. 8). We put forth six cases, as follows.

Case A: input of coarse sediment (Ta) was associated with an overall increase in oxygenation from the anaerobic laminated interval (Te) below to dysaerobic, conditions in Ta, probably related to a relatively high amount of food, allowing a disperse colonization of larger Phycosiphon trace-makers.
Case B: deposition of the cross-laminated interval was not associated with an increase in oxygen content sufficient to allow a colonization by *Phycosiphon* trace-makers; anaerobic conditions during deposition of the upper part of the previous laminated interval (Tb) continued until the base of the next bioturbated interval (Td) above the cross-laminated interval was deposited.

Case C: this case, similar to case B, is characterized by a thin cross-laminated interval that could be related to rapid deposition of the cross-laminated interval or to minor erosion. Absent or only scarce *P. incertum* can be interpreted as reflecting persistent low-oxygen conditions (as in case B) or as evidence of a short time of deposition for the thin cross-laminated interval before deposition of the next interval (Td) which impeded colonization by *Phycosiphon* trace-makers.

Case D: this case could be associated with a gradual decrease in oxygen content from dysaerobic to finally nearly anaerobic conditions, documented by the colonization by large *Phycosiphon* trace-makers, then by small *Phycosiphon* and finally lacking bioturbation.

Case E: this case reflects a gradual decrease in oxygenation from dysaerobic conditions, allowing a colonization by large *Phycosiphon* trace-makers, to nearly anaerobic conditions generally preventing burrowing activities which are, registered only during short-time improvements in oxygenation.

Case F: in this case, the oxygen content decreased from dysaerobic conditions, allowing a colonization by large *Phycosiphon* trace-makers, to nearly anaerobic sediments strongly reducing the activity of *Phycosiphon* trace-makers. However, patches of *Phycosiphon* could be associated with short-time improvements in oxygenation together with localized patches of available benthic food, probably related to minor flows that brought oxygen and food from shallower zones.

**Conclusions**

Ichnological, sedimentological and geochemical analyses of sediments from the Eocene Frysjaodden Formation allow an interpretation of palaeoenvironmental conditions during prodelta development in the Central Basin of Spitsbergen.

Ichnological analysis reveals the exclusiveness of *P. incertum* of different size showing variations in distribution, abundance and density: (a) large *P. incertum* are dispersed, isolated and randomly distributed throughout weakly/non-laminated intervals, with variable orientation with respect to the bedding; (b) small *P. incertum* occur sporadically in single, isolated, laminated intervals or in patches of variably oriented burrows; (c) densely packed small *P. incertum*, mainly horizontal to the bedding plane, are present in laminated horizons; and (d) densely distributed small *P. incertum* are found in fully bioturbated horizons which are several centimetres thick. These overall patterns reflect variable relationships of *P. incertum* and the distinguished laminated/bioturbated intervals. Variations in size, abundance and distribution of *P. incertum* can be related to oxygen contents fluctuating between anaerobic and dysaerobic conditions, short-time local inputs of benthic food and an episodic sedimentation affecting all these factors.

Several cases of distal turbiditic sequences conforming prodelta sediments could be distinguished according to the sedimentological and ichnological features found, mainly related to variable oxygenation and rates of sedimentation, determining variations in the Bouma sequence.

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