Evidence for a diachronous Late Permian marine crisis from the Canadian Arctic region

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ABSTRACT

A high-resolution chem stratigraphic study of a 24-m-thick section at West Blind Fiord on Ellesmere Island (Canadian Arctic) documents stepwise environmental deterioration in the marine Sverdrup Basin during the late Changhsingian (late Late Permian) as a result of volcanic disturbances to surrounding landmasses. A horizon within the upper Lindström Formation (datum A) is characterized by increased Fe-oxyhydroxide fluxes and weathering intensity as well as modest shifts toward more reducing water-mass conditions and higher marine productivity, recording an initial disturbance that washed soils into the marine environment. The contact between chert of the Lindström Formation and silty shale of the overlying Blind Fiord Formation, which is 1.6 m higher and ~50 k.y. younger than datum A, records a large increase in detrital sediment flux, more strongly enhanced marine productivity, and a regional extinction of siliceous sponges, therein termed the “Arctic extinction event.” The horizon equivalent to the latest Permian mass extinction of Tethyan shallow-marine sections is 5.6 m higher and ~100 k.y. younger than the Arctic extinction event, demonstrating the diachronous nature of the marine biotic and environmental crisis at a global scale; it is associated with intensified anoxia and possible changes in phytoplankton community composition in the study section. Marine environmental deterioration in the Sverdrup Basin, probably triggered by terrestrial ecosystem deterioration and elevated detrital sediment fluxes, was under way by the early part of the late Changhsingian, well before the onset of main-stage Siberian Traps flood basalt volcanism. The event sequence at West Blind Fiord may record the deleterious effects of early-stage explosive silicic eruptions that affected the Boreal region, possibly through deposition of toxic gas and ash within a restricted latitudinal band, while having little impact on marine ecosystems in the peri-equatorial Tethyan region.

INTRODUCTION

The latest Permian mass extinction was the largest mass extinction in Earth history, during which ~90% of marine and a large fraction of terrestrial taxa died out (Erwin, 1994; Retallack, 1995; Alroy et al., 2008). In shallow-marine carbonate facies from South China and elsewhere across the Tethyan region, the latest Permian mass extinction was generally an abrupt event of large scale; it is associated with intensified anoxia and possible changes in phytoplankton community composition in the study section.

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Late Permian marine crisis

at Meishan D, the global stratotype section and point (GSSP) for the Permian-Triassic boundary (Yin et al., 2001). A similar pattern of stepwise extinction was reported from the Chaotian section in northern Sichuan Province, where the main radiolarian crisis is located 3.5 m below the latest Permian mass extinction (Isozaki et al., 2007a). At both Dongpan and Chaotian, the extinction sequence was probably the result of depth-related environmental changes. As the oxygen-minimum zone intensified and expanded, taxa occupying progressively shallower habitats were adversely affected (Shen et al., 2012). These studies are significant in showing that (1) major biotic and environmental changes were under way at intermediate water depths in the Tethyan region prior to the latest Permian mass extinction defined by shallow-marine facies, and (2) pre–latest Permian mass extinctions among deeper-dwelling organisms such as radiolarians are a manifestation of a globally diachronous extinction event with a minimum duration of about 100 k.y.

Although less well documented, major environmental and biotic changes were under way prior to the latest Permian mass extinction in intermediate-depth marine settings on the northern Pangean margin as well. The marine biota in the Sverdrup Basin of the Canadian Arctic was in decline from the early part of the Late Permian, culminating in a major extinction of sponges in the early part of the late Changhsingian (Henderson and Baud, 1997; Beauchamp and Baud, 2002; Gates et al., 2004; Beauchamp et al., 2009), which we herein designate the “Arctic extinction event.” The uppermost 50 m section of the Black Stripe Formation at Buchanan Lake is characterized by a gradual ~3‰ negative shift in δ13Corg that was punctuated by three episodes of fly ash loading inferred to represent deposition of airborne combustion products of magmatic intrusions into Siberian coalfields (Grasby et al., 2011). This negative δ13Corg shift commenced during the early late Changhsingian Clarksina bachmanni zone (Korte and Kozur, 2010), and it predates the ~3‰~4‰ negative shift in marine carbonate δ18O records globally (Luo et al., 2011). Based on average sedimentation rates, Grasby et al. (2011) estimated that the onset of the δ13Corg shift at Buchanan Lake preceded the latest Permian mass extinction by ~500–750 m.y.; however, Shen et al. (2010) correlated the C. bachmanni zone to the lower C. yini zone and estimated its age as only ~200–300 k.y. older than the latest Permian mass extinction. Collectively, these studies show that major changes in marine biotas and environmental conditions were under way in the Boreal region from the early part of the late Changhsingian, well in advance of the latest Permian mass extinction, and that such changes may have been caused by early (pre–main-stage) eruptions of the Siberian Traps volcanic province (cf. Kozur and Weems, 2011).

The goals of the present study are to better understand (1) environmental changes in intermediate-depth marine facies of the Sverdrup Basin prior to the latest Permian mass extinction, (2) the timing of these changes in relation to marine extinction events in the Canadian Arctic and elsewhere globally, and (3) the causes of environmental stress and regional biotic decline prior to the latest Permian mass extinction. For this purpose, we undertook a multiproxy chemostratigraphic analysis of a 24-m-thick interval of the uppermost Permian at West Blind Fiord on southwestern Ellesmere Island. We generated chemostratigraphic profiles at a high level of stratigraphic resolution, with 56 samples from an interval representing ~500 k.y., or an average of one sample per ~10 k.y. These data were used to assess the hypothesis that environmental changes were under way in intermediate-depth marine settings at mid-northern paleolatitudes (~40°N) at least 100 k.y. prior to the latest Permian mass extinction, as defined by extinction patterns in per-equatorial (Tethyan) shallow-marine facies.

**GEOLOGIC SETTING—SVERDRUP BASIN**

**Paleogeography and Climate**

The Sverdrup Basin is a >1000-km-long, 400-km-wide, northeast-trending pericratonic successor basin underlying the northernmost islands of the Canadian Arctic region (Embry, 1991; Embry and Beauchamp, 2008; Fig. 1A). The basin developed through rifting and collapse of the Franklinian geosyncline following the Late Devonian–Mississippian Ellesmerian orogeny (Beauchamp et al., 1989a). During the Permian–Triassic, it was surrounded by landmasses to the north, east, and south but linked to the global ocean via a shallow strait on its western margin (Reid et al., 2007; Embry, 2009). This strait existed throughout the late Paleozoic except for transient episodes of closure in the early Pennsylvanian and, possibly, in the Late Permian (Beauchamp et al., 1989b). The basin achieved its present configuration when the northern landmass (termed “Crockerland” by Berg, 1994; Symons and McCausland, 2006). Climatic conditions in the Sverdrup Basin region evolved from warm and dry in the Early Permian to cool and humid in the Late Permian (Beauchamp, 1994; Beauchamp and Baud, 2002; Reid et al., 2007). Around the margin of the basin, sedimentary facies included glauconitic sandstone, siltstone, and red shale of marginal marine to nonmarine character containing a low-diversity, temperate-zone fossil assemblage dominated by sponges, bryozoans, echinoderms, and brachiopods. Concurrently, the basin center accumulated a >5-km-thick succession of dark-gray shale, cherty siltstone, and chert. The high concentration of sponge-derived chert in Upper Permian beds has been interpreted as evidence of vigorous thermohaline circulation and nutrient upwelling in the Panthalassic Ocean, and its absence in Lower Triassic beds has been interpreted as evidence of sluggish circulation and nutrient-poor surface waters (Beauchamp, 1994; Beauchamp and Desrochers, 1997; Beauchamp and Baud, 2002). Water depths in the basin center were probably on the order of a few hundred meters, a depth range favored by modern hexactinellid sponges. The water column was stratified, resulting in enhanced burial of organic carbon in deeper areas and contributing to 13C enrichment of Middle and early Late Permian carbonates in the Sverdrup Basin (Beauchamp et al., 1987) and globally (Isozaki et al., 2007b; Grossman et al., 2008).

**Basinal Sequence Stratigraphy**

Sedimentation in the center of the Sverdrup Basin was continuous from the Pennsylvanian through the Early Cretaceous, whereas thinner, unconformity-punctuated successions are present on the basin margins (Fig. 1B: Beauchamp et al., 1989b; Beauchamp et al., 2009). The Permian–Triassic is represented by a series of large-scale (third-order, or multimillion-year) transgressive-regressive (T-R) cycles, with Sequence 7 made up of a >200-m-thick upper Capitanian to lower Changhsingian succession, and Sequence 8 represented by an ~350-m-thick middle Changhsingian to Olenekian succession. The study section at West Blind Fiord (section 24 of Beauchamp et al., 2009) was deposited in a slope setting approximately midway between the basin center and southeastern basin margin (Fig. 1B). In this area, the uppermost ~100 m section of the Upper Permian has been assigned to the Black Stripe and Lindström Formations, which are composed of medium-bededded black to white spiculitic cherts passing upward into increasingly argillaceous and thinly bedded siliceous shale, representing the regressive systems tract (RST) of Sequence 7 (Gates et al., 2004; Beauchamp et al., 2009). The sequence boundary between Sequences 7 and 8 is conformable and located ~19 m below the top of the Lindström Formation at West Blind Fiord, although...
Figure 1. (A) Sverdrup Basin facies map showing the location of the West Blind Fiord (WBF) study section, and the Otto Fiord South (OFS) and Buchanan Lake (BL) sections of Grasby and Beauchamp (2008) and Grasby et al. (2011). The dashed line represents the erosional edge of the basin. The inset map shows global paleogeography at ca. 252 Ma; M—Meishan, PA—paleo-Arctic Ocean, S—Siberian Traps, SC—South China, Sv—Sverdrup Basin, T—Trøndelag, W—West Blind Fiord. (B) Basin-center to basin-margin cross section and sequence stratigraphy of Middle Permian—Lower Triassic units of southern Ellesmere Island. Abbreviations: Wuchia.—Wuchiapingian; Chang.—Changhsingian; Olenek.—Olenekian; Sequ.—sequence; DW-SB—deep-water sequence boundary; RST—regressive systems tract; TST—transgressive systems tract. Note that RST is equivalent to the “highstand systems tract” of some authors, but we retain the sequence terminology of Beauchamp et al. (1989b, 2009) for the Sverdrup Basin succession.
Permian–Triassic Biotic Changes

The marine invertebrate biota of the Sverdrup Basin underwent major changes during the Permian. Basin-margin sediments of Early Permian age (Sakmarian and younger) are dominantly cool-water carbonate ramp deposits composed of bryozoans, echinoderms, brachiopods, foraminifers, and solitary rugose corals with a minor fraction of sponge spicules (Beauchamp, 1989; Beauchamp and Desrochers, 1997; Beauchamp and Baud, 2002; Reid et al., 2007). Replacement of this prolific carbonate factory by a siliceous sponge-dominated biota occurred between the Artinskian (late Early Permian) and Guadalupian (late Middle Permian) as a consequence of climatic cooling and an increase in nutrient levels (Beauchamp and Henderson, 1994; Reid et al., 2007; Bensing et al., 2008).

The lower beds of the Upper Permian Lindström and Black Stripe Formations contain rare macrofaunal remains, but productid and spiriferid articulate brachiopods, trepostome and fenestrate bryozoans, and echinoderms all disappear near the base of the Mesogondolella rosenkrantszi–Mesogondolella sheni zone, around the lowest Changhsingian (Beauchamp, 1994; Henderson and Baud, 1997). Thereafter, fossil remains consist almost exclusively of siliceous sponge spicules (Gates et al., 2004). The spicules are 10–150 μm wide and 100–2000 μm long and include monactine monaxons and tetraxons, generally associated with the classes Demospongia and Hyalospongia, respectively (Rigby, 1987). The spicules were derived from sponges lacking a fused skeleton, and the resulting state of disaggregation makes generic identification impossible. The top of the Mesogondolella sheni zone, dating to the mid–late Changhsingian, marks the point where demosponges, hyalosponges, and some soft-bodied taxa (as noted by trace fossils) went extinct, culminating the progressive reduction in biodiversity that characterized the Middle to Late Permian of the Sverdrup Basin (Beauchamp, 1994). Since both sponge classes range from the Cambrian to the Holocene, their demise in the Sverdrup Basin during the Arctic extinction event represents either extinction at a lower taxonomic level or local rather than global taxonomic disappearances. Grasby and Beauchamp (2008, 2009) have correlated this extinction (Arctic extinction event) with the global mass extinction (latest Permian mass extinction), but this correlation is inaccurate because the succeeding Clarkina hauschkei zone represents a pre–latest Permian mass extinction level (see Biostratigraphic control section).

The basal 10–20 m of the Blind Fiord Formation (of latest Permian age) contain no biota except for microbes, rare ammodiscid foraminifers, and conodonts (Beauchamp and Baud, 2002) and exhibit very limited ichnofossil abundance (Beatty et al., 2008). The Griesbachian (earliest Triassic) is characterized by a limited macrofauna including several species of the ammonite genera Otoceras, Ophiceras, and Bukkenites and the bivalve Claraia (Henderson and Baud, 1997; Baud et al., 2008). Greater bioturbation is apparent from ~60 m above the base of the Blind Fiord Formation (Beatty et al., 2008), and a more diverse benthic macrofaunal assemblage that includes bryozoan biostromes is found in the Dienerian-Smithian, ~200–300 m above the Permian-Triassic boundary (Baud et al., 2008). However, sponges did not reappear in the Sverdrup Basin until the Middle Triassic.

STUDY SECTION

History of Investigation

The 24-m-thick study section at West Blind Fiord consists of the uppermost 6 m of the Lindström Formation and the lowermost 18 m of the Blind Fiord Formation (Fig. 2A). This section...
was visited by Benoit Beauchamp and Charles Henderson first during the 1985 field season, when conodont samples were collected from the base of the Blind Fiord Formation (Fig. 2B). Beauchamp and Henderson revisited this section in 1989, when Henderson collected a larger suite of samples over a wider stratigraphic interval. The results of this biostratigraphic work were first reported at the Pangea meeting in Calgary (Henderson, 1993), when it was suggested that events in the Permian-Triassic boundary interval of the Canadian Arctic were diachronous with respect to the Tethyan Permian-Triassic boundary record. Subsequently, numerous additional sections were collected at sites on several Arctic islands by Beauchamp, Henderson, and co-workers, culminating in a summary paper by Beauchamp et al. (2009). Grasby and Beauchamp (2008, 2009) investigated the C-isotope stratigraphy and other geochemical signatures of a number of Arctic Permian-Triassic boundary sections, but these studies utilized the biostratigraphic framework of Beauchamp and Baud (2002) and did not fully integrate new biostratigraphic information developed by Henderson and reported in the present study (see next section).

Biostratigraphic Control

Research on conodont biostratigraphy of the Permian-Triassic boundary has advanced substantially in the past few years owing to studies in China (Jiang et al., 2007; Zhang et al., 2007; Chen et al., 2009) and Iran (Kozur, 2005; Shen and Mei, 2010). As a result of these studies, biostratigraphic age assignments and correlations of Tethyan and boreal conodont faunas now can be undertaken with greater confidence. A long-lived boreal Mesogondolella lineage, culminating in the Sverdrup Basin with M. rosenkrantzi and M. sheni, was replaced during the late Changhsingian age. This zone includes Clarkina cf. changxingensis, Clarksina hauschkei, Clarksina cf. jolensis, and very rare Hindeodus typicalis. Clarkina cf. changxingensis (Figs. 3E, 3F, 3H, and 3I) represents a new subspecies of C. changxingensis similar to C. yini (Kozur, 2005); the latter was also described originally as a subspecies of C. changxingensis from Beds 23–24e at Meishan D (Yin et al., 2001). Our specimens of C. hauschkei (Figs. 3M and 3N) have been compared to those specimens described by Kozur (2005) as Clarkina hauschkei borealis from a zone below both the Permian-Triassic boundary clay and the microblaste facies at Abadeh, Iran. Clarkina cf. jolensis (Figs. 3J and 3K) has the relatively large, posteriorly directed cusp of Clarkina meishanensis, but it has a fused carina that is more comparable to C. jolensis from Iran. Clarkina jolensis is associated with C. yini and C. hauschkei in Iran (Kozur, 2005). These three species all exhibit the flattening of the posterior platform, reduction in the width and depth of the adcarinal furrows, and a mostly small, posteriorly directed cusp that is typical of prelatest Permian mass extinction conodonts of the C. yini zone of South China (Jiang et al., 2007; Zhang et al., 2007; Chen et al., 2009). At West Blind Fiord, this zone is located below the base of the present study section (Fig. 4A).

Mesogondolella sheni Zone

Specimens of M. sheni (Figs. 3C and 3D) from the TST at the base of Sequence 8 (upper Lindström Formation) indicate a Changhsingian age, but the exact relationship between these specimens and the type material from the Selong, Tibet, which has been assigned to the Selong, Tibet, when it was suggested that events in the Permian-Triassic boundary interval of the Canadian Arctic were diachronous with respect to the Tethyan Permian-Triassic boundary record. Subsequently, numerous additional sections were collected at sites on several Arctic islands by Beauchamp, Henderson, and co-workers, culminating in a summary paper by Beauchamp et al. (2009). Grasby and Beauchamp (2008, 2009) investigated the C-isotope stratigraphy and other geochemical signatures of a number of Arctic Permian-Triassic boundary sections, but these studies utilized the biostratigraphic framework of Beauchamp and Baud (2002) and did not fully integrate new biostratigraphic information developed by Henderson and reported in the present study (see next section).

Clarkina hauschkei Zone

This zone includes Clarkina cf. changxingensis, Clarksina hauschkei, Clarksina cf. jolensis, and very rare Hindeodus typicalis. Clarkina cf. changxingensis (Figs. 3E, 3F, 3H, and 3I) represents a new subspecies of C. changxingensis similar to C. yini (Kozur, 2005); the latter was also described originally as a subspecies of C. changxingensis from Beds 23–24e at Meishan D (Yin et al., 2001). Our specimens of C. hauschkei (Figs. 3M and 3N) have been compared to those specimens described by Kozur (2005) as Clarkina hauschkei borealis from a zone below both the Permian-Triassic boundary clay and the microblaste facies at Abadeh, Iran. Clarkina cf. jolensis (Figs. 3J and 3K) has the relatively large, posteriorly directed cusp of Clarkina meishanensis, but it has a fused carina that is more comparable to C. jolensis from Iran. Clarkina jolensis is associated with C. yini and C. hauschkei in Iran (Kozur, 2005). These three species all exhibit the flattening of the posterior platform, reduction in the width and depth of the adcarinal furrows, and a mostly small, posteriorly directed cusp that is typical of prelatest Permian mass extinction conodonts of the C. yini zone of South China (Jiang et al., 2007; Zhang et al., 2007; Chen et al., 2009). At West Blind Fiord, this zone is located by the interval from datum B (5.6 m) to datum C (9.0 m; Fig. 4A). Based on the age model of Age model and sedimentation rates, this zone has a duration of ~70 ± 10 k.y. (Fig. 4B).

Clarkina meishanensis Zone

Although C. meishanensis (Figs. 3G, 3K, and 3L) is restricted to Beds 25–28 at Meishan (Jiang et al., 2007), a similar species (C. zhangi) ranges from Bed 23 to Bed 24e. C. zhangi was originally described as a subspecies of C. meishanensis (Mei et al., 1998), and it is possible that these taxa represent the same species. Clarkina meishanensis migrated into Iran during the latest Changhsingian, correlative with Beds 23 and 24 at Meishan (Kozur, 2005), but its occurrence in various sections globally seems to be diachronous. This zone is interpreted to correlate with the C. yini and C. meishanensis zones of latest Changhsingian age, equivalent to Beds 24–27b at Meishan D (Henderson and Baud, 1997; cf. Chen et al., 2009). Associated species, at least in the lower part of the zone at Otto Fiord South (Fig. 4A), include C. hauschkei and C. cf. changxingensis, indicating an overlap with taxa characteristic of the C. hauschkei zone, a pattern also noted by Kozur (2005) from Iranian sections. Furthermore, Henderson and Baud (1997) reported the occurrence of the ammonoid Otoceras concavum within this zone, which has been accepted as a Late Permian species by Yin et al. (1996). At West Blind Fiord, this zone is represented by the interval from datum C (9.0 m) to datum E (= Permian-Triassic boundary, ~18 m; Fig. 4A). Based on the age model (see age model and sedimentation rates), this zone has a duration of ~140 ± 10 k.y. (Fig. 4B).

Clarkina taylorae–Hindeodus parvus Zone

Clarkina taylorae (Figs. 3Q and 3R) appears above the maximum flooding surface at Otto Fiord South closely associated with the ammonoid Otoceras boreale. Very rare specimens of H. parvus (Fig. 3O) appear a few meters higher in the same section (Henderson and Baud, 1997). The first occurrence of C. taylorae at its type locality in Selong, Tibet (Orchard et al., 1994), coincides with that of H. parvus, the first appearance datum (FAD) of which defines the Permian-Triassic boundary in the Meishan GSSP. Jiang et al. (2007) suggested that C. taylorae ranged below the Permian-Triassic boundary into Bed 27a at Meishan, but we interpret their figured specimens from Beds 27a and 27b as C. zhejiangensis and comparable to specimens referred to C. praeutaylorae from Iran (Kozur, 2004; Chen et al., 2009). Clarkina taylorae represents a good index for the Permian-Triassic boundary when species of Hindeodus are rare; it has an upright cusp.
and better developed adcarinal furrows that are typical of most immediately post–latest Permian mass extinction species of *Clarkina*. The stratigraphic position of the Permian-Triassic boundary was narrowly constrained at Otto Fiord South by the first appearances of *C. taylorae* and *H. parvus*, but this datum represents a local first occurrence (FO) of *H. parvus* that is younger than its FAD at Meishan. The correlative horizon at West Blind Fiord is ~13 m above the base of the Blind Fiord Formation. At West Blind Fiord, this zone approximately spans the uppermost 6 m of the study section (~18–24 m), between datums E and F (Fig. 4A). Based on the age model (see age model and sedimentation rates), this zone has a duration of ~70 ± 10 k.y. (Fig. 4B).

**Clarkina taylorae–C. carinata Zone**

Clarkina carinata (*Neogondolella* of some authors; for discussion, see Henderson and Mei, 2007) is the most commonly identified Lower Triassic conodont species (Clark, 1959; Sweet, 1970; Orchard and Krystyn, 1998). A sample from higher in the Blind Fiord Formation (~35 m above its base) yielded *C. taylorae*, *Clarkina tulongensis* (Fig. 3S), and *C. carinata* (Figs. 3T, 3U, 3V, and 3W), which are characteristic of the early Griesbachian (lowermost Triassic), equivalent to Beds 27c–30 at Meishan D (Jiang et al., 2007; Zhang et al., 2007; cf. Chen et al., 2009). Although conodont data are available only for these few horizons at West Blind Fiord, an age-equivalent section at Otto Fiord South on northern Ellesmere Island was the subject of a more detailed conodont study by Henderson and Baud (1997), and the two sections can be correlated accurately on the basis of C-isotope profiles (Grasby and Beauchamp, 2008) and sequence stratigraphic relationships (Beauchamp et al., 2009). At West Blind Fiord, this zone begins approximately at the top of the 24-m-thick study

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**Figure 3.** Scanning electron microscope (SEM) photomicrographs of conodonts from West Blind Fiord (WBF) and Otto Fiord South (OFS) sections. (A–B) *Mesogondolella rosenkrantzi*, WBF C-125990 (22 m below formation contact): (A) upper and (B) oblique upper view of same specimen. (C–D) *Mesogondolella sheni*, WBF C-125990 (22 m below formation contact), upper view; and (D) WBF C-125991 (21 m below formation contact), upper view. (E–F, H–I) *Clarkina cf. changxingensis*, WBF C-125970 (0.2 m above formation contact): (E) oblique upper and (G) upper view of same specimen; (H–I) WBF C-125969 (0.1 m above formation contact), upper views. (G, K–L) *Clarkina meishanensis*, OFS (31.5 m above sequence boundary): (G) upper view; WBF C-125569 (50 m above formation contact), (K) upper and (L) oblique upper view of same specimen. (J, P) *Clarkina cf. jolfensis*, J) WBF C-125969 (0.1 m above formation contact), upper view; and (P) WBF C-125970 (0.2 m above formation contact), upper view. (M–N) *Clarkina hauschkei*, WBF C-125969 (0.1 m above formation contact), upper views. (O) *Hindeodus parvus*, OFS (31.5 m above sequence boundary), lateral view. (Q–R) *Clarkina taylorae*, OFS (31.5 m above sequence boundary), upper views. (S) *Clarkina tulongensis*, OFS (31.5 m above sequence boundary), upper view. (T–W) *Clarkina carinata*, WBF C-125569 (50 m above formation contact): (T) upper view; (U) OFS (31.5 m above sequence boundary), upper view; (V–W) WBF C-125569 (50 m above formation contact), (V) oblique upper and (W) upper view of same specimen.
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Figure 4. (A) Correlation of West Blind Fiord to Meishan D and Otto Fiord South based on biostratigraphic and δ¹³C₉₈ constraints. Datum B represents the Arctic extinction event (AEE), datum D is the latest Permian mass extinction (LPME), and datum E the Permian-Triassic boundary (PTB); the positions of datum A at Meishan and datums E and F at West Blind Fiord are approximate. For Meishan D, conodont zonation is from Jiang et al. (2007) and Zhang et al. (2007); δ¹³C₉₈ profile is from Cao et al. (2002); age model is based on Shen et al. (2010, 2011); note stratigraphic condensation in proximity to the Permian-Triassic boundary. For Otto Fiord South, conodont data are from Henderson and Baud (1997); correlations to West Blind Fiord were refined on the basis of C-isotope profiles in Grasby and Beauchamp (2008) and the sequence stratigraphic model of Beauchamp et al. (2009); note that the 10-cm-thick unit below the sequence boundary represents the feather edge of the Lindström Formation, which thins proximally. Abbreviations: SSTs—sequence systems tracts; MFS—maximum flooding surface; RST—regressive systems tract; SB—sequence boundary; TST—transgressive systems tract. For West Blind Fiord, δ¹³C₉₈ data are original to this study, except for uppermost two δ¹³C₉₈ points (circled; from Grasby and Beauchamp, 2008); note change in vertical scale at level of sequence boundary represented by the two 5 m scale bars. For both Canadian Arctic sections, productive samples and samples barren of conodonts are shown by solid and open dots, respectively, to right of lithologic columns; lowercase letters on or adjacent to biostratigraphic range bars refer to conodont specimens in Figure 3; note that Clarkina hauschkei and C. cf. changxingensis are equivalent, respectively, to C. cf. subcarinata and C. aff. changxingensis of Henderson and Baud (1997), and that C. cf. changxingensis is a new subspecies of changxingensis (C.M. Henderson). Abbreviations: Ch.—Changhsingian; chg.—changxingensis; gond.—gondolellids; hind.—hindeodids; mei.—meishanensis; parv.—parvus; tay.—taylorae. Recovery of ammonoids shown to right of lithologic columns: O-v—Otoceras concavum; O-b—Otoceras boreale; O-c—Ophiceras commune. (B) Sedimentation rate model for West Blind Fiord (WBF) based on an age model for the Meishan global stratotype section and point (GSSP; Shen et al., 2010) and correlation datums A–F of main figure; average sedimentation rates are shown at right.
section and includes stratigraphically younger beds analyzed by Grasby and Beauchamp (2008) and Beauchamp et al. (2009).

**Carbon Isotope Correlations**

Organic C-isotope records have utility for chemostratigraphic correlation, especially when biostratigraphically constrained (e.g., Gröcke, et al., 1999; Heimhofer et al., 2003; Weissert et al., 2008). Organic C-isotope profiles assisted in refining the correlations between West Blind Fiord and Meishan D already developed on the basis of conodont biostratigraphy (see Biostratigraphic control section). The two sections show similar patterns of variation in the precrisis interval of the Upper Permian. A gradual trend toward more ¹³C-enriched values is observed in the upper *C. changxingensis* zone at Meishan, peaking in the upper Bed 22–lower Bed 23 interval (Cao et al., 2002), while a correlative shift is observed in the upper *M. sheni* zone at West Blind Fiord, peaking at the Lindström–Blind Fiord formation contact (Fig. 4A). Above the formation contact (= datum B), a sharp reversal initiated a trend toward more ¹³C-depleted values that culminated at datum C within the *C. yini* zone (Bed 24) at Meishan and the *C. hauzechkei* or lower *C. meishanensis* zone at West Blind Fiord. Between datums C and D, both sections show a gradual trend toward more ¹³C-enriched values, a trend defined by eight samples over 0.3 m at Meishan and by six samples over 2.2 m at West Blind Fiord (and, thus, not a single-sample anomaly). At the level of datum D (= latest Permian mass extinction), both sections are characterized by a large (~4‰–5‰) and abrupt negative shift in δ¹³Corg (Fig. 4A). The correlative character of the two organic C-isotope profiles breaks down above datum D, with Meishan showing relatively ¹³C-enriched, but highly variable values and West Blind Fiord showing relatively ¹³C-depleted and stable values.

The C-isotopic composition of organic matter is subject to global as well as local influences, and both may be apparent in the West Blind Fiord δ¹³Corg profile (Fig. 5A). The pre–latest Permian mass extinction patterns exhibited by the sections at West Blind Fiord and Meishan are similar to those in other Permian-Triassic boundary sections globally, e.g., the Buchanan Lake section from the Sverdrup Basin (Grasby and Beauchamp, 2008; Fig. 5B), and the Trøndelag core from Norway (Hermann et al., 2010; Age Zones West Blind Fiord Buchanan Lake Beds Zones Meishan (Corg) Meishan (Ccarb) Trøndelag

<table>
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<th>Age</th>
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<th>Buchanan Lake</th>
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<td>22</td>
<td><em>C. changxingensis</em></td>
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Figure 5. Comparison of organic C isotopic profiles for (A) West Blind Fiord (this study), (B) Buchanan Lake (Grasby and Beauchamp, 2008), (C) Meishan global stratotype section and point (GSSP; Cao et al., 2002; carbonate δ¹³C profile also shown), and (D) Trøndelag core, Norway (Hermann et al., 2010). The Meishan profiles were plotted on a linear depth scale, and the other records were adjusted vertically within biostratigraphic constraints. The Arctic extinction event (AEE) coincides with the onset of a gradual negative shift at the base of Bed 23 in the Meishan Corg record, whereas the global latest Permian mass extinction (LPME) is located at a sharp negative-positive couplet at the base of Bed 25. The youngest interval of heavy fly ash deposition at Buchanan Lake (Grasby et al., 2011, their fig. 4) is indicated in B.
Fig. 5D). All four sections show a trend toward more $^{13}$C-depleted values beginning around the M. sheni/C. hauschkei or C. changxingensis/C. yini zonal boundary (= datum B), followed by a reversal toward more $^{13}$C-enriched values a little below the latest Permian mass extinction at datum C. These similarities suggest a common response at a global scale to carbon-cycle forcings, e.g., increased photosynthetic fractionation by eukaryotic marine plankton (as recorded by increasing $\Delta ^{13}$C$_{carb}$ at Meishan; Fig. 5C) in response to rising atmospheric CO$_2$ levels triggered by early-stage Siberian Traps volcanism. Above the latest Permian mass extinction (= datum D), the Tromsø core exhibits a large and sustained shift toward more $^{13}$C-enriched values similar to that at Meishan, whereas the Buchanan Lake section mirrors the lighter and more stable compositions shown by West Blind Fiord (Fig. 5). These divergent patterns may reflect regional differences in the degree to which marine primary production following the latest Permian mass extinction was taken over by prokaryotic green sulfur bacteria (Chlorobiaceae), which fractionate C isotopes during photosynthesis to a substantially smaller degree than eukaryotic algae and, hence, generate more $^{13}$C-enriched organic matter (Riccardi et al., 2007). The inference of low abundances of green sulfur bacteria at West Blind Fiord based on light $\delta ^{13}$C$_{org}$ values is confirmed by low concentrations of Chlorobiaceae-derived biomarkers (see Organic fraction section). Conversely, high abundances of green sulfur bacteria at Meishan are indicated both by heavy $\delta ^{13}$C$_{org}$ values and by high concentrations of relevant biomarkers (Grice et al., 2005; Cao et al., 2009).

Marine carbonate C-isotope profiles of a specific age are often similar owing to the long residence time of dissolved inorganic carbon in seawater (~80 k.y.), which exerts a strong global influence. Carbonate $\delta ^{13}$C profiles for Permian-Triassic boundary sections can be readily correlated globally (e.g., Algeo et al., 2007; Tong et al., 2007; Luo et al., 2011). The Meishan $\delta ^{13}$C$_{carb}$ profile is typical of these records, showing minor variation in the range of $+3\%e$ to $+4\%e$ through most of the Changhsingian, followed by a gradual $-3\%e$ to $-4\%e$ shift beginning in the mid-C. yini zone and culminating around the latest Permian mass extinction (Fig. 5C). Carbonate $\delta ^{13}$C$_{carb}$ records exhibit important differences in detail from organic $\delta ^{13}$C records, as shown by the paired profiles for Meishan (Cao et al., 2002). The onset of the negative shift in $\delta ^{13}$C$_{carb}$ (= base of Bed 24 at Meishan) postdates that of the negative shift in $\delta ^{13}$C$_{org}$ (= lower Bed 23 at Meishan) by an interval of ~100 k.y. (Fig. 4A).

Further, the positive shift in the $\delta ^{13}$C$_{org}$ profiles between datums C and D is not matched in $\delta ^{13}$C$_{carb}$ profiles, in which the negative trend begun in the mid-C. yini zone continues to the level of the latest Permian mass extinction (Fig. 5C). Owing to such differences, correlations based on matching of $\delta ^{13}$C$_{carb}$ and $\delta ^{13}$C$_{org}$ profiles can result in erroneous inferences concerning temporal relationships. For example, Grasby and Beauchamp (2008, 2009) and Grasby et al. (2011) equated a negative shift in the $\delta ^{13}$C$_{org}$ profiles of West Blind Fiord and Buchanan Lake (datum D) with the negative excursional in carbonate C-isotopes at the base of Bed 25 at Meishan. On this basis, Grasby and Beauchamp (2008) conflated the regional significant isotope excursions event represented by datum B (Arctic extinction event) with the global mass extinction event (latest Permian mass extinction) at Meishan and other Tethyan sections (datum D; Fig. 5). As discussed in Permian-Triassic biotic changes, the Arctic extinction event and latest Permian mass extinction are separate events, and it is important to recognize them as such because this relationship reveals the globally diachronous character of the Late Permian biotic crisis.

Placement of Permian-Triassic Boundary, Latest Permian Mass Extinction, and Maximum Flooding Surface Horizons

The Permian-Triassic boundary was originally placed at the base of the Blind Fiord Formation (Nassichuk et al., 1973; Thorsteinsson, 1974), but recent definition of the Permian-Triassic boundary based on the FAD of the conodont Hindeodus parvus in the Meishan D GSSP (Yin et al., 2001) dictates a higher placement. The lowermost few meters of the Blind Fiord Formation have yielded the conodonts C. hauschkei, C. cf. changxingensis, and C. cf. jof Jensis on the basis of which an uppermost Permian age has been inferred (Beauchamp et al., 2009). H. parvus was recovered 31.75 m above the base of the Blind Fiord Formation south of the head of Otto Fiord (Henderson and Baud, 1997), associated with the Early Triassic ammonoid Otoceras boreale and just below specimens of Ophioceras commune, although this horizon probably represents a local first occurrence of H. parvus rather than a true FAD. The stratigraphic correlation framework developed in the study (Fig. 4A) suggests that the Permian-Triassic boundary is ~13 m above the base of the Blind Fiord Formation at West Blind Fiord, with an uncertainty of about ±5 m.

The end-Permian mass extinction became the “latest Permian mass extinction” upon redefinition of the system boundary. In their analysis of the West Blind Fiord section, Grasby and Beauchamp (2008) placed the latest Permian mass extinction at the contact between the Lindström and Blind Fiord Formations (their fig. 4). However, we interpret the formation contact to represent a regional extinction event affecting the sponge biota of the Sverdrup Basin, herein termed the “Arctic extinction event” (see Introduction section). As shown by the correlation framework developed in Figures 4 and 5, the Arctic extinction event is ~5.6 m below the latest Permian mass extinction at West Blind Fiord and, hence, ~100 k.y. older than the latest Permian mass extinction as defined at Meishan D and elsewhere in the Tethyan region. At West Blind Fiord, the latest Permian mass extinction is located at ~10.2 m within a uniform interval of dark-gray shale lacking any macrofossils or trace fossils. This placement is supported by both biostratigraphic data and C-isotopic correlations (Figs. 4A and 5).

A maximum flooding surface represents the surface of maximum water depth in a given sequence and, thus, the transition from a transgressive systems tract (TST) to a highstand or regressive systems tract (RST) (Allen and Allen, 2005; Catuneanu et al., 2009). The maximum flooding surface of Sequence 8 of Beauchamp et al. (1989b) was assumed to be correlative with the Lindström–Blind Fiord Formation contact (FC) by Grasby and Beauchamp (2008, 2009). However, several features, including thinner bedding, increased total organic carbon (TOC) concentrations, and maxima in pyrite abundance and degree of pyritization values (see Organic fraction, and S-Fe system sections), suggest that the maximum flooding surface should be placed ~5 m above the formation contact at West Blind Fiord (Fig. 4A). Thus, the TST of Sequence 8 of Beauchamp et al. (1989b) extends through the lower ~5 m of the Blind Fiord Formation. The transition from the TST to the RST of Sequence 8 may have been due to an increasing influx of terrigenous detrital material resulting from elevated subaerial weathering rates during the Permian-Triassic boundary crisis interval (see Detrital fluxes and weathering rate changes section), rather than from a fall in global sea-level elevations. In fact, eustatic elevations probably continued to rise into the Early Triassic due to climatic warming (Hallam and Wignall, 1999), so the RST of Sequence 8 at West Blind Fiord is likely to represent the point in time at which rates of sediment influx into the Sverdrup Basin outpaced rates of eustatic deepening.

Age Model and Sedimentation Rates

We constructed an age model for the West Blind Fiord section (Fig. 4B) based on the correlation framework developed previously herein (see Biostratigraphic control, and Carbon iso-
Late Permian marine crisis

Detrital Siliciclastic Fraction

In total, 56 samples from the 24-m-thick West Blind Fiord section were analyzed in the present study (see Methods supplement for details of preparatory and analytical procedures\(^1\)).

The formation contact at 5.6 m is marked by abrupt shifts in major-element concentrations associated with the detrital fraction but relatively smaller changes in trace elements.

The upper part of the Lindström Formation (0–5.6 m at West Blind Fiord) contains abundant SiO\(_2\) (80%–90%) but only limited Al\(_2\)O\(_3\) (2%–4%) and K (<0.5%; Figs. 6A–6C). The silica is present mainly as chert derived from siliceous sponges and, hence, is of biogenic origin (Beauchamp and Desrochers, 1997; Beauchamp and Baud, 2002). Al is present mainly in the clay fraction, in which the dominant mineral is chlorite (Fig. 7A).

The overlying Blind Fiord Formation (5.6–24 m at West Blind Fiord) contains smaller quantities of SiO\(_2\) (50%–60%) but larger amounts of Al\(_2\)O\(_3\) (15%–18%) and K (3%–4%; Figs. 6A–6C). The latter elements reside mainly in the clay fraction, whereas silica is present both in the clay fraction and as quartz silt. Major minerals in the clay fraction include illite, illite-smectite, and chlorite (Figs. 7B–7D). Trace metals (Figs. 6D–6F) exhibit concentrations that are mostly close to those of post-Archean average shale (PAAS; Taylor and McLennan, 1995), suggesting residence primarily in the detrital fraction. Although V is somewhat enriched relative to PAAS in the Blind Fiord Formation, strong covariance with Al and K indicates residence mainly in the detrital fraction. In contrast, Mo shows local enrichments, especially between datum A (4.0 m) and the formation contact (5.6 m), which may represent a redox control (see Paleoredox conditions and environmental changes section). Other components of the sediment, such as calcite, organic matter, and pyrite, are present in only small quantities (<5%).

The chemical index of alteration (CIA) is a commonly used proxy for the degree of weathering of siliciclastic sediments, with higher values signifying more intense weathering (Nesbitt and Young, 1982; Price and Velbel, 2003). This ratio generally reflects the extent of conversion of feldspars to clay minerals, especially catation-poor clays such as kaolinite (Maynard, 1992; Sutton and Maynard, 1993). In the Lindström Formation, CIA values decrease from 0.65 to 0.70 at the base of the section to 0.50–0.55 just below datum A at 4.0 m (Fig. 6G). CIA values then rise sharply to >0.80 and remain generally high to the formation contact at 5.6 m. CIA values fall abruptly to ~0.72 at the base of the Blind Fiord Formation and then decline gradually to ~0.62 toward the top of the section but without a measurable shift at the latest Permian mass extinction horizon.

Cross plots of high field strength elements (e.g., Nb/Y vs. Zr/Ti) were discriminated among magma types (Winchester and Floyd, 1977; Pearce, 1996), but they have also been employed to assess the provenance of siliciclastic sediments (Umazano et al., 2009). At West Blind Fiord, the lower part of the Lindström Formation (0–4.0 m) exhibits the lowest Nb/Y and the highest Zr/Ti ratios, whereas the upper Lindström Formation (4.0–5.6 m) exhibits a shift toward higher Nb/Y and lower Zr/Ti ratios (Fig. 8). All Lindström Formation samples plot within the field of rhodacite to dacite magmas, suggesting derivation from source rocks of intermediate to slightly felsic composition, a pattern typical of coarser siliciclastics derived from old cratons (cf. Fralick, 2003). Samples of the Blind Fiord Formation plot mostly within the field of PAAS, indicating a mixed provenance, as is typical of shales. This graph reveals a systematic up-section change in the composition of the detrital fraction beginning at datum A (4.0 m) in the upper Lindström Formation—a change that is not apparent in the major- and trace-element profiles of Figure 6.

Major-element concentration data can be used to estimate the concentrations and fluxes of the major mineral fractions, including clay minerals, detrital quartz, biogenic chert, and calcite (see Methods supplement [see footnote 1]). The upper part of the Lindström Formation (0–5.6 m at West Blind Fiord) consists of 15%–30% clay minerals and detrital quartz, 70%–85% biogenic chert, and <2% calcite (Fig. 9A). The mineral-fraction composition of samples changed abruptly at the formation contact, and the overlying Blind Fiord Formation (5.6–24 m at West Blind Fiord) generally consists of ~80% clay minerals, 16%–18% detrital quartz, 2%–4% calcite, and 0% biogenic chert.

A few samples (at 7 m, 14 m, and 22–23 m) contain larger amounts of calcite (to 9%) and measurable concentrations of “biogenic” silica (Fig. 9A); however, these samples were identified as siltstones in the field and are likely to have a higher siliciclastic to clay ratio than that used in estimating detrital quartz concentrations (see Methods supplement [see footnote 1]); hence, all of the quartz/chert in these samples is presumed to be of detrital origin. Mineral-fraction fluxes also changed abruptly at the formation contact (Fig. 9B). The flux of biogenic chert decreased from ~5 g cm\(^{-2}\) k.y.\(^{-1}\) in the Lindström Formation to ~0 g cm\(^{-2}\) k.y.\(^{-1}\) in the Blind Fiord Formation, while the flux of clay minerals and detrital quartz increased from <0.5 g cm\(^{-2}\) k.y.\(^{-1}\).
to >10 g cm\(^{-2}\) k.y.\(^{-1}\). Thus, the bulk sediment flux of the Lindström Formation (~6 g cm\(^{-2}\) k.y.\(^{-1}\)) was dominated by biogenic silica, whereas that of the Blind Fiord Formation (~12–21 g cm\(^{-2}\) k.y.\(^{-1}\)) was dominated by detrital components (Fig. 9B). Bulk sediment flux is directly proportional to linear sedimentation rate, changes in which occur at stratigraphic levels corresponding to age tie points in our correlation framework (Fig. 4B). Thus, the stepwise increases in bulk sediment flux shown in Figure 9B may be a procedural artifact. In view of the relatively continuous increase in sedimentation rates at West Blind Fiord during the latest Permian (Fig. 4B), it is reasonable to infer that bulk sediment flux accelerated at the formation contact and thereafter increased steadily during deposition of the Blind Fiord Formation (heavy dashed line, Fig. 9B).

**Organic Fraction**

TOC values are low (0.2%–0.5%) throughout the study section but show a distinct pattern of stratigraphic variation (Fig. 10A). TOC increases sharply at datum A, peaking at 0.5%, and then falls toward the latest Permian mass extinction interval. Above the latest Permian mass extinction (10.2 m), TOC remains relatively elevated (0.4%–0.5%) to near the top of the study section. Biomarkers confirm the presence of thermally mature marine organic matter in all samples, as shown by C\(_{21}/C_{29}\) hopane ratios of 0.26–0.53 and C\(_{21}/C_{21}\) hopane/C\(_{29}\) hopane ratios of 0.15–0.30, 0.25–0.60, and 0.65–1.05, respectively (cf. Peters et al., 2004). However, these compounds do not provide any information regarding the presence or absence of highly weathered (and thus relatively inert) terrestrial organic matter. A petrographic study revealed considerable amounts of organic matter of both marine and terrestrial provenance in two samples close to the formation contact (Fig. 11). Clumps of marine organic matter tend to be larger than terrestrial fragments of vitrinite and inertinite, so the former probably dominates volumetrically. The absence of a relationship between clay content (Fig. 9A) and TOC (Fig. 10A) is noteworthy, indicating that grain-size variation did not serve as a dominant control on organic-matter accumulation (cf. Kennedy et al., 2002). More generally, changes in the C-N-S proxies at West Blind Fiord are largely independent of lithologic changes (which are confined mainly to the formation contact at 5.6 m), indicating that the former provide a record of biogeochemical influences in the West Blind Fiord depositional system rather than of lithologic controls.
Late Permian marine crisis

Organic carbon $\delta^{13}C$ values are between $-26\%e$ and $-27\%e$ below datum A (0–4.0 m), but they show a slight shift toward more $^{13}C$-enriched values ($-25\%e$ to $-26\%e$) between datum A and the formation contact (Fig. 10B). The $\delta^{13}C_{org}$ profile shifts abruptly toward more negative values at the formation contact (5.6 m) and then exhibits a gradual negative trend (from $-27.5\%e$ to $-29.5\%e$) as the latest Permian mass extinction horizon is approached. At the latest Permian mass extinction horizon (10.2 m), the profile shows an abrupt excursion toward a heavier composition ($-25.5\%e$), followed by an equally rapid shift toward lighter values ($-30\%e$ to $-32\%e$), which are sustained to the top of the section. N isotopes are mostly +4.0‰ to +5.0‰ below datum A (0–4 m), but they shift toward heavier values (+4.8‰ to +5.7‰) between datum A and the formation contact (Fig. 10C). Just below the formation contact (5.6 m), the N isotope profile shows an abrupt excursion toward a lighter composition (+2.3‰), followed by a shift toward heavier values (+5.5‰ to +6.5‰) between the formation contact and the latest Permian mass extinction horizon. Above the latest Permian mass extinction (10.2 m), the N isotope profile exhibits a shift back toward lighter values (+4.5‰ to +5.2‰), which remain relatively uniform to the top of the section.

The concentration profiles of all biomarkers measured in the West Blind Fiord section fluctuate over relatively narrow ranges. Aryl isoprenoids (AI) are present in all samples analyzed, although at relatively low concentrations (<12 ng/g TOC; Fig. 10D). AIs increase gradually upward throughout the 24-m-thick section but show a local maximum in the 2 m interval just below the latest Permian mass extinction. The ratio of 28,30-disthenoane (DNH) to C30 hopane (28,30-DNH ratio) ranges from 0.15 to 0.65, with higher values above the formation contact (Fig. 10E). The C27/C29 sterane ratio shows a similar pattern, ranging between 0.6 and 1.2, with higher values above the formation contact (Fig. 10F). Both the 28,30-DNH and C27/C29 sterane ratios show small, transient decreases at the latest Permian mass extinction horizon (10.2 m). The 2-methylhopane index (2-MeHI), i.e., the ratio of 2-methylhopane to (2-methylhopane + hopane), shows generally higher values below the formation contact (mostly 2%–5%) and lower values above the formation contact (mostly <2%; Fig. 10G), but the range of values is muted. The homohopane index (HHI), i.e., the ratio of C35 to C34 homohopanes (Peters et al., 2004), also shows little variation (0.5–0.9), except for a peak (2.7) at the latest Permian mass extinction horizon (Fig. 10H). The Ts/(Ts + Tm) profile, which is the ratio of two...
Permian mass extinction; PTB—Permian-Triassic boundary.

4B. Abbreviations: AEE—Arctic extinction event; FC—formation contact; LPME—latest Permian mass extinction.

Values in both panels are cumulative. Mineral fractions were calculated per Equations 2–6 of Methods supplement (see text footnote 1). Bulk (total) sediment flux is given by the rightmost solid curve in B; the stepped character of this curve is a function of assuming constant linear sedimentation rates between age tie points in Figure 9. Stratigraphic trends in (A) mineral fractions and (B) mineral-fraction fluxes. Figure 9. Stratigraphic trends in (A) mineral fractions and (B) mineral-fraction fluxes. Values in both panels are cumulative. Mineral fractions were calculated per Equations 2–6 and fluxes per Equations 7–8 of Methods supplement (see text footnote 1). Bulk (total) sediment flux is given by the rightmost solid curve in B; the stepped character of this curve is a function of assuming constant linear sedimentation rates between age tie points in Figure 4B. Abbreviations: AEE—Arctic extinction event; FC—formation contact; LPME—latest Permian mass extinction; PTB—Permian-Triassic boundary.

$C_{27}$ hopanes, shows a limited range (0.64–0.67) through most of the study section but modestly lower values below datum A (Fig. 10J). The $T_s/(T_s + T_m)$ ratio commonly varies as a function of organic source, sediment lithology, and thermal maturity (Moldowan et al., 1986); relatively uniform values in the West Blind Fiord section may reflect the similar maturity of all samples. The indigenous character of the biomarkers in the host rock is confirmed by parallel changes in multiple biomarkers in concert with other geochemical proxies, as well as by the observation that multiple parameters for source rock depositional facies change in conjunction with the lithologic transition from chert to shale at the formation contact. The generally low abundances of AI and other biomarkers, as well as the carbon-number distributions in the samples, are consistent with the intermediate level of thermal maturity of the study area (see Basinal sequence stratigraphy section).

Cross plots of TOC versus TN (Fig. 12A) and $\delta^{13}C_{org}$ versus $\delta^{15}N$ (Fig. 12B) reveal marked changes in the composition of the organic fraction with stratigraphic position. In both cross plots, sample groups for the Lindström (0–5.6 m), lower Blind Fiord (5.6–10.2 m), and upper Blind Fiord formations (10.2–24 m) show little overlap, and the differences in sample population means are highly significant ($p(\alpha) < 0.001$). Lindström Formation samples yield $C_{27}/N$ ratios of ~20–30, $\delta^{13}C_{org}$ values of −25.5%e to −27.5%e, and $\delta^{15}N$ values of +4.0%e to +5.8%e. Blind Fiord Formation samples yield uniformly lower $C_{27}/N$ ratios (mostly 2–5) and almost uniformly lower $\delta^{13}C_{org}$ values (mostly −27.0%e to −29.5%e). Although lower and upper Blind Fiord samples exhibit similar $C_{27}/N$ ratios (Fig. 12A), these two groups are readily distinguished on the basis of $\delta^{13}C_{org}$ and $\delta^{15}N$ values (Fig. 12B). Upper Blind Fiord samples yield more $^{13}$C-depleted (−30.0%e to −31.7%e) and $^{15}$N-depleted (+4.5%e to +5.3%e) compositions relative to the $\delta^{13}C_{org}$ (−27%e to −30%e) and $\delta^{15}N$ (+5.1%e to +6.4%) compositions of lower Blind Fiord samples.

**S-Fe System**

Total sulfur (S) concentrations in the Lindström Formation are uniformly low (<0.2%) below datum A (4.0 m) but rise to somewhat higher values (0.5%–0.6%) between datum A and the formation contact (4.0–5.6 m; Fig. 13A). Total S concentrations in the Blind Fiord Formation are uniformly low (<0.2%) between the formation contact and the latest Permian mass extinction horizon (5.6–10.2 m) but rise to substantially higher values (mostly >1.0%) above the latest Permian mass extinction horizon (10.2 m). More than 95% of total S is pyrite S (data not shown), and most of this S (especially in the high-S intervals) is “excess S” (Fig. 13A), i.e., S that exceeds the amount of reduced S expected relative to TOC for oxic-suboxic marine facies (Berner and Raiswell, 1983; Leventhal, 1983). Petrographic study showed that pyrite was present both as irregular masses and frambooids, and that pyrite was associated almost exclusively with organic clumps of probable marine origin (Fig. 11). In contrast, organic macerals of undoubted terrestrial provenance show virtually no pyrite overgrowths.

A TOC-S cross plot reveals that Lindström Formation samples from below datum A (0–4.0 m) and Blind Fiord samples from between the formation contact and latest Permian mass extinction horizon (5.6–10.2 m) cluster along the oxic-suboxic marine trend of Berner and Raiswell (1983; Fig. 14). Lindström Formation samples from between datum A and the formation contact (4.0–5.6 m) define a vector toward higher S concentrations relative to TOC (trend 1), and Blind Fiord Formation samples from above the latest Permian mass extinction (10.2–24 m) define a second vector toward higher S concentrations relative to TOC (trend 2). These trends reflect increasing
Figure 10. Chemostratigraphy (organic fraction): (A) total organic carbon (TOC), (B) $\delta^{13}$C$_{org}$, (C) $\delta^{15}$N, (D) total AI (aryl isoprenoids), (E) 28,30 dinorhopane (DNH) ratio, (F) C$_{27}$/C$_{29}$ sterane ratio, (G) 2-MeHI (methylhopane index), (H) HHI (homohopane index), and (I) Ts/(Ts + Tm) ratio. In H, values >0.5 (dashed vertical line) are indicative of anoxic conditions (Peters et al., 2004). The positions of samples WBF-303 and WBF-306 used in petrographic analysis (Fig. 11) are shown to right of lithologic column. Abbreviations: AEE—Arctic extinction event; FC—formation contact; LPME—latest Permian mass extinction; PTB—Permian-Triassic boundary.
amounts of excess S (see previous) that may be related to the accumulation of syngenetic pyrite in the sediment. The observation that excess S increases with higher TOC may have significant implications for the relationship between paleoredox conditions and marine productivity (see Paleoredox conditions and environmental changes, and Plankton community and changes in primary productivity sections).

Degree of pyritization is a commonly used paleoredox proxy that represents the ratio of pyrite Fe to total reactive Fe (i.e., pyrite Fe plus extractable Fe that is potentially pyritizable; Raiswell et al., 1988). Degree of pyritization values are low (<0.2) in the lower part of the study section but increase abruptly at the latest Permian mass extinction horizon (10.2 m) to 0.40–0.45 (Fig. 13B). Degree of pyritization values fluctuate above the latest Permian mass extinction but episodically reach values >0.2 through the remainder of the section. Total Fe concentrations, ranging from 2% to 8%, yield a profile quite similar to magnetic susceptibility: low values below datum A, a transient rise between datum A and the formation contact, a sustained rise above the formation contact and peaking at ~7.5 m (except for one outlier higher
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Figure 12. (A) TOC (total organic carbon) versus TN (total nitrogen) and (B) δ¹³C-org versus δ¹⁵N. Crosses enclosed by ovals show the mean ± the standard error of the mean for each of four sample groups.

in the section), followed by a gradual decrease through the remainder of the section (Fig. 13C). Reactive Fe concentrations are nearly equivalent to total Fe in the Lindström Formation but decline to 50%–70% of total Fe values in the Blind Fiord Formation. Pyrite Fe covaries positively with total Fe but comprises no more than 25% of total Fe in the pyrite-rich intervals and <10% of total Fe in the pyrite-poor intervals.

The pyrite δ³⁴S profile, while varying over an ~40‰ range through much of the study section, exhibits marked changes in character in conjunction with event horizons (Fig. 13D). In the Lindström Formation, samples below datum A (4.0 m) show a systematic increase in δ³⁴S up section from ~34‰ to ~1‰. Above datum A, δ³⁴S values become highly variable, fluctuating between extremes of ~40‰ and 0‰ from sample to sample. In the Blind Fiord Formation, a marked reduction in δ³⁴S variance is observed at the formation contact (5.6 m), such that δ³⁴S values are limited to the range of ~40‰ to ~28‰ in the 2 m of section just above the formation contact. Variance increases up section toward the latest Permian mass extinction horizon (10.2 m) but decreases again above the latest Permian mass extinction horizon, where most samples yield δ³⁴S values between ~40‰ and ~30‰ (Fig. 13D).

The magnetic susceptibility signal in marine sediments is generally dominated by ferrimagnetic Fe-oxyhydroxides and paramagnetic clay minerals such as illite, i.e., minerals of detrital origin, rather than authigenic origin (Ellwood et al., 2000, 2007). Magnetic susceptibility values in the Lindström Formation are about average for marine sedimentary rocks (<0.5 × 10⁻⁷ m³ kg⁻¹) below datum A (4.0 m), but they exhibit a transient positive excursion (to ~1.0 × 10⁻⁷ m³ kg⁻¹) between datum A and the formation contact (Fig. 13E). Magnetic susceptibility values in the Blind Fiord Formation rise sharply above the formation contact (5.6 m) to a maximum of ~3 × 10⁻⁷ m³ kg⁻¹ at 7.5 m and then decline gradually upward through the remainder of the section. There is little or no change in the magnetic susceptibility profile at the level of the latest Permian mass extinction horizon (10.2 m). Relationships between magnetic susceptibility and other proxies can assist in identifying the main carriers of the magnetic susceptibility signal. Magnetic susceptibility correlates most strongly with non-pyrite Fe (r = +0.89; p[α] < 0.001; n = 56; Fig. 15A) and more weakly with clay minerals (r = +0.61; Fig. 15B) and other sediment fractions. This observation suggests that the magnetic susceptibility signal is carried primarily by detrital ferrimagnetic phases such as fine-grained Fe-oxyhydroxides, although clay minerals (mainly illite) may make a secondary contribution to the magnetic susceptibility signal (cf. Ellwood et al., 2007). Magnetic susceptibility also exhibits positive covariation with P concentrations (r = +0.60), possibly because of absorption of P onto Fe-oxyhydroxides (Föllmi, 1996).

DISCUSSION

Paleoredox Conditions and Environmental Changes

Paleoredox conditions in the West Blind Fiord study section can be inferred from a combination of petrographic, chemostratigraphic, and biomarker proxies. Although not entirely consistent, these proxies document a general change from less-reducing to more-reducing conditions upward within the section. Differences in redox interpretations among proxies may reflect differences in the time scales at which each proxy imparts its characteristic signature to the sediment. Such differences can result when redox conditions in the depositional system vary at time scales shorter than the depositional time represented by individual samples, which is ~1 k.y. for the Lindström Formation and ~0.1 k.y. for the Blind Fiord Formation (for 1- to 2-cm-thick samples; see Age model and sedimentation rates section; cf. Bond and Wignall, 2010). For example, if a largely oxic facies experiences short episodes of anoxia, it may not accumulate sufficient amounts of pyrite and redox-sensitive trace metals to document these events yet is likely to yield biomarkers with an anoxic signature, because even low concentrations of compounds such as aryl isoprenoids are taken as evidence of photic-zone euxinia (Summons and Powell, 1987; Grice et al., 2005; Cao et al., 2009). A similar phenomenon in which short intervals of oxygenation occur within an
otherwise anoxic facies can result in juxtaposition of bioturbation with geochemical signatures of anoxia (Kenig et al., 2004).

The lower part of the Lindström Formation (0–4 m) was deposited under largely oxic conditions. This inference is based on low degree of pyritization values (<0.1; Fig. 13B) and U and Mo concentrations close to PAAS (Figs. 6E–6F). In addition, uniformly low S concentrations (<0.2%; Fig. 13A) and relatively 34S-enriched pyrite S-isotopic values (Fig. 13D) provide no evidence for the presence of syngenetic pyrite (i.e., pyrite formed from H2S-bearing waters) and, thus, of water-column euxinia. The upper part of the Lindström Formation (4.0–5.6 m) was deposited under conditions that were mainly suboxic, but ranged to weakly euxinic (i.e., marked by low H2S concentrations). This inference is based on the observations that degree of pyritization values (<0.2; Fig. 13B) and U concentrations (mostly <4 ppm; Fig. 6E) remain low, but that Mo concentrations rise sharply above background detrital levels to ~8 ppm (Fig. 6F). S concentrations show a modest increase (to ~0.5%) associated mainly with excess S (Fig. 14) that may be present in the form of syngenetic framboidal pyrite (cf. Leventhal, 1983), an inference supported by a shift toward somewhat more 34S-depleted pyrite sulfur (Fig. 13D; Lyons, 1997; Wilkin and Arthur, 2001). Finally, δ15N values shift toward more 15N-enriched compositions (Fig. 10C), consistent with enhanced denitrification under suboxic conditions (Jenkyns, 2010).

The lower part of the Blind Fiord Formation (5.6–10.2 m) was deposited under redox conditions similar to or somewhat more oxidizing than those of the underlying upper Lindström Formation. This interval exhibits low values for (1) DOP (<0.2; Fig. 13B), (2) redox-sensitive trace metals such as U and Mo (Figs. 6E–6F), and (3) excess S concentrations (Fig. 13A). However, pyrite sulfur compositions continue to show the 34S-depleted values characteristic of syngenetic pyrite (Fig. 13D), and relatively high δ15N values (Fig. 10C) are consistent with denitrification under suboxic conditions. Beginning at the latest Permian mass extinction horizon (10.2 m), redox proxies imply that the Blind Fiord Formation was deposited under more strongly reducing conditions. Degree of
pyritization shows a sharp increase at the latest Permian mass extinction and generally higher values to the section top (Fig. 13B). Although the actual values (mostly 0.2–0.5) are not representative of euxinic conditions (cf. Raiswell et al., 1988; Lyons and Berner, 1992), we infer that euxinia was a transient phenomenon at West Blind Fiord that failed to yield the reactive Fe enrichments that typify persistently euxinic basins (Lyons and Severmann, 2006). It should be noted that elevated sedimentation rates, as developed at West Blind Fiord above the formation contact (see Age model and sedimentation rates section), can lead to lower-than-expected degree of pyritization values in euxinic facies, e.g., values of ~0.4 along the modern Black Sea margin (Lyons and Severmann, 2006). S increases locally to >2% above the latest Permian mass extinction horizon (Fig. 13A) and consists mainly of excess S that cannot be accounted for through degradation of sedimentary organic matter and, hence, is likely to represent syntogenic pyrite (Fig. 14). Values of δs4S pyrite become more consistently negative (~30‰ to ~40‰, with a single outlier; Fig. 13D), supporting the hypothesis that the pyrite fraction is composed largely of frambooids formed in the water column rather than authigenic pyrite formed in the sediment (cf. Wilkin and Arthur, 2001; Nielsen and Shen, 2004; Algeo et al., 2008). Pyrite frambooids were observed petrographically at a similar stratigraphic level at Buchanan Lake, a section located further toward the center of the Sverdrup Basin (Grasby and Beauchamp, 2009). These observations are consistent with an increase in the frequency and/or duration of water-column euxinia, although the relatively low degree of pyritization values and absence of U and Mo enrichment (Figs. 6E–6F) may indicate that euxinia continued to develop sporadically in a water mass of generally oxic or suboxic character.

Biomarker data provide some additional information about redox conditions in the West Blind Fiord section, although subject to certain caveats. Aryl isoprenoids (AI) with a 2,3α,6-methyl substitution pattern are derived from the diagenetic breakdown of the carotenoid isorenieratene and chlorobactene, part of the light-harvesting complex of green sulfur bacteria (Chlorobiaceae), which are photosynthetic microorganisms that require hydrogen sulfide as an electron donor and, thus, are obligate anaerobes found only in strongly reducing diagenetic environments, isorenieratene is reduced to isorenieratene. Increasing thermal maturation leads to progressive rupture of the isoprenoid chain, yielding a homologous series of monoaromatic compounds between C₂₅ and C₃₀ (Summons and Powell, 1987). AIs were identified in at least trace quantities in all samples from the West Blind Fiord section (Fig. 10D), but the quantities are smaller relative to TOC (<12 ng/g) compared to other Permian-Triassic boundary sections (Grice et al., 2005; Cao et al., 2009) and do not show any systematic stratigraphic trend. The AI profile is consistent with sporadic but transient development of euxinic conditions at West Blind Fiord throughout the study interval.

Of the biomarker proxies measured in the saturated hydrocarbon fraction, the ratio of 28,30-dinorhopane to C₂₅ hopane (28,30-DNH ratio) and the homohopane index (HII) have been used as indicators for water-column anoxia (Curiale and Odermatt, 1989; Peters et al., 2004). In the West Blind Fiord samples, the 28,30-DNH ratio increases from the Lindström Formation to the Blind Fiord Formation, but it is generally high in all the samples measured (Fig. 10E). Homohopanes, a series of C₂₅–C₃₀ hopane hydrocarbons, are formed as breakdown products of bacteriohopanepolyols (Ourisson et al., 1979; Rohmer et al., 1984). The ratio of the C₂₅/C₂₃ homohopanes (or HII) records the preferential preservation of the C₂₅ precursor hydrocarbon. Values >0.5 can be interpreted as evidence of anoxic conditions (Peters et al., 2004), although this proxy is also subject to a maturity influence. The homohopane index shows only modest variation through the study section, with values between 0.5 and 1.0, with the exception of a single outlier (2.7) just above the latest Permian mass extinction horizon at 10.2 m (Fig. 10H). Given the likely uniform thermal maturity of all samples in the study, these values suggest that reducing conditions prevailed throughout deposition of the West Blind Fiord section, with moderate intensification of anoxia in conjunction with the latest Permian mass extinction. The biomarker data are consistent with the hypothesis that photic-zone euxinia developed intermittently and for intervals of short duration at West Blind Fiord, although with an increase in the frequency and/or duration of such episodes following the global mass extinction event.

Plankton Community and Changes in Primary Productivity

Fossil and molecular evidence of major changes in the composition of phytoplankton communities can be observed in some stratigraphic sections following the latest Permian
Evaluating changes in primary productivity in paleomarine systems is difficult, but qualitative inferences may be possible for the West Blind Fiord study section. $C_{org}/N$ ratios of 20–30:1 in the Lindström Formation (Fig. 12A) may record comparatively high concentrations of terrestrial organic matter (Meyers, 1994), an inference supported by petrographic observations of higher-plant–derived organic macerals in samples close to the formation contact (Fig. 11). $C_{org}/N$ ratios are mostly ≤5:1 in the Blind Fiord Formation (Fig. 12A), suggesting dominance of marine algal over terrestrial organic matter (Meyers, 1994). Although control of the shift in $C_{org}/N$ ratios by changes in organic matter preservation related to more reducing conditions higher in the section cannot be excluded, several factors argue against such a mechanism: (1) The largest shift in $C_{org}/N$ ratios is located at the formation contact (5.6 m), not at the latest Permian mass extinction (10.2 m), where the most pronounced intensification of reducing conditions occurred (see Paleoredox conditions and environmental changes section), and (2) the interval of the Blind Fiord Formation deposited under more oxidizing conditions (5.6–10.2 m) shows lower $C_{org}/N$ ratios than the interval deposited under more reducing conditions (10.2–24 m; Fig. 12A). Thus, it seems likely that a large increase in the flux of marine algal material to the sediment occurred at the formation contact, probably in response to an increase in marine primary productivity.

**Detrital Fluxes and Weathering Rate Changes**

Major changes in subaerial weathering processes may have played an important role in the Permian-Triassic marine biotic crisis (Algeo et al., 2011). Globally, marine sections exhibit an average approximately sevenfold increase in sediment flux across the latest Permian mass extinction horizon (Algeo and Twitchett, 2010). Within the latest Permian interval of the West Blind Fiord study section, sedimentation rates increased by a factor of 3.5x, from 25 to 88 m m.y.⁻¹ (Fig. 4B), and even higher sediment fluxes characterize the Lower Triassic of the northern Pangean margin (Algeo and Twitchett, 2010, their fig. 2). However, even before the major increase in sediment flux that occurred at the formation contact (Fig. 9B), the composition of the detrital fraction at West Blind Fiord had begun to change: High field strength elements show a gradual shift from a rhodacitic composition in the Lindström Formation to an average upper-crustal composition (PAAS) in the Blind Fiord Formation (Fig. 8). Although this pattern could be interpreted in terms of changing sediment

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**Figure 15.** (A) Magnetic susceptibility (MS) versus nonpyrite Fe and (B) magnetic susceptibility versus clay minerals. In A, nonpyrite Fe was calculated as $F_{\text{total}} - F_{\text{pyrite}}$ (see Fig. 13C). Magnetic susceptibility exhibits strong positive covariation ($r = +0.61$; $p \ll 0.001$; $n = 56$) with nonpyrite-Fe concentration (Fig. 13C) but weaker covariation ($r = +0.61$) with clay-mineral abundance (Fig. 9A). These relationships suggest that the magnetic susceptibility signal is carried mainly by Fe-oxyhydroxides and secondarily by clay minerals such as illite.

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mass extinction. In general, eukaryotic marine algae declined (Knoll et al., 2007; Chen et al., 2011), although prasinophyte and acritarch “disaster taxa” appear to have thrived (Payne and van de Schootbrugge, 2007). Photosynthetic prokaryotes, namely, green sulfur and N-fixing cyanobacteria, also thrived (Grice et al., 2005; Xie et al., 2005, 2007; Hays et al., 2007; Cao et al., 2009). Biomarker data from the West Blind Fiord study section suggest that changes in the phytoplankton community of the Sverdrup Basin during the latest Permian may have been relatively minor. First, aryl isoprenoids indicate the presence of green sulfur bacteria throughout the study interval, with a consistently high abundance just below the latest Permian mass extinction horizon (Fig. 10D). The short-term shifts toward higher δ¹³C organ values at 7.5 m and 10.2 m (Fig. 10B) might be due to transient increases in biomass coming from Chlorobiaceae owing to the relatively ¹³C-enriched composition of these bacteria (Riccardi et al., 2007). Second, the 2-MeHI profile documents a limited and relatively uniform presence of diazotrophic (N-fixing) cyanobacteria throughout the study section (Fig. 10G). Concentration increases at 1.0 m, 5.0 m, and 10.2 m suggest transient increases in cyanobacterial productivity (Summons et al., 1999; Xie et al., 2007). Each peak coincides with a shift toward lower δ¹⁵N values (Fig. 10C), consistent with enhanced N fixation by diazotrophic cyanobacteria (Jenkyns et al., 2001; Kuypers et al., 2004). Third, shifts in the C₃₇/C₃₉ sterane ratio (Fig. 10F) may reflect variability in the types of photosynthetic eukaryotes, specifically, red versus green algae, contributing organic matter to the sediments (Volkman et al., 1998; Peters et al., 2004). If correct, the increase in C₃₇/C₃₉ sterane ratios at the formation contact at West Blind Fiord suggests enhanced proportions of red algae, which have a lower requirement for iron, making them better adapted to euxinic conditions than green algae. However, the ratios for this proxy are all close to 1.0, indicating that any changes in the proportions of the two groups of algae were modest at most. It can be observed that trends in the C₃₇/C₃₉ sterane profile are opposite those in the 2-MeHI profile (Fig. 10G), suggesting that algal communities varied in tandem with abundances of diazotrophic cyanobacteria.
provenance (Dickinson, 1985), we infer that it reflects a gradual increase in subaerial weathering intensity. Secular changes in weathering intensity are supported by the CIA profile, which increases from 0.52–0.70 at 0–4.0 m to 0.80–0.85 at 4.0–5.6 m (Fig. 6G). The large increase in CIA values just below the formation contact (5.6 m) may represent a short-term erosional event, e.g., a rapid stripping of highly weathered soils from adjacent landmasses (cf. Sephton et al., 2005). The subsequent decline in CIA across the formation contact to intermediate values (~0.68–0.73) may reflect influx of less fully weathered material following the loss of terrestrial soils (cf. Algeo and Twitchett, 2010). This interpretation is supported by the magnetic susceptibility profile (Fig. 13E): The transient increase at 4.0–5.6 m is consistent with a major influx of soil-derived material (especially from the Fe-enriched B horizon), while the subsequent sustained increase above the formation contact (5.6 m) reflects an elevated flux of Fe-bearing clay minerals produced through intensified chemical weathering. Intensified soil erosion and chemical weathering around the latest Permian mass extinction have been inferred in earlier studies of both terrestrial (Retallack, 2005; Sheldon, 2006) and marine sections (Sephton et al., 2005; Xie et al., 2007; Wang and Visscher, 2007).

**Relationship to Early Eruptions of Siberian Traps?**

The Siberian Traps flood basalt eruptions are thought to have caused major global environmental changes. Release of a combination of volcanic CO₂ and volanogetic methane produced when magma intruded Siberian coal basins triggered strong climatic warming (Retallack, 1999), as shown by oxygen isotope studies (Korte et al., 2005a, 2005b; Kearsey et al., 2009). Higher levels of atmospheric CO₂ along with volcanic SO₂ emissions are likely to have increased the acidity of precipitation (Wignall, 2001, 2007), which, in combination with warmer temperatures, resulted in elevated rates of chemical weathering (Retallack, 1999; Sheldon, 2006). These changes may have been important factors in the latest Permian destruction of terrestrial ecosystems, when gymnosperm-dominated forests representing a mature successional stage yielded to rapidly growing lycopsid-fern communities representing an early successional stage (Wang, 1996; Looy et al., 2001; Grauvogel-Stamm and Ash, 2005; Hochuli et al., 2010; Hermann et al., 2011). The destruction of terrestrial vegetation had profound consequences for weathering processes, nutrient fluxes, and marine environmental conditions. A combination of climate warming and increased nutrient fluxes (eutrophication) resulted in greatly expanded oceanic anoxia (Winguth and Maier-Reimer, 2005; Meyer et al., 2008; Algeo et al., 2010). Shallow-marine ecosystems were harmed by episodic expansion of the oxygen-minimum zone into the ocean-surface layer as well as by elevated particulate fluxes (siltation) in paralic environments. High rates of terrestrial erosion persisted widely for up to ~2 m.y. during the Early Triassic and were potentially a factor in the delayed recovery of marine biotas (Algeo and Twitchett, 2010; Algeo et al., 2011), although stable refugia are known to have existed (Beaty et al., 2008).

We propose that changes in the marine environment and biota of the Sverdrup Basin during the Late Permian were driven by perturbations to adjacent land areas in response to Siberian Traps volcanism (Fig. 16). Silicic eruptions and magmatic intrusions into Siberian coalfields are

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**Figure 16.** Time line of events in the Sverdrup Basin and Tethyan region during the Late Permian and Early Triassic. Late Changhsingian extinctions of sponges in the Sverdrup Basin and radiolarians in the Tethyan region predate main-stage Siberian Traps eruptions by ~100 k.y. and may have been associated with early stage explosive silicic eruptions (cf. Grasby et al., 2011). Note expanded time scale between 252.5 and 252.0 Ma. A to F refer to correlation datums in Figures 4A and 5. MP—Middle Permian, MTr—Middle Triassic; LPME—latest Permian mass extinction; AEE—Arctic extinction event; FAD—first appearance datum; OMZ—oxygen minimum zone; LIP—large igneous province.
likely to have resulted in deposition of toxic gases and ash that degraded or destroyed terrestrial floras (Grasby et al., 2011; cf. Self et al., 2008), leading to large-scale erosion of soils with consequent detrimental effects on marine ecosystems (Algeo et al., 2011). Pre-main-stage eruptions are thought to have begun during the early late Changhsingian (Kozur and Weems, 2011; Grasby et al., 2011), up to several hundred thousand years prior to the latest Permian mass extinction and thus roughly contemporaneous with marine environmental changes in the Sverdrup Basin (this study) and with changes in intermediate-depth marine systems globally (see Introduction section). In the following, we summarize the pattern of marine environmental changes discussed above in reference to the main event horizons at West Blind Fiord.

Datum A (4.0 m) in the upper Lindström Formation is the first horizon recording major environmental changes within the study section. At this level, the detrital fraction underwent a compositional change toward higher Fe-oxhydroxide concentrations and a CIA signature indicative of greater weathering intensity. High CIA values may be an indication of a large particulate contribution from soils, where strong cation leaching is common (Maynard, 1992), and the pulse of Fe-oxhydroxides may reflect erosion of the Fe-rich B horizons of soils. This event was accompanied by a modest shift toward more reducing conditions, probably caused by elevated fluxes of soil-derived nutrients that triggered an increase in marine productivity. Concurrently, the abundance of siliceous sponges, the most prominent remaining component of the Late Permian biota in the Sverdrup Basin, was reduced modestly, probably in response to changing nutrient and/or redox conditions.

The formation contact (5.6 m) between the Lindström and Blind Fiord formations records the final extinction of siliceous sponges in the Sverdrup Basin, herein termed the “Arctic extinction event.” A concurrent large increase in detrital sediment flux may have been the primary cause of this event. A shift toward lower Corg(NO3)N ratios is indicative of a larger fraction of marine organic matter, and higher Corg/Ca ratios may reflect greater abundance of algal- or bacterial-consuming ciliated protozoans. Both patterns are consistent with an increase in marine productivity, which is a likely consequence of enhanced terrestrial inputs to marine systems (Algeo et al., 2011). A shift toward less-reducing conditions across the formation contact may have been related to enhanced freshwater runoff into the Sverdrup Basin in conjunction with higher detrital sediment fluxes.

The latest Permian mass extinction (10.2 m) is not characterized by any apparent change in the fossil biota (since all macrofauna had disappeared earlier) or marine productivity, nor did it have any noticeable effect on the composition of the detrital fraction or subaerial weathering intensity. The most significant changes at this level are related to benthic redox conditions, which became more persistently and/or more intensely euxinic, as recorded by multiple redox proxies, although changes in phytoplankton community composition may have occurred at the same time, as proxied by δ13Corg and δ15N. Persistently euxinic conditions reflect a sustained shallowing of the Sverdrup Basin chemocline to the water depth of the West Blind Fiord study site. One possibility is that chemocline shallowing was driven by enhanced marine productivity, which had accelerated at the formation contact (~100 k.y. earlier) and may have coincidentally peaked at the latest Permian mass extinction. An alternative is that the chemocline shallowed abruptly at the latest Permian mass extinction due to global climatic warming triggered by release of volcanic CO2 or coalfield methane (Retallack, 1999), resulting in a rapid reduction of the solubility of dissolved oxygen in seawater (but see Bond and Wignall [2010] for an opposing view). The latter process could have been triggered by the onset of the main stage of Siberian Traps volcanism (Reichow et al., 2009).

An important issue is the apparent difference in timing of terrestrial disturbances in the Boreal and Tethyan regions. An increased flux of detrital siliciclastics to the Sverdrup Basin commenced in the early part of the late Changhsingian (this study), whereas the Tethyan region did not experience this influx until the latest Changhsingian, just prior to the latest Permian mass extinction (Sephton et al., 2005; Xie et al., 2005, 2007; Algeo et al., 2007; Wang and Visscher, 2007). We suggest that the key difference was paleolatitude. More northerly sites may have been more vulnerable to deposition of acidic gases and ash from early stages of the Siberian Traps eruptions (Fig. 16; Grasby et al., 2011; cf. Self et al., 2008). In the latest Permian, the West Blind Fiord study site was located at ~40°N, which was much closer to the Siberian Traps volcanic centers at 60–70°N in terms of both paleolatitude and absolute distance (~3000–4000 km) than sites in the Tethyan region (Fig. 1, inset). Following explosive silicic eruptions, fallout is commonly initially constrained to a narrow latitudinal belt, and the intensity of fallout diminishes considerably as it spreads to higher and lower latitudes (cf. McCormick et al., 1995; Robock, 2002). The environmental effects of volcanic emissions can be regionally and/or latitudinally constrained, as following the 1783 Laki eruption in Iceland (Thordarson and Self, 2003). Sulfate emissions, in particular, can have environmentally harmful effects related to acidification of soils and surface waters and metabolic toxicity (Thordarson and Self, 2003; Ward, 2009; Bao et al., 2010). Such effects may have been “felt” in the boreal Sverdrup Basin sooner than in the peri-equatorial Tethyan region.

CONCLUSIONS

The West Blind Fiord section on southwestern Ellesmere Island provides a record of deteriorating environmental conditions in the Sverdrup Basin during the late Changhsingian (late Late Permian). Environmental deterioration proceeded stepwise, with most change associated with a few stratigraphic horizons. Datum A, of early late Changhsingian age, records increased Fe-oxhydroxide fluxes and weathering intensity as well as modest shifts toward more-reducing water-mass conditions and higher marine productivity, recording an initial disturbance that washed soils into the marine environment. The contact between the Lindström Formation and the overlying Blind Fiord Formation, ~50 k.y. younger than datum A, records a large increase in detrital sediment flux, elevated marine productivity, and the disappearance of siliceous sponges during the Arctic extinction event. The horizon equivalent to the global latest Permian mass extinction, ~100 k.y. younger than the Arctic extinction event, records strong intensification of anoxia and possible changes in phytoplankton community composition. Compared to some sections, Meishan in particular, the phytoplankton community remained relatively stable throughout the study interval, although characterized by recurrent increases in the relative abundances of green sulfur bacteria and diazotrophic cyanobacteria and, perhaps, an increase in the ratio of red to green algae above the Arctic extinction event. These patterns are consistent with a fluctuating but generally rising chemocline in the Sverdrup Basin during the latest Permian, probably triggered by an increase in seawater nutrient levels and, at the latest Permian mass extinction, a reduction in dissolved oxygen solubility due to climatic warming. These changes in the marine environment were ultimately driven by increasing detrital sediment fluxes resulting from terrestrial ecosystem degradation, possibly related to early stage explosive silicic eruptions of Siberian Traps volcanic centers. The deleterious effects of these eruptions may have been felt sooner in the Boreal region than in the peri-equatorial Tethyan region owing to latitudinally concentrated fallout of toxic gas and ash.
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Addendum

Evidence for a diachronous Late Permian marine crisis from the Canadian Arctic region

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The samples from the West Blind Fiord section analyzed for geochemistry in this study were collected in 1999 by the Geological Survey of Canada, namely by Steve Grasby and Benoit Beauchamp, and are the property of the Government of Canada.