



A Basin Redox Transect at the Dawn of Animal Life

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A basin redox transect at the dawn of animal life

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44 **Multiple eukaryotic clades make their first appearance in the fossil record between**
45 **~810 and 715 Ma. Molecular clock studies suggest that the origin of animal**
46 **multicellularity may have been part of this broader eukaryotic radiation. Animals**
47 **require oxygen to fuel their metabolism, and low oxygen levels have been**
48 **hypothesized to account for the temporal lag between metazoan origins and the**
49 **Cambrian radiation of large, ecologically diverse animals. Here, paleoredox**
50 **conditions were investigated in the Fifteenmile Group, Ogilvie Mountains, Yukon,**
51 **Canada, which hosts an 811 Ma ash horizon and spans the temporal window that**
52 **captures the inferred origin and early evolution of animals. Iron-based redox**
53 **proxies, redox-sensitive trace elements, organic carbon percentages and pyrite**
54 **sulfur isotopes were analyzed in seven stratigraphic sections along two parallel**
55 **basin transects. These data suggest that for this basin, oxygenated shelf waters**
56 **overlay generally anoxic deeper waters. The anoxic water column was dominantly**
57 **ferruginous, but brief periods of euxinia likely occurred. These oscillations coincide**
58 **with changes in total organic carbon, suggesting euxinia was primarily driven by**
59 **increased organic carbon loading. Overall, these data are consistent with proposed**
60 **quantitative constraints on Proterozoic atmospheric oxygen being greater than 1%**
61 **of modern levels, but less than present levels. Comparing these oxygen levels against**
62 **the likely oxygen requirements of the earliest animals, both theoretical**
63 **considerations and the ecology of modern oxygen-deficient settings suggest that the**
64 **inferred oxygen levels in the mixed layer would not have been prohibitive to the**
65 **presence of sponges, eumetazoans or bilaterians. Thus the evolution of the earliest**
66 **animals was probably not limited by the low absolute oxygen levels that may have**

67 **characterized Neoproterozoic oceans, although these inferred levels would constrain**

68 **animals to very small sizes and low metabolic rates.**

69

70 **1. Introduction**

71 A number of eukaryotic groups first appear in the fossil record between the Bitter
72 Springs isotope excursion at ~810 Ma and the Sturtian glaciation at ~715 Ma (Macdonald
73 et al., 2010). This apparent radiation includes the first unequivocal appearances of groups
74 such as the vase-shaped microfossils, interpreted to be related to lobose, and perhaps
75 filose, testate amoebae (Porter and Knoll, 2000; Porter et al., 2003), scale microfossils of
76 uncertain phylogenetic affinity (Cohen et al., 2011; Cohen and Knoll, 2012), and simple
77 multicellular and coenocytic green algae (Butterfield et al., 1994). Interestingly,
78 molecular clock studies suggest that the origin of animal multicellularity may have been
79 part of this broader radiation. Studies utilizing different taxa, genes, calibration points
80 and clock models have converged on an estimated divergence of ~800 Ma for the last
81 common ancestor of animals (Berney and Pawlowski, 2006; Lartillot et al., 2009;
82 Sperling et al., 2010; Erwin et al., 2011; Parfrey et al., 2011). Similar results in these
83 studies, despite broad methodological differences, suggest this divergence estimate is
84 approximately correct. This age finds further support in the appearance of presumed
85 demosponge-specific biomarkers beneath ca. 635 Ma Marinoan glacial deposits (Love et
86 al., 2009; Kodner et al., 2008); as demosponges represent a derived lineage within
87 animals, the origin of the animal crown group must be even deeper in time. If the
88 molecular clock ages and biomarker data are accurate, however, the lack of metazoan
89 body and trace fossils throughout the Cryogenian and early Ediacaran periods presents a
90 conundrum (Erwin et al., 2011). It has been hypothesized that animal body size and
91 diversity may have been limited by relatively low levels of oxygen in the Proterozoic
92 atmosphere and oceans. In such oceans, it is posited that animals could have been

93 restricted to small and thin body plans that did not fossilize well, with the explosion of
94 larger and ecologically diverse organisms in the late Ediacaran and Cambrian related in
95 part to increasing O₂ levels (Cloud, 1968; Rhoads and Morse, 1971; Runnegar, 1982a;
96 Knoll and Carroll, 1999). Consistent with this hypothesis, different geochemical redox
97 proxies support a directional change towards more oxygenated conditions in the latest
98 Proterozoic (reviewed by Och and Shields-Zhou, 2012; Kah and Bartley, 2011).

99 What remains highly uncertain, however, are the atmospheric and oceanic oxygen
100 concentrations prior to and during earliest animal evolution, specifically during the
101 Cryogenian period (850-635 Ma). Oxygen levels are generally assumed to have been
102 relatively low in Cryogenian oceans (e.g. Kump, 2008), but given the lack of widespread
103 paleoenvironmental documentation, the extent to which early animals were limited by
104 low oxygen levels remains unknown. Specifically, the physiological requirements of
105 small animals with low-energy lifestyles that may have characterized the Cryogenian
106 Period were likely different from the larger, more active and muscular organisms
107 preserved in Cambrian rocks. This difference needs to be considered when comparing
108 physiological requirements against the constraints provided by geochemical proxies.

109 Here, we investigate the environmental context of early animal evolution and
110 compare inferred redox constraints with the likely physiological requirements associated
111 with different grades of organization in early animal evolution. Previous iron speciation
112 and sulfur isotope studies of the pre-Sturtian Chuar Group (Canfield et al., 2008; Nagy et
113 al., 2009; Johnston et al., 2010) provide insight into Cryogenian environments, but are
114 limited to a single section deposited between ca. 770 and 742 Ma (Karlstrom et al.,
115 2000). Here we report geochemical redox proxies through seven sections along two

116 parallel platform-to-basin transects in the early Cryogenian Fifteenmile Group in the
117 Tatonduk and Coal Creek inliers, Ogilvie Mountains, Yukon, Canada (Figs. 1 and 2). The
118 Fifteenmile Group was deposited in a basin that originated during an episode of
119 continental extension (Macdonald et al., 2012) prior to 811.51 ± 0.25 Ma, the U-Pb
120 zircon date on a tuff in the upper portion of the Reefal Assemblage (green line in Mt.
121 Harper Section, Figs. 3 and 6; Macdonald et al., 2010). Thus the Fifteenmile Group spans
122 a time period that significantly preceded the earliest macroscopic multicellular forms in
123 the Ediacaran Period (Narbonne, 2011) but overlaps with molecular-clock estimates for
124 the divergence of crown-group animals (Erwin et al., 2011, and references above).

125 The paleoredox state of shale samples collected from measured stratigraphic
126 sections was investigated using a multi-proxy approach. Specifically, iron speciation data
127 are integrated with major-element and redox-sensitive trace element abundances, total
128 organic carbon (TOC) percentages, and pyrite sulfur isotope values to obtain an estimate
129 of overall water-column redox profiles. Together, the geochemical data from these
130 stratigraphic sections provide the first early Neoproterozoic basin redox transect and give
131 insight into paleoenvironmental conditions in this basin at the dawn of animal life. These
132 data can then be placed in the context of other information constraining Mesoproterozoic
133 and early Neoproterozoic oxygen levels and compared to the likely physiological
134 requirements of early animals.

135

136 **2. Geologic Background**

137 Neoproterozoic strata in the northern Canadian Cordillera are exposed in
138 erosional windows ('inliers') separated by Phanerozoic cover (Rainbird et al., 1996;

139 Thorkelson et al., 2005) (Fig. 1). In the Coal Creek inlier, the focus of this study,
140 geological mapping (Fig. 2) and stratigraphic analysis indicate that Neoproterozoic
141 extension produced a series of NNW-side down normal faults, such that the basin, at least
142 locally, deepened towards the northwest in present-day coordinates (Macdonald et al.,
143 2012). The Fifteenmile Group consists of lagoonal, tidal, and supertidal carbonates of the
144 informal Gibben formation, tidal flat and deltaic deposits of the Chandindu formation,
145 and mixed carbonates and siliciclastics of the Reefal Assemblage, which is characterized
146 by km-scale stromatolitic reefs that transition laterally into shale-dominated, deeper water
147 sub-basins (Macdonald et al., 2012). Shales were sampled from two parallel transects
148 across the basin (Fig. 2), including a shorter transect passing a short distance from a
149 stromatolite reef complex into the shale basin (Fig. 4), and a longer transect stepping
150 further into the basin (Fig. 3). Shales were also investigated from exposures of the Reefal
151 Assemblage ~75 km to the northwest in the Tatonduk inlier that have yielded distinctive
152 scale microfossils (Cohen et al., 2011; Cohen and Knoll, 2012). As Fifteenmile Group
153 strata in the Tatonduk inlier are represented only by shale interbedded with re-deposited
154 carbonate (and no evidence for shallow-water sedimentation), these exposures are
155 interpreted to have formed in a deeper, more distal environment than correlative sections
156 in the Coal Creek inlier (Macdonald et al., 2012); however, displacement along poorly
157 exposed post-Jurassic faults between the two inliers precludes precise paleogeographic
158 reconstruction.

159

160 **3. Materials and Methods**

161 234 shale samples from logged stratigraphic sections were crushed to flour and
162 analyzed for major and minor-element concentrations, iron speciation systematics,
163 percent carbonate carbon and organic carbon, and pyrite sulfur isotope composition. Iron
164 sequential extraction followed standard protocols for iron carbonate, iron oxide and
165 magnetite extractions (Poulton and Canfield, 2005), while pyrite iron content was
166 quantified using the chromous chloride extraction method (Canfield et al., 1986). Pyrite
167 sulfur isotopes were determined through combustion via a Costech Elemental Analyzer
168 linked to a Thermo Scientific Delta V in continuous flow mode (measured as SO-SO₂)
169 using Ag₂S from the chromous chloride extraction. Major- and minor-element
170 abundances were determined following a standard acid digestion (hydrofluoric,
171 perchloric, hydrochloric and nitric) and measurement with ICP-AES at SGS Laboratories,
172 Canada. Percent carbonate carbon was quantified by percent loss on acid dissolution.
173 Total organic carbon values were determined on acidified samples by combustion within
174 a Carlo Erba NA 1500 Analyzer attached to a Thermo Scientific Delta V Advantage
175 isotope ratio mass spectrometer. Complete materials and methods and precision estimates
176 for each analysis are contained in Supplementary Information.

177

178 **4. Results**

179 All geochemical measurements are reported in Supplemental Information Tables
180 1 and 2. Iron speciation data are plotted against the sequence stratigraphic framework for
181 the Coal Creek inlier (Macdonald et al., 2012) in Figs. 3 and 4. Full redox proxy data are
182 plotted against stratigraphy for the principal investigated sections including the short
183 transect at Reefer Camp (Fig. 5), the long transect at Mt. Harper (Fig. 6), and the deepest-

184 water section at Mt. Slipper (Fig. 7). Similar plots for sections with more limited data in
185 the Coal Creek inlier (Mine Camp, East Harper and Mt. Gibben) can be found in
186 Supplemental Figs. 1-3, respectively.

187

188 4.1 *Multi-proxy estimation of paleo-redox state*

189 An estimate of water-column redox state was determined using a multi-proxy
190 approach based on iron speciation chemistry, redox-sensitive trace elements (especially
191 Mo and V) and pyrite sulfur isotope values. In iron speciation chemistry, the highly-
192 reactive pool (FeHR) consists of iron in pyrite (FeP) plus iron that is reactive to sulfide
193 on early diagenetic timescales (iron carbonates such as siderite and ankerite, and iron
194 oxides, including magnetite). The remaining unreactive pool (FeU) consists mainly of
195 iron in sheet silicates; the sum of the two pools is total iron (FeT). Key to the geological
196 application of this proxy is the observation that modern sediments deposited under oxic
197 water columns have a $\text{FeHR}/\text{FeT} < 0.38$, while those deposited beneath anoxic water
198 masses generally have $\text{FeHR}/\text{FeT} > 0.38$ [Raiswell and Canfield (1998); see also Farrell,
199 (2011), and Supplementary Information for further discussion]. The proxy can also
200 distinguish the nature of an anoxic water column based on the proportion of highly
201 reactive iron that has been sulfidized, with FeP/FeHR ratios >0.80 indicating an euxinic
202 water column, and lower ratios pointing towards ferruginous conditions (Anderson and
203 Raiswell, 2004; Poulton and Canfield, 2011).

204 Like all proxies, iron speciation has acknowledged caveats. For instance, dilution
205 by turbidites or rapid sedimentation can result in low FeHR/FeT ratios, imparting a false
206 oxic ‘signature’ to sediments deposited under an anoxic water column (Raiswell and

207 Canfield, 1998; Lyons and Severmann, 2006). Near-shore or estuarine sediments can trap
208 large amounts of iron oxides, leading to an anoxic FeHR/FeT signature for sediments
209 deposited under oxic conditions (Poulton and Raiswell, 2002). Weathering can oxidize
210 Fe²⁺ phases to Fe³⁺ phases, potentially skewing the interpretation of euxinic versus
211 ferruginous conditions (see below), although the FeHR term should remain constant
212 (Canfield et al., 2008). Consistency between independent proxies is the best test of an
213 inference, and consequently we integrated the iron speciation chemistry with other redox
214 proxies and sedimentological constraints. Redox-sensitive trace elements such as
215 vanadium and molybdenum are soluble under oxic conditions but are less soluble under
216 reducing conditions, and are scavenged by organic and inorganic particles or complex
217 with sulfide, leading to enrichments compared to average shale values (Tribovillard et al.,
218 2006). Pyrite sulfur isotope values can further inform paleoenvironmental reconstruction,
219 because sulfate reduction within a water-column where sulfate is not limiting allows
220 expression of the biological preference for lighter ³²S, and consequently depleted isotopic
221 compositions in the resulting pyrite with respect to seawater sulfate. Sulfate reduction
222 within sediments, on the other hand, where sulfate availability is often diffusion limited,
223 results in Rayleigh distillation, leading to pyrite values that approach seawater sulfate
224 (Johnston and Fischer, 2012).

225 In our multi-proxy framework, samples were considered likely to have been
226 deposited under an oxic water column when they showed FeHR/FeT < 0.38 (Raiswell
227 and Canfield, 1998), no enrichment in Mo and V with respect to average shales (Gromet
228 et al., 1984), and relatively enriched $\delta^{34}\text{S}$ pyrite sulfur isotope values (or not enough
229 sulfide present in the rock for measurement). Samples were considered to have been

230 deposited under an anoxic, ferruginous water column when they showed FeHR/FeT >
231 0.38, little to no Mo enrichment but often with V enrichment, and relatively enriched $\delta^{34}\text{S}$
232 pyrite sulfur isotope values. Finally, samples were considered to have been likely
233 deposited under an anoxic, euxinic water column when they showed FeHR/FeT > 0.38,
234 relatively high FeP/FeHR ratios, Mo and V enrichments, and depleted $\delta^{34}\text{S}$ pyrite sulfur
235 isotope values.

236 Euxinic water columns are usually distinguished by FeP/FeHR > 0.80 or 0.70
237 (Poulton and Canfield, 2011), a ratio which few of these samples surpasses. Samples
238 interpreted here as euxinic, though, clearly have much higher FeP/FeHR ratios than
239 samples interpreted as ferruginous (see Figs. 6 and 7), and essentially no iron carbonate
240 or magnetite. Further, these shales have very depleted $\delta^{34}\text{S}$ pyrite sulfur isotopes (to -34
241 ‰) and high molybdenum abundances relative to other Neoproterozoic samples (Scott et
242 al., 2008). Two likely possibilities exist to explain these patterns: 1) these shales were
243 deposited beneath an euxinic water column, with subsequent oxidation of pyrite to iron
244 oxides, or (2) the shales were deposited beneath a ferruginous water column, with the
245 zone of free sulfide accumulation essentially at the sediment-water interface.

246 Petrographic examination of selected shales did not show widespread evidence for
247 oxidation of pyrite, although because the samples are from outcrop and surely have
248 suffered some alteration, it is possible that micron-scale pyrite grains beneath the limits
249 of routine petrographic detection have been wholly or partially oxidized. In the second
250 possibility, full access to seawater sulfate and molybdenum pools could explain the
251 isotopic and abundance patterns for these two elements, while the shorter time interval
252 exposed to high sulfide levels compared to a fully euxinic water column would result in

253 less sulfidization of highly-reactive iron phases. Recognizing that the development of
254 truly euxinic conditions is ambiguous and these data may represent sulfide production at
255 the sediment-water interface, inferences of euxinia in Figs. 5-7 should be treated with
256 caution.

257

258 4.2 *Sedimentary Geochemistry of the Fifteenmile Group*

259 4.2.1 *Reefer Camp transect, Coal Creek inlier*

260 Near Reefer Camp, samples from the shallow-water Chandindu formation show
261 the hallmarks of deposition under an oxic water column (Fig. 5A). Within the
262 stromatolite reef core of the Reefal Assemblage, thin black shales show high FeHR/FeT,
263 but because they show no redox-sensitive trace element enrichment, and the FeHR signal
264 is entirely dominated by iron oxides, this may represent nearshore trapping of oxides, as
265 occurs in modern settings (Poulton and Raiswell, 2002) rather than a true ocean redox
266 signal. Samples above the flooding surface atop the stromatolite reef tract have iron
267 speciation values persistently above 0.38, moderate amounts of iron carbonate, no Mo
268 enrichment and enriched pyrite sulfur isotope values, pointing to deposition under
269 ferruginous conditions. Samples from the upper part of the Reefal Assemblage signal an
270 apparent return to oxic deposition. In Fig. 5B (See Fig. 4 for the stratigraphic relationship
271 of these sections), the Chandindu formation samples again show evidence for oxic
272 deposition. Continuing upsection into shale of the Reefal Assemblage, all available
273 evidence points to deposition under a generally oxic water column.

274

275 4.2.2 *Mt. Harper, Coal Creek inlier*

276 Near Mt. Harper, shallow-water sediments of the Chandindu formation also show
277 evidence for oxic deposition (Fig. 6). The Mt. Harper transect steps much farther
278 westward into the Reefal Assemblage shale basin than the short transect at Reefer Camp,
279 and records a thick package of black shale and foreslope carbonate that fill
280 accommodation space associated with tectonic extension (Macdonald et al., 2012). In
281 section GO134, the stratigraphically lowest exposed shales of the Reefal Assemblage,
282 there is evidence for euxinic deposition. Many of these samples do not show $Fe_{HR}/Fe_T >$
283 0.38, but as this succession contains many siltstone turbidites and redeposited carbonates
284 and was likely deposited rapidly during active extension (Macdonald et al., 2012), it is
285 possible that the highly-reactive iron was diluted by high sedimentation rates. Thus a
286 threshold for anoxia of 0.22 may be more appropriate (Raiswell and Canfield, 1998; see
287 also discussion in Supplementary Information). These samples show relatively high
288 Fe_P/Fe_{HR} , high Mo (~10-32 ppm; high for Neoproterozoic shales—Scott et al., 2008),
289 and depleted pyrite sulfur isotope values, indicating sulfide production very near to the
290 sediment-water interface, if not in the water column (see above). The upper half of
291 section GO134 shows lower Mo and less depleted pyrite sulfur isotope values, potentially
292 suggesting ferruginous or even oxic conditions. Samples at the base of section S1103
293 have $Fe_{HR}/Fe_T < 0.38$, no redox-sensitive trace element enrichments, and no pyrite,
294 possibly recording deposition under oxic conditions. This is followed by a second pulse
295 of euxinic deposition, showing similar characteristics to the samples in GO134, with
296 elevated Fe_P/Fe_{HR} , high Mo abundances, and depleted pyrite sulfur isotope values.
297 There is little evidence for euxinia above this level, with ferruginous conditions dominant
298 in the upper Reefal Assemblage. In contrast to samples from the lower Reefal

299 Assemblage at Mt. Harper, where the FeHR pool is almost entirely in pyrite and iron
300 oxides, samples from the upper Reefal Assemblage contain moderate quantities of iron
301 carbonate. In combination with low Mo, and enriched and variable pyrite sulfur isotope
302 values, this suggests that the upper half of the Reefal Assemblage accumulated under an
303 anoxic, ferruginous water column. Brief and fluctuating water column oxygenation may
304 have occurred, as evidenced by stratigraphically-variable iron speciation signatures.

305

306 4.2.3 *Mt. Slipper, Tatonduk inlier*

307 Iron speciation values from the Reefal Assemblage in the Tatonduk inlier (Figure
308 7), which represent the deepest-water setting studied (Macdonald et al., 2012), generally
309 show $\text{FeHR}/\text{FeT} > 0.38$, indicating persistent deposition under an anoxic water column.
310 Samples from the base of the section have relatively high FeP/FeHR , high Mo and
311 depleted pyrite sulfur isotope values, pointing to euxinic deposition (or at least
312 fluctuating euxinia). At ~90m, these proxy values decrease, indicating a transition to
313 ferruginous conditions. A possible return to euxinia is seen at the top of the section, from
314 strata that have yielded scale microfossils (Cohen et al., 2011).

315

316 4.2.4 *Sediment total iron contents*

317 The total iron to aluminum ratio is another informative redox proxy, because
318 sedimentary iron is authigenically enriched under anoxic water columns (Lyons and
319 Severmann, 2006). An interesting feature of shale samples from the Fifteenmile Group is
320 that even samples considered to have been deposited under anoxic conditions have Fe/Al
321 ratios lower than average shale (Gromet et al., 1984). Given the general concordance in

322 these samples of FeHR/FeT, redox-sensitive trace element abundances and pyrite sulfur
323 isotope data, the inconsistency with Fe/Al likely indicates an intrinsic bias to either total
324 iron or total aluminum in the Reefal Assemblage, rather than this representing oxic
325 deposition. Total Al abundances in all shales investigated (average = 7.03 wt %) are
326 slightly depressed relative to the North American Shale Composite (NASC; Gromet et
327 al., 1984; 8.94 wt %). In contrast, total Fe (average = 2.43 wt %) is significantly reduced
328 relative to NASC (4.43 wt %), especially considering that basinal samples interpreted as
329 anoxic should be enriched in iron. Some Reefal Assemblage shales are exceptionally low
330 in total iron (<1 weight percent), and have very high FeHR/FeT ratios indicating a near-
331 absence of detrital iron-silicates. Dilution by carbonate may explain some of the low iron
332 contents, as some samples are slightly calcareous (to ~30-40%, average 9.45% ± 9.50),
333 Supplementary Table 1) but low iron contents persist in shale samples that have
334 essentially no carbonate (e.g. GO134 and S1103 sections). Open-system diagenesis could
335 have potentially affected these rocks, although even the marls would have had very low
336 permeability. Further, the main effect of diagenesis in carbonates is to add iron (Brand
337 and Veizer, 1980), which is unlikely given the low overall amounts of acetate-extractable
338 iron (average 0.13 weight percent) and the lack of a relationship between percent
339 carbonate and acetate-extractable iron ($R^2 = 0.062$). Another possibility is that the
340 provenance was extremely weathered, iron-free material. However, Chemical Index of
341 Alteration (CIA; Nesbitt and Young, 1982) values average ~70 throughout the dataset,
342 indicating a fairly unweathered provenance. A few values in the 75-85 range suggest a
343 weathered source for those samples, but overall there is no obvious correlation between
344 CIA and total iron. Thus, while several factors may explain some low iron values, none

345 can explain all low values. We note that some other Neoproterozoic sections show
346 anomalously low FeT/Al (e.g. Sahoo et al., 2012); further study is needed to determine if
347 these are local, basin-level effects or an as-yet-unexplained aspect of the Neoproterozoic
348 iron cycle.

349

350 *4.2.5 Redox proxy data and sediment organic carbon contents*

351 Sediment TOC results vary consistently compared to multi-proxy inferences of
352 redox state (Fig. 8). Sediments likely deposited under an oxic water column have low
353 organic carbon abundances (average = 0.31% ± 0.49 weight percent; median = 0.19%),
354 whereas those inferred to have been deposited beneath a ferruginous water column have
355 higher sediment TOC values (average = 0.66% ± 1.37; median = 0.28%). And sediments
356 likely deposited beneath euxinic conditions show much greater TOC (average = 2.87% ±
357 1.49; median = 2.63. Thus, these data are consistent with the hypothesis that the
358 development of euxinic conditions in Neoproterozoic basins is primarily driven by the
359 degree of organic carbon loading and the exhaustion of more energetically-favorable
360 electron acceptors than sulfate (e.g. Fe³⁺) (Johnston et al., 2010).

361

362 *4.2.6 Redox proxy data and water depth*

363 Redox proxy data show a consistent pattern with respect to sedimentological
364 structures that indicate relative water depth. In particular, all occurrences of hummocky
365 cross-stratified sandstones encased within shale, which indicate deposition above storm
366 wave base, show evidence for oxic deposition (Fig. 3-7). In other words, the surface
367 mixed layer in the basin appears to be oxygenated, at least during storms. While the depth

368 of storm wave base varies among basins (Peters and Loss, 2012), these data indicate that
369 in this basin, the water column in direct contact with the atmosphere remained oxic.
370 Oxygenated conditions may extend slightly deeper, as some sediments likely deposited
371 below storm wave base (such as shales basinward of the stromatolite reef at Reefer
372 Camp, Fig. 5B) still indicate oxic conditions. A few brief intervals of oxygenated
373 conditions, or fluctuating anoxia, persist deeper into the Coal Creek inlier shale basin as
374 recorded in the Mt. Harper (Fig. 6) and Mine Camp (Supplementary Fig. 1) sections, but
375 the majority of these deeper-water sediments record anoxic conditions. The deepest-water
376 section at Mt. Slipper in the Tatonduk inlier (Fig. 7), which shows no evidence for wave
377 activity, is persistently anoxic. Thus, there is a clear redox structure to the basin, with an
378 oxygenated surface layer where the sediments are in contact with the atmosphere (storm
379 wave base), and anoxic conditions below this depth.

380

381 **5. Discussion**

382 5.1 *Fifteenmile Group redox structure in a global context*

383 Quantitative constraints on Proterozoic oxygen levels are notoriously difficult to
384 obtain (Kump, 2008). O₂ levels must have been above 0.001% present atmospheric levels
385 (PAL), the limit imposed by the disappearance of mass-independent fractionation of
386 sulfur isotopes at ~2.45 Ga (Farquhar et al., 2000; Pavlov and Kasting, 2002). Two other
387 constraints have been proposed for mid-Proterozoic O₂ (Kump, 2008). First, anoxic deep
388 oceans likely require atmospheric O₂ to be less than 40% PAL (Canfield, 2005). Second,
389 it has been proposed that iron is only retained in lithified soil horizons, and it has been
390 since the Paleoproterozoic, when O₂ is greater than 1% PAL (Holland and Beukes, 1990).

391 These limits on Proterozoic O₂ have caveats, and it has even been hypothesized that
392 levels may not have been dramatically different from the Phanerozoic (Butterfield, 2009).
393 Nonetheless, it is notable that the basin redox transect of the Fifteenmile Group is
394 consistent with proposed quantitative limits (Kump, 2008). Indeed, the basin redox
395 structure of the Fifteenmile Group is similar in many ways to that of the Mesoproterozoic
396 Roper Group in Australia (Shen et al., 2003), with an oxygenated shelf overlying anoxic
397 basinal waters. Although there are local drivers for anoxia (Tyson and Pearson, 1991), the
398 available basin redox transects point to extensive subsurface anoxia in the Proterozoic
399 oceans, sustained over hundreds of meters of stratigraphic section. This clearly differs
400 from Phanerozoic ocean anoxic events (Campbell and Squire, 2010), indicating a
401 different driver and implying lower atmospheric O₂ than the modern. Placing minimum
402 constraints on global atmospheric pO₂ levels from local iron speciation data is difficult,
403 but shallow-water facies in the Fifteenmile Group record oxic deposition, as do samples
404 from the shale basin just off the reef margin at Reefer Camp (Fig. 5), and some samples
405 from deeper in the shale basin at Mt. Harper (Fig. 6) and Mine Camp (Supplementary
406 Fig. 1), implying enough atmospheric oxygen to counteract strong benthic reductant
407 fluxes in a basin otherwise prone to euxinia (cf. Kump et al., 2005).

408 In sum, although there is clear need to study more basins, and develop new global
409 redox proxies and models, the basin redox transect of the Fifteenmile Group is consistent
410 with proposed constraints on Proterozoic oxygen levels as being <40% and >1% PAL
411 (Kump, 2008). We apply these bounds for comparison with the physiological
412 requirements of early animals.

413

414 5.2 *Physiological requirements of early animals*

415 The consistency of previously proposed constraints on atmospheric oxygen with
416 the basin redox transect of the Fifteenmile Group prompts the question of whether such
417 oxygen levels would have prohibited the evolution of animal, eumetazoan or bilaterian
418 body plans. A common assumption in attempts to link late Precambrian oxygenation and
419 biospheric evolution is that animals have high respiratory demands. While metazoans do
420 have a clear and definite requirement for oxygen, they are not a monolithic group, and the
421 oxygen requirements for any given organism varies widely based on size, metabolism,
422 and the presence or absence of a circulatory system (Vaquer-Sunyer and Duarte, 2008).
423 Hypotheses relating geochemical change to early animal evolution must therefore
424 compare inferred changes against the explicit body plans, ecological strategies and
425 taxonomic groups presumed to be affected. Determining the physiological requirements
426 of ancient organisms has obvious uncertainty, but can be accomplished through analogy
427 with living representatives (Knoll et al., 2007), and thus it is possible to make general
428 statements about the likely oxygen requirements of Precambrian animals.

429

430 5.2.1 *Diploblastic metazoans*

431 Whether sponges are monophyletic (Philippe et al., 2009) or paraphyletic
432 (Sperling et al., 2009), they are certainly the sister group or grade of all other animals
433 (Philippe et al., 2011). Moving up the metazoan phylogenetic tree, the exact relationships
434 of cnidarians, ctenophores and placozoans to bilaterians are unclear, but all are likely
435 more closely related to bilaterians than they are to sponges (Philippe et al., 2011).
436 Importantly, all these animals (diploblasts) are characterized by only two epithelial cell

437 layers, with the space between layers filled largely with metabolically-inert material (e.g.
438 mesohyl in sponges, mesoglea in cnidarians). From a respiratory point of view, then,
439 essentially every cell in a diploblastic metazoan is in direct contact with seawater
440 (Ruppert et al., 2004). Thus, the theoretical oxygen limit for diploblastic animals will not
441 differ from that of a single-celled eukaryote, barring two minor differences. First, for
442 unicellular eukaryotes, diffusion of oxygen into the cell can occur across the entirety of
443 its surface, whereas diffusion into a sheet of cells cannot occur at cell-cell contacts.
444 Second, animals have a collagenous extracellular matrix, and molecular oxygen is
445 required for the formation of hydroxyproline in collagen (Fujimoto and Tamiya, 1962;
446 Prockop et al., 1962). Using the *K_m* for the proline hydroxylase system of chick embryos,
447 Towe (1970) suggested oxygen levels of ~3% PAL would be required for collagen
448 synthesis. However, Rhoads and Morse (1971) cogently noted that collagen-rich
449 invertebrates are found at oxygen levels beneath this value (see also discussion below on
450 modern oxygen minimum zones), suggesting that the oxygenase requirements of a
451 terrestrial vertebrate cannot be applied to marine invertebrates. Further, collagen is now
452 known to exist in fungi (Celerin et al., 1996; Wang and St. Leger, 2006) and
453 choanoflagellates (King et al., 2008--although the homology of both to metazoan
454 collagens remains uncertain), which suggests collagen may have been present in the last
455 common ancestor of opisthokonts. If so, any oxygen requirement for collagen synthesis
456 was met far earlier than the origin of animals.

457 In the fossil record, clear eukaryotic organisms are found at ~1800 Ma, and
458 several lineages of multicellular eukaryotes, which would also have been subject to the
459 same constraint as early animals of limited diffusion at cell-cell contacts, are found in

460 Mesoproterozoic rocks (Runnegar, 1991; Knoll et al., 2006). The presence of these
461 organisms long before the Cryogenian implies that any physiological oxygen threshold
462 for the body plans that characterized the earliest (diploblastic) period of early animal
463 evolution must have been surpassed far prior to the origin of animals themselves.

464

465 5.2.2 *Bilaterian metazoans- theoretical lower oxygen limits*

466 In contrast to diploblasts, which have sheets of cells separated by inert material,
467 bilaterian (triploblastic) organisms have metabolically-active cells in three-dimensions
468 (Knoll, 2011). Body size (and the ability of the organism to exist at a given oxygen
469 concentration) is consequently limited by the ability to maintain functional internal
470 oxygen levels, either through pure diffusion or through a blood vascular system (BVS).
471 The implications of this constraint under hypothetical Precambrian oxygen levels have
472 been extensively discussed (e.g. Raff and Raff, 1970; Runnegar, 1982a,b; 1991; Catling
473 et al., 2005; Payne et al., 2010). Using a theoretical framework for the diffusion of
474 oxygen into an idealized animal (Alexander, 1971), these studies have demonstrated that
475 low oxygen levels will restrict bilaterians to small, thin body plans. What has not been
476 asked in these theoretical calculations is what oxygen levels will prohibit the existence of
477 bilaterian body plans.

478 Superficially, this question would seem to hinge on the nature of the last common
479 ancestor of bilaterians (consider Carroll et al., 2001, versus Erwin and Davidson, 2002),
480 specifically whether this ancestor was a complex, coelomate organism with a heart and
481 BVS, or a much simpler organism that transported oxygen through pure diffusion.
482 However, as noted by Budd and Jensen (2000), due to structural size requirements,

483 notably the physical space required to fit a functional BVS, this transport system is not
484 present in modern organisms less than ~3mm in size. Following the framework of
485 Alexander (1971; see Supplementary Information for details), we estimate that the most
486 likely minimal oxygen requirement for a 3 mm-long x 67 μm -wide worm with a
487 circulatory system, such as an annelid, is ~0.14% PAL (Fig. 9). The most likely minimal
488 oxygen requirements for a 600 x 25 μm diameter worm limited by pure diffusion, such as
489 a nematode, is ~0.36% PAL (Fig. 9)—note that these values are with respect to ambient
490 dissolved oxygen concentrations and do not consider temperature or salinity effects on
491 the dissolution of oxygen in water. The estimated oxygen requirements for these two
492 hypothetical ancestors differ slightly, but their broad similarity and the overlap in
493 sensitivity analyses (Fig. 9) suggests that pure diffusion and a BVS likely represent
494 optimal designs below and above this size threshold.

495 Although there are uncertainties in the optimal values for the parameters in the
496 equations governing oxygen requirements (see Supplementary Information), three facts
497 suggest the values described above represent conservative estimates for the minimum
498 oxygen concentrations necessary to sustain bilaterians. First, for the bilaterian limited by
499 pure diffusion, a sensitivity analysis (Fig. 9 and Supplementary Table 7) demonstrates
500 that one of the most important terms at very low oxygen levels is the minimum cellular
501 oxygen concentration. This will be a small, but non-zero, number (Raff and Raff, 1970).
502 Raff and Raff (1970) used a value of $1/10^{\text{th}}$ the shared K_m of yeast and mammalian
503 cytochrome oxidase. Here, rather than adopting an arbitrary but likely more accurate
504 fractional value, we use the shared yeast-mammal K_m (Chance, 1957) as our ‘most
505 likely’ value for this parameter (note that investigated invertebrate cytochrome oxidases

506 have a similar value (e.g. Gnaiger et al., 2000)). This ensures that the most important
507 parameter in the model is an over-estimate. Second, for the hypothetical ancestor with a
508 circulatory system, we assumed the organism did not have respiratory pigments.
509 Although the homology of metazoan respiratory pigments is unclear (Terwilliger, 1998),
510 their presence in this hypothetical last common bilaterian ancestor would greatly increase
511 diffusion rates. Thus, our assumption that respiratory pigments were absent again results
512 in a conservative estimate. Finally, and most importantly, these theoretical calculations
513 assume a perfectly tubular organism (Alexander, 1971). Such an organism does not exist,
514 as real animals have body wall rugosities, gills, and other structures that dramatically
515 increase diffusive surface area with respect to volume; even the gut is a gas-exchange
516 organ. Consequently, these ‘most likely’ values and the sensitivity analyses are not
517 intended to yield a precise number. Rather, these models provide an indication of the
518 lower bound of oxygen levels necessary to preclude the bilaterian body plan from
519 Proterozoic oceans. No matter the complexity of the last common ancestor of bilaterians,
520 theoretical modeling suggests the bilaterian body plan was unlikely to have been
521 prohibited unless O₂ levels were < 0.4% PAL.

522

523 5.2.3 *Bilaterian metazoans- empirical lower oxygen limits*

524 These theoretical calculations can be tested with empirical observations of the
525 oxygen limits of bilaterians in modern oxygen-minimum zones (OMZs). Unlike the biota
526 on shelves or in regions of anthropogenic eutrophication that show deleterious oxygen
527 responses at relatively minor oxygen depletions (Diaz and Rosenberg, 1995; Levin et al.,
528 2009), OMZs have experienced geologically long-lasting dysoxic- to anoxic conditions,

529 allowing the fauna to adapt to these levels and providing an excellent analogue for
530 Precambrian oceans with persistently low oxygen levels. It should be noted that
531 organisms in modern OMZs have likely secondarily adapted to these environments rather
532 than originating in them. Thus, the type of adaptations allowing organisms to inhabit
533 these environment must be considered. For example, organisms with extreme metabolic
534 adaptations, such as amitochondriate loriciferans living in an euxinic Mediterranean basin
535 (Danovaro et al., 2010), cannot inform us about Precambrian animal evolution, as the
536 transformation of the mitochondria into a hydrogenosome was certainly not a primitive
537 feature. Most of the adaptations allowing bilaterians to inhabit modern low-oxygen
538 environments, though, appear to lie in their very small, thin body plans (with high
539 surface-area to volume ratios for increased diffusion) and enlarged respiratory organs
540 (Levin, 2003; Gooday et al., 2010; Jeffreys et al., 2012; Lamont and Gage, 2000; Neira et
541 al., 2001)—that is, with morphological adaptations that would have been possible, and
542 perhaps likely, in early bilaterians. Consequently these animals can provide a useful
543 analogue for Precambrian animal life in low-oxygen conditions.

544 In using OMZs as Precambrian analogues, it has long been recognized that the
545 faunas are characterized by such small, thin, body plans (Rhoads and Morse, 1971). What
546 has emerged in the four decades of oceanographic research since Rhoads and Morse's
547 seminal paper is just how little oxygen is actually required by bilaterian animals. It is now
548 clear that non-chemosymbiotic benthic macrofaunal (retained on 0.3 mm sieves)
549 bilaterians can and do live in Rhoads and Morse's 'azoic' zone of <0.10 mL/L oxygen¹

¹ A difficulty in interdisciplinary research on the biological effects of differing oxygen levels is the use of different units by different research communities (Hofmann et al.,

550 (Levin, 2003; Gooday et al., 2010; Levin et al., 2000; Palma et al., 2005; Zettler et al.,
551 2009; Levin et al., 1991; Levin et al., 2002; Ingole et al., 2010), often with densities of
552 hundreds to thousands of animals per square meter. Bilaterian faunas can even be found
553 as low as 0.02 ml/L O₂, equivalent to ~0.3% of modern surface ocean levels (assuming a
554 normal surface ocean concentration of ~6 mL/L) in the OMZ off Chile (Palma et al.,
555 2005), Peru (Levin et al., 2002) and the Bay of Bengal (R. Akkur, pers. comm.). The
556 exact oxygen concentrations actually required to exclude bilaterians are likely even
557 lower, these oxygen measurements are determined from O₂ sensors or seawater samples
558 from CTD casts collected several meters (~5 m) above the seafloor. The oxygen levels at
559 which bilaterians are recorded (namely 0.02 mL/L) approach the detection limit of the
560 Winkler titration technique (Paulmier et al., 2006), and CTD cast values generally over-
561 estimate in-situ benthic conditions (Breur et al., 2009). Thus, both theoretical calculations
562 and empirical observations in modern OMZs suggest the presence of bilaterians would
563 not have been limited unless atmospheric O₂ was considerably less than 1% PAL, and
564 likely less than 0.4% PAL.

565

566 7. Conclusions

567 Geochemical transects of the ~800 Ma Fifteenmile Group in the Ogilvie
568 Mountains document shallow-water facies characterized by low FeHR/FeT, a lack of
569 enrichment in redox-sensitive trace elements, and relatively heavy and variable pyrite
570 sulfur isotope values. Deeper-water facies (those deposited below storm wave base) are
571 characterized by FeHR/FeT > 0.38, enrichment in redox-sensitive trace elements, and

2011). For consistency here we report oxygen levels as in the benthic ecology literature (mL/L). For reference 0.01 mL/L \approx 0.44 μ mol/kg \approx 0.014 mg/L \approx 0.4 atm.

572 more depleted pyrite sulfur isotope values. Overall, this points towards an oxygenated
573 surface layer, down to storm wave base, overlying a generally anoxic deep basin.
574 Fluctuations between euxinic and ferruginous conditions sub-storm wave base appear to
575 have been controlled by variations in organic carbon loading. As proxies like iron
576 speciation and redox-sensitive trace elements provide evidence of local environments,
577 more geochemical studies from other basins are necessary to begin building the global
578 picture of redox heterogeneity. Further, the development of quantitative global redox
579 tracers and better modeling are needed to place tighter constraints on the history of
580 oxygen on Earth. Nonetheless, the Fifteenmile Group redox structure is comparable to
581 that of a well-characterized Mesoproterozoic basin (Shen et al., 2003)—albeit with more
582 evidence for ferruginous conditions-- and both basins are consistent with broad estimates
583 for atmospheric oxygen levels between 1 and 40% PAL (Kump, 2008).

584 Comparing these likely O₂ levels with the estimated physiological requirements of
585 early animals suggests that sufficient atmospheric oxygen, even for mobile bilaterians,
586 was present well in advance of the origin of animals. Unless early Neoproterozoic oxygen
587 levels were substantially < 1% PAL, and likely < 0.4% PAL, atmospheric oxygen levels
588 would not have prohibited the sponge, eumetazoan and bilaterian body plans. This
589 conclusion does not imply that animals necessarily lived in the Fifteenmile basin, but
590 rather that global O₂ levels were likely adequate for the presence of animals. Notably, this
591 does not negate the possibility of an oxygenation event around the Sturtian glaciation
592 (Planavsky et al., 2010; Frei et al., 2009), or the use of oxygenated ‘oases’ beneath
593 photosynthetic mats by the earliest trace makers in the geological record (Gingras et al.,

594 2011), but it does suggest that such conditions were not necessary for the origin of either
595 animals or bilaterians.

596 It is important to remember, though, that while low Precambrian oxygen levels
597 would not have prohibited animals, including bilaterians, the environmental milieu would
598 still have exerted a strong effect on life. Most importantly, low oxygen certainly would
599 have constrained these organisms to very small and thin body plans with little metabolic
600 scope (Raff and Raff, 1970; Runnegar, 1982a, b; Payne et al., 2010). Faunas in modern
601 low-O₂ OMZ analogues have very small body sizes, reduced diversity, and simple food
602 webs (Levin, 2003; Gooday et al., 2009; Sperling et al., in review). In other words,
603 although all available data suggests bilaterians *can* live down to 1% PAL or less, the
604 fauna would be limited to a select few—those organisms that were a couple millimeters
605 in length and had low-energy lifestyles. Thus, while no oxygenation event need be
606 invoked to explain the origin of animals or bilaterians themselves, the hypothesized end-
607 Neoproterozoic oxygenation event (the timing and magnitude of which remains
608 debated—Kah and Bartley, 2011; Och and Shields-Zhou, 2012) may still have played a
609 role in the later Ediacaran diversification of macroscopic animals and the Cambrian
610 ‘Explosion’ (e.g. Runnegar, 1982a; Rhoads and Morse, 1971; Knoll and Carroll, 1999).
611 Although Cambrian diversification was certainly multifaceted (Erwin et al., 2011), a
612 latest Proterozoic increase in oxygen levels could have allowed for an increase in both
613 size and metabolic scope, including potentially the advent of predation, a metabolically-
614 costly feeding strategy.

615

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628

629 **FIGURE CAPTIONS**

630 **Fig. 1-** Location map of the Coal Creek and Tatonduk inliers, Yukon Territory, Canada,
631 with stars marking the location of the inliers.

632

633 **Fig. 2-** Geological map of the Coal Creek inlier, Ogilvie Mountains, Yukon Territory,
634 showing sections (in red) studied in this paper. The stratigraphic framework for basin
635 transects A – A' and B – B' are found in Fig. 3 and 4. The units studied as part of this
636 paper are the informal Gibben formation, Chandindu formation and Reefal Assemblage
637 of the Fifteenmile Group. Geological mapping by Macdonald et al. (2012).

638

639 **Fig. 3-** Stratigraphic framework and iron speciation chemistry for transect A – A' at Mine
640 Camp, Mt. Harper and Mt. Gibben, Coal Creek inlier. Iron speciation chemistry
641 (specifically the ratio of highly reactive iron (FeHR) to total iron (FeT)) from fine-
642 grained siliciclastic rocks is plotted against the stratigraphic columns. Vertical line on
643 iron speciation plots denotes a ratio of 0.38, with samples having higher ratios considered
644 to have been deposited under an anoxic water column, and samples with lower ratios
645 likely to have been deposited under an oxic water column. Complete redox proxy data for
646 individual sections is found in Fig. 6 (Mt. Harper), Supplemental Fig. 1 (Mine Camp),
647 Supplemental Fig. 2 (East Harper) and Supplemental Fig. 3 (Mt. Gibben). *m-* mud; *si-*
648 silt; *ms-* medium sand; *cs-* coarse sand; *cg-* conglomerate. *Gib.* = Gibben formation.

649

650 **Fig. 4-** Stratigraphic framework and iron speciation chemistry for transect B – B' at
651 Reefer Camp, Coal Creek inlier. Sections are located 5 km apart and record a transition in
652 the Reefal Assemblage from a stromatolite reef complex into a deeper-water shale basin.
653 Iron speciation chemistry (specifically the ratio of highly reactive iron (FeHR) to total
654 iron (FeT)) from fine-grained siliciclastic rocks is plotted against the stratigraphic
655 columns. Vertical line on iron speciation plots denotes a ratio of 0.38, with samples
656 having higher ratios considered to have been deposited under an anoxic water column,
657 and samples with lower ratios likely to have been deposited under an oxic water column.
658 Complete redox proxy data for individual sections at Reefer Camp is found in Fig. 5.
659 Legend for stratigraphic columns and sediment type abbreviations as in Fig. 3.
660 Abbreviations: *W.* = Wernecke Supergroup, *Gib.* = Gibben formation, *Cha.* = Chandindu
661 formation.

662

663 **Fig. 5-** Redox proxy data from sections B' (Fig. 5A) and B (Fig. 5B) at Reefer Camp,
664 Coal Creek inlier. From left to right, proxy data plotted and their respective relevant
665 baseline data denoted by vertical red lines are: highly reactive to total iron (FeHR/FeT ;
666 0.38), pyrite iron to highly reactive iron (FeP/FeHR ; 0.80), total iron to total aluminum
667 (FeT/Al ; 0.50), molybdenum (2.6 ppm), vanadium (130 ppm), pyrite sulfur isotope
668 values (0 ‰), and weight percent total organic carbon. Relative base level curve from
669 sequence stratigraphic study of Macdonald et al. (2012). The far right column is a
670 subjective estimate of water column redox state based on the multi-proxy data, and is
671 meant to represent general trends rather than an estimate for every point. Euxinia(?)
672 denotes uncertainty regarding whether these samples represent deposition under a truly
673 euxinic water column or a ferruginous water column with sulfide production at or near
674 the sediment-water interface (see text). Legend for stratigraphic column and formation
675 name abbreviations as in Fig. 3 and 4.

676

677 **Fig. 6-** Redox proxy data from the Mt. Harper section, Coal Creek inlier. This figure is a
678 composite section from the Mt. Harper area (Fig. 2). Redox proxy data and their relevant
679 baseline values (marked by vertical red lines) as in Figure 5. Relative base level curve
680 from sequence stratigraphic study of Macdonald et al. (2012). The far right column is a
681 subjective estimate of water column redox state based on the multi-proxy data, and is
682 meant to represent general trends rather than an estimate for every point. Euxinia(?)
683 denotes uncertainty regarding whether these samples represent deposition under a truly
684 euxinic water column or a ferruginous water column with sulfide production at or near

685 the sediment-water interface (see text). Legend for stratigraphic column and formation
686 name abbreviations as in Fig. 3 and 4.

687

688 **Fig. 7-** Redox proxy data from the Mt. Slipper section, Tatonduk inlier. Redox proxy data
689 and their relevant baseline values (marked by vertical red lines) as in Figure 5. Relative
690 base level curve from sequence stratigraphic study of Macdonald et al. (2012). The far
691 right column is a subjective estimate of water column redox state based on the multi-
692 proxy data, and is meant to represent general trends rather than an estimate for every
693 point. Euxinia(?) denotes uncertainty regarding whether these samples represent
694 deposition under a truly euxinic water column or a ferruginous water column with sulfide
695 production at or near the sediment-water interface (see text). Legend for stratigraphic
696 column and formation name abbreviations as in Fig. 3 and 4. Scale microfossils described
697 by Cohen et al. (2011) are found at the top of the Reefal Assemblage at this locality.

698

699 **Fig. 8-** Boxplot analysis of total organic carbon weight percentages for samples
700 determined likely to have been deposited under a ferruginous and euxinic water column.
701 Bottom-water redox state for each sample was estimated using a multi-proxy framework
702 including iron speciation data, redox-sensitive trace elements and pyrite sulfur isotope
703 values (see text for details). Samples designated as euxinic may represent sulfide
704 accumulation at the sediment-water interface rather than true water-column euxinia (see
705 text). The box represents the 25th and 75th percentiles, the thick horizontal line represents
706 the median, and the whiskers represent minimum and maximum values. An extreme

707 outlier in the ferruginous set (E1002- 470.4; 9.87 wt %) was included in the boxplot
708 calculations but not graphed.

709

710 **Fig. 9-** Theoretical minimum oxygen requirements for the last common ancestor (LCA)
711 of bilaterians, following the equations governing the diffusion of oxygen into an
712 organism from Alexander (1971) and modified by Payne et al. (2010). Estimates were
713 made for two potential body plans characterizing the bilaterian LCA, a 600 μm long
714 worm limited by pure diffusion, and a 3-mm long worm with a circulatory system. ‘Most
715 likely’ values represent values estimated from optimal values for all parameters (see
716 Supplemental Information). Minimum and maximum values were derived from the
717 literature for each parameter and global minimum and maximum values were estimated.
718 A sensitivity analysis was then conducted for each parameter by varying that parameter
719 between minimum and maximum values while keeping all other parameters at their ‘most
720 likely’ values. All estimates for oxygen requirements are far less than the 1% of Present
721 Atmospheric Levels indicated by canonical views of atmospheric oxygen levels in the
722 Proterozoic (Kump, 2008). Lowest row for extant mobile bilaterians shows the current
723 lower oxygen limit at which bilaterians are found in the modern ocean ($0.02 \text{ mL/L O}_2 \approx$
724 0.33% of modern surface ocean levels assuming a normal surface ocean level of 6 mL/L).
725 Bilaterians are found at these levels off the coasts of Peru, Chile, and in the Bay of
726 Bengal. This oxygen level may represent an overestimate due to the methodology used to
727 measure oxygen in most benthic ecology studies (Breur et al., 2009 and see text).

728

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Highlights for Sperling *et al.*

- 1) We present the first early Neoproterozoic basin redox transect
- 2) Redox proxy data are consistent with quantitative constraints on Proterozoic O₂
- 3) Inferred Proterozoic oxygen levels would not prohibit the presence of animals

Figure 1

[Click here to download Figure: Figure 1 Locality map.pdf](#)



Figure 2
[Click to download Figure: Figure 2 Coal_Creek_Erik.19910500.W](#)

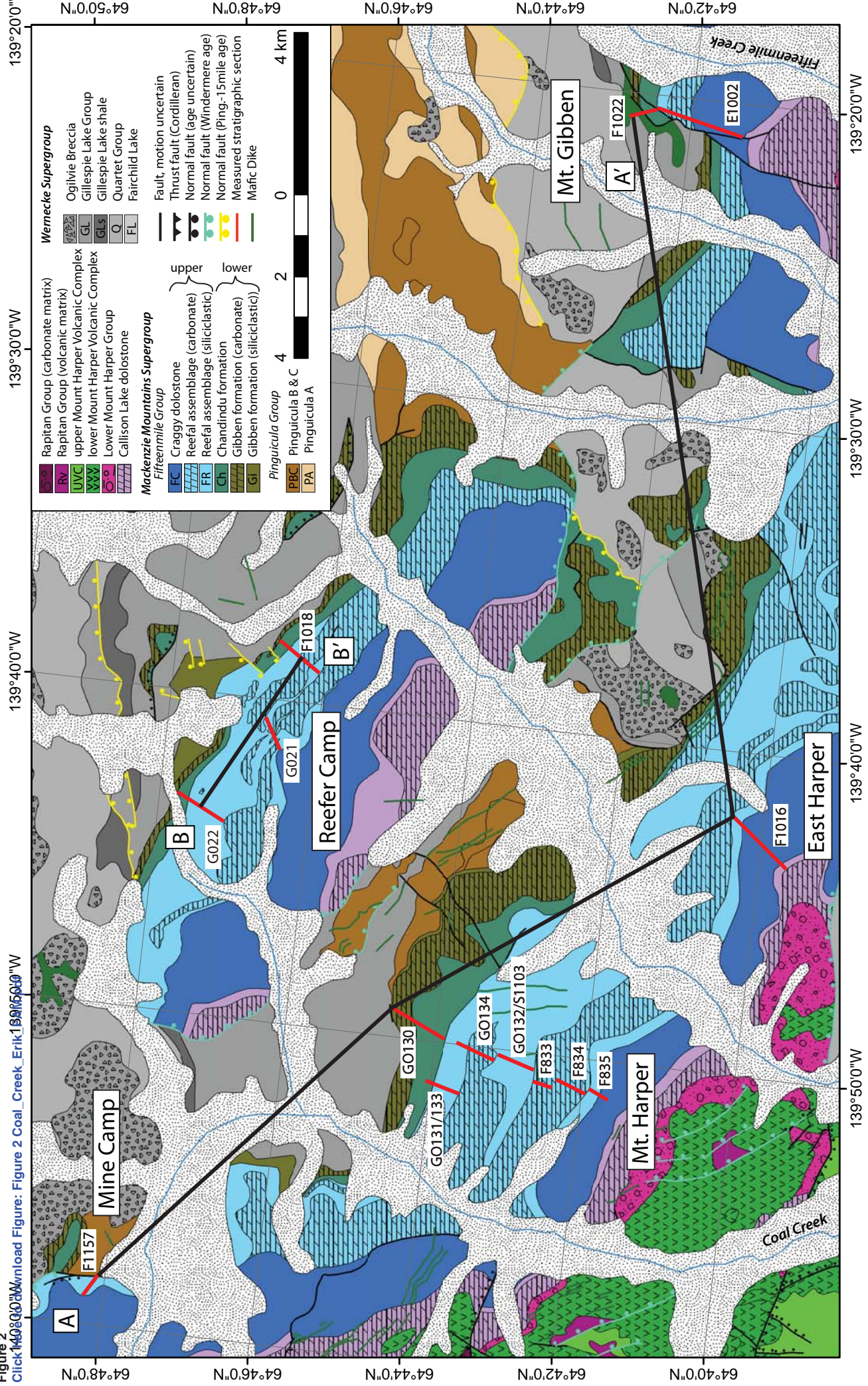
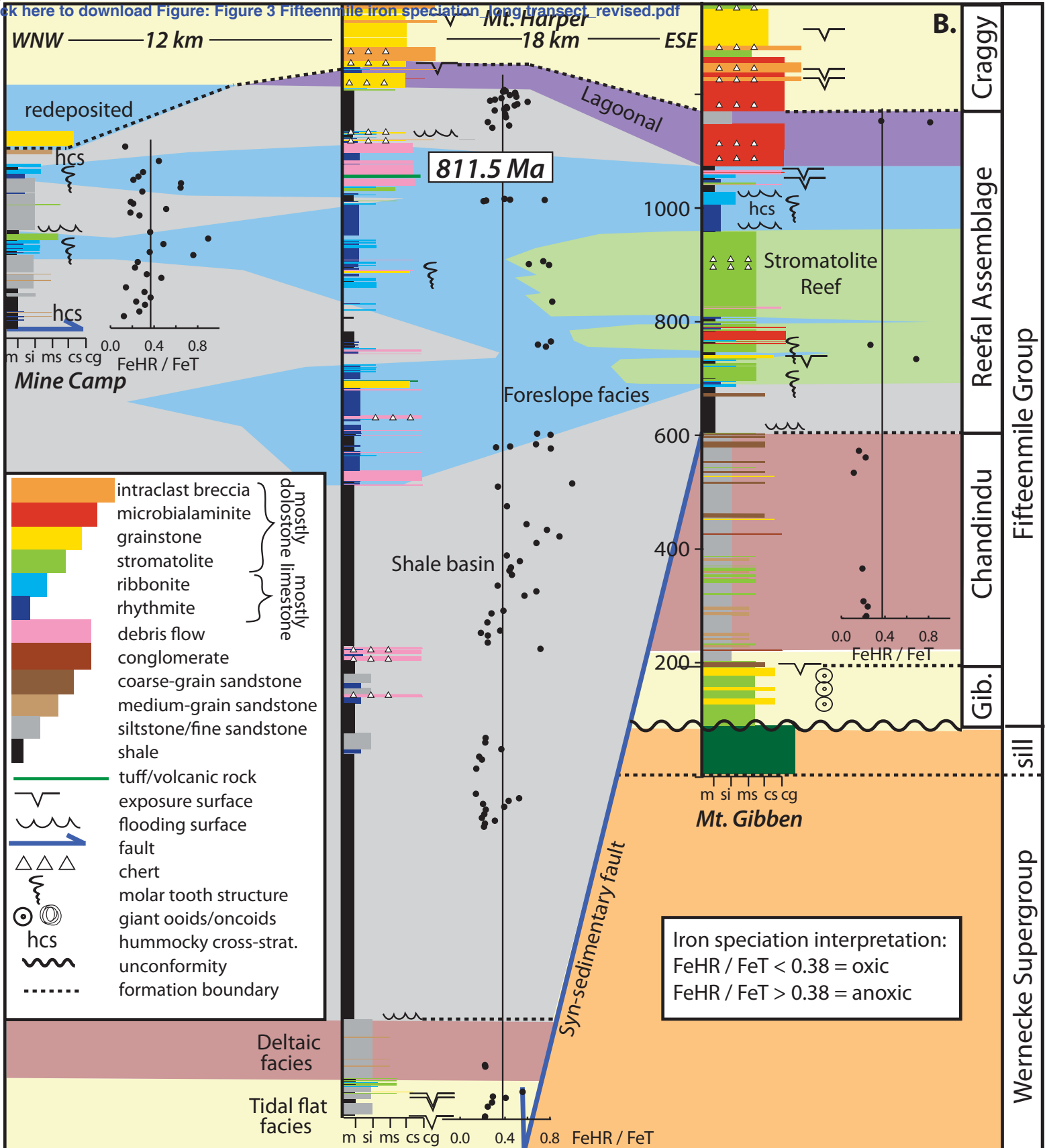


Figure 3

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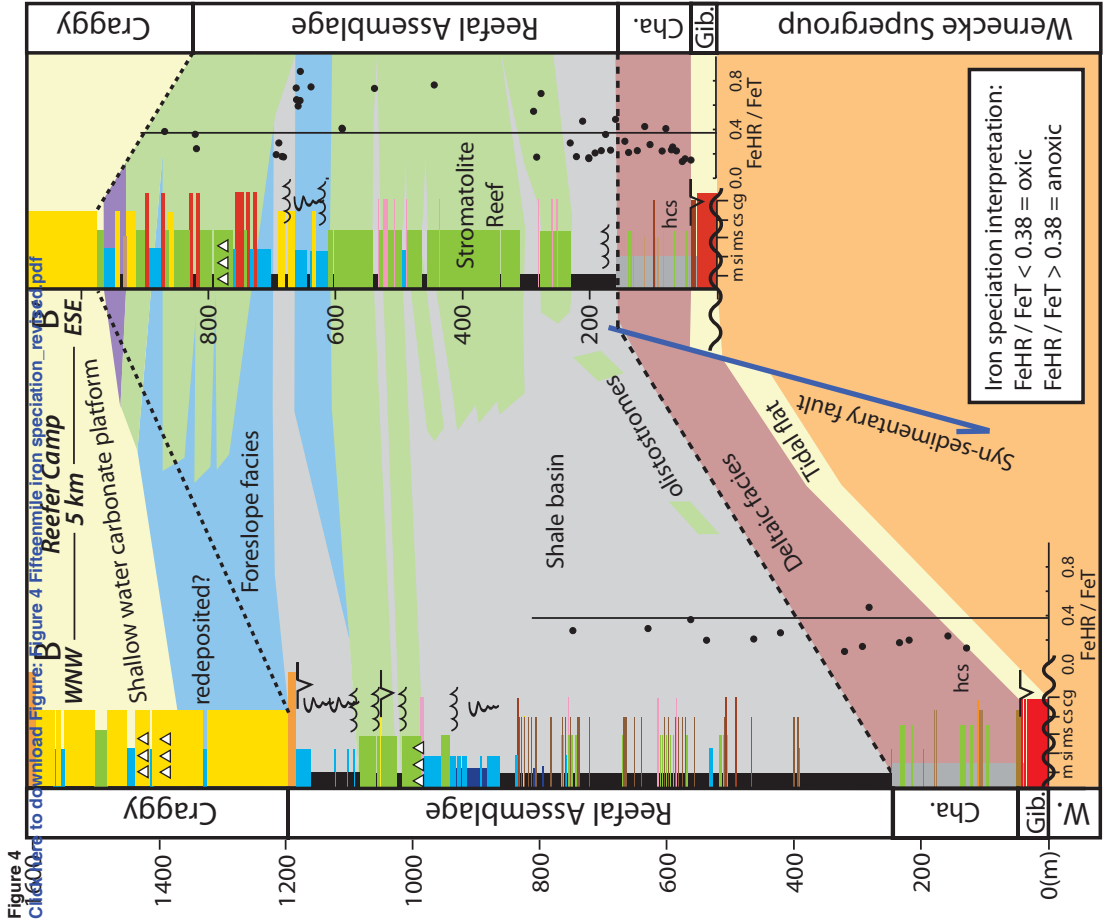


Figure 5

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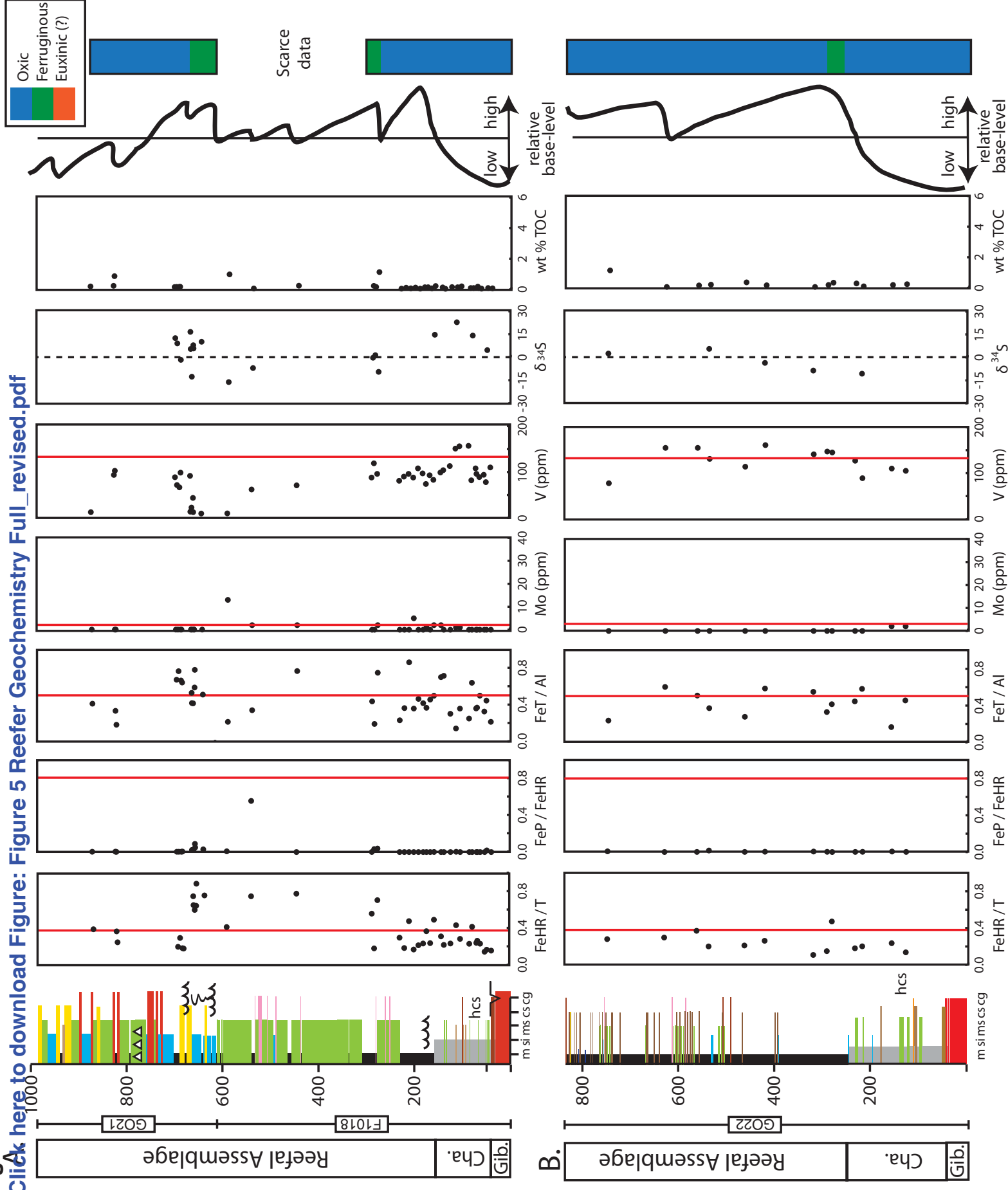


Figure 6

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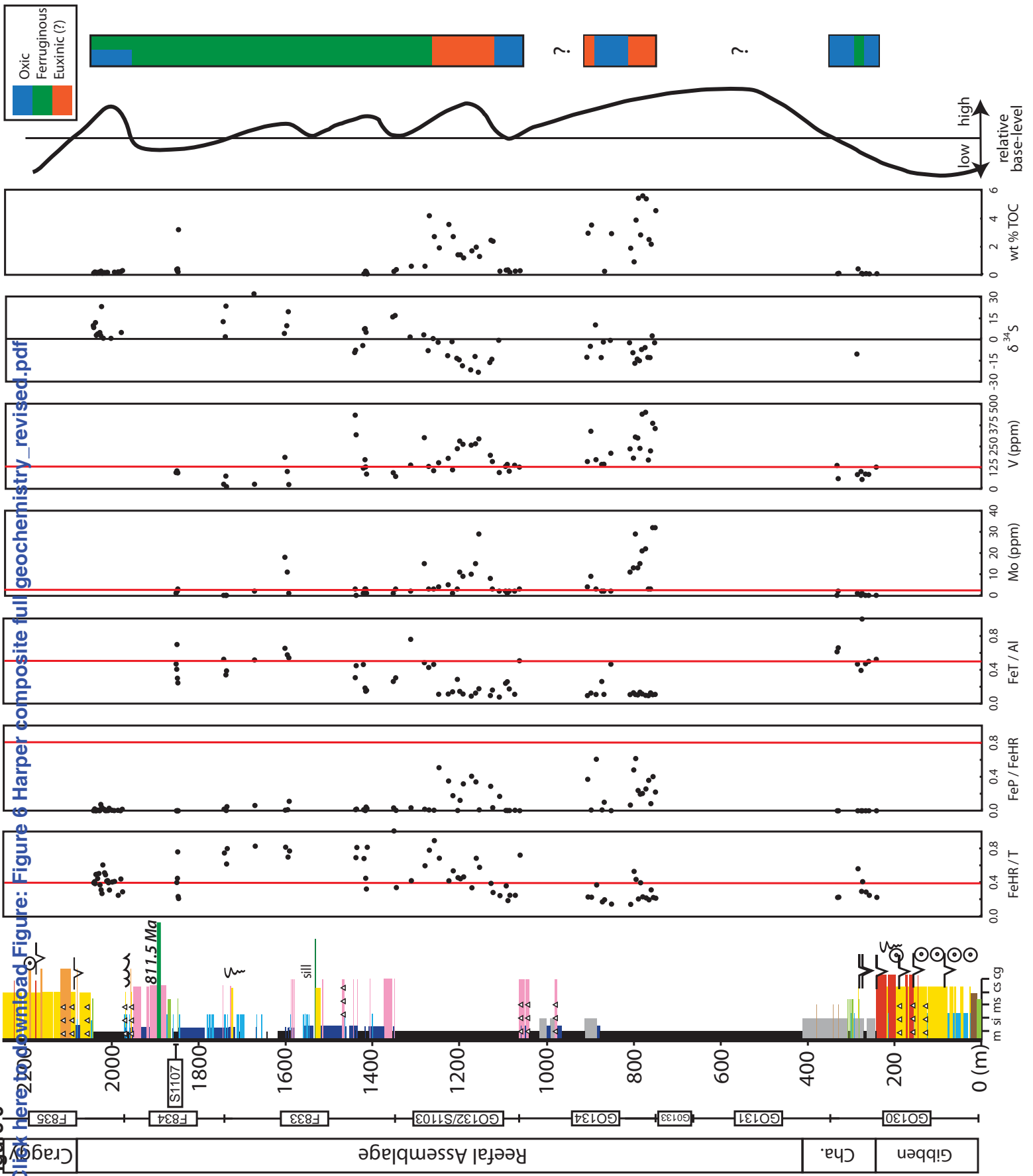


Figure 7

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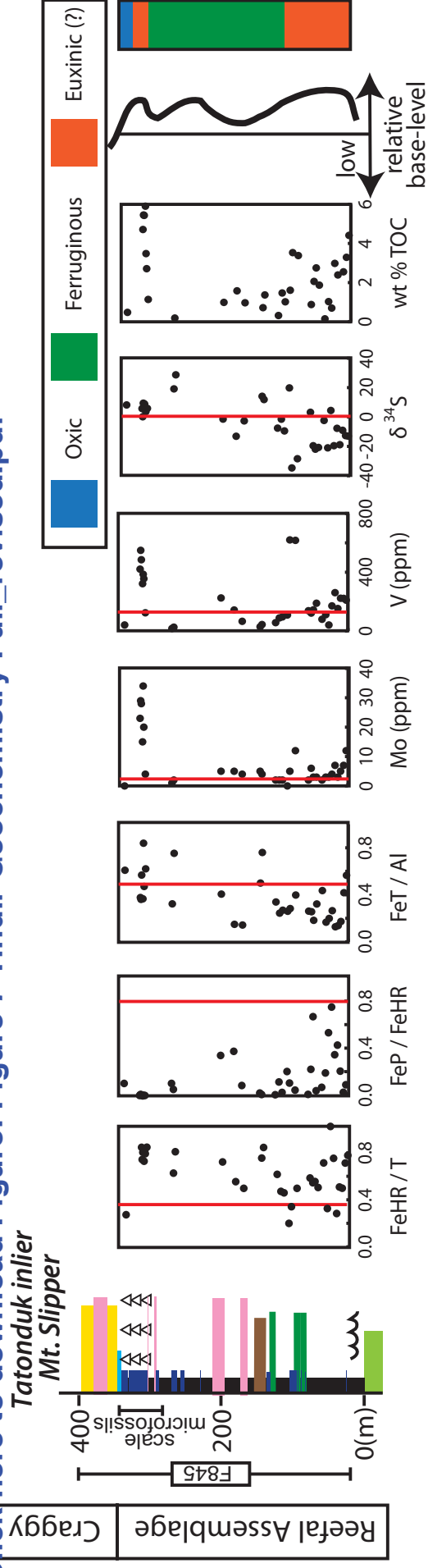


Figure 8
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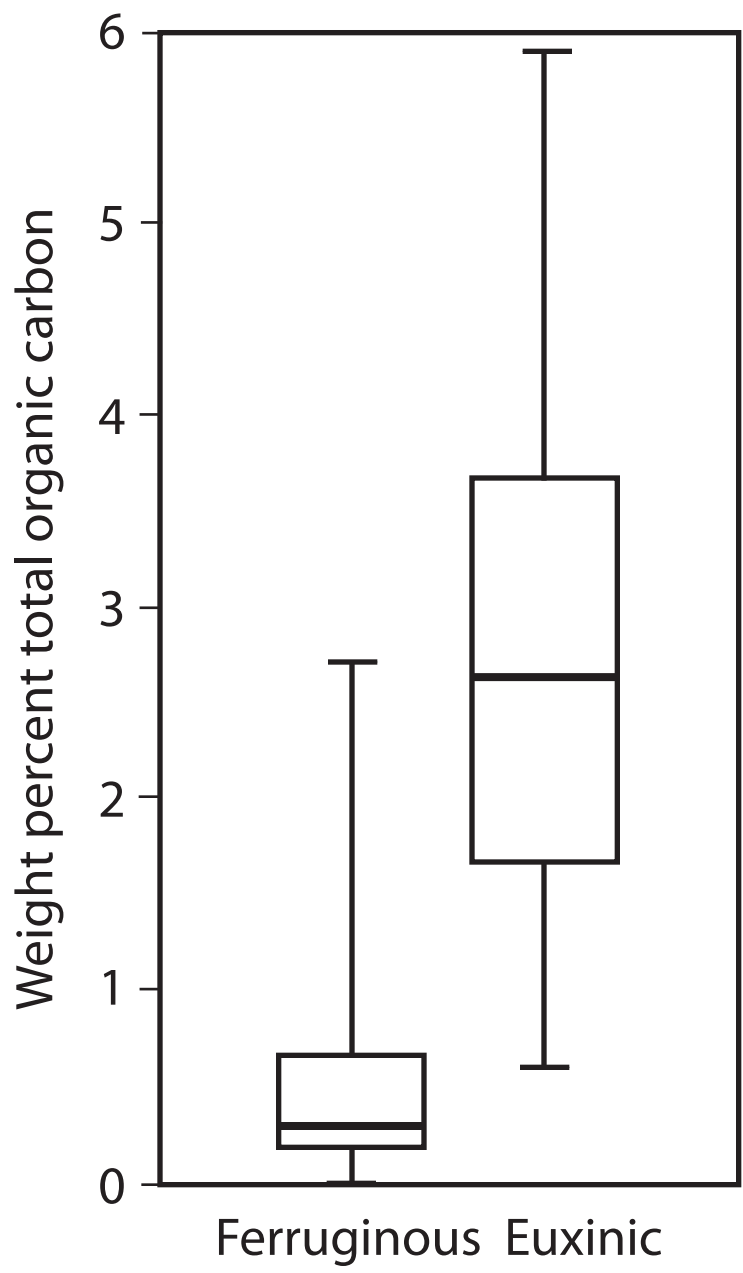


Figure 9

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