Chromium evidence for protracted oxygenation during the Paleooproterozoic


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Abstract

It has commonly been proposed that the development of complex life is tied to increases in atmospheric oxygenation. However, there is a conspicuous gap in time between the oxygenation of the atmosphere 2.4 billion years ago (Ga) and the first widely-accepted fossil evidence for complex eukaryotic cells <1.7 Ga. At present the gap could either represent poor sampling, poor preservation, and/or difficulties in recognizing early eukaryote fossils, or it could be real and the evolution of complex cells was delayed due to relatively low and/or variable O₂ levels in the Paleooproterozoic. To assess the extent and stability of Paleooproterozoic O₂ levels, we measured chromium-based oxygen proxies in a >2400-m core from the Onega Basin (NW-Russia), deposited ~2.1–2.0 billion years ago—a few hundred million years prior to the oldest definitive fossil evidence for eukaryotes. Fractionated chromium isotopes are documented throughout the section.
(max. 1.63±0.10‰, $\delta^{53}$Cr), suggesting a long interval (possibly >100 million years) during which oxygen levels were higher and more stable than in the billion years before or after. This suggests that, if it is the case that complex cells did not evolve until after 1.7 Ga, then this delay was not due to O$_2$-limitation. Instead, it could reflect other limiting factors—ecological or environmental—or could indicate that it simply takes a long time—more than the tens to more than 100 million years recorded in Onega Basin sediments—for such biological innovations to evolve.

**Keywords**

Cr isotopes, oxygen overshoot, isotope geochemistry, eukaryotes

**Introduction**

Free O$_2$ in the atmosphere, the result of oxygenic photosynthesis, shapes ecology on the global scale. Understanding Earth’s oxygenation is therefore paramount to understanding the evolution and history of life. Given that O$_2$-dependent metabolisms are highly energetic and that atmospheric O$_2$ concentrations ($p$O$_2$) correlate with increased nutrient richness (Reinhard et al., 2020), high oxygen levels are often, though controversially, thought to be necessary for ‘complex’ (e.g., eukaryotic) life (Cole et al., 2020). A distinct, but related question is whether high $p$O$_2$ alone is sufficient for this purpose—or in other words, was oxygenation the trigger for shifts towards more complex life. In this view, removing the O$_2$-limitation will quickly lead to the evolution of traits thought to underpin complexity, including eukaryote-grade cells, large size, and multicellularity (e.g., Payne et al., 2009; Zhang et al., 2018), implying that $p$O$_2$ above all else modulated the evolution of complex life on Earth (cf. Catling et al., 2005).

One challenge to this view is the mismatch between the appearance of unambiguous eukaryotic fossils in rocks younger than ~1.7 Ga (e.g., Agić et al., 2015, 2017; Adam et al., 2017; Javaux et al., 2001; Javaux and Knoll, 2017; Miao et al., 2019), and the initial oxygenation of the atmosphere 700 million years earlier (Farquhar et al., 2000; Warke et al., 2020). While this could be interpreted to suggest that O$_2$ levels were not the primary modulator of eukaryote emergence, others have hypothesized that O$_2$ levels simply were not stable or high enough to support substantial eukaryotic communities (Javaux and Lepot, 2018) before the latest Paleoproterozoic (Zhang et al., 2018).

A way to test this hypothesis is to examine the record of redox changes across an extended period of time in the early Proterozoic Eon prior to the first unambiguous eukaryotic fossils. The
early Proterozoic witnessed the disappearance of mass-independent sulfur isotope fractionations by ~2.5 to 2.32 billion years ago (Ga) (Farquhar et al., 2000; Warke et al., 2020), signifying the permanent oxygenation of the atmosphere. This change is colloquially known as the Great Oxidation Event (GOE) (Holland, 2002). The Rhyacian Period (2.3–2.05 Ga) is notable for high positive carbon isotope fractionations in marine carbonates (δ¹³C_carb) (Karhu and Holland, 1996). As δ³⁴S_carb has been closely coupled to the burial of biomass and the release of O₂ accumulated in the atmosphere, this led Bekker and Holland (2012) to hypothesize that atmospheric O₂ levels may have reached 50% of the modern in this period. This is in accordance with signals of highly elevated seawater sulfate concentrations, including the deposition of massive evaporites (e.g., Blättler et al., 2018) and sedimentary sulfur isotope signatures (Planavsky et al., 2012), that point to a robust marine sulfate pool that is only stable in oxygenated conditions, but pO₂ remains difficult to robustly constrain.

A more recent approach has been to use the concentrations and isotopes of redox-sensitive trace metals to infer the amount of O₂ required to mobilize those elements from the crust to the oceans where they are subsequently incorporated into sedimentary rocks. However, this approach has led to conflicting results. For instance, Mänd et al. (2020) used the concentrations of molybdenum (Mo), rhenium (Re) and uranium (U), as well as U isotopes, in marine sediments to demonstrate that highly oxidized conditions continued up to around 2 Ga. Yet, the same Orosirian Period (2.05 to 1.8 Ga), and more generally the mid-Proterozoic (2 to 0.8 Ga), is generally thought to be characterized by lower levels of oxygen, as evidenced by muted sedimentary U concentrations (Partin et al., 2013), absent cerium (Ce) anomalies (Bellefroid et al., 2018), and reduced Mo isotope fractionations (Ossa Ossa et al., 2018, among others). In essence, different proxies present varying views of Earth’s oxygenation due to their different spatial scales and sensitivities to pO₂ that, along with the fragmentary nature of studied sedimentary successions, hinder attempts to link oxygenation to preserved shifts in fossil complexity.

A continuous record of chromium isotope fractionations (δ⁵³Cr) presents the opportunity for probing the stability of pO₂ in Earth’s atmosphere, as substantial δ⁵³Cr fractionations are inherited from continental weathering environments in direct contact with atmospheric oxygen (Frei et al., 2009; Wei et al., 2020), although see Daye et al. (2019), Liu et al. (2020), and Lyons et al. (2020). Crucially, although δ⁵³Cr fractionations record mainly oxygen-poor weathering settings in the middle Proterozoic (Cole et al., 2016; Colwyn et al., 2019; Planavsky et al., 2014), with potential episodes of oxygenation (Canfield et al., 2018; Gilleaudeau et al., 2016; Wei et al., 2021), there is intriguingly no Cr isotope evidence for weathering under oxygen-rich environments in the middle
Paleoproterozoic—when it has been commonly proposed as a high-oxygen interval based on other proxies (Bekker and Holland, 2012; Blättler et al., 2018; Planavsky et al., 2012).

Amongst the best locations to establish a long, continuous Paleoproterozoic Cr record is the Onega Basin in the Karelian Republic, Russia (Melezhik et al., 2013). This is a >3-km-thick Paleoproterozoic succession of evaporites, carbonates, and siliciclastics, including organic-rich mudstones, intercalated with mafic igneous rocks (Figure 1). In this study, we present new bulk-rock $\delta^{53}$Cr data from a ~2400-m-thick volcano-sedimentary drill core from the Onega Basin, obtained from the Onega Parametric Hole (OPH; drilled in 2008–2009 in the southern Onega Basin at 62.1559 N, 34.4073 E; Figure 1). The drill core intersects ~800 m of the Tulomozero Formation, ~1500 m of the Zaonega Formation, and ~500 m of the Suisari Formation. We find that substantially fractionated sedimentary $\delta^{53}$Cr values persist throughout the Onega Basin succession, across several facies transitions. This suggests that the Onega Basin records a protracted time period—possibly more than 100 million years (Myrs)—that was host to a fundamentally stable and oxygen-rich ocean-atmosphere system providing a platform to revisit the links between oxygen and the development of complex life.

The Onega Basin

The Onega Basin succession begins with the deposition of coarse-grained siliciclastics and lava flows onto a basement of Archean granites and gneisses. These are overlain by the ~800-m-thick Tulomozero Formation (~2100–2050 Ma; see below for age references) made up of evaporites and dolomites. The Zaonega Formation (~2050–2000 Ma) follows as a ~1500-m-thick volcano-sedimentary package consisting of dolostones, siliciclastic turbidites, and mudstones intercalated with mafic igneous rocks. On top of that lies the ~500-m-thick Suisari Formation (~2000–1970 Ma) comprising mainly tuffs and mafic lavas/sills. The section is capped by fluvial-lacustrine siliciclastics of the Kondopoga Formation. Following deposition, the Onega Basin underwent greenschist-facies metamorphism during the ~1.89–1.79 Ga Svecofennian orogeny, which deformed the succession into a series of northwest–southeast trending folds (Melezhik et al., 2013, and references therein).

This study focuses on the Tulomozero, Zaonega, and Suisari formations that constitute the middle and upper part of the Onega Basin succession. There is significant lithofacies variation in the Tulomozero Formation—in the northern part of the basin, along with the western and eastern margins, the formation consists mainly of dolostone with varying amounts of magnesite and calcium sulphate pseudomorphs, and siliciclastic layers. By contrast, the formation in the southern
part of the basin, as recorded by the drill core of the Onega Parametric Hole, begins with a ~600-
meters-thick halite and anhydrite-magnesite unit that grades upwards into a dolomite dominated
unit. It is likely that in its present northern extent the formation was deposited in a low-energy
intertidal, sabkha or playa lake environment with fluvial influences, whereas the southern part of the
basin was more restricted and evaporitic, but still experienced periodic seawater influx that
sustained the accumulation of hundreds of meters of evaporites (Blättler et al., 2018; Melezhik et
al., 2013). Carbonates of the Tulomozero Formation are characterized by strongly positive $\delta^{13}$C_carb
values, reaching as high as 18‰, that potentially reflect local amplification of a globally $^{13}$C-
enriched bicarbonate pool (e.g., Melezhik et al., 1999). As such, it is one of the type sections for the
Lomagundi-Jatuli carbon isotope excursion (Karhu and Holland, 1996). The paragenesis of halite
and calcium sulfate, together with their S and Ca isotope composition, have been taken as evidence
of a large marine sulfate pool, constituting $>30\%$ of modern levels (Blättler et al., 2018).

The overlying Zaonega Formation was, in contrast, deposited in alternating shallow and deeper
water conditions, possibly in a rift basin developed on a highly active continental margin. Between
35–70% of the succession is composed of mafic sills and lavas emplaced into unconsolidated
sediments (Melezhik et al., 2013). The lowermost Zaonega Formation is dominated by mudstones
with common dolostone interbeds which transition into deeper-water mixed mudstones, dolostones,
and siliciclastic rhythmites further up section. The formation is notable for containing extremely
high organic carbon content (up to 70 wt.%), the earliest known oil field (Qu et al., 2012), highly
elevated redox-sensitive element concentrations (Mänd et al., 2020), and one of the earliest
sedimentary phosphorus enrichments (Lepland et al., 2014). Variable and often high pyrite S
isotope ratios have been interpreted as recording a diminished global marine sulfate pool or,
alternatively, anomalously high local sulfate demand by microorganisms (Paiste et al., 2020b, and
references therein). In the lower part of the Zaonega Formation, dolostones enriched in $^{13}$C record
the Lomagundi-Jatuli excursion (Melezhik et al., 2015), but the best-preserved carbonates of the
upper Zaonega Formation have normal-marine $\delta^{13}$C values (Kreitsmann et al., 2020, 2019).
Strongly negative $\delta^{13}$C values in carbonates and organic matter from the middle part of the Zaonega
Formation were earlier thought to reflect a global negative C isotope excursion but are now viewed
as artifacts of hydrothermal de-dolomitization (Kreitsmann et al., 2020, and references therein) and
the incorporation of methanotrophic biomass (Qu et al., 2012).

The deposition of the Zaonega Formation was followed by the Suisari Formation. The latter
signifies a slowdown of subsidence and the infilling of the basin with hundreds-of-meters-thick
subaqueous mafic-to-ultramafic lavas and sills interlayered with relatively thin, tuffaceous
mudstone beds (Melezhik et al., 2013).
The age of the Onega Basin remains imprecisely constrained (Figure 1a). The most robust anchor for the older age boundary is given by a Pb–Pb age of 2449 ± 1.1 million years ago (Ma) from a pluton crosscutting the Archean basement (Amelin et al., 1995). The Tulomozero Formation is additionally constrained by an imprecise dolomite Pb–Pb age of 2090 ± 70 Ma (Ovchinnikova et al., 2007) and by virtue of it recording the Lomagundi-Jatuli isotope excursion (along with the lowermost Zaonega Formation), which terminated in Fennoscandia at ~2060 Ma (Martin et al., 2013). A single zircon from a tuff layer in the lower Zaonega Formation yielded a U–Pb age of 1982 ± 4.5 Ma (Martin et al., 2015). A suite of cross-cutting dykes and sills in the Onega Basin succession provide younger age boundaries. A mafic sill in the Jangozero Formation, below the Tulomozero Formation, has yielded U–Pb zircon and baddeleyite ages of 1976 ± 9 Ma (Puchtel et al., 1998) and 1975.3 ± 2.8 (Martin et al., 2015). Zircons in dolerite and kimberlite sills in the Zaonega Formation have been U–Pb dated to 1919 ± 18 Ma (Priyatkina et al., 2014), 1956 ± 5 Ma (Stepanova et al., 2014), and 1961 ± 5.1 Ma (Martin et al., 2015); some of these sills have peperitic contacts, implying that the sediments were still wet and unconsolidated at the time of intrusion. Gabbro sills of the Suisari Formation, interpreted as coeval to the platformal lavas, are dated 1975 ± 24 (whole-rock and clinopyroxene Sm–Nd), 1980 ± 57 Ma (leach residue, plagioclase, and clinopyroxene Pb–Pb, Puchtel et al., 1998), 1988 ± 34 Ma (whole-rock and clinopyroxene Sm–Nd), 1985 ± 57 Ma (whole-rock and plagioclase Pb–Pb), and 1969 ± 18 Ma (whole-rock, ilmenite, and ulvöspinel Re–Os, Puchtel et al., 1999). Re–Os dates on Zaonega Formation mudstones have yielded a preliminary, and not yet peer-reviewed, age of ~2050 Ma (Bauer et al., 2019). Finally, a sandstone in the overlying Kondopoga Formation yielded a detrital zircon Pb–Pb age of 1967 ± 3.5 Ma (Martin et al., 2015). In summary, while dates are imprecise, deposition of the ~2400-m-thick succession comprising the Tulomozero, Zaonega, and Suisari formations likely spanned several tens of millions and possibly over a hundred million years. Provisionally, we constrain the age of the succession between ~2.1–2.0 Ga.

Materials and methods

Drill core

Drill core samples were obtained from the ~3500-m-long Onega Parametric Hole (OPH), drilled in 2008–2009 in the southern Onega Basin (62.1559 N, 34.4073 E; Figure 1). The drill core intersects ~800 m of the Tulomozero Formation, ~1500 m of the Zaonega Formation, and ~500 m of the Suisari Formation. Samples for this study were obtained from each of the aforementioned formations and constitute three sample sets: “NGU” (140 samples), “ACME” (40 samples), and
Care was taken to remove any macroscopically visible veins or monocrystalline mineral grains from rock slabs, especially for the “Yale” samples from which Cr isotope compositions were measured.

**X-ray diffraction**

The mineralogical composition of 147 samples was studied using X-ray diffractometry (XRD) at the University of Tartu, Estonia. Rock samples were ground to a fine powder, pressed into unoriented tablets and scanned on a Bruker D8 Advance diffractometer, where a copper Kα X-ray source and a LynxEye positive sensitive detector were used to generate diffractograms in a 2–70° 2θ range. Quantitative mineral abundances were modeled using the Rietveld algorithm-based Topaz software suite. The relative error for major mineral components (>5 wt.%) was ~10% and ~20% for minor mineral components (<5 wt.%).

Based on the results, the samples were divided into three main lithotypes: (1) evaporites (containing >10 wt.% halite, magnesite, and/or anhydrite) which mainly occur below 2330 m depth; (2) carbonates (containing >50 wt.% dolomite, calcite, magnesite, and/or siderite) above 2330 m depth that are dominated by dolomite and magnesite in the Tulomozero Formation, and dolomite-calcite in the Zaonega Formation; and (3) mudstones which first appear above 2330 m, but become dominant in the Zaonega Formation, where they contain mostly quartz, feldspars, micas and chlorite. See Supplementary Text for a more detailed description. For some mudstone and carbonate samples, mineralogical data was not available. For classification, the carbonate content of these samples was, instead, estimated using elemental abundances and the formula (Ca + Mg) / (Si + Al), with values >0.5 being considered carbonates.

**Element abundances**

Total organic carbon (TOC) content was measured at the Geological Survey of Norway (NGU) using a LECO SC-444 analyser (“NGU” samples) (Paiste et al., 2020a) and at the University of Alberta via loss on ignition (“ACME” and “Yale” samples). For the latter, sample aliquots weighing between 0.5 and 2 g were loaded into ceramic beakers and combusted at 500 °C for 8–12 h, and the change in mass was recorded. Element concentrations for sample set “NGU” were determined at NGU with a Philips PW 1480 X-ray fluorescence spectrometer equipped with a rhodium X-ray tube. For major elements, 0.6 g of powdered sample aliquots and 4.2 g of Li₂B₄O₇ were heated to 1000 °C and fused into a bead using a CLAISSE FLUXER-BIS. For minor elements, samples were prepared by mixing 9.6 g of powdered sample with 2.4 g of Hoechst wax in a Spex Mixer/Mill and pressing them into a pellet using a Herzog press. The detection limits for major elements were
generally below 0.02% and the typical precision (1σ) was ~2%. For minor elements, detection limits were ≤10 μg g$^{-1}$. Calibration of the XRF was done using a set of ~120 internationally certified natural rock standards as well as ~20 artificial standards provided by the XRF manufacturer.

Element concentrations for sample set “ACME” were determined at ACME Laboratories, Bureau Veritas Commodities Canada Ltd. The samples were powdered and then heated to mineralize organic carbon. Major elements were analyzed via inductively coupled plasma optical emission spectroscopy (ICP-OES) from aliquots fused into a LiBO$_2$ bead. Minor elements were digested using a mix of HNO$_3$, HClO$_4$, and HF and analyzed via inductively coupled plasma mass spectrometry (ICP-MS). Average relative standard deviation was less than 5% for all elements; accuracy was monitored using the OREAS 25a and OREAS 45e reference materials.

Element concentrations of sample set “Yale” were measured at the Yale Metal Geochemistry Center (YMGC). The samples were pulverized in an agate mill, combusted in ceramic crucibles at 500 °C for 8 h to remove organic carbon and digested in a class ten Pico-trace clean laboratory using a succession of concentrated, Teflon-distilled acids. Digestions were initiated with the addition of 3 ml of HNO$_3$ and 1 ml of HF, then heated at 100 °C for 24 h in closed Teflon vials, before being evaporated to dryness. Once dry, 3 ml HCl and 1 ml of HNO$_3$ were added to each sample, and samples were heated at 95 °C for 24h before being dried down. Final residues were then taken up in 5 mL of 3N HNO$_3$ solution at 70°C for 1h. Prior to analyses by ICP-MS, samples were diluted 200:1 in 5% Teflon-distilled HNO$_3$ spiked with 1 ng g$^{-1}$ indium (In). All samples were measured on a Thermo Scientific Element XR high-resolution ICP-MS. The In spike was used to correct for instrumental drift over the analytical session. Five dilutions of a customized internal laboratory standard were measured at the end and beginning of the analyses and several times throughout the run. Repeat analysis on the laboratory standard yielded a relative standard deviation (1σ) for all elements (excepting Na) of ≤8%, averaging ~4% across the analytical session. External accuracy was assessed by measuring the geostandard BHVO-2; relative difference between measured and known values for all elements was 8.3% on average and generally below 10%. In particular, the Cr and Ti values for BHVO-2 were within the 95% confidence interval for accepted values (Jochum et al., 2005). For all sample sets, Cr was normalized to the detrital tracer element Ti to assess authigenic enrichments, as suggested by Cole et al. (2017).

**Isotope ratios**

Oxygen isotope ratios of carbonates from micro-drilled samples were measured at the University of Tartu using a Thermo Scientific Delta V Advantage continuous flow isotope ratio mass
spectrometer and include data published in Kreitsmann et al. (2019). The precision of the measurements was 0.2‰ (2σ) and the long-term reproducibility exceeded ±0.2‰ (2σ). The compositions are expressed as the ratio of $^{18}$O over $^{16}$O, normalized to the Vienna Pee Dee Belemnite (VPDB) reference standard as $\delta^{18}$O:

$$\delta^{18}O = \left( \frac{^{18}O/^{16}O_{\text{sample}}}{^{18}O/^{16}O_{\text{VPDB}}} - 1 \right) \times 1000‰ \quad (1)$$

A correction was applied for the differing phosphoric acid fractionation factor of dolomite and calcite (Rosenbaum and Sheppard, 1986) according to the mineralogy of individual samples. Oxygen isotope compositions of bulk samples are presented here as an average of several micro-drilled sub-samples (1–7 micro-drilled measurements per bulk sample).

Chromium isotope compositions were determined for the “Yale” sample set at the YMGC from solutions prepared for elemental concentration analysis (see above). Sample preparation adapted the methods of Schoenberg et al. (2008) and Reinhard et al. (2014). An aliquot of each sample, containing ~200 ng of Cr, was diluted in 2 ml of 6N HCl and a $^{50}$Cr–$^{54}$Cr double spike was added at a spike/sample ratio of ~0.5, in order to correct for isotope fractionation during sample processing and measurement (Schoenberg et al., 2008). The samples were purified chromatographically to remove mass interference from $^{54}$Fe, $^{50}$Ti, and $^{50}$V (see Supplementary Information). First, samples were passed through columns containing AG1-X8 anionic resin, where the matrix was eluted by addition of 0.2N HCl and Cr was subsequently collected with 2N HNO$_3$. Second, samples were loaded on microcolumns filled with AG1-X8 resin to remove all remaining Fe, and Cr was eluted with 6N HCl. In the final step, Ti was removed using the cationic resin AG50W-X8: the matrix was eluted with 0.5N HNO$_3$, 0.5N HF, and 1N HCl and Cr was subsequently released with 1.8N HCl. Residues were taken up in 5% HNO$_3$ and analyzed on a Thermo-Finnigan Neptune Plus multicollector ICP-MS. To correct for remaining interferences, the samples were run in high-resolution mode and Fe, Ti, and V were monitored. Standard-sample bracketing with the NIST SRM 979 reference standard was employed to correct for instrumental drift. Instrumental mass bias and fractionation during sample preparation was corrected for using the Cr double-spike. Chromium isotopic compositions are normalized to the composition of the reference material NIST SRM 979, and reported as:

$$\delta^{53}\text{Cr} = \left( \frac{^{53}\text{Cr}/^{52}\text{Cr}_{\text{sample}}}{^{53}\text{Cr}/^{52}\text{Cr}_{\text{SRM979}}} - 1 \right) \times 1000‰ \quad (2)$$

The internal precision was calculated based on 20 duplicate analyses of NIST SRM 979 and reported as 2σ uncertainty. Geostandards BHVO-2B and Nod-1-a were used to monitor the external precision; the measured values were between −0.13‰ to −0.10‰ and 0.04‰ to 0.09‰, respectively, which are within error of previously reported measurements (Cole et al., 2016).
In-situ element mapping

In-situ element mapping of two polished samples (OPH-1572 and OPH-4081) from representative lithologies was conducted at the University of Tartu. Mapping involved parallel laser scans perpendicular to bedding from areas containing secondary silicate veins and anhydrite. Scans were performed with a Cetac LSX-213 G2+ laser system with a HelEx II fast-washout two-volume large-format cell using 800 ml min$^{-1}$ helium as carrier gas. A square 65 μm laser spot moving left-to-right at 65 μm s$^{-1}$ at 10 Hz and a power of 3.45 J cm$^{-2}$ was used. The total duty cycle was 0.2 s. For sample OPH-4081, $^{24}$Mg, $^{27}$Al, $^{28}$Si, $^{31}$P, $^{34}$S, $^{43}$Ca, $^{49}$Ti, $^{51}$V, $^{52}$Cr, $^{55}$Mn, $^{56}$Fe, $^{60}$Ni, $^{63}$Cu, $^{66}$Zn, $^{88}$Sr, $^{89}$Y, $^{137}$Ba, $^{140}$Ce, and $^{238}$U were measured at a dwell time of 7 ms. For sample OPH-1572, $^{13}$C, $^{27}$Al, $^{28}$Si, $^{31}$P, $^{34}$S, $^{43}$Ca, $^{47}$Ti, $^{51}$V, $^{52}$Cr, $^{55}$Mn, $^{56}$Fe, $^{60}$Ni, $^{63}$Cu, $^{66}$Zn, $^{95}$Mo, $^{140}$Ce, and $^{238}$U were measured at a dwell time of 8 ms. The composition of the volatilized material was analyzed using an Agilent 8800 quadrupole ICP-MS in single quad mode. Maps are presented as relative abundances in counts per second.

Chromium as a paleoredox proxy

The utility of Cr as a paleoredox proxy is due to its contrasting solubilities in oxygen-rich and oxygen-poor environments and the distinctive isotope fractionations associated with redox transitions. Chromium occurs in igneous rocks as insoluble Cr(III), with an isotopic composition of $-0.12\pm0.10\%\delta^{53}$Cr (e.g., Schoenberg et al., 2008), but can be converted to soluble Cr(VI) through reactions with manganese(IV) oxides in terrestrial weathering environments (e.g., Frei et al., 2009). Since the main pathway for Mn(IV) oxide formation requires molecular oxygen (although see Supplementary Text and Daye et al., 2019; Liu et al., 2020), and Cr(III) oxidation with Mn(IV) oxides induces a strong positive fractionation— theorists $\sim+6\%$, but lower in natural environments (see references in Wei et al., 2020)—the presence of fractionated Cr in aqueous environments indirectly attests to the availability of free oxygen in the atmosphere (see Supplementary Text and Miletto et al., 2021; Saad et al., 2017 for alternative takes). Additional Cr isotope fractionations can occur during redox reactions in rivers (e.g., Ellis et al., 2002) and oceans (e.g., Miletto et al., 2021), producing $\delta^{53}$Cr values that vary between $+0.3\%$ and $+1.6\%$ in modern seawater. Such fractionated $\delta^{53}$Cr signals can be captured by sediments accumulating under a reducing water column that scavenge dissolved Cr, so that the presence or absence of fractionated Cr in sediments can be used to track atmospheric oxygen levels through Earth history (Frei et al., 2009; Wei et al., 2020). Similarly, since concentrations of soluble Cr(VI) in seawater scale with the generation of Mn(IV) oxides in terrestrial settings and with the abundance of oxygen-rich settings in
the oceans, Cr concentration trends in anoxic sediments (which readily scavenge dissolved Cr) have also been tied to fluctuations in the extent of anoxic and euxinic seafloor area. For example, Reinhard et al. (2013) reported a shift from ~0.017 μg g\(^{-1}\)/μg g\(^{-1}\) Cr/Ti ratios in mid-Proterozoic mudstones to ~0.05 in Neoproterozoic mudstones, corresponding to expected shifts in atmospheric oxygen levels over the Proterozoic.

Organic-rich mudstones, such as those of the Zaonega Formation, can quantitatively capture marine δ\(^{53}\)Cr due to the fast kinetics of Cr(VI) reduction in anoxic waters, as has been demonstrated in the anoxic Cariaco Basin (e.g., Reinhard et al., 2014). Carbonates, while a much less efficient sink, capture Cr with a negative fractionation in biogenic carbonates but possibly minimal fractionation in abiogenic ones (Rodler et al., 2015). Therefore, δ\(^{53}\)Cr values in Tulomozero Formation carbonates can be considered a minimum boundary on seawater values. Compared to mudstones and carbonates, evaporites are much less studied in terms of their Cr isotopic composition. Yet, existing work on groundwater-derived evaporites in the Atacama Desert suggests that Cr is captured quantitatively in chromate salts (Pérez-Fodich et al., 2014). Likewise, the presence of bittern salts in the lower Tulomozero Formation implies cycles of almost complete seawater evaporation in the basin, which strongly support nearly quantitative capture of all seawater components. This is confirmed by element mapping of the most \(^{53}\)Cr-enriched evaporite sample (OPH-4081, 2305.25 m depth) which shows that chromium is hosted in diffuse, likely authigenic, reddish Fe-rich phases that are disseminated throughout the magnesite-dolomite matrix (Figure 2a).

In our Onega Basin samples, bulk Cr concentrations range between 0.3 and 2283.1 μg g\(^{-1}\), with a median of 86.1 μg g\(^{-1}\) (n = 253). Authigenic Cr abundances, expressed as Cr/Ti (Cole et al., 2017), range between 0.002 and 0.694 (μg g\(^{-1}\)/μg g\(^{-1}\); Figure 3), with a median of 0.039 (n = 253). Due to different Cr drawdown affinities of different lithologies (see Methods for classification), the median Cr/Ti value in Tulomozero Formation evaporites is 0.075 (range 0.002–0.366; n = 16); in Tulomozero and Zaonega Formation carbonates, 0.040 (range 0.008–0.694; n = 93); and in mudstones throughout the succession, 0.037 (range 0.003–0.554; n = 144). Significantly fractionated δ\(^{53}\)Cr values are present throughout the entire studied section, varying between −0.43±0.03‰ and +1.64±0.10‰ δ\(^{53}\)Cr (errors are 2σ), with an average of +0.24‰ (n = 41; Figure 3). In evaporites, δ\(^{53}\)Cr ranges between −0.34 and +1.17‰ (average +0.42‰; n = 6); in dolostones, between −0.20 and +1.17‰ (average +0.20‰; n = 14); and in mudstones, from −0.43 to +1.64‰ (average +0.22‰; n = 21) (Figure 4). These values, being the highest reported so far for the Paleoproterozoic (Reinhard et al., 2013; Wei et al., 2020), are difficult to explain through currently known oxygen-independent processes. For instance, ligand-based Cr(III) solubilization occurs at significantly lower rates than proton-driven solubilization (Saad et al., 2017), UV-based Mn(II)
photo-oxidation was, by the time of the middle Paleoproterozoic, hampered by the ozone layer (Liu et al., 2020), and Mn(IV) oxides produced through anoxygenic photosynthesis in reduced surface settings (Daye et al., 2019) would have been vulnerable to back-reduction prior to diffusing to subsurface Cr weathering environments (Anbar and Holland, 1992). Therefore, the data most likely reflect elevated redox potentials during the deposition of the Tulomozero and Zaonega formations (see Supplementary Text for more discussion on alternatives).

**Post-depositional alteration of chromium**

Due to its complex geological history, secondary processes within the Onega Basin succession may have altered some geochemical signals. For example, stromatolitic carbonates in the Tulomozero Formation have undergone pervasive dolomite/magnesite recrystallization with micritic fabrics being destroyed, but most samples have retained recognizable bedding and lamination (Melezhik et al., 2015, 1999). Zaonega Formation carbonates, where dolomite is the primary phase, have in large part been secondarily de-dolomitized to calcite, with a concomitant decrease in both their $\delta^{18}O$ and $\delta^{13}C$ values, though this has primarily affected the margins of dolomite beds (Kreitsmann et al., 2019). Furthermore, there are pervasive quartz-mica veins in the mudstones of the Zaonega Formation that were emplaced due to syn-depositional hydrothermal activity triggered by the emplacement of mafic lavas and sills or during later greenschist facies metamorphism (Paiste et al., 2018). Elemental mapping of the most $^{53}$Cr-enriched mudstone sample (OPH-1572, 1112.08 m depth) shows that Cr is enriched both in C$_{org}$-rich laminae, as well as in microscale mica veins, suggesting some Cr mobilization into percolating fluids (Figure 2b).

Despite the evidence above for post-depositional fluid alteration, several factors suggest that this process cannot explain the positively fractionated $\delta^{53}$Cr values in the Onega Basin. First, conspicuous veins, mono-mineral clusters, and altered margins of carbonate beds were carefully screened and avoided during sample selection. Second, igneous-derived Cr from hydrothermal sources is expected to host negative or crustal $\delta^{53}$Cr ratios (Schoenberg et al., 2008), and remobilization of primary Cr(III) is likely to drive residual sediment $\delta^{53}$Cr values more negative (Ellis et al., 2002), yet our samples have positive $\delta^{53}$Cr values. The lack of alteration is also evident by Cr mapping—concentrations in mica veins are highest in the areas where they cross Cr-rich laminae (Figure 2b), suggesting adjacent laminae as the source of Cr. Finally, the evaporite section of the lower Tulomozero Formation is composed of minerals, such as halite and bittern salts, which are highly susceptible to fluid alteration. Their persistence implies that substantial fluid movement has not occurred in these rocks.
Carbonates in the Onega Basin deserve further assessment since this lithology has a high propensity for diagenetic recrystallization. Oxygen isotope values provide a means of tracking the influence of diagenetic processes, given that more pervasive diagenesis typically leads to lower $\delta^{18}O$ values (Klaebe et al., 2021, and references therein). Carbonate $\delta^{18}O$ values in our Onega Basin samples range between $-17.2$ and $-4.8\%\text{o}$, with an average of $-11.0 \pm 2.7\%\text{o}$ (VPDB; $n = 45$; Figure S1). However, Figure 5 shows that the samples with the highest $\delta^{18}O$ values, hence being the ‘least altered,’ tend to have more fractionated Cr compositions than those with low $\delta^{18}O$ values, suggesting, in the standard framework, that secondary processes likely decreased, not increased $\delta^{53}\text{Cr}$ values (cf. Klaebe et al., 2021).

Another illustration that Cr geochemistry in the Onega Basin reflects primary processes is Figure 6a, which relates Cr/Ti to $\delta^{53}\text{Cr}$ in all three lithologies. That nearly all samples lie on a general positive trend suggests that $\delta^{53}\text{Cr}$ variance throughout the Onega Basin can largely be explained through the mixing of two components—an unfractionated detrital source and a fractionated seawater source (e.g., Planavsky et al., 2014)—without needing to invoke secondary overprint.

**Basin controls on the dissolved chromium pool**

Even if the authigenic $\delta^{53}\text{Cr}$ reflects a seawater signal, it is possible that elevated $\delta^{53}\text{Cr}$ was only a local phenomenon, and not representative of global Cr cycling. There is strong evidence for basin restriction in the Onega Basin that would allow such a scenario—evaporite mineralogy and sedimentary textures indicative of shallow deposition abound in the Tulomozero Formation (Blättler et al., 2018; Melezhik et al., 1999). Furthermore, the sulfur and iron isotope records in the Zaonega Formation have been interpreted to reflect varying levels of restriction (Mänd et al., 2021; Paiste et al., 2020b). In this case, distillation of the dissolved Cr pool by preferential drawdown of $^{52}\text{Cr}$ (i.e., Rayleigh fractionation) could have driven $\delta^{53}\text{Cr}$ more positive. Alternatively, positive values may have resulted from an anomalously positive riverine source of Cr.

However, Cr concentrations consistently reach hundreds of $\mu\text{g g}^{-1}$ over the hundreds of meters of mudstone-carbonate stratigraphy in the Zaonega and Suisari formations making it unlikely that a single point source could have provided this Cr. Distillation of the Cr pool is also unlikely, as that implies a substantial diminution of the dissolved Cr—it is far more parsimonious to invoke a fractionated open marine Cr source (e.g., Mänd et al., 2020). Conversely, in the lower Tulomozero Formation, the precipitation of a thick evaporite succession including bittern salts implies that seawater evaporation and Cr drawdown likely proceeded nearly to completion during numerous
evaporative cycles (Blättler et al., 2018), again precluding a distillation effect on the $\delta^{53}$Cr signal. While the shallow-water upper Tulomozero Formation carbonates present the highest likelihood of $\delta^{53}$Cr distillation, the low efficiency of Cr incorporation into carbonates (Reinhard et al., 2013) again precludes strong distillation.

Variable Cr abundances and isotope ratios attest to differences in Cr cycling and drawdown across the Onega Basin succession (Figure 3). For example, the highest Cr/Ti ratios (up to 0.353) and $\delta^{53}$Cr values (up to 1.64‰) are consistently present in the ~1115–1108 m interval that hosts a distinctive P-rich mudstone-dolostone contact—a marker horizon within the Onega Basin (Paiste et al., 2020a). This same interval displays extremely high TOC content (up to >70 wt.%), authigenic apatite, and fossilized sulfur cycling microbial ecosystems which are interpreted as having been formed in a highly biologically productive setting with fluctuating sulfidic–suboxic redox boundaries close to sediment surface that are especially conducive to redox-sensitive metal drawdown (Lepland et al., 2014; Mänd et al., 2020). Additionally, since the interval was host to a degree of hydrocarbon migration (Qu et al., 2012), secondary Cr accumulation may have contributed to these maximum values (though as a high-temperature process, this is unlikely to have caused isotope fractionations). On the other hand, marl samples from the transition zone of the Tulomozero and Zaonega formations (between 2100–1920 m) and in the Suisari Formation (above 640 m) host unfractionated Cr ($\delta^{53}$Cr below −0.13‰ and −0.12‰, respectively) at low authigenic concentrations (Cr/Ti below 0.065 and 0.196, respectively). In these latter cases, high levels of plagioclase (up to 53.2%), and the presence of amphibole and titanite, suggest that the sediments may have experienced substantial volcanic ash input that delivered unfractionated, igneous Cr. Since almost all samples lie on a generally positive trend between $\delta^{53}$Cr and Cr/Ti (Figure 6a), variation in both of these parameters can be sufficiently explained through variable Cr drawdown efficiencies or detrital input, leading to differing ratios of authigenic to detrital Cr. While it remains possible that waters in the Onega Basin experienced basin-specific Cr cycling, the presence of substantial volumes of isotopically fractionated Cr demand an oxygenated ocean–atmospheric system that can host Cr(VI)-cycling.

**Atmospheric–oceanic redox in the Paleoproterozoic**

The Rhyacian period has been proposed as a time of elevated oxygen abundance in the middle Paleoproterozoic, potentially sandwiched between times of comparatively lower oxygen abundance (Bekker and Holland, 2012). Oxygenated conditions are evidenced by evaporite mineralogy together with calcium and sulfur isotope records that indicate elevated marine sulfate levels (Blättler et al., 2018), consistent with high redox-sensitive element concentrations and isotope ratios...
This period of O$_2$ abundance was originally tied to the Lomagundi-Jatuli carbon isotope excursion through the mechanism of excess organic carbon burial (Bekker and Holland, 2012; Karhu and Holland, 1996), although this interpretation has become increasingly challenged (e.g., Mänd et al., 2020).

Intriguingly, the Rhyacian rock record is characterized by very minor sedimentary Cr isotope anomalies, despite this proxy being frequently cited as evidence for atmospheric oxygenation (Wei et al., 2020). Previous reports of positively fractionated Cr in the Paleoproterozoic do exist (Figure 7), particularly in the ~1.85 Ga Gunflint iron formation (Fralick et al., 2017; Frei et al., 2009), coupled with subtly negatively fractionated Cr in the ~1.9 Ga Schreiber Beach paleosol (Frei and Polat, 2013), both consistent with Cr(VI) solubilization and transport to the oceans. However, with the exception of meteorically derived diagenetic carbonate cements in the Gunflint Formation (Fralick et al., 2017), these fractionations barely extend beyond the crustal mean (maximum of +0.21‰), and are even surpassed by $\delta^{53}$Cr data from the Neoarchean that are indicative of relatively minor amounts of Cr(IV) weathering and cycling in a predominantly anoxic world (Frei et al., 2009). Furthermore, $\delta^{53}$Cr values of Rhyacian iron formations are even less fractionated (up to +0.03‰) (Frei et al., 2009). The anomalous feature of the later Paleoproterozoic $\delta^{53}$Cr record is high positive fractionations (up to +2.34‰) in the ~1.9 Ga Flin Flon and Beaverlodge Lake paleosols (Babechuk et al., 2017; Toma et al., 2019), which constitute the opposite of the expected isotope effect of oxic Cr weathering and have, thus, been linked to redox-independent Cr solubilization, instead of oxidative Cr cycling (e.g., Konhauser et al., 2011).

Collectively, these data have been used to suggest a late Paleoproterozoic to Mesoproterozoic Cr cycle highly distinct from the modern—variable atmospheric pO$_2$ produced, potentially in an alternating fashion, localized positively fractionated Cr runoff related to Mn(IV) oxide cycling (Fralick et al., 2017; Frei and Polat, 2013) or negatively fractionated runoff related to ligand- or acid-based solubilization (Babechuk et al., 2017; Toma et al., 2019). The oceans, meanwhile, are implied to have remained predominantly anoxic, with relatively minor fractionated Cr runoff diluted within an unfractionated marine reservoir.

In contrast to this mid-Proterozoic scenario, illustrated by a general lack of correlation between Cr/Ti and $\delta^{53}$Cr at this time (Figure 6b), our data from the ~2.1–2.0 Ga Onega Basin suggest a modern-type Cr cycle in the Rhyacian to the early Orosirian: nearly all samples follow a positive trend between Cr/Ti and $\delta^{53}$Cr (Figure 6a) that is a telltale sign of oxidative Cr(VI) cycling coupled to Mn(II) oxidation (Planavsky et al., 2014). Whereas Fe(II) oxidation occurs at circumneutral pH at near 0 mV potential, high-potential redox reactions ($\geq$500 mV) are required to oxidize Mn(II) under the same conditions. Although it remains unresolved how much, and for how long, oxygen...
levels would have to rise above ~0.1–1% of the present atmospheric level (PAL), as is required for
substantial Mn(IV)-driven Cr(III) oxidation without quantitative Cr(VI) back-reduction by Fe(II)
phases (Planavsky et al., 2014), highly fractionated $\delta^{53}\text{Cr}$ values coupled to high authigenic Cr
enrichments imply globally elevated $pO_2$ levels (see discussion on alternative Mn oxidation and Cr
fractionation mechanisms in Supplementary Text). This Cr(VI) then accumulated in relatively
oxxygen-replete oceans, where the oxyanion was stable. The large, positively fractionated marine Cr
pool was then further subjected to biochemical Cr redox cycling in the photic zone (e.g., Miletto et
al., 2021).

Our key finding is that, despite downcore variability in magnitude, the fractionated $\delta^{53}\text{Cr}$
values persist for nearly the entirety of the ~2400-m-thick sampled succession, across facies that
transition from a shallow evaporative setting to a deeper water turbidite system, and most likely
representing several tens to potentially more than a hundred Myrs of time (Figures 3, 7). Most
parsimoniously, the data suggest that Earth’s atmospheric oxygenation remained above the ~0.1–
1% $pO_2$ threshold for Cr(III) oxidation and transport to the oceans (Planavsky et al., 2014) for the
entirety of this time period. These findings provide a striking contrast to what are increasingly seen
as highly variable mid-Proterozoic redox conditions (e.g., Tang et al., 2017). In this light, the
Orosirian period perhaps witnessed a momentous transition from a well redox buffered to a poorly-
buffered atmospheric–oceanic redox state. This provides support for recent triple O isotope studies
that suggest elevated bioproductivity and $O_2$ production in the middle Paleoproterozoic (Crockford
et al., 2018), possibly due to increased nutrient fluxes (Konhauser et al., 2011; Bekker & Holland,
2012).

**Implications for the evolution of biological complexity**

Oxygen is often cited as the most important factor modulating the evolution and diversification of
complex life on our planet, driving, for example, increases in body size through time, the
diversification of eukaryotes, the origin of animals, and ultimately the invasion of land by
vertebrates and arthropods (e.g., Falkowski et al., 2005; Ward et al., 2006; Berner et al., 2007;
Payne, 2009; Sperling et al., 2013; Knoll, 2014; Planavsky et al., 2014; Cole et al., 2020). More
difficult to assess is the connection between oxygen and the origin of complex cells, in particular,
the eukaryotic cell, with its organelles, endomembrane system, relatively large size, and
sophisticated cytoskeleton. It is reasonable to suggest that free oxygen is a necessary precondition
for the evolution of crown group eukaryotes—the clade comprising the last common ancestor of all
living eukaryotes (LECA) and all of its descendants—given that LECA possessed mitochondria
and, presumably, sterols (Desmond and Gribaldo, 2009), both of which would have required
oxygen, albeit at very low levels (Waldbauer et al., 2011) or, possibly, only intermittently (Müller et al., 2012). But it is an open question whether an increase in oxygen was the driver for the origin of crown group eukaryotes—whether very low oxygen levels acted as a brake, that, once released, resulted in the immediate, or at least inevitable, evolution of cellular complexity.

One challenge in answering this question is that it is hard to pin down when the complex cells of eukaryotes evolved. In part this is because much of early eukaryotic evolution probably occurred in species that would not have been preserved, or, if preserved, would not have exhibited preservable characters that would allow us to recognize them as eukaryotes. Molecular clock studies permit the possibility that stem group eukaryotes emerged in the early Paleoproterozoic (e.g., Betts et al., 2018; Gold et al., 2017), but by definition these first eukaryotes would not have possessed any of the complex traits that characterize crown group eukaryotes; they would have been nearly or completely indistinguishable from their prokaryotic relatives (Porter, 2020). Rocks younger than 1700 Ma (Agić et al., 2015, 2017; Adam et al., 2017; Javaux et al., 2001; Javaux and Knoll, 2017) preserve fossils that indicate that eukaryotes had evolved a sophisticated cytoskeleton, the ability to form resistant organic walls, and, possibly, a Golgi apparatus and endomembrane system (Javaux and Knoll, 2017), but whether other traits like mitochondria and the nucleus were present by this time—or had evolved long before—is not clear. Molecular clock estimates for LECA are wide ranging (Porter, 2020), and, in any case, only provide minimum age constraints for the origin of complex cells: an age of 1.8 Ga for LECA, for example, does not tell us whether the nucleus, sophisticated cytoskeleton, and mitochondria were present much earlier (see Shih and Matzke, 2013 for a younger age estimate for the last of these). Thus, we do not know when complex eukaryotic cells emerged. Furthermore, even if we were able to constrain the timing of their origin to an interval when oxygen levels increased, it is still difficult to assess a causal connection given that it’s a single biological event (and therefore a just-so story). More robust evidence would be multiple independent acquisition of complex cells across numerous clades associated with a rise in oxygen (Butterfield, 2009). But just as with the early eukaryotic fossil record, it seems likely that if other, now-extinct lineages had evolved complex, eukaryote-grade cells, they would not be easily preserved, or if preserved, they would be difficult to recognize as such. In fact, it is worth considering whether some of the puzzling structures reported from Paleoproterozoic rocks could represent these extinct lineages (e.g., El Albani et al., 2010, 2019; Rasmussen et al., 2002; Bengtson et al., 2007).

What we can say, however, is that if there had been a delay in the appearance of complex cells—if complex eukaryotic cells did in fact arise ca. 1.7 Ga or later, and not ca. 2.1 Ga—it is unlikely to be due to a lack of oxygen. Results from the Onega Basin indicate that oxygen levels were high
enough and stable enough to support modern eukaryotic life for tens to >100 million years, from at least 2.1 to 2.0 Ga and perhaps earlier. Thus, if there was a delay, then we are forced to seek other reasons to explain it. Maybe other physical or ecological barriers prevented the evolution of complex cells during the Paleoproterozoic; maybe oxygen was an important driver but it simply takes a very long time for complex cells to evolve—longer than the tens to >100 million years recorded in the Onega Basin; or maybe the evolution of cellular complexity is not an inevitable result of rising oxygen (or any other environmental factor). In the case of the latter, the appearance of complex life on this planet might reflect, in large part, a series of contingent events.

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Figure captions

Figure 1: Onega Basin geology. (a) Simplified stratigraphy of the Onega Basin sedimentary succession showing formation boundaries, age constraints (explained in text; superscripts are references: 1 – Amelin et al. (1995); 2 – Puchtel et al. (1998); 3 – Martin et al. (2015); 4 – Ovchinnikova et al. (2007); 5 – Stepanova et al. (2014); 6 – Priyatkina et al. (2014); 7 – Bauer et al. (2019); 8 – Puchtel et al. (1999)), predominant geological features, and the location of the Onega Parametric Hole (OPH, indicated with a triangle). Modified from Melezhik et al. (2015) and Paiste (2018). (b) The exposures of the Tulomozoero, Zaonega, and Suisari formations within the Onega Basin, NW Russia are shown by color. (c) Mafic pillow lavas of the Suisari Formation. Hammer head is ~15 cm across (outcrop on the western shore of Lake Onega, Melezhik et al., 2013). (d-i) Polished drill core photographs of common rock types of the Tulomozoero and Zaonega formations. Scale bars are 1 cm across. (d) Laminated fine-grained dolostone in the upper Zaonega Formation (OnZaP drill cores, northern Onega Basin, 13 m, Paiste et al., 2018). (e) Laminated organic- and sulphide-rich mudstone in the upper Zaonega Formation displaying soft-sediment deformation as well as quartz-mica and pyrobitumen veining (OnZaP, 63 m, Paiste et al., 2018). (f) Rhythmically bedded greywacke-siltstone in the lower Zaonega Formation (FAR-DEEP 12AB drill core, northern Onega Basin, 282 m, Melezhik et al., 2013). (g) Pale pink dolostone of the upper Tulomozoero formation with red-brown stromatolitic laminae (FAR-DEEP 11A drill core, northern Onega Basin, 107 m, Melezhik et al., 2013). (h) Massive, coarse-crystalline anhydrite in the middle Tulomozoero Formation (OPH, 2510 m, Melezhik et al., 2015). (i) Brown and pink, massive, coarse-grained halite of the lower Tulomozoero Formation with inclusions of anhydrite (white), magnesite (yellow), and shale (pale grey) (OPH, 2902 m, Melezhik et al., 2015).

Figure 2: In-situ element maps for representative Onega Basin samples. Optical micrographs and relative carbon, aluminum, iron, and chromium abundance maps (reported in counts per second) were determined through laser-ablation mass-spectrometry from representative lithologies in the OPH core. (a) Evaporitic carbonate sample OPH-4081 (depth 2305.25 m, δ⁵³Cr = 1.16±0.08‰), composed mainly of magnesite–dolomite, hosts disseminated Fe-rich phases, in which Cr is preferentially concentrated. (b) Corg-rich mudstone sample OPH-1572 (depth 1112.08 m, δ⁵³Cr 1.63±0.11‰) displays Corg-rich sedimentary laminae and Al-rich cross-cutting mica veins. Cr is concentrated both in the laminae and veins. Scale bars are 1 mm. Note difference in color scale.
Figure 3: Chromium geochemistry in the Onega Parametric Hole drill core. Columns show the lithology, abundance of authigenic chromium (Cr/Ti), and its isotopic composition ($\delta^{53}$Cr). Color of datapoints corresponds to lithology. Crustal averages, vertical grey bars, are taken from Cole et al. (2017) and Schoenberg et al. (2008).

Figure 4: Chromium isotope ratios per lithology. Boxplots show medians, 25/75th quantiles, and the full range; superimposed black circles are individual samples. Grey bar denotes average crustal composition from Schoenberg et al. (2008).

Figure 5: Relationships between chromium and oxygen isotope ratios in carbonates. Chromium isotope ratios ($\delta^{53}$Cr) are plotted against oxygen isotope ratios ($\delta^{18}$O VPDB); in the latter case, lower values are indicative of stronger alteration of carbonates. Grey bar denotes average crustal composition from Schoenberg et al. (2008).

Figure 6: Authigenic chromium abundances (Cr/Ti) plotted against isotope ratios ($\delta^{53}$Cr). (a) Onega Basin data from this study, coloured per lithology. (b) Published data from mid-Proterozoic sections of various lithologies, coloured according to age (1950–1000 Ma). A positive trend, as opposed to a horizontal one, suggests Cr(VI) cycling and the presence of atmospheric oxygen (e.g., Planavsky et al., 2014). Data for panel b is from Frei and Polat (2013), Planavsky et al. (2014), Cole et al. (2016), Gilleaudeau et al. (2016), Canfield et al. (2018), and Wei et al. (2021).

Figure 7: Fossil evidence for eukaryotes (a), authigenic chromium abundances (using Cr/Ti as proxy) in shales (b), and $\delta^{53}$Cr values in various geological archives (c) through time. References for (a) are Miao et al. (2019), Bengtson et al. (2017a, 2017b), Gibson et al. (2018), Brocks et al. (2017) and Bobrovskiy et al. (2021). Differently coloured pastel dots on panels b and c represent previously published data of different lithological affinities. Saturated cross symbols are data from this study. Cr/Ti data is from Reinhard et al. (2013) and Robbins et al. (2016); $\delta^{53}$Cr data is from the compilation of Wei et al. (2020) and Wei et al. (2021). Since Cr concentrations in the crust have decreased through geological time due to a decreasing prevalence of Cr-rich ultramafic igneous rocks, samples have been normalized to changes in crustal abundance according to Condie (1993). Grey bars represent average crustal compositions as in Figure 2. Errors are smaller than the data symbols.
Figures

Figure 1:
Figure 2:
Figure 4:
Figure 5:
Figure 6:
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