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The Österplana Fossil Meteorites and . . . What Else? Terrestrial Cr-Spinels and Zircons in the Ordovician Limestones of the Thorsberg Quarry (Sweden)

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Abstract: In the Ordovician limestone of the Thorsberg quarry (South Sweden), about 130 meteorites have been found. Among the extraterrestrial material, several terrestrial Cr-spinels and zircons have been found too. In particular, in the interval 416–447 cm above the Arkeologen bed, terrestrial Cr-spinels, compositionally different from previous studied Cr-spinels of the same sequence, are present. Previous studies on zircon provided depositional ages that range from 464.22 ± 0.37 Ma to 465.01 ± 0.26 Ma. The trace element content of zircons suggests different possible source rocks. In fact, zircons from the oldest ash layer resemble those from dolerite, while those in the youngest layers are similar to zircons commonly found in granitoids, with more than 65% wt. SiO2. The chemistry of Cr-spinels suggests a strong alteration, so that it is difficult to assign them to a specific area, however they recall the chemistry of altered spinels from ophiolitic occurrences (among other possibilities). The geological setting of the Laurentia and Baltica areas, including the description of basalts to rhyolite association and the presence of ophiolitic slices, makes us confident about the derivation of these zircons and Cr-spinels from those areas.

Keywords: Cr-spinel; zircon; Ordovician; Sweden; Laurentia; Baltica

1. Introduction

About a quarter of all meteorites falling on Earth today originate from the breakup of the L-chondrite parent body ~466 Ma ago, the largest documented breakup in the asteroid belt in the past ~3 Ga. A systematic search for fossil meteorites was initiated in the Thorsberg quarry in southern Sweden in the 1990s [1,2]. In the quarry, since then, about 130 meteorites, 1–21 cm in diameter, have been found. The meteorites fell over an area of less than ca. 30,000 m² of the seafloor during ~2 Ma. About 100 of the recovered meteorites have been analysed for chrome-spinel composition and classified so far [3]. All except one of these are ordinary chondrites, and are almost certainly L chondrites, based on the more detailed oxygen isotope and chondrule-size studies of a representative subset of fossil meteorites [4–6]. One of the recovered meteorites, once called Mysterious Object (now Österplana 065), is an 8 cm winonaite-related meteorite of a type not known among present-day meteorite falls and finds [7]. The Österplana 065 seems to be a fragment of the impactor that broke up the L-chondrite parent body that, according to [8], was shown to be the first documented example of a meteorite type that does not fall on Earth today, because its parent body has been consumed by collisions. In the sediments containing fossil meteorites, there are also abundant L-chondritic chromite grains [9].

The Österplana 065 was found in the Glaskarten 3 bed that spans the interval from 427–467 cm above the base of the Arkeologen bed. In the same interval (416–447 cm),
the so-called Mysterious Layer (ML), characterized by exceptionally high concentrations of Cr-spinel grains, can be found. The Cr-spinels of the Österplana 065 have nothing in common with the Cr-spinels from the ML that are terrestrial. Moreover, the chemistry of the terrestrial Cr-spinels of ML is very different from that of the other terrestrial Cr-spinels in the rest of the section, and occurring in the same levels as zircons.

Zircons from different ash-bearing layers provide stratigraphically consistent depositional ages that range from 464.22 ± 0.37 Ma to 465.01 ± 0.26 Ma [10].

Among the several heavy minerals that can be found in sediments, Cr-spinel is of particular significance because, unlike silicate minerals, it is resistant to low-grade alteration and mechanical breakdown. Furthermore, it is a widespread accessory mineral in rocks that are potential sources of sediments as ultramafic intrusive, cumulate and volcanic suites, or metamorphic rocks (see review in [11,12]). When derived from mantle peridotites and primitive rocks, this is indicative of magmatic and tectonic evolution in the source areas [13–19]. Spinel chemistry is diagnostic of parental melt composition and crystallisation conditions [11,12,20,21], and it often contains melt inclusion carrying direct information on magma chemistry [12,17,22–24].

Zircons forms in a wide variety of igneous and metamorphic rocks, as they are ultrastable, are often used in paleogeographic reconstruction for single-grain U-Pb dating. However, [25–27], among others, they showed that zircon displays variations in trace elements that could reflect host rock compositions, the most significant being those related to REE, U, Th, Y, and Nb. Owen (1987) [28] suggested the Hf content as a possible discriminant in provenance. Metamorphic zircons can be distinguished by magmatic ones according to Th/U ratios [29–31].

Here, we will study the chemistry of both spinels and zircons found in the Ordovician limestones of the Thorsberg quarry, trying to estimate their possible provenance.

2. Geological Setting

In 1993, following two finds of fossil meteorites in mid-Ordovician marine limestone in Sweden, a systematic search for meteorites was initiated in the Thorsberg quarry in southern Sweden [1,2]. During most of the Ordovician, this area was situated several hundreds of kilometres from the mainland, and constituted a tectonically stable depositional environment. The lithified sediments formed slowly, 2–4 mm Ka⁻¹. It is in the industrial process of limestone extraction that the meteorites are recovered, and until today, that around 130 meteorites have been found, representing ~98% of all fossil meteorites known to science. Although the recovered meteorites fell during ~2Ma, the limestone beds being sawed and yielding meteorites, however, represent <1 Ma of time [1].

3. Materials and Methods

A total of 1320 kg of limestone samples (about 50 rock pieces) from the Hällelekis-Thorsberg section were processed at the Astrogeobiology Laboratory at Lund University, using techniques developed for the separation and extraction of extraterrestrial minerals from sediments [2]. The sample depths range from −6.18 to +10.30 m [2,9] relative to the base of the Arkeologen bed, with no visible bentonite layer identified (Figure 1). The bulk samples were dissolved in HCl (6 M) and HF (11 M) at room temperature after thorough cleaning (see [9], for details). After sieving at mesh sizes 32 and 63 µm, residues were carefully searched under the binocular microscope for chrome-spinel and zircon grains. Only samples from three levels, i.e., +1.80 to +1.90 m (Likhall), +4.18 to +4.35 m (HKIS 4c), and +5.09 to +5.26 m (HKIS 2a) contained well-preserved zircons with a likely volcanic ash origin; a lower layer seemingly not related to ash is also present (No Ash).

Zircons are colourless, euhedral grains of up to 500 µm in length. They are elongated to acicular morphologies, with aspect ratios greater than 3, and have well-defined crystal faces and sharp edges, with no signs of transport abrasion. According to [32,33], these zircons are probably from volcanic ash or tuff, as they show no signs of transport...
abrasion, and share common features with the elongated acicular morphologies typical of rapid crystallization.

Figure 1. Stratigraphic column.

The grains were mounted in epoxy, polished flat with 6 and 1 μm diamond paste, and coated with carbon. Elemental composition was determined with an INCA Oxford systems energy dispersive X-ray device (EDS) attached to a Hitachi S-3400N scanning electron microscope (SEM) at the Department of Geology, Lund University. Electron voltage was set to 15 keV and working distance to 10 mm. Counting time was set to 60–80 s per analysis. For the standardization of the instrument, a cobalt standard was used. For the standardization of the spectrum peaks, we used synthetic pure spinel for Mg and Al, chromite for Cr, hematite for Fe, and titanite for Ti. Analytical accuracy was controlled by the USNM 117075 (Smithsonian) reference standard [34].

In situ chemical analyses on zircons have been performed by the LA-ICP-MS (Laser Ablation Inductively Coupled Plasma Mass Spectrometry) hosted by the Department of Physics and Geology of the University of Perugia. The laser ablation device was a Teledyne/Photon Machine G2 (ATL-I-LS-R solid state-triggered excimer 193 nm laser) equipped with a two-volume ANU (Australian National University) HelEx 2 cell. The spectrometer was a quadrupole Thermo Fisher Scientific iCAP Qc ICP-MS. The analytical
setup and relative figures of merit (precision, accuracy, and detection limits) are described in [35,36]. The spot size for all the analyses has been set at 12 µm, and helium has been used as carrier. The zircon trace elements (\(31P, 42Ca, 47Ti, 53Mn, 57Fe, 71Ga, 85Rb, 89Y, 90Zr, 93Nb, 118Sn, 137Ba, 139La, 140Ce, 141Pr, 146Nd, 147Sm, 153Eu, 157Gd, 159Tb, 163Dy, 165Ho, 166Er, 169Tm, 173Yb, 175Lu, 178Hf, 181Ta, 208Pb, 232Th and 238U) have been analysed using the reference standard STD 91500, and following the method suggested by [37].

4. Results and Discussions
4.1. Cr-Spinels

The analysed spinels are essentially chromites in composition. The Cr\(_2\)O\(_3\) amount is usually high; in fact, more than 75% of the studied spinels present Cr\(_2\)O\(_3\) higher than 30 wt. % (Figure 2). It is followed by iron that, in the FeO form, usually comprises from 20–30 wt. % (about 85% in frequency), while in the Fe\(_2\)O\(_3\) form (calculated according to stoichiometry), it is variable with the highest frequency 20–40 wt. % (more than 75% of the analysed spinels fall into this field). About 90% of the spinels have an Al\(_2\)O\(_3\) content lower than 5 wt. %, while MgO is usually below 5 wt. % (about 85% of the spinels). Apart from a couple of samples, TiO\(_2\) is below 3 wt. %, MnO below 0.35 wt. % and V\(_2\)O\(_3\) below 0.5 wt. %.

![Figure 2](image-url). Frequency distribution of Cr\(_2\)O\(_3\), Al\(_2\)O\(_3\), FeO, Fe\(_2\)O\(_3\), and MgO in the studied spinels.

Commonly, detrital spinels are related to ophiolitic material. Lenaz et al. (2000) [17] and [12] provided the spinel TiO\(_2\) vs. Al\(_2\)O\(_3\) diagram (Figure 3). This diagram allows the discrimination of ophiolitic spinels from different tectonic settings, such as island arc, ocean island, back-arc basin, and MORB. The studied spinels show compositions
different from any represented field. In the same figure, the composition of Cr-spinels from the Shetland, Ballantrae, and Leka ophiolites is also represented. The composition of the detrital Cr-spinels found in Kinnekulle resembles that of the ferrianchromite rims of Cr-spinels from those ophiolites, but the detritals are usually richer in TiO₂. This fact, the low Mg#, i.e., Mg/(Fe²⁺ + Mg) and the enrichment of MnO could suggest a possible secondary process [38].

Alwmark and Schmitz (2007) [39] noticed that in extraterrestrial chromites from the Lockne crater in central Sweden, there were Zn enrichment and oxidation. They suggested that it was a result of the hydrothermal system induced by the impact. However, the extraterrestrial Cr-spinels from Kinnekulle are not considered altered, due to the absence of Zn enrichment [40].

According to the chemical compositions, the Cr#, i.e., Cr/(Cr + Al), is in the range 0.78–0.94 for the majority of spinels, while the Fe#, i.e., Fe²⁺/(Fe²⁺ + Mg), is in the range 0.67–0.97. The Fe³⁺/(Fe³⁺ + Al + Cr) is usually higher than 0.2 (maximum 0.7). This value is particularly interesting because, in meteoritic spinels, the Fe³⁺ is virtually absent, so that it allows the possibility to discriminate between terrestrial and extraterrestrial occurrences.

To understand the likely provenance for these spinels, we compared them with the database of [11]. If we consider the Cr# vs. Mg# diagram, our spinels fall into the field or close to the field of alkali basalts, lamprophyres, and related rocks and worldwide Alaskan zoned ultramafic complexes (Figure 4). Anyway, they differ from the closest Alaskan complexes of the Urals, that, by the way, are also younger, showing a comparable Cr# in the range 0.60–0.85, but lower Fe# in the range 0.2–0.8 [44,45]. When inserted into the ternary Cr-Fe³⁺-Al diagram, they perfectly fall into the field of alkali basalts, lamprophyres, and related rocks, and are similar to the worldwide Alaskan ultramafic complexes [11] (Figure 5). However, we were unable to find source areas with similar compositions.

Figure 3. TiO₂ vs. Al₂O₃ diagram. Red circles: Cr-spinels from Kinnekulle (this study); yellow fields: Cr-spinels from Ballantrae ophiolites [41]; red dotted lines: Cr-spinels from Shetland ophiolites [42]; red dot and dash lines: ferrianchromite rims of Cr-spinels from Shetland ophiolites [42]; blue dotted lines: Cr-spinels from Leka ophiolites [43]; blue dot and dash lines: ferrianchromite rims of Cr-spinels from Leka ophiolites [43]. Fields for Arc, OIB (ocean island basalts), BABB (back arc basin Basalts), and MORB (mid ocean ridge basalts) after [12,19].
In the Scandinavian area, there are scattered occurrences of chromites and/or Cr-spinels. Among other things, in central Sweden, chromites occur in the Proterozoic metallogenic province of Bergslagen [46], in the Kittelfjäll spinel peridotite; a fragment of lithospheric mantle, which occurs as an isolated body within high grade metamorphic crustal rocks of the Seve Nappe Complex [47]. In Norway, they are usually associated with ophiolites, as in the case of Leka [43,48], while in Finland, they occur in Paleoproterozoic primitive volcanic rocks [49], or in layered deposits such as Kemi [50]. Other ophiolites related to the closure of the Iapetus can be found in Newfoundland [51,52]. Anyway, none of these occurrences fit our analyses. Considering that their chemistry is similar to the one of the altered rims of Shetland and Leka ophiolites [42,43], they may be the product of alteration of an ophiolitic
segment, or they could be supplied as an aeolian contribution, as happened, for example, for spinels in the Maiolica formation in Italy [53], deriving from unknown alkaline basalt, lamprophyre, or Alaskan ultramafic occurrences.

4.2. Zircons

In zircons, Hf and Y, which can be considered as minor elements for zircons, are in the ranges 5800–20,000 ppm and 100–7200 ppm, respectively. In particular, Hf is in the range 5800–14,700 ppm for the No Ash sample (mean 9195 ± 2329), 7860–12,500 ppm for LIK (mean 9626 ± 992), 6300–18,900 ppm for 4C (mean 9693 ± 1769), and 8600–20,260 ppm for 2A (mean 12,351 ± 2597). Yttrium is in the range 260–2220 ppm for the No Ash sample (mean 1029 ± 510), 540–2800 ppm for LIK (mean 1711 ± 601), 104–5300 ppm for 4C (mean 2562 ± 1058), and 1060–7170 ppm for 2A (mean 2717 ± 1347). The ionic radius of Zr is most closely matched by the smaller radii HREE than the larger LREE [31]. Given this, the total content of LREE (La to Pr) is lower than about 100 ppm (No Ash: 4–34 ppm; LIK: 6–45 ppm; 4C: 9–103 ppm; 2A: 7–53 ppm), that of MREE (Nd to Gd) is lower than about 250 ppm (No Ash: 8–41 ppm; LIK: 12–106 ppm; 4C: 7–248 ppm; 2A: 18–179 ppm), and that of HREE (Tb to Lu) is in the range 55–4055 ppm (No Ash: 354–1922 ppm; LIK: 421–1972 ppm; 4C: 110–3865 ppm; 2A: 770–4288 ppm). The values of P can be very different in zircons according to the host rocks. This is in the range 59–2615 ppm for the studied zircons (No Ash: 125–814 ppm; LIK: 185–838 ppm; 4C: 59–1167 ppm; 2A: 418–2615 ppm). According to [31], Th and U are somewhat compatible with Zr (ionic radii: Th⁴⁺ 1.05 Å, U⁴⁺ 1.00 Å) and occur in crustal zircon with abundances typically ranging from tens- to thousands-of-ppm. Thorium is in the range 30–529 ppm for the studied zircons (No Ash: 50–263 ppm; LIK: 40–349 ppm; 4C: 38–529 ppm; 2A: 30–510 ppm), while U is in the range 57–447 ppm (No Ash: 144–383 ppm; LIK: 66–348 ppm; 4C: 57–377 ppm; 2A: 61–447 ppm).

To discriminate between possible sources, some provenance diagrams have been used. According to the Th vs. U diagram [31] (Figure 6), zircons with Th/U ratios lower than 0.1 are metamorphic, and those with values above 0.1 are magmatic, even if [54] suggested that recrystallized and newly grown metamorphic zircons often have high Th/U ratios. The zircons studied here show values close to or greater than 1, so they can be considered as magmatic. It is interesting to notice that all the occurrences of zircons show subparallel trends and those from the No Ash level are usually lower in Th for a given U content.
Another effective method to discriminate between pristine magmatic zircons and the “metamorphic” ones is to verify their REE patterns, mainly focusing on HREE ratio \( (\text{Gd}_{\text{CN}}/\text{Yb}_{\text{CN}}) \), and secondarily, Ce and Eu anomalies \[27,31,55,56\]. Generally, the analysed crystals show quite strong positive Ce and negative Eu anomalies, together with a steep HREE trend, typical of magmatic crystals (Figure 7). Only one zircon (4c-53) shows no Eu anomaly, and a slightly flatter \( \text{Gd}_{\text{CN}}/\text{Yb}_{\text{CN}} \) ratio \( (\text{Gd}_{\text{CN}}/\text{Yb}_{\text{CN}} = 0.14) \) that could suggest an eclogitic origin \[56\]. Grimes et al. (2007) \[57\] divided oceanic (MORB) and continental zircons by using \( \text{U/Yb} \) vs. \( \text{Hf} \) and \( \text{U/Yb} \) vs. \( \text{Y} \) diagrams (Figure 8). These diagrams show that at least some of the zircons from the No Ash level present a MORB affinity. All the others are related to a continental arc. The same results can also be obtained by using the \( \text{U/Yb} \) vs. \( \text{Nb/Yb} \) \[55\] (Figure 9). Recently, to discriminate zircons from different tephras, \[58\] introduced some discriminant functions, taking into consideration several trace and rare earth elements in zircons. The applications of the so-called DF1 and DF2 parameters show that the No Ash level falls in a field different from the one where the other samples fall (Figure 10). Moreover, the \( \text{Yb/Dy} \) vs. \( \text{Dy} \) diagram is characterized by a relatively low \( \text{Yb/Dy} \) ratio and a wide variation in \( \text{Dy} \) consistent with dominant plagioclase crystallization accompanied with minor allanite, zircon, and amphibole fractionation for the No Ash, 4C and 2A levels, whereas the LIK zircons have high \( \text{Yb/Dy} \) ratios and limited \( \text{Dy} \) variation, which can be explained by dominant amphibole fractionation, in addition to plagioclase crystallization.

Different sources could also be differentiated by employing the \( \text{Gd}_{\text{CN}}/\text{Yb}_{\text{CN}} \) vs. \( \text{Nb/Yb} \) (Figure 11) diagram \[55\], which emphasizes the increase of garnet signature, zircon (cooling of the melt), and titanite-apatite-ilmenite crystallization. The investigated samples plot in a quite small area in which they partially overlap, but it is possible to recognize an increase in the garnet signature from No Ash-2a to Likhall, and finally, to 4c. However, if we consider the stratigraphic continuity, it is possible to imply that there was an increase of the garnet signature occurring from No Ash to Likhall, and then 4c. Successively, a fractionation of titanite-apatite-ilmenite in 2a occurred, plotting the sample near No Ash.

**Figure 7.** Range of REE content in the studied samples.
Figure 7. Range of REE content in the studied samples.

Figure 8. (A) U/Yb vs. Hf and (B) U/Yb vs. Y diagrams according to [57]. Symbols as in Figure 7.

Figure 9. Log$_{10}$U/Yb vs. Log$_{10}$Nb/Yb diagram according to [58]. Symbols as in Figure 7.

Figure 10. (A) DF1 vs. DF2 and (B) Yb/Dy vs. Dy diagrams according to [58]. Symbols as in Figure 7.
Different sources could also be differentiated by employing the GdCN/YbCN vs. Nb/Yb diagram \[55\] (Figure 11) which emphasizes the increase of garnet signature, zircon (cooling of the melt), and titanite-apatite-ilmenite crystallization. The investigated samples plot in a quite small area in which they partially overlap, but it is possible to recognize an increase in the garnet signature from No Ash-2a to Likhall, and finally, to 4c. However, if we consider the stratigraphic continuity, it is possible to imply that there was an increase of the garnet signature occurring from No Ash to Likhall, and then 4c. Successively, a fractionation of titanite-apatite-ilmenite in 2a occurred, plotting the sample near No Ash.

**Figure 11.** Gd/Yb vs. Nb/Yb diagram according to \[57\]. Symbols as in Figure 7.

Belousova et al. (2002) \[27\] studied the trace and rare earth elements in igneous zircons from several lithologies. Their classification trees based on recursive partitioning techniques provide a rapid means of relating parent rock type to zircon trace element analysis at confidence levels of 75% or more. By applying those criteria, zircons from the No Ash level are predominantly (about 70%) from doleritic affinity rocks; LIK zircons are almost equally subdivided between granitoids with SiO$_2$ < 65%, and granitoids with SiO$_2$ higher than 70–75%, in 4C and 2A levels with more than 50% of zircons, are assigned to granitoids with SiO$_2$ greater than in the 70–75% group (Figure 12). In the most recent levels, there are also about 10% of zircons with a syenitic/monzonitic affinity (Figure 12).
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Figure 12. Frequency distributions of zircons according to the possible sources, as determined using the classification tree diagram by [27]. GR70–75: granitoids with about 70–75% SiO$_2$; GR < 65: granitoids with less than 65% SiO$_2$; SY/MONZ: syenites and monzonites; DOL: dolerites; CARB: carbonatites.

The temperature of crystallization has been calculated using the Ti-in-zircon geothermometer [59], and a relative density curve has been plotted (Figure 13). No Ash zircons show a smaller gap, and the lowest crystallization temperature among the samples (673–776 °C), while both are similar for Likhall and 4c (680–839 °C and 685–857 °C, respectively). Finally, 2a shows a gap similar to No Ash, but at higher temperatures (703–815 °C). The errors employed for building the relative density curve are arbitrary, and a value of 2$\sigma$ = ±10 has been chosen, taking into account the number of analyses and the temperature distribution. Even though the diagram shows that the patterns are quite different from each other, many peaks are shared by two or more samples, but with different heights. In particular, the most shared ones are those at 770 °C (shared by all the samples), and at about 708 °C (No Ash, Likhall, and 2a), the major parts of the others are shared by two samples (Figure 13). Other than these, 4c shows little exclusive peaks at 719 °C and 856 °C, while the one at 815 °C is rather high, and belongs only to 2a.
Torsvik and Rehnström (2003) [60] suggested that, during the Late Ordovician, suturing between the Avalonia and Baltica plates closed the Tornquist Sea that had separated these palaeocontinents during most of the Ordovician. This suture zone, the so-called Trans-European Suture Zone, during the Variscan and Alpine orogenesis, further developed and/or rejuvenated, and is now largely concealed by deep sedimentary basins of the late Palaeozoic to Tertiary age. Avalonia separated from Gondwana during the Arenig–Llanvirn, however, the exact nature and time of collision between Avalonia and Baltica are not clear in the geological record [60]. In Avalonia, Late Ordovician folding, faulting, and igneous activity is known from the Welsh Borderlands (Shelve area), and may relate to Avalonia–Baltica docking, being more or less coeval with Taconic deformation (455–422 Ma), along the Laurentian margin [61]. Large K-bentonite beds are widespread in Baltica, and the most regionally important is the Kinnekulle bed (Mid-Caradoc age, according to the D. multidensgraptolite zone). The Kinnekulle bed has been dated 452 ± 2 Ma (U/Pb zircon age cited in [62]) and 455 ± 2 Ma (40Ar/39Ar biotite; [63]). According to Huff et al. (1992) [64], this unit shows affinity with granites from the volcanic arc, indicating generation along convergent continental plate margins. Based on the climatic zones, Ref. [60] argued that the Kinnekulle K-bentonite beds could be associated with a Mid-Caradoc calc-alkaline magmatic event in Avalonia. The thickness distribution of these ash beds requires a palaeo-west source area, and the Avalonian magmatic arc is the best candidate to supply Baltica with massive ash falls. According to this reconstruction, Ref. [65] suggested, for zircons found in the “Täljsten” interval at Kinnekulle, that the source volcanoes were probably situated at the west or southwest (present-day directions), where the magmatic activity occurred during the subsequent amalgamation of Baltica, Avalonia, and Laurentia, and [10] endorses the interpretation of [65]. Haynes et al. (2011) [66] studied the biotites present in the Kinnekulle bentonites, and suggested that they are the product of explosive volcanism, with the source magmas having passed through the continental crust on their way to large, single vent eruptions associated with subduction along continental margins like Laurentia–Baltica margins during the Ordovician. However, both the zircons from the ash layer and those from the No Ash layer are slightly older than those previously described [10]. Woodcock et al. (2012) [67] described voluminous back-arc basin volcanism, typically with a bimodal basalt–rhyolite composition, beginning in Welsh basins during the Llanvirn hiatus in arc activity. The back-arc basin volcanism continued after the arc rejuvenated outboard on the microcontinent. The study of [68,69] on the Fishguard Volcanic

![Figure 13. Density curve of zircon crystallization temperatures. Blu line: 2a; grey line: 4c, black line: Likhall; red line: No-ash.](image-url)
Group in South Wales showed that it represents an example of a volcanic sequence linked to the closure of the Iapetus Ocean. Its age spans in the range 467–458 Ma and reveals that the Fishguard Volcanic Group represents a closely related series of basalts, basaltic andesites, dacites, and rhyolites originating in a back-arc environment. Zircon U–Pb geochronological data for felsic volcanic rocks indicate deposition over a short interval between 465 and 462 Ma. Fritschle et al. (2018) [70] reported arc-related rhyolites from the Avoca volcanic sequence in SE Ireland dated at 463.6 ± 2.6 Ma. At the same time, Refs. [71–74] showed that the Canadian Appalachians comprise a tectonic collage of Ordovician peri-Laurentian terranes that formed during the closure of the Iapetus Ocean and imbricated beneath the composite margin of Laurentia, and is comprised in the 470–460 Ma interval.

5. Conclusions

The literature on detrital zircon presents a lot of age determination, but few analyses of trace elements. Many trace element analyses of non-detrital zircons can be found, however, to our knowledge, there are no analyses of trace elements from what we consider the possible source areas. Commonly, authors using detrital zircons compare the ages while, in past years, Hf isotopes have been used too [75–78]. We think that trace elements could be a new discriminant, because in the same possible source areas, there could be zircons with the same age, but bearing a different trace element fingerprint related to different source rocks. However, the geological setting of the Laurentia and Baltica areas, including the description of basalts to rhyolite association and the presence of ophiolitic slices, makes us confident about the derivation of Kinnekulle zircons and Cr-spinels from those areas.

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