1 Developing luminescence analysis of Icelandic volcanic glass: a case study

2 using the Þórsmörk Ignimbrite

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Abstract

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Large volcanic eruptions from Iceland can produce significant volumes of glass-rich rhyolitic tephra, which are then deposited across NW Europe and the North Atlantic-Arctic region, forming time-parallel marker horizons useful to palaeoenvironmental studies. Here we investigate new ways of improving the tephrochronological record of Iceland using (thermo)luminescence analysis of rhyolitic volcanic glass shards that dominate airfall ash deposits of the Þórsmörk Ignimbrite (ÞIG), tephra from the Askja 1875 AD, Öræfi 1362 AD eruptions, and the Ópoli tephra from NW Iceland. Following screening experiments, which showed that pure volcanic glass samples retained age-related TL signals, we undertook glassphase TL dating of the PIG and Óboli tephra. Our TL age estimate of c. 40 ± 10 ka for the PIG supports phenocryst-based radiometric ages of c. 50 ka rather than older age estimates of c. 200 ka. Results from the Óboli tephra were consistent with the fission track age established at c. 2 Ma age, but further investigations of high dose sensitivity changes and longer-term stability factors such as athermal fading are required for quantitative dating of volcanic glass deposits >100 ka. However, as thermoluminescence signals from purified glass fractions of Icelandic tephra can be obtained over 100-1,000,000-year time scales, luminescence characterisation of glass shards can be used alongside geochemical and morphological analysis to distinguish between distal tephras with similar geochemical signatures, and assist with tephrochronological investigations beyond the limits of radiocarbon dating.

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Keywords: Tephra; geochronology; geochemistry; Arctic; palaeoclimate

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Introduction

Large explosive rhyolite-producing eruptions from Iceland, such as the one responsible for the Þórsmörk Ignimbrite (PIG) in southern Iceland (Fig. 1), are comparatively rare, but particularly useful as they form time-parallel marker horizon, which provide chronological constraints for palaeoenvironmental records. As recent eruptions from Iceland have shown, volcanic ash from even comparatively small eruptions can be distributed across the North Atlantic and Arctic regions, and north-western Europe. Numerous studies have used of tephra layers to link ice, marine and terrestrial records of past environmental and climate change [1–30], but glacial erosion has removed many of the largest eruptions of the pre-Holocene, Quaternary era (>11.7 ka–2.5 Ma) from the Icelandic terrestrial record and some key large Pleistocene eruptions remain relatively poorly characterised and dated near their sources.

To be useful as time-parallel marker horizons, tephra deposits need to be fully characterised. Major and trace element geochemistry and phenocrystic composition are commonly used alongside shard morphology, texture and chronological data to distinguish between different tephra deposits, and correlate those that are similar [24, 27–42]. However, geochemical correlation based on glass major element geochemistry can sometimes be problematic. Multiple eruptions from the same magma body can produce geochemical heterogeneity within the glassy products of individual deposits [43, 44], but different eruptions from the same magma chamber over a period of time can also produce glass with broadly similar major element geochemical composition. Recent studies have shown there are significantly more visible and non-visible (crypto-tephra) Icelandic ash deposits in sediment cores extracted from the Arctic Sea and North Atlantic than previously thought, and not all can be distinguished by shard-specific major or trace element analysis [33, 37]. Widely dispersed

tephra deposits can also be re-deposited a long time after the eruption event by post eruption processes such as ice-rafting [45, 46] and several studies have confirmed that glass is not geochemically stable in all burial environments or able to withstand harsh laboratory digestion procedures intact [47, 48].

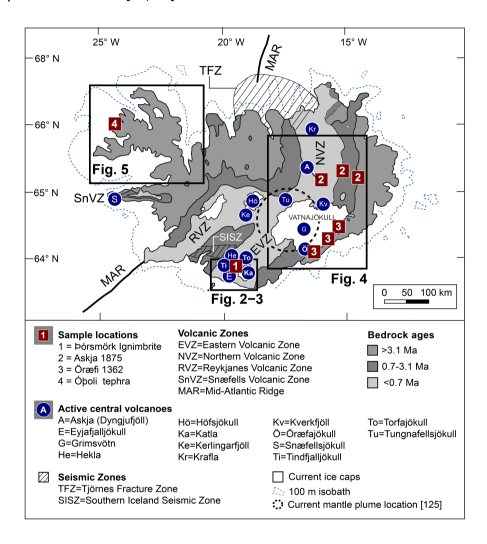


Figure 1. Summary geological map of Iceland showing active central volcanoes and sample locations [adapted from 11].

More than 300 historical eruptions have been documented or identified by geochemical analysis of tephra layers in Iceland in the last c. 1,100 years, on average, one every four years [7, 10, 11, 29, 49-59]. Tephra is the only product of more than 130 of these eruptions, and

over 75 % of these have produced a visible tephra layer on the Icelandic mainland [60]. However, rhyolitic tephra and large ignimbrite-forming eruptions are relatively uncommon in the Icelandic tephrochronological record [13, 15, 37, 53, 61].

Since rhyolitic magmas produce the most geochemically evolved and distinctive tephra, geochemical discrimination between different volcanic systems and/or multiple eruption events from the same volcanic system can be achieved. Moreover, large volumes of rhyolitic volcanic glass, which are unequivocally associated with the eruption event, form explosively on contact with water and/or ice during magma-quenching. For example, volcanic glass formed from the rhyolitic magma of the Öræfi 1362 AD eruption is geochemically homogenous and has <1% phenocrystic content [62]. Glass-rich volcanic ash ejected into the stratosphere can be distributed over a wide geographical area, forming long-distance chronological markers [63, 64]. Advantageously, selective removal of heavier phenocrystic components during long-distance stratospheric transport of ash provides a natural glass-purification process.

Radiation induced thermoluminescence (TL) is a well-established property of crystalline minerals such as quartz and feldspars [65]. Non-crystalline materials including synthetic glass [66,67] and naturally occurring volcanic glass (*e.g.*, obsidian) [68] also exhibit TL, though with lower sensitivities than most crystalline materials. A study of 55 archeaological silicate glass slices [66] showed high temperature TL sensitivities ranging from 10^{-1} to 10^{2} photon counts per mg per Gy in 10 degree centrigrade bands. By contrast TL sensitivities from quartz samples from diverse lithologies have been reported [111] as ranging from 10^{3} -2x10⁴ photon counts

per mg per Gy over similar temperature intervals (centred on 380 degrees centigrade), and from feldspars from 10^4 - 10^6 phioton counts per mg per Gy.

Therefore, though volcanic glass is a metastable material [72], luminescence signals from the glass phase of tephra provide a useful additional provenance and, potentially, chronological tool, especially in the c. 50-75 ka age-range, which is difficult to date using other radiometric methods (e.g., fission-track) and when suitable phenocrysts are absent [68–72]. Despite the relatively low signal levels, and variability in signal outputs due to retention of some fine crystalline components, pioneering luminescence studies from the early 1980s on fine-grained volcanic glass-rich fractions from tephra deposits from North America and New Zealand showed considerable potential [73–74].

Here, first, we investigate new ways of improving the tephrochronological record of Iceland bycharacterising the geochemical and TL properties of the (airfall) rhyolitic glass component that dominates tephra (>95%) from four key rhyolitic Icelandic eruptions of Late Quaternary age: the Þórsmörk Ignimbrite (ÞIG), the Óþoli tephra (ÓT), the Askja 1875 AD (A1875) and Öræfi 1362 AD (Ö1362) eruptions and (Figs. 1–6; Fig. S2, Table S1). Second, we focus on producing quantitative ages from the ÞIG and compare our results to published age estimates.

Based on geochemical correlation to potassium-argon dated phenocrysts from ignimbrite deposits outside of the Pórsmörk area [1, 2], the PIG was originally considered to be c. 200 ka. An argon-argon (Ar-Ar) age of 54.5 ± 2 ka [3] has been geochemically linked to the c. 55 ka North Atlantic Ash Zone-II (NAAZ-II) II-RHY-II deposit, a widespread and geochemically homogenous tephra deposit found in North Atlantic and Arctic Sea marine sediment cores

Greenland ice core records [3–5]. The eruption that formed the PIG is the same order of magnitude as the thickest rhyolitic tephra layers in the Holocene and historical tephrochronological record of Iceland, *e.g.*, Hekla 3, Hekla 4, Öræfi 1362 AD and Askja 1875 AD [11, 76, 88, 89].

The Ópoli tephra was found in plateau-top ice-dammed lake deposits of NW Iceland. Following initial TL analysis which suggested an early Quaternary age and has been fission-track dated to 2.26 ± 0.11 Ma [75]. In this study, the tephra deposit acts as an 'older' age-comparison for quantitative TL dating of purified volcanic glass from the PIG.

The A1875 and Ö1362 tephra deposits examined are from two of the largest rhyolite forming Plinian-style eruptions in the Icelandic historical record. These eruptions distributed large volumes of tephra across eastern Iceland and NW Europe. In this study, we use them as 'recent' TL age-controls because they have similar glass-phase rhyolitic geochemistry to the Óþoli tephra and the PIG.

Study sites

Pórsmörk Ignimbrite: The Pórsmörk Ignimbrite (PIG), first described by Thórarinsson [1] and named by Jørgensen [76, 77], is located in the transitional-alkalic province of the Eastern Volcanic Zone (Fig. 1). It is one of Iceland's largest eruptive bodies, covering an area of about 80 km² (Figs. 2, 3). Outcrops in the Pórsmörk area are the only known record of the PIGforming eruption in the Icelandic terrestrial tephra record. Intermittent volcanic aggradation and glacial erosion have exposed well-preserved stratigraphic sections along the Krossa and

Markarfljót valleys, with deposits north of the Krossa that are typically 10–50 m thick but can be up to 200 m thick [76, 77] (Figs. 2, 3). Accessible exposures, up to 30 m thick, are located on a 4–5 km stretch along the northern border of the Krossa (Figs. 2, 3). No basal contact is visible at most sites and only slope deposits are accessible at some sites [1, 76] (Fig. 3).

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The Þórsmörk area is surrounded by three ice-capped, active central volcanoes: Eyjafjallajökull to the south, Katla to the east, Tindfjalljökull to the north (Fig. 2b). Torfajökull, Iceland's largest active rhyolite volcano, is located further to the NE and Hekla is located further north beyond Tindfjalljökull. Torfajökull is amongst the most active rhyolitic volcanoes in the world with at least ten rhyolitic eruptions in the post-glacial period alone with the last occurring in ca. 1480 AD [78]. Tindfjalljökull is the least active central volcano in this region and classified as a dormant volcano because there have only been a few eruptions along its margins in post-glacial times [1, 79, 80]. Eruptions from Tindfjalljökull are typically dominated by slightly alkaline basalts and minor intermediate rocks and abundant sub-alkaline to slightly peralkaline rhyolites [76, 80-83]. Thórarinsson suggested that the summit of Tindfjalljökull is collapse caldera connected with the formation of the PIG [1, 76, 84]. All structural and depositional features of the PIG preserved in and around Pórsmörk are associated with emplacement from pyroclastic flows or surges [76, 77] (Fig. 3). Jørgensen [76, 77] found no evidence for a Plinian phase of activity, but abundant airfall ash deposits (e.g., Fig. 3i-l) could have been formed by explosively-generated pyroclastic flows, created by instabilities in the lower parts of the eruption column [85]. These combined into an ignimbrite-forming surge that mantled the existing topography, with welded and unwelded pumice and ash forced into valley troughs and sides. Given an average thickness of 20-25 m, Thórarinsson [1] calculated that the equivalent volume of the PIG as 1.5–2.0 km³. At least a further 2–3 km³ of freshly fallen tephra was probably dispersed over a wider area, but this has since been eroded by glacial activity.

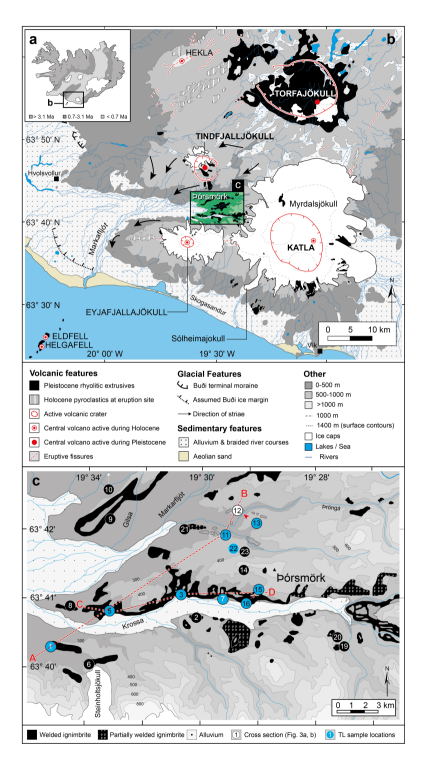


Figure 2. Regional setting and sampling locations for the Pórsmörk Ignimbrite. (a) Location map of Iceland; (b) Location map of Pórsmörk in relation to active volcanoes, pyroclastic and extrusive deposits and main glacial features of the Eastern Volcanic Zone (EVZ), Iceland. The

geology of area covered by the Þórsmörk box is basic and intermediate hyaloclastites and tuffaceous sediment younger than 0.7 Ma (based on the 1:250,000 Geological Map no. 6 of the Iceland Geodetic Survey). (c) Sketch map of main outcrops of the Þórsmörk Ignimbrite, and its broad classification into welded and unwelded outcrops [adapted from 76, 77] and thermoluminescence sampling locations (blue circles) investigated in this study. Some sample sites (e.g., PIG10) were located off-map, while others (e.g., PIG12) were inaccessible.

Thórarinsson [1] and Jørgensen [77] suggested a c. 200 ka age for the PIG by correlating deposits in Pórsmörk with potassium-argon (K-Ar) ages from geochemically similar deposits north of Pórsmörk [2, 4]. Zielinski et al. [9] later proposed that the PIG was the source of the rhyolitic component of North Atlantic Ash Zone-II (NAAZ-II), a widespread tephra deposit that has been found in numerous marine and ice cores across the North Atlantic and the Arctic Sea (Fig. S2). Dated to c. 50 ka, based on its correlation-based age in the GISP2 Greenland ice core of 53.5-55 ka [9, 86, 87], NAAZII has been geochemically linked with an argon-argon (Ar-Ar) dated PIG deposit of 54.5 ± 2.0 ka, sampled from an unspecified location [3].

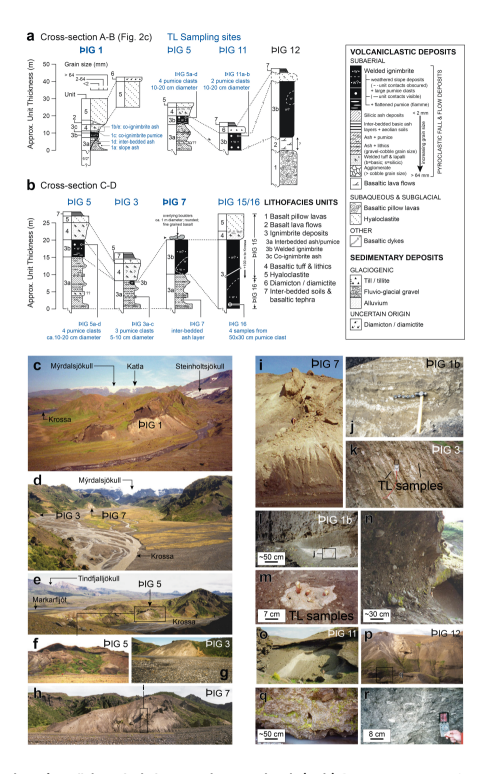


Figure 3. The Þórsmörk Ignimbrite, southern Iceland. (a–b) Summary composite schematic stratigraphic logs of main lithofacies units at principal sites and schematic cross section sketches along lines A-B and C-D; (c–r) Photographs of the Þórsmörk Ignimbrite sampling locations and luminescence sampling methods: (c) Site ÞIG 1, taken from the lateral moraine of Gigjökull, looking approximately E up the Krossa towards Mýrdalsjökull; (d) Sites ÞIG 3 & 7,

taken from Valahnjúkur looking approximately E up the Krossa towards Mýrdalsjökull; (e) Site PIG 5, taken from the top of site PIG 6, looking approximately NNE towards Tindfjalljökull in the distance; (f) Site PIG 5; (g) Site PIG 3 looking approximately W down the Krossa; (h) Site PIG 7 taken from the Krossa sandur looking N at sampling site, and erratics near the top of the deposit (circled); (i) TL sampling at site PIG 7 composed of inter-bedded ash, pumice and lithic layers beneath an outcrop of welded ignimbrite, similar to those at PIG 3, shown in (k); (j) TL sampling at PIG 1b in ash rich co-ignimbrite ash and pumice layers; (k) TL sampling of inter-bedded ash layers at PIG 3; (l) Co-ignimbrite ash, black tuff and capping hyaloclastites at site PIG 1b (Units 3c, 4, 5); (m) TL sampling of a large pumice clast, c. 25–30 cm in diameter, embedded in a welded ash and lithic matrix at site PIG 16; (n) Poorly sorted, rounded boulders and pebbles in loose soil and fine-grained matrix (diamicton) contact with welded ignimbrite at site PIG 11 (hammer for scale); (o) Site PIG 11 looking NNE (person at the bottom edge of the shadow for scale); (p) Site PIG 12 taken from the top of the hill at site PIG 13 looking approximately north; (q) Pillow lavas at the base of PIG 12; (r) Fiamme at the base of PIG 5 (Unit 3b).

Askja 1875 AD (A1875) tephra: Post-glacial volcanism in Iceland is concentrated in four separate areas within the fissure system of Dyngjufjöll (Askja), located approximately 100 km North of Vatnajökull in the Eastern Highlands of the tholeitic Northern Volcanic Zone (NVZ) of Iceland (Figs. 1, 4) [83]. Askja, the name of the main caldera and central volcano, covers an area of c. 45 km² (Fig. 4) and is bounded by steep cliffs, with Pleistocene-age rocks and predominantly basaltic composition forming the basement bedrock inside the crater (Fig. 4) [89]. Post-glacial lavas contain several rhyolitic ash layers, the oldest dated to c. 9,800 years BP [90]. Between 9,800-4,500 years BP, following rapid early Holocene deglaciation in Iceland,

volcanic productivity was 30 times greater than during the historical era [90]. The last Plinian eruption in the Dyngjufjöll volcano took place from within the Askja caldera a few hours before 8 a.m. on 29th March 1875 AD [90, 91]. The explosive phase of the eruption produced large volumes of fine-grained, grey-white, non-vesicular ash, which was dispersed across eastern Iceland by strong westerly winds forming visible layers 20-30 cm thick up to 50 km away [6]. The A1875 eruption is a classic example of how tephra from explosive Icelandic eruptions can be rapidly distributed across the North Atlantic and NW Europe [92, 93]. Tephra layers dominated by glass-shards invisible to the naked eye with a similar eruption-related geochemistry to A1875 deposits in Iceland have been found in Scandinavia [90, 94].

Öræfi AD1362: The Öræfajökull volcano, located in the Eastern Volcanic Zone on the southern margins of the Vatnajökull ice cap (Figs. 1, 4b), is Iceland's highest (2119 m) and most active composite stratovolcano during the post-glacial and Holocene era. There have only been two rhyolite-producing eruptions from Öræfajökull in the historical period: in 1362 AD and 1727 AD [11]. The 1362 AD eruption (henceforth referred to as Ö1362) was particularly explosive, producing >10 km³ of freshly fallen rhyolitc ash and pumice, and accompanied by two massive jökulhlaups from Falljökull and Rótarfjallsjökull. Although the eruption took place mainly in the caldera, and most of the fallout occurred over the sea, it is still the second most voluminous tephra deposit in Iceland in recorded history, after the more effusive and basaltic tephra producing Veiðivötn eruption of 1477 AD [58]. Up to 2 km³ of highly evolved rhyolitic (SiO2>70%) tephra fell on land with prevailing westerly winds transporting most of the ejecta ESE across the North Atlantic [11, 62, 95]. Tephra layers up to 10 cm thick have been found in soils covering an area of 4300 km² surrounding the main edifice [96].

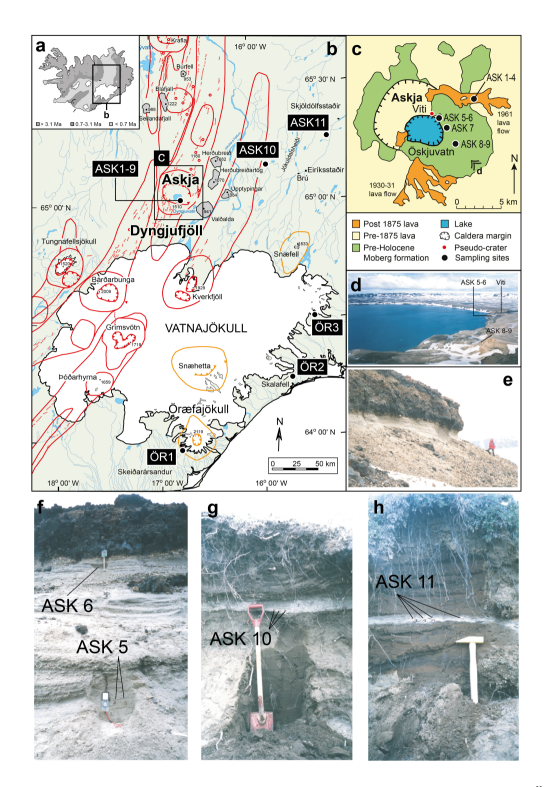


Figure 4. Historical eruption age-control study sites and sampling of A1875 and Ö1362 tephra deposits. (a) Location map; (b) Regional setting of the Askja (Dyngjufjöll) (red) and the Öræfi (orange) volcanic systems and Askja 1875 (ASK1-11) and Öræfi 1362 (ÖR1-3) sampling locations. Solid red & orange lines are outlines of the main volcanic systems; broken lines red and orange lines are fissure swarms; triangulated lines are the caldera margin; red circles are

craters and grey areas outlines of the main table mountains ('mobergs'). (c) Summary geological map of Askja adapted from [89] and showing the location of samples ASK 1-9. (d) Askja caldera from SE rim and ASK 5-9 sampling sites; (e) A1875, Layer 'D' [89] and capping lava flow; (f) TL sampling of Askja 1875 - ASK 5 (Layer 'C' [89] and ASK 6 (Layer 'D' [89]); (g) ASK 10 at Fiskidalur, 55 km NNE of Askja; (h) ASK 11 at Arnórsstaðamuli, 74 km NNE of Askja.

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Ópoli Tephra: The Ópoli tephra (ÓT) is a rhyolitic tephra deposit within a glacio-lacustrine sedimentary sequence near the plateau surface of Skagafjall [75, 97] (Fig. 5a-c). The ÓT sedimentary unit consists of 8-10 cm of airfall tephra overlain by up to c. 10 m of reworked volcanic ash (Fig. 5d). The airfall ash sub-unit is composed of more than 99% rhyolitic glass, with very minor basaltic-andesite/andesitic altered glass and sparse traces of feldspar, iron ore and pyroxene of uncertain origin [75, 97]. The 8-10 cm thick airfall ash is unusual for a tephra deposit on the NW (Vestfírðir) Peninsula, where visible tephra layers in Holocene peat deposits are uncommon, and usually less than a centimetre thick (Fig. 5d). Based on its stratigraphic position within an ice-dammed lake sequence in NW Iceland, the ÓT was originally thought to have been deposited c. 17-23 ka [97], but has since been dated to 2.26 ± 0.11 Ma using the isothermal fission-track method [75]. Prior to fission-track dating analysis, preliminary luminescence experiments on the airfall rhyolitic glass fraction revealed the ÓT to be older, with a minimum age estimate of at least c. 400 ka (Fig. 6f) [75]. The ÓT is most likely the product of a large Plinian-style eruption somewhere in the Húsafell region, a now inactive part of the SW Iceland axial rift (RVZ in Fig. 1) from near-central Iceland [75]. Prevailing north/north-north-westerly winds at the time of the eruption transported ash c. 200 km, depositing it on the heavily glaciated landscape of the Vestfírðir Peninsula during the

Late Pliocene [75]. The A1875 tephra is the closest geochemical match to the Óboli tephra (Figs. 1, S1) [75].

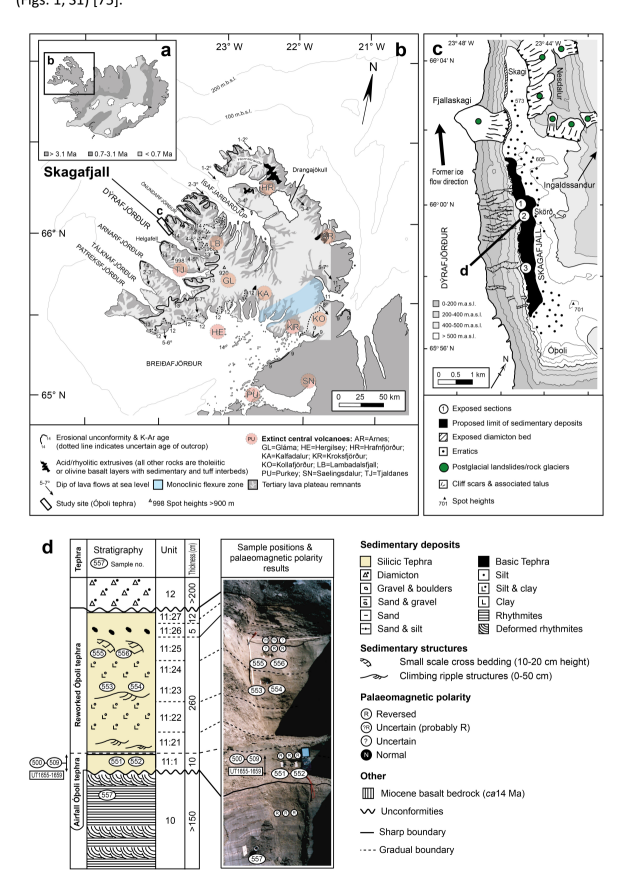


Figure 5. Regional setting, stratigraphic logs and sampling of the Óþoli tephra. (a–c) Location of Skagafjall and main geological features of the Western Troughs of the Vestfírðir Peninsula (based on [75, 97]). (d) Detailed stratigraphic log of and summary of palaeomagnetic results from Skagafjall section B units 10–12 and sampling strategy/sample numbers for the air fall and reworked components of the Óþoli tephra. Sample numbers 500-510 and 551-557 are SUTL laboratory numbers (SUERC-luminescence dating laboratory) sampled from the airfall tephra deposits in section B. SUTL558 is from section C. SUTL500-509 were sampled at 1 cm intervals from the basal 10 cm of Óþoli tephra above the rhythmite bed/tephra contact; UT sample numbers are University of Toronto fission-track laboratory numbers [97].

Methods

Luminescence dating relies on the laboratory reconstruction of the radiation dose absorbed by sensitive materials in the natural environment since formation or since the last heat- or light- induced resetting or 'zeroing' event [98–100]. For volcanic glass, the event being dated is the formation event, *i.e.*, cooling from magma, which due to its extremely high temperature, may be assumed to be free from residual thermoluminescence signals. Here, this assumption was verified, for practical purposes, using 19^{th} Century material from the A1875 eruption. A laboratory determination of the total radiation dose experienced since the formation or last bleaching event is referred to as the equivalent or palaeodose, D_e . This is obtained, following measurement of the "natural" luminescence signal in the laboratory by reconstructing a stepwise luminescence dose-response curve by laboratory irradiation, and remeasurement of stimulated luminescence from purified sub-samples (or aliquot discs). If D_T , the rate of radiation dose received by decay of radioactive elements, and cosmic radiation, *in-situ* in the natural environment, can be reliably assessed [99, 100], an age

estimate can be determined, simply, as the quotient of palaeodose (D_e) and environmental dose rate (D_T) or [D_e / D_T].

Sample collection and purification

We examined the luminescence characteristics of undisturbed airfall deposits to avoid possible complications associated with post-depositional reworking and light exposure. Samples were collected by hammering light tight 22–27 mm diameter, 20–50 cm long copper tubes into airfall tephra and pumice deposits that had been cut back 10–30 cm. All samples were transported and stored under light-tight conditions and 2–5 cm of material at the end of the tubes that could have been exposed to light was removed.

Contaminant mineral phases that also retain a luminescence signal, such as quartz, feldspar and zircon, have a higher density (>2.45 g cm⁻³) than rhyolitic glass (2–2.2 g cm⁻³). Therefore, double-purified sieved 90–150 µm volcanic glass fractions were prepared by density separation techniques [48, 73, 102]. Samples were cleaned ultrasonically in 10% hydrochloric acid (HCl) and density separated 2.3–2.4 or <2.4 g cm⁻³ using sodium polytungstate solution (Figs. S3). Harsh chemical treatments and etching were avoided due to the fragile nature of platey and pumiceous glass shards; hence, no attempt was made to clean the glass surfaces with hydrogen peroxide (H₂O₂) or etch the shards with hydrofluoric acid (HF) to remove their thin outer layer influenced by external alpha-dose rates. Contamination by non-glassy phases was assessed using TL sensitivity response, and visually confirmed by light microscopy and Scanning Energy Dispersive-Scanning SEM (EDS-SEM) geochemical mapping of discs used in TL analysis and by examination of preliminary N/50 Gy dose response curves (Fig. 6, S4). A summary

SAR-TL experiment

Based on findings from our preliminary screening experiments (Fig. 6), and after testing multiple aliquot methods [101], it was noted that the relatively high radiation doses needed to match natural signals from the early Quaternary Ópoli tephra analysis had potential to induce radiation colouring in the samples. Moreover, sensitivities in response to repeated 50 Gy irradiation and readout cycles tended to decline, suggesting that such radiation colouring was not being fully annealed by the TL heating cycle to 400 degrees. In recognition of this, to quantify equivalent doses and constrain ages for the Þórsmörk Ignimbrite and the Ópoli tephra, we adapted the Single-Aliquot Regeneration OSL (SAR-OSL) technique [103, 104] for TL analysis (SAR-TL). In the SAR method, the same disc is exposed to a series of laboratory radiation doses prior to preheating and readout, followed by a test dose (TD) measurement to assess laboratory induced sensitivity changes across the measurement cycle. The natural TL was measured from purified glass on each disc, and TL was then artificially regenerated beyond the naturally accumulated TL by applying a series of laboratory radiation doses to each disc.

Stainless steel discs were cleaned with acetone and Electolube SCO200D silicone grease then loaded with 4–6 mg of purified volcanic glass concentrated in the central area of each disc. Read-outs were undertaken using a Scottish Universities Environmental Research Centre (SUERC) TL Research Reader consisting of an Electron Tubes 9883 QB Photo-multiplier Tube (PMT) fitted with Schott 7/59 and KG1 filters and continuously flushed with nitrogen to prevent oxidising reactions that might create spurious TL. Measurements were undertaken manually with real time analysis to monitor glow shapes, assess integrated photon count

rates to define successive regeneration doses levels required, and check for signal quality. Discs were heated from room temperature to 400 °C at 5 °C s⁻¹. Blanks and blank contamination discs were cleaned with acetone and sprayed with Electolube SCO200D silicone grease, were placed on the work surface and read out at the end of each session. Discs for the N/50 Gy and SAR-TL experiment were irradiated using a shielded ELSEC 9022 irradiator fitted with a 1.85 GBq 90 Sr β -ray source with a source to sample distance of 15 mm and a silicate dose rate of 3.353 \pm 0.015 Gy min⁻¹ for the N/50 Gy experiment and 3.270 \pm 0.015 Gy min⁻¹ for the later SAR-TL experiments. Reconstructing the large natural signal retained in the Óþoli tephra proved particularly time consuming due to its large stored natural dose. For this experiment a higher dose rate irradiator incorporating a 11.1 GBq 90 Sr β -ray source at a working distance of 7.5 mmm was used, yielding a mean dose rate in the central areas of the sample of 32.0 \pm 0.6 Gy min⁻¹ Both irradiators had been calibrated to high precision relative to the UK air kerma standards at the National Physical Laboratory (NPL) when commissioned in 1997, before this study.

SAR-TL Palaeodose reconstruction (D_e)

After the natural TL signal had been thermally removed and measured (read-out), the TL and test dose and read-out measurement cycle for each disc consisted of: (1) Individual disc irradiation (PIG: 50, 75, 100, 125, 150, 200, 300 Gy; ÓT: 500, 1000, 1500, 2000, 3000, 4000, 5000, 6000 Gy); (2) A short, high temperature (within-chamber) pre-heat of 220 °C for two minutes and 0–400 °C TL read-out; (3) Test-dose (TD), pre-heat, as stage 2, and 0–400 °C TL read-out. Disc-specific changes in sensitivity were in stage 3 were corrected by applying a test-dose (TD) after each successive regeneration dose (PIG = 25 Gy; ÓT = 100 Gy). Palaeodose (De) values for the Pórsmörk Ignimbrite PIG7 and the Óþoli tephra SUTL551 samples were

calculated from standard error weighted regression of 200-400 °C, 10 °C integrated data (Tables 1, 2, S2, S4). The final weighted mean palaeodose values used in age estimates were calculated from the most thermally stable peak intensity regions of the glow curve, determined by a D_e plateau plot. In summary, D_e values were integrated over three temperature regions that correspond to the wide (280-360 °C), median (290-330 °C) and narrow (290-310 °C) TL intensity peak region of the glow curves (Figs. 7-8; Tables 1, 2). Uncorrected (UC) and test-dose (sensitivity) corrected (TDC) regeneration curves, were modelled using best mathematical fit (highest adjusted r² values, p-value <0.05) linear regression (y=ax+b) and non-linear least squares saturating exponential regression analysis (y=a[1-e^(-bx)]) applied to 10 °C interval datasets for both samples (Tables S2, S4). The nonlinear and non-saturating response of the Óboli Tephra was also modelled using saturating exponential + linear regression analysis $(y=a[1-e^{(-bx)}] + cx)$ [following 66]. Regression results shown in Tables 1 and 2 represent methods with the highest r² and lowest RMSE values (see also Table S2, S4). Other regression models (e.g., saturating exponential, 2nd and 4th order polynomial) were investigated (e.g., Table S4). Regression and statistical analysis was undertaken SigmaPlot Software California in (Systat Inc., San Jose USA, www.systatsoftware.com), XLSTAT (version 2010.3.09, www.xlstat.com).

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Fading tests were conducted at the end of the measurement cycle using multiple aliquot grouped approaches as suggested by Sanderson 1988 [116]. For this, two groups of discs were defined for each sample. As these had same prior radiation and TL readout histories as each other, both groups were expected to be in the same state, in respect of sensitivity and sensitisation behaviour. One of the groups was irradiated to a 50 Gy dose, at the start of the test period, and the other irradiated at the end of it -5×10^6 seconds. Both were preheated

together, and then readout, in their 9th readout cycle since the start of the experiment. These results were normalised to those obtained from a prompt readout in their 8th cycle. By double normalising the data sets and comparing the results from the two groups, the fading quotient associated with the dark storage for the period between prompt and stored readings was obtained. This approach provides for replication across many discs, and importantly is not affected by sensitivity changes in the samples (based on sensitisation between successive irradiation and readout cycles), or in the equipment (since the readout sessions for both groups of samples are conducted together both for the prompt and stored measurements). The fading observed crosses log time cycles 4-5 and 6-7.

A similar approach was adopted by Ward [67], who studied TL dose response, from 200 Gy to 160 kGy, and stability of highly reproducible mass produced and fully melted silicate glass slices (i.e., microscope cover slips). Fading tests using high dose (44 kGy) covered 11 logarithmically spaced post-irradiation intervals from 45 minutes after irradiation up to 35 days. After an initial rapid fading over the first few hours following irradiation, subsequent losses diminished in rate as seen on a logarithmic time scale and appeared to be dominated by thermal fading processes. On this basis, the procedure adopted in the study aimed to avoid the immediate post-irradiation period. It is recognised that fading rates over longer Quaternary timescales represent an extrapolation over any observable laboratory storage test. Therefore, the results of the study were also be assessed relative to independent age controls of the Ópoli tephra and PIG in this study.

Dose rate reconstruction (D_T)

The overall dose rate to the tephra shards includes internally generated alpha and beta contributions, and externally delivered alpha, beta, gamma components from the bulk sediment matrix, which are attenuated by water, plus an external cosmic dose rate component (Tables 3, 4). Bulk sample geochemical data, obtained by X-ray fluorescence (XRF) and Inductively Coupled Plasma-Mass Spectrometry (ICP-MS) analysis, was used to assess 'external' or 'bulk' matrix dose-rates. Bulk dose rate calculations were based on the infinite matrix assumption [100], which states that samples are essentially homogeneous on a macroscale. Shard-specific electron probe microanalysis (EPMA) potassium (K) data and ICP-MS analysis of the purified glass fraction U, Th and Rb data were used for (internal) glass-shard dose rate calculations, (D_Tshard_(in-situ-sat)) in Table 4, following [98, 100]. See Supplementary Note 1 for experimental procedures. We also compared dose rates derived from bulk and glass shard geochemistry with data derived from spectrometry measurements (Tables 3, 4, S5).

For the most realistic assessment of dose rate, we partitioned the shard alpha dose into an internal contribution, which is not subject to water-content attenuation, and an external alpha contribution, delivered across the pore space of the bulk matrix, which is attenuated by pore-water. For the final weighted mean glass-bulk dose rate model, $D_T^{glass-bulk}$ (in-situ-sat) (Table 4), we partitioned the dose rate into internal glass shard and external matrix components according to water content attenuation as follows. Samples for this study were collected during the warmest and driest parts of the year. Since the Icelandic climate is significantly colder and wetter outside our sampling months of June-August, we defined the mean water content as the mid-point of the collected and saturated water content and applied a 15% error to cover possible fluctuations in water content during burial [following 105].

Dry dose rates were attenuated by appropriate factors before calculation of final 'wet' dose rates using equations in [100], as follows: α -wet = α -dry / (1 + 1.5WF); β -wet = β -dry / (1 + 1.25WF); γ -wet = γ -dry / (1 + 1.14WF), where W and F are as defined W = (saturation wet weight - dry weight) / (dry weight). F = average fractional water content during burial history and was determined for two scenarios: as received water content (*in-situ*) and average palaeowater content, i.e., the mid-point between *in-situ* and saturated water content, with error limits that encompass the measured F_{in-situ} and W_{saturation} values. Errors in F_{in-situ} are mean \pm 1 σ of three representative measurements, while W_{saturation} errors are measurement errors from six duplicate measurements.

Internal (glass) and external (bulk) dose rates were based on geochemical and/or spectrometry data according to the following equations: $D\alpha_T = k[\varphi_g D\alpha_g + (1-\varphi_M)WFD\alpha_M]$ and $D_{\beta T} = \varphi_g D_{\beta} + (1-\varphi_M)WFD_{\beta M}]$, where: φ_g , φ_M are published absorbed dose factors and $(1-\varphi_M)$ attenuation factors [107–109] that correspond to the median grain size for glass shards and matrix, respectively. Spectrometry-based dose rates were calculated from *in-situ* portable gamma-spectrometry, thick source beta counting (TSBC) and alpha spectrometry (Table 4). Geochemical-based dose rates were calculated from bulk and shard-specific geochemical data in Tables 4 and S1 using well-established conversion factors [109]. No thoron or radon loss was assumed in conversion calculations because samples are believed to be sufficiently compact. External attenuation factors for U, Th and K taken from [108] were based on a median grain-size of 100 μ m. Equilibrium between U and Th decay was assumed as similarities between elemental and TSBC dose rates suggested that disequilibrium between decay products is minimal.

Results

Sample collection, purification & preliminary dose response screening

We collected light-tight TL samples from 17 out of 23 locations visited in the Pórsmörk region (Figs. 2, 3). Samples covered a wide range of rhyolitic material, including pumice, airfall ash layers dominated by glass shards, inter-bedded rhyolitic ash layers associated with pyroclastic flow deposits, and reworked ash deposits overlying the pumice-rich horizons (Fig. 3; Table S1). Sampled pumice clasts were unwelded and typically c. 10–50 cm in diameter (Fig. 3m). Ten light-tight samples were collected from airfall and reworked Ópoli tephra deposits (Fig. 5). Light-tight samples of A1875 tephra were collected from three locations: 0.5 km SE of Víti within the caldera (ASK 5) (ash-rich layers C & D [90]), 55 km to the ENE of Víti at Fiskidalur (ASK 10), and 74 km NNE of Víti at Arnórsstaðamuli (ASK 11), where visible layers, up to 1 cm thick exist (Fig. 4). Ö1362 tephra samples were collected from three sites at 9 km (Kvísker), 65 km (Hólmur), 86 km (Fóssdalur) to the East of the volcano along the principal eruption plume axis (Fig. 4). The major element geochemical composition of the rhyolitic glass-shards in the Ö1362 tephra is most similar to those of the Pórsmörk Ignimbrite (Fig. S2).

Visual inspection under the microscope revealed the rhyolitic ash and pumice Þórsmörk Ignimbrite samples were >95% glass, and the Óþoli tephra samples were >99% clear, platey glass shards of rhyolitic composition (Fig. 6). The bulk A1875 and Ö1362 samples were composed of >90–95% platey or pumiceous volcanic glass shards (Fig. 6). Þórsmörk Ignimbrite samples had the highest glass-shard alkali (Na+K) content of samples examined (Fig. S2). No purified fractions or SEM-EDS geochemical maps of discs used in SAR-TL experiments had

exceptionally high K, Na, Ca, indicative of possible feldspar contamination (K, Na, Ca)AlSi₃O₈,) (Fig. S5).

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Using the purified 90-150 micron volcanic glass shard fraction of the four deposits, we thermally read out the natural signal and examined their response to a 50 Gy radiation dose. We also undertook TL bleaching, TL inducement and pre-heating tests (Figs. 6, S5; Supplementary Notes 2, 3). Results showed an age-related signal was retained by volcanic glass (Fig. 6), and that a fairly consistent volcanic glass-phase response to 50 Gy laboratory radiation dose existed across all samples. This was characterised by a broad peak intensity region of up to 10³–10⁴ counts per second (cps) between 200–400 °C (Fig. 6a-d, f). The natural TL signal retained by the A1875 and Ö1362 samples was too small for further quantitative analysis using the existing set up (Fig. 6). Bleaching and inducement tests showed the TL signal could not be fully removed by optical exposure and that a TL signal of similar magnitude to the natural Ópoli tephra signal could not induced by optical exposure. Manual pre-heats of 220 °C for two minutes, 155 °C for five hours or 135 °C for 16 hours all successfully removed the laboratory-induced low temperature (c. 110 °C) TL peak observed in the natural and 50 Gy dose response experiment (Figs. 6, S5; see Supplementary Note 2 for experimental details).

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SAR-TL experiments & palaeodose reconstruction

The Þórsmörk Ignimbrite PIG 7 purified glass sample was significantly more sensitive (defined as TL intensity per unit applied dose) than the Óþoli Tephra sample and had a straightforward linear dose response in the 200–400 °C temperature region (Figs. 7a, 8a). Notably, SAR-TL regression curves used for final analysis of the PIG7 and the Óþoli Tephra samples had mean

 r^2 values consistently >0.98 with p< 0.0001 in the 240–360 °C peak intensity region (Tables 1,

520 2).

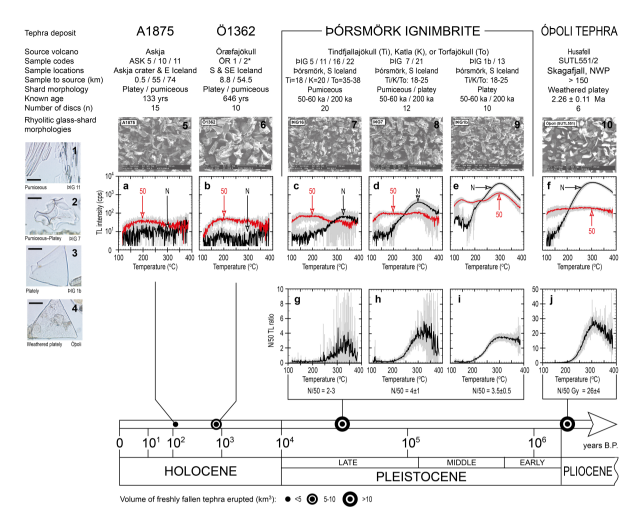


Figure 6. Preliminary natural versus 50 Gy (N/50 Gy) dose response tests for purified volcanic glass samples. (a)—(f) black lines represent the mean natural TL intensity (N) in counts per second (cps), at 1 °C intervals. Dark (red) lines represent the mean TL intensity produced by applying a 50 Gy laboratory dose. Light grey lines are ± 1σ errors from n discs shown at 1 °C intervals. Pre-heats were deliberately not applied prior to the N or 50 Gy TL read outs to allow examination of the natural and dose response glow curve shape. This creates a low temperature peak visible in some 50 Gy response profiles (e.g., PIG 1b and PIG 13). As no pre-heats or regression analysis were undertaken, the information contained in this figure is not linked to age estimates obtained from the SAR-TL experiment. (g)—(j) Natural and 50 Gy ratio plateau plots, showing the mean ratio and single standard deviation of the N/50 Gy TL response; *=ÖR 3 was visibly contaminated by minerals unrelated to the eruption

event and is not included for clarity. Microphotographs 1–4, taken at x400 magnification, illustrate the difference between pumice and platey glass shard morphologies; black scale bar is 25 mm; purified samples were mounted in Canada Balsam. Scanning Electron Microscope (SEM) microphotographs 5–10 show selected Ö1362, A1875, PIG7 and ÓT purified glass shards on discs used in the preliminary N/50 Gy experiment. Discs were carbon coated for SEM-EDS geochemical mapping causing low image resolution.

For PIG7, the uncorrected natural TL lies within an approximately linear growth phase of the regeneration curve, enabling reliable D_e values to be obtained by interpolation (Fig. 7b). Neither the type of regression curve fitted or the temperature integral chosen (280–360 °C , 290–330 °C or 290–310 °C peak intensity regions of the glow curve) made a significant impact on the reconstructed D_e value (Figure 7, Table 1). The PIG7 290–310 °C peak intensity region had the most stable D_e plateau region with the lowest measurement errors (Fig. 7c) and was chosen for D_e analysis. Uncorrected weighted linear (LIN) regression applied to 290–310 °C data produced a D_e value of 169 ± 4 Gy, while weighted saturating exponential (SE) regression produced a D_e of 165 ± 4 Gy (Fig. 7b; Table 1).

There was an approximately linear increase in TL intensity for PIG7 25 Gy test-doses TD1–TD9 (Fig. 6a–b). The PIG7 test-dose sensitivity-corrected 290–330 °C peak intensity weighted mean D_e values were 221 ± 6 Gy (LIN) and 217 ± 5 Gy (SE) (Fig. 7e). Peak region (290–310 °C) D_e values were similar (227 \pm 8 Gy (LIN); 225 \pm 8 Gy (SE)). Athermal fading did not appear to be a significant for PIG7, with 96% of the signal retained in the 290–330 °C and 99% in the 290–310 °C stable peak temperature regions after 60-day storage (Tables 1, S3). Test-dose (sensitivity) and 60-day storage loss-corrected SAR-TL D_e values across the 280–360 °C region

range from 226 \pm 7 Gy to 234 \pm 11 Gy (Table 1). The most accurate assessment of the PIG7 palaeodose is considered to be the weighted mean sensitivity and 60-day storage-corrected SAR-TL D_e value of 230 \pm 11 Gy, obtained by weighted linear regression from the 290–310 °C peak intensity temperature region (Table 1).

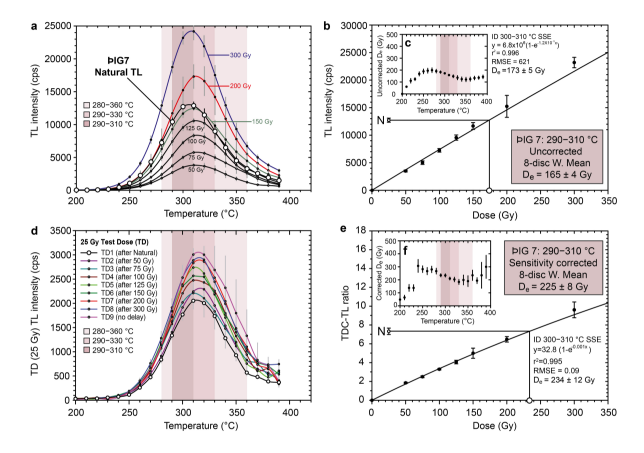


Figure 7. SAR-TL palaeodose (D_e) data for PIG 7 of the Pórsmörk Ignimbrite: (a) Natural and applied dose mean TL intensity glow curves. Weighted mean TL intensity standard measurement errors were $10.48 \pm 1.32\%$ in the 280-360 °C peak intensity region, $5.93 \pm 0.66\%$ for 290-330 °C and $5.95 \pm 0.95\%$ for 290-310 °C; **(b)** Example of a saturating exponential (SE) weighted regression curve fitting of the form $y=a[1-e^{(-bx)}]$ in the peak TL intensity 300-310 °C region. Similar SSE weighted regression was undertaken on 10-degree interval data between 200-400 °C to generate the D_e plateau plot shown in (c) (Tables 1, S2); **(c)** 200-400 °C D_e and temperature plateau plot; **(d)** 25 Gy test-dose (TD) mean TL intensity

glow curves; **(e)** Saturating exponential (SE) weighted regression of 300–310°C temperature test-dose (sensitivity)-corrected data (see also Fig. S6); **(f)** 200–400 °C D_e temperature test-dose (sensitivity) corrected D_e plateau plot. The weighted mean 290–310 °C sensitivity corrected D_e = 225 ± 8 Gy. Results are based on 10 °C integration of original 1 °C data (Tables 1, S2).

| Þórsmörk Igni | mbrite Palaeodose | Statistics & Age | estimates | | Weighted mean age (no fading | 60-day storage | 60-day storage | 60-day storage loss | 60-day storage loss corrected weighted | |
|---|-------------------|--------------------------|--|------------|---|---------------------------------|---|---|---|--|
| ÞIG7 SAR-TL | | Weighted mean palaeodose | 8-Disc Mean Regression Error Analysis (all | | correction) (D _T ^{glass-bulk}) _{(in-} | loss correction (Fraction of | loss corrected weighted mean palaeodose | mean age (D _T ^{glass-bulk}) _(in-situ-sat) | mean age $(D_T^{glass-bulk})_{(in\text{-situsat})}$ | |
| | | D _e ± SE | Mean | Mean RMSE | situ-sat) 5.82 ± 1.48 | signal retained) | D _e ± SE | 5.82 ± 1.48 mGy a ⁻¹ | 6.25 ± 1.51 mGy a ⁻¹ (ka) | |
| | | (Gy) | $r^2 \pm 1\sigma$ | ±1σ | (ka) | | (Gy) | (ka) | | |
| - | Linear regression | n | | | | | | | | |
| | (a) 280-360 °C | 157 ± 3 | 0.973 ± 0.025 | 639 ± 253 | 27 ± 7 | 0.94 ± 0.04 | 167 ± 8 | 29 ± 7 | 27 ± 7 | |
| | (b) 290-330 °C | 166 ± 4 | 0.991 ± 0.009 | 527 ± 244 | 29 ± 7 | 0.96 ± 0.02 | 173 ± 5 | 30 ± 8 | 28 ± 7 | |
| Uncorrected | (c) 290-310 °C | 169 ± 4 | 0.997 ± 0.003 | 352 ± 112 | 29 ± 7 | 0.99 ± 0.03 | 171 ± 7 | 29 ± 8 | 27 ± 7 | |
| Oncorrected | Saturating expon | ential regression | | | | | | | | |
| | (d) 280-360 °C | 151 ± 3 | 0.987 ± 0.009 | 511 ±93 | 26 ± 7 | 0.94 ± 0.04 | 160 ± 8 | 28 ± 7 | 26 ± 6 | |
| | (e) 290-330 °C | 158 ± 3 | 0.995 ± 0.003 | 543 ±62 | 27 ± 7 | 0.96 ± 0.02 | 165 ± 5 | 28 ± 7 | 26 ± 6 | |
| | (f) 290-310 °C | 165 ± 4 | 0.997 ± 0.003 | 528 ±65 | 28 ± 7 | 0.99 ± 0.03 | 167 ± 7 | 29 ± 7 | 27 ± 7 | |
| | Linear regression | n | | | | | | | | |
| | (a) 280-360 °C | 220 ± 5 | 0.984 ± 0.018 | ### ± 0.05 | 38 ± 10 | 0.94 ± 0.04 | 234 ± 11 | 40 ± 10 | 37 ± 9 | |
| | (b) 290-330 °C | 221 ± 6 | 0.996 ± 0.004 | ### ± 0.03 | 38 ± 10 | 0.96 ± 0.02 | 231 ± 8 | 40 ± 10 | 37 ± 9 | |
| Test dose- (Sensitivity-) corrected | (c) 290-310 °C | 227 ± 8 | 0.999 ± 0.001 | ### ± 0.03 | 39 ± 10 | 0.99 ± 0.03 | 230 ± 11 | 39 ± 10 | 37 ± 9 | |
| | Saturating expon | ential regression | | | | | | | | |
| | (d) 280-360 °C | 215 ± 5 | 0.993 ± 0.009 | ### ± 0.06 | 37 ± 9 | 0.94 ± 0.04 | 229 ± 11 | 39 ± 10 | 37 ± 9 | |
| | (e) 290-330 °C | 217 ± 5 | 0.999 ± 0.001 | ### ± 0.04 | 37 ± 10 | 0.96 ± 0.02 | 226 ± 7 | 39 ± 10 | 36 ± 9 | |
| | (f) 290-310 °C | 225 ± 8 | 0.999 ± 0.001 | ### ± 0.05 | 39 ± 10 | 0.99 ± 0.03 | 228 ± 10 | 39 ± 10 | 36 ± 9 | |

Table 1. Summary palaeodose statistics and age estimates for the PIG7 sample. The weighted mean test-dose (sensitivity) and 60-day storage-loss corrected palaeodose and age estimates shaded in dark grey and in bold are considered to be reliable. Best fit regression analysis was determined from highest adjusted r^2 values. Ten-degree integrated ranges for the 280–360 °C, 290–330 °C and 290–310 °C TL peak intensity regions are shown. The 290–310 °C and 290–330 °C ranges (dark grey shading) have the lowest TL intensity measurement errors, the highest 8-disc mean r^2 , lowest RMSE values and lowest p-values. This produced sensitivity and 60-day storage loss corrected palaeodoses of c. 230 \pm 10 Gy and ages of 40 \pm 10 ka. Data have been rounded to the nearest 10 Gy and 5 ka to reflect calculation errors,

In the 50 Gy dose response screening experiment, the natural TL signal from the Óþoli tephra formed 'broad' unimodal peak of 10³–10⁴ cps °C⁻¹ between 280–360 °C (Fig. 6), indicating a

TL signal that is predominantly stored in the more stable, longer-term energy traps evicted at high temperature. In the SAR-TL experiment, the uncorrected natural TL peak intensity region of the glow curves from the Óþoli tephra (ÓT) covered a broader temperature range, 290–330 °C, than PIG 7 (Fig. 8a). The broader shape of the regeneration dose response curve alone implied that the ÓT was significantly older than the PIG7 sample (Fig. 8b). Regression results show that the ÓT natural signal was located in an approximately linear growth phase *after* an initial non-linear growth phase, best modelled by a saturating exponential plus linear regression fit (Fig. 8b; Table S4). We chose this type of regression because it had a consistently higher r^2 and lower RMSE values than single saturating exponential or polynomial (quadratic) regression models (290-330 °C region mean \pm 1 σ r^2 = 0.995 \pm 0.001; p<0.0001; Figs. 8b, S6; Tables 2, S4). The 290–330 °C peak intensity region had the most stable D_e plateau, and produced a weighted mean D_e value of 4581 \pm 350 Gy (Fig. Table 2).

Assessing sensitivity change for the ÓT was more complex than for PIG7, with TL intensity reversals between some adjacent 100 Gy test-dose runs (Fig. S6c–d). These non-linear effects could be due to the (age-related) highly weathered and comparatively hydrated state of the ÓT glass shards, the high laboratory dose rate, the higher magnitude of applied doses (>1 kGy), and/or cumulative radiation induced sensitivity changes (such as radiation 'colouring'). If the ÓT test-dose corrected response curve continued to grow in the same approximately linear manner beyond 6 kGy, an applied dose in excess of 10 kGy would be required to produce an interpolated sensitivity-corrected palaeodose estimate (Fig. 8e; Table 2). It was not possible to apply such a high dose in the time available. Moreover, athermal fading appeared to be a significant factor for the ÓT sample examined, with 73% the signal retained

in the 290–330 °C and 79% retained in the 290–310 °C stable peak temperature regions after 60-day storage (Tables 2, S3).

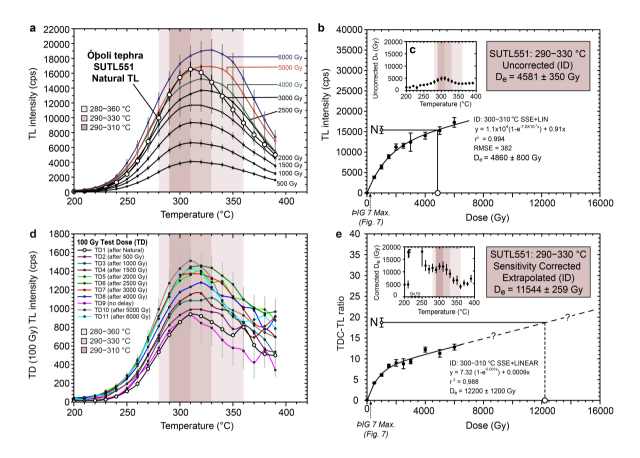


Figure 8. SAR-TL D_e data for the Ópoli tephra airfall ash sample SUTL551: (a) Natural and applied dose mean TL intensity glow curves. Weighted mean uncorrected TL intensity measurement errors were 6.40 \pm 0.11 % for the 280–360 °C peak intensity region, 6.37 \pm 0.06% for 290–330 °C and 6.40 \pm 0.13% for 290–310 °C; (b) Example of a saturating exponential (SE) plus linear (LIN) weighted regression curve fitting y = a[1-e^(-bx)]+cx in the peak TL intensity 300–310 °C region. Similar SSE+LIN weighted regression was undertaken on 10 degree interval data between 200-400 °C to generate the D_e plateau plot shown in (c) (Tables 2, S4); (c) 200–400 °C D_e vs temperature plateau plot; (d) 100 Gy test-dose (TD) mean TL intensity glow curves; (e) SE+LIN weighted regression of 300-310°C temperature peak region, highlighting how the reconstructed natural test-dose (sensitivity) corrected is greater than

the highest applied dose of 6000 Gy (see also Fig. S6); **(f)** 200–400 °C (D_e) test-dose (sensitivity) corrected palaeodose (D_e) plateau plot. Results shown are based on 10 °C integrals with standard measurement error from measured 1 °C data (see also Tables 1, S3, S4).

| Óþoli Tephra | Palaeodose St | atistics & Age e | stimates | | Weighted mean age (no fading | 60-day storage loss correction | 60-day storage loss corrected | 60-day storage loss corrected | 60-day storage loss corrected | |
|-----------------------------------|----------------|--------------------------------|--|------------------|--|--------------------------------|----------------------------------|--|--|--|
| Óþoli Tephra (SUTL 551) SAR-TL | | Weighted mean palaeodose | 8-Disc Mea Regression Err (all p<0.0 | or Analysis | correction) (D _T ^{glass-bulk}) _{(in-} | (Fraction of signal retained) | weighted mean palaeodose | weighted mean age estimate | weighted mean age estimate | |
| | | D _e ± SE (Gy) | ` . | Mean RMSE ±1σ | (D _T ⁹¹⁸⁸⁸ 283) _(in-situ-sat) 2.83 ± (ka) | | D _e ± SE (Gy) | (D _T ^{glass-bulk}) _(in-situ-sat) 3.97 ± 0.47 (ka) | (D _T ^{glass-bulk}) _(in-situ-sat) 2.83 ± (ka) | |
| | (a) 280-360 °C | | 0.995 ± 0.0015 | 331 ± 26 | 1178 ± 100 | 0.74 ± 0.03 | 4505 ± 278 | 1135 ± 90 | 1592 ± 149 | |
| Uncorrecte | (b) 290-330 °C | 4581 ± 350 | 0.995 ± 0.0012 | 342 ± 29 | 1619 ± 168 | 0.73 ± 0.02 | 6294 ± 511 | 1585 ± 151 | 2224 ± 239 | |
| ~ | (c) 290-310 °C | 4549 ± 524 | 0.995 ± 0.0002 | 355 ± 38 | 1607 ± 217 | 0.79 ± 0.01 | 5764 ± 668 | 1452 ± 184 | 2037 ± 277 | |
| Test-dose | (a) 280-360 °C | 8173 ± 153 | 0.986 ± 0.003 | 0.68 ± 0.13 | 2888 ± 211 | 0.74 ± 0.03 | 11040 ± 493 | - | - | |
| | (b) 290-330 °C | 11544 ± 259 | 0.988 ± 0.003 | 0.58 ± 0.09 | 4079 ± 302 | 0.73 ± 0.02 | 15861 ± 562 | - | - | |
| corrected | (c) 290-310 °C | 11539 ± 302 | 0.986 ± 0.003 | 0.63 ± 0.09 | 4077 ± 307 | 0.79 ± 0.01 | 14622 ± 426 | - | - | |

Table 2. Summary palaeodose statistics and age estimates for the Óþoli tephra, sample.

Ten-degree integrated intensity ranges for the three peak temperature plateau regions 280–360 °C, 290–330 °C and 290–310 °C are shown. The weighted mean sensitivity and 60-day storage loss-corrected palaeodose and age estimates highlighted in light grey overlap the fission-track age of 2.26 ± 0.11 Ma [75].

Dose rates and age estimates

Environmental dose rates for PIG7, PIG1b and Óþoli tephra samples shown in Tables 3, 4 and S5 were produced using well-established equations and dose rate conversion factors. For PIG7, the spectrometry-based dose rate of 5.82 ± 1.48 mGy a⁻¹ and geochemistry-based dose rate of 6.25 ± 1.51 mGy a⁻¹ in combination with the sensitivity and 60-day storage loss-corrected palaeodose values of c. 230 ± 10 Gy produced age estimate of 40 ± 10 ka. High errors of c. 25% are mainly due to large uncertainties associated with estimating palaeowater content fluctuations. Due to erratic changes in sensitivity and unresolved, but significant, athermal signal losses, we cannot currently reconstruct a reliable SAR-TL age estimate for the Óþoli tephra sample. Interestingly, though, the minimum uncorrected age shown in Table 2

is 1178 ± 100 ka or 1.18 ± 0.1 Ma, and the 60-day storage loss-corrected SAR-TL D_e value of 6294 ± 511 Gy and a geochemical dose-rate of 2.83 ± 0.41 mGy a^{-1} produces an age estimate of $2,224\pm239$ ka (2.22 ± 0.24 Ma) (Table 2), which is similar to the Óþoli tephra glass-shard fission-track age of 2.26 ± 0.11 Ma [75].

| Geochemistry | | Latitude | Longitude | Sampling | D _{cosmic} ¹ | Elemental concentrations | | | | | | | |
|--------------|-----------|----------|-----------|------------|----------------------------------|--------------------------|-------------------------------|-------------|------------|----------|--|--|--|
| Tephra | Matrix | (°N) | (°W) | Depth (m) | dose rate | _ | K or K ₂ O* (wt %) | U (ppm) | Th (ppm) | Rb (ppm) | | | |
| Lab. Code | | | | | (mGy a ⁻¹) | n | | | | | | | |
| Þórsmörk | (Ignimbr | ite | | | | | | | | | | | |
| ÞIG 7 | Bulk | 63°40.37 | 19°34.37 | 1 ± 0.5 | 0.16 ± 0.06 | - | 3.03 ± 0.15 | 3.80 ± 0.19 | 13 ± 0.7 | 107 ± 5 | | | |
| | Glass | | | | | 20 | 4.07 ± 0.19* | 4.37 ± 0.22 | 14.3 ± 0.7 | - | | | |
| ÞIG 1b | Bulk | 63°40.35 | 19°36.22 | 20 ± 5 | 0.04 ± 0.01 | - | 3.28 ± 0.16 | 3.13 ± 0.16 | 10 ± 0.5 | 73 ± 4 | | | |
| | Glass | | | | | 24 | 4.11 ± 0.61* | 3.39 ± 0.17 | 11.6 ± 0.6 | - | | | |
| Óþoli Tephra | | | | | | | | | | | | | |
| SUTL551 | /2 Bulk | 65°59.78 | 23°45.40 | 5 ± 1 | 0.11 ± 0.04 | - | 2.16 ± 0.11 | 2.26 ± 0.03 | 8.5 ± 0.4 | 55 ± 3 | | | |
| Glass | | | | | | 137 | 2.75 ± 0.14* | 2.60 ± 0.10 | 10.8 ± 0.5 | - | | | |

Table 3. Location and geochemical data used to calculate dose rates for the Þórsmörk Ignimbrite and Óþoli tephra samples. 1 The cosmic dose rate calculation assumes no change in sediment depth throughout burial and was calculated from $0.21e^{\left[-0.07(dr)+0.0005(dr)^2\right]}$ mGy a^{-1} [following 130] where d is current depth of burial and r is the density of the attenuating medium where r = 0.998 g cm⁻³ for water and 2.6 g cm⁻³ were used for sediment; n = number of shards analysed (Supplementary Note 1 and Table S1 for details).

| Dose rate data | | Glass shard dose rates | | | | | | Bu | lk mat | Weighted mean | | |
|----------------|-------------------------|------------------------|-----------------|-----------------|------------------|--|------------------------|-----------------|-----------------|------------------|--|---|
| Tephra | Measurement | $D\alpha_T$ | Dβ _T | Dγ _T | D _{cos} | (D _T glass) _(in-situ to sat) | $D\alpha_T$ | Dβ _T | Dγ _T | D _{cos} | (D _T bulk) _(in-situ-sat) | (D _T glass-bulk) _(in-situ to sat) |
| Lab ID | technique | (mGy a ⁻¹) | | | | (mGy a ⁻¹) | (mGy a ⁻¹) | | | | (mGy a ⁻¹) | (mGy a ⁻¹) |
| Þörsmó | rk Ignimbrite | | | | | | | | | | | |
| ÞIG 7 | α -dose excluded | - | - | - | - | - | - | 2.81 | 1.07 | 0.16 | 2.85 ± 0.60 | 2.85 ± 0.60 |
| | Spectrometry* | 1.84 | 2.12 | 1.09 | 0.16 | 5.22 ± 2.11 | 3.04 | 2.12 | 1.07 | 0.16 | 6.39 ± 2.07 | 5.82 ± 1.48 |
| | Geochemical | 1.84 | 2.09 | 1.09 | 0.16 | 5.18 ± 2.38 | 3.04 | 3.32 | 1.09 | 0.16 | 6.97 ± 1.96 | 6.25 ± 1.51 |
| ÞIG 1b | α-dose excluded | - | - | - | - | - | - | 2.55 | 1.09 | 0.05 | 2.96 ± 0.57 | 2.96 ± 0.57 |
| | Spectrometry* | 1.72 | 1.91 | 1.11 | 0.04 | 4.78 ± 2.12 | 2.43 | 1.91 | 1.09 | 0.05 | 5.48 ± 2.03 | 5.14 ± 1.47 |
| | Geochemical | 1.71 | 2.30 | 1.11 | 0.05 | 5.17 ± 2.64 | 2.43 | 3.12 | 1.11 | 0.05 | 6.71 ± 2.14 | 6.10 ± 1.66 |
| Óþoli Tephra | | | | | | | | | | | | |
| | α-dose excluded | - | _ | - | - | - | - | 1.59 | 0.56 | 0.11 | 1.83 ± 0.17 | 1.83 ± 0.17 |
| | Spectrometry* | 1.30 | 1.11 | 0.30 | - | 2.70 ± 0.62 | 1.51 | 1.11 | 0.30 | - | 2.92 ± 0.54 | 2.83 ± 0.41 |
| | Geochemical | 1.30 | 1.65 | 0.85 | 0.11 | 3.90 ± 0.66 | 1.51 | 1.58 | 0.85 | 0.11 | 4.04 ± 0.67 | 3.97 ± 0.47 |

Table 4. Summary dose rate data for the Pórsmörk Ignimbrite and Óþoli tephra samples. Summary dose rate values were calculated with α -dose rate excluded, using spectrometry data (α -dose included) and using geochemical data (α -dose included) for three scenarios:

glass shard, bulk matrix, and weighted mean (glass shard–bulk) matrix. Values in bold are considered most likely. For geochemical calculations, the glass shard matrix dose rate was calculated using well-established equations and conversion factors [107–109, 130–133]. Since the biggest source of uncertainty is variations in palaeowater content, using more recent conversion factors does not significantly alter these calculated dose rates or errors. Total alpha, beta and gamma dose rates ($D\alpha_T$, $D\beta_T$, $D\gamma_T$) incorporate attenuation factors based on *in situ*–saturation water content values [105, 132], which we consider to be most representative of the average water content through burial history at both the PIG 7 and Óþoli tephra sites (see Methods, Table S5). D_{cos} is the cosmic dose rate shown in Table 3. Errors for dose rate components are not shown for clarity but are typically $\pm 10\%$.

Discussion

We discuss the accuracy of the new PIG7 TL age estimate, examine implications of the age obtained, and make suggestions for future TL analysis of glass-rich tephra deposits based on the results from the PIG7 and Óþoli tephra experiments.

Experimental Implications

Our experiments on the Pórsmörk Ignimbrite (PIG7) produced age estimates that were similar orders of magnitude to published ages, highlighting that a glass-formation age-related TL signal is retained by the rhyolitic volcanic glass component of Icelandic tephra. In palaeodose reconstruction, even the smallest laboratory doses are applied at rates which are several orders of magnitude greater than those experienced in the natural environment. This creates two problems. First, high laboratory doses tend to fill or saturate shallow, thermally unstable traps, a process which does not occur during slower natural irradiation processes and may

result in differences in deep trap sensitivity [99, 110, 111]. Second, although laboratory dose rates less than c. 10 Gy min⁻¹ are not thought to be problematic [100], studies have shown that laboratory irradiation at 10⁸–10¹¹ (c. 0.1–100 Gy min⁻¹) times greater than that experienced in the natural environment can lead to defect creation, migration and, ultimately, defect complex creation, which does not occur naturally [121]. Glass should be one of the materials least affected, because localised charge transport is not favourable for shallow-deep trap competition. The erratic sensitivity changes observed for the Ópoli tephra in the SAR-TL experiment suggest the comparatively higher laboratory dose rates (an order of magnitude greater than for the PIG7 sample) were detrimental in this respect. The obvious solution is to reduce the applied dose rate, but this would be extremely time consuming for multiple discs with several applied doses runs >1000 Gy.

Further investigation into the influence of sample age and applied dose on athermal signal loss is recommended. This effect was minimal for the PIG sample but appears to be a significant issue for the much older Óþoli tephra, which received applied doses in the c. 1-6 kGy range. From our investigations, the athermal signal loss characteristics of 'younger' volcanic glass deposits appears to be similar to those of commercial glasses, which do not exhibit significant athermal losses at 'low' applied doses i.e., <100 kGy [66]. Compositional differences, additional lattice defects and/or weathering could explain the lower dose threshold observed in our experiments.

Athermal fading of stored TL in phenocrysts of volcanic origin is thought to be responsible for significant D_e underestimates in OSL dating of quartz phenocrysts present in some tephra deposits from Japan [112, 114, 115] and infra-red stimulated luminescence (IRSL) dating of

volcanic feldspar and other phenocrysts in the Old Crow Tephra, Alaska [113]. 'Young' volcanic glass shards appear less susceptible to long-term athermal signal loss at low irradiation doses [65, 66, 110-115], perhaps because fewer opportunities for long-range charge transport exist in volcanic glass compared to mineral-based luminescence systems. The relatively small TL signals per unit dose and low short-range order of glasses may also result from increased opportunities for competing non-radiative relaxation. At low doses, recombination is most likely to take place in hole centres populated from non-correlated ionising events, leading to the type of reproducible dose response curves observed in the PIG7 sample. At higher doses (e.g., >1 kGy) [66], the increased proximity between donors and acceptors pair may eventually favour tunnelling recombination and limit stability, as seen by the greater athermal losses in the Óþoli tephra.

By calculating the ionisation density as a function of dose and partitioning the dose (energy per unit mass) amongst the atoms in feldspar, Sanderson [116] showed that the mean spacing of randomly positioned charge-carriers in a lattice irradiated at low doses was greater than 1000 lattice units. The random spatial model for fading in calcite [117], which has been subsequently applied to feldspars [116, 118], other minerals [119], and volcanic products [120] is perhaps less relevant for volcanic glass since it cannot support high rates of tunnelling at low dose without some form of charge-defect clustering, leading to deviations from the logarithmic (1/t) decay behaviour predicted by the Visocekas [117] model. Ward [67] showed that fading from glass cover slips across a range of doses up to 160 kGy exhibits logarithmic (1/t) decay behaviour only at the highest doses used. At very high doses, the proximity of neighbouring ion pairs becomes sufficiently close to enable anomalous signal losses from the glass matrix by quantum mechanical tunnelling [cf. 116], as observed by high dose

experiments from glass slices [66, 67, 111]. In fact, all materials should exhibit (quantum mechanical) tunnelling as the mean concentration of electron-ion pairs reaches the percent level. If 1 kGy corresponds to 10 ppm initial ionisation, the mean concentration of electron-ion pairs reaches the percent level at doses in the 10's-100's kGy region. At low doses, thermal fading occurs from the continuum of traps where charge is located, and is presumably lost to non-luminescence processes, but this is not necessarily a significant barrier to successful dating providing the stability of the signal can be demonstrated by, for example, the existence of stable thermal plateau plots.

In early work examining a range of glass-rich tephras Berger [73,74], expressed concern that feldspar phenocrysts had potential to dominate fading behaviour for impure samples. He noted the apparent stability of the glass rich tephra from the Mazama, tephra and suggested that the glass phase experienced long-term athermal fading. Following the demonstration that tephra showed weak OSL and IRSL response [75], Auclair [113] studied the potential of IRSL dating from Crow Lake tephra from Alaska, in conjunction with stratigraphically associated loess. The tephra samples, which had not been subject to flotation, had IRSL signals which were an order of magnitude lower that the glass rich fractions, but, unlike samples used in this study, retained plagioclase as well as volcanic glass, pyroxene and iron oxides. Three approaches to fading analysis over a timescale ranging from 16 hours to 2000 hours were examined, with differing results. When aliquots were irradiated and staggered intervals and read together at the end of the storage period (to minimise the effects of sensitivity change in readout equipment), scattered results were obtained, with a mean value of 2% fading per decade of time. The use of short IRSL measurements, which scarcely depleted individual aliquots, allowed multiple measurement from individual samples, and produced more

coherent data. Fading rates of 4% and 6% per decade were obtained from loess and tephra samples, which are within uncertainties of each other. These values are very similar to values obtained from feldspars. Auclair [113] noted the possibility that they may relate to the phenocrysts in the sample, rather than the glass phase, and also noted the possible ambiguity between fading and sensitivity change in the experimental design.

While our experiments suggest that the purification of coarser material from both PIG7 and the Ópoli tephra was more successful, at this stage, it is not clear if storage signal losses observed from the Ópoli tephra were due to insufficient thermal stability in the temperature range examined, the onset of proximity effects or possibly related to the increased level of weathering experienced by this sample over time. Despite low luminescence sensitivities, the signal obtained from synthetic glasses continued to grow in a similar approximately linear fashion after near-saturation to the Ópoli tephra [67]. This implies that dating of volcanic glass up to at least two million years old could be possible, but several technical refinements are needed to obtain accurate and reliable ages. Further high-dose experiments on volcanic glass of Quaternary age and different geochemical composition are recommended to determine if signal loss is related to burial age, geochemical composition and/or the scale of laboratory doses or dose rates.

The other key factor in assessing the reliability of age estimates is the accuracy of calculated dose rates. The large age errors in this study relate to the large dose rate errors and are primarily due to long-term variations in palaeowater content during burial. However, our dose rate calculations also had to account for variations in the alpha dose contribution since it was not possible to etch the thin plately glass shards without destroying them. The alpha

dose contributed as much as 50% of the total dose rate in this study and creates uncertainties that are as significant as those associated with changes in palaeowater content (Table 4). The observed similarity between bulk and internal glass-shard compositions in both samples examined suggests that sharp micro-dosimetric discontinuities probably do not exist at shard boundaries. Nevertheless, whilst we constrained alpha dose-related uncertainties as far as currently feasible, alpha dose attenuation in irregularly shaped glass shards could be significantly different to published dosimetric equations, which are based on spherical grains [100, 108, 122, 123]. Numerical modelling, beyond the scope of this paper, is therefore recommended to fully evaluate the dosimetric properties of glass shard shapes that are not perfect spheres. An in-depth comparison with dose rates from tephra deposits where volcanic glass constitutes less than c. 95% of the bulk tephra matrix might also be useful.

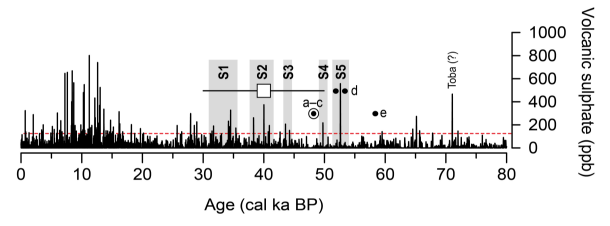
In summary, we conclude that sensitivity and long-term storage tests are critical components that should be included in all future analyses. Further reducing uncertainties associated with our age estimates would also require extensive *in-situ* dose rate measurements, including for example, a better assessment of seasonal to multi-decadal fluctuations in water content at each site. Automation, single-grain glass shard analysis (where sensitivity allows) and pulsed-laser infra-red stimulated luminescence (IRSL) methods could be applied and compared to ages obtained from feldspar-phenocryst phases in glass-rich deposits. This could result in reliable TL ages from tephra across a broad age-spectrum, up to and beyond the radiocarbon dating upper limit of c. 50 ka.

Palaeoenvironmental Implications

The c. 40 ± 10 ka age estimate of the PIG7 sample is similar in magnitude to existing age estimates obtained from the Þórsmörk region. It is also a closer approximation to 'known' ages for this deposit than, for example, the glass-phase TL age of 23.4 ± 2.4 ka obtained from deposits associated with the c. 75 ka Toba eruption [124]. Post depositional thermal annealing of the PIG7 sample could have reduced its measured age, but there are no currently active thermal areas in the Þórsmörk valley. Our c. 40 ± 10 ka age estimate also eliminates the possibility of a c. 200 ka eruption age for deposits at the PIG7 site. Following the SAR-TL experiments, we examined glass shards in the PIG1b and PIG7 samples for the presence of spontaneous fission-tracks. None were observed. The absence of natural fission-tracks in volcanic glass with a c. 3–4.5 ppm uranium and c. 10-14.5 ppm thorium content provides an additional age constraint of < c.100 ka, and rules out the possibility of a c. 200 ka eruption age in more than one location within the Þórsmörk valley.

The 40±10 ka age is potentially younger than the established age of NAAZ2 and raises the intriguing possibility of multiple ignimbrite- or rhyolitic-ash forming eruptions in the area between 60-30 ka. Lacasse et al. [2, 4] first suggested the PIG is the terrestrial geochemical equivalent of North Atlantic Ash Zone 2 (NAAZ-2). NAAZ-2 was originally dated in marine cores to around 64 ka [46]. Later, correlation ages of 48.5-58 ka were proposed by comparing the astronomically calibrated oxygen isotope time scale from marine cores with the Greenland GISPII ice core incremental timescale [9, 86]. The widespread distribution of the Vedde Ash and NAAZ-II in marine cores of the North Atlantic has been linked with ice rafting [9, 46], raising doubts over the precision of c. 64-48 ka correlation ages for NAAZ-II in marine cores. The spread of ages within NAAZ-II could also indicate more than one eruption event. Zielinski et al. [9] proposed that NAAZ-II was the result of two simultaneous eruptions from Katla and

Torfajökull based on two distinct tephra deposits with similar chemical composition to NAAZ-II and the PIG found in Greenland ice cores at 53.5 ka and 57.3 ka [86, 87] (Fig. 9). The c. 40 ka age estimate for the PIG7 sample is approximately 23% younger these ice core ages, but coeval with an elevated volcanic sulphate peak in the GISP2 ice core record centred on 40 ka (Fig. 9). Moreover, the tephrochronology of 60-30 ka eruptions from Iceland has recently been geochemically mapped in unprecedented detail and supports the possibility of multiple large rhyolite-forming eruptions from volcanic complexes in and around Pórsmörk during this period.



□ PIG7 SAR-TL age
 S1-S5 Volcanic sulphate >126 ppb (99% percentile) in SAR-TL range
 Closest geochemical correlation match
 Geochemical correlation match

Figure 9 The GISP2 Volcanic Sulphate record of palaeovolcansim for the last 80,000 years compared to new SAR-TL age estimates for Þórsmörk Ignimbrite. Notes: IC ka = ice core years before present; References: (a) [53]; (b), (c) [2, 4]; (d) [9]; (e) [46].

To summarise, TL screening and dating of volcanic glass in tephra holds great potential, particularly for the rapid classification of geochemically similar volcanic systems with widely separated eruption events. Whether all the ignimbrite deposits in the Þórsmörk area are from the same eruption event remains uncertain and warrants further investigation. Luminescence

and radiometric dating of other deposits in the Þórsmörk area is recommended to determine if more than one ignimbrite-forming eruption event occurred between 60-30 ka and further improve the tephrochronological record of Iceland.

Conclusions

1) To improve the Late Pleistocene tephrochronology of Iceland, NW Europe, and the North Atlantic and Arctic regions, we investigated the thermoluminescence (TL) dose response characteristics of rhyolitic volcanic glass produced by four large Plinian eruptions from Iceland: the c. 50 ka Þórsmörk Ignimbrite, Askja 1875 AD, Öræfi 1362 AD and the Late Pliocene Óþoli Tephra,

2) Our screening experiments showed an unequivocal age-related increase in naturally retained glass-phase TL. Single aliquot regeneration-TL analysis of volcanic-glass from the Pórsmörk Ignimbrite produced an age estimate of c. 40 \pm 10 ka, supporting evidence for a major eruption in the Pórsmörk area of Southern Iceland 30–60 ka rather than c. 200 ka.

3) The application of combined glass-phase thermoluminescence and radiometric dating at multiple sites in the Pórsmörk area could reveal if more than one ignimbrite-forming eruption occurred between 30-60 ka. Results from the Ópoli tephra were a similar order of magnitude to its established c. 2 Ma age, but further investigation of sensitivity change and signal loss from 'older' volcanic glass deposits is required. The tephrochronological record of Iceland could be improved using TL analysis of volcanic glass deposits.

Acknowledgements

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Data availability

Datasets are summarised in the Supplementary Material, are available from the corresponding author (siro@bas.ac.uk) and these links:

Supplementary material: https://doi.org/10.6084/m9.figshare.13634666.v1

Datasets: https://doi.org/10.6084/m9.figshare.13633877.v1,

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