A Holocene history of climate, fire, landscape evolution, and human activity in Northeast Iceland

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9 Abstract

10 Paleoclimate reconstructions across Iceland provide a template for past changes in climate across the northern North Atlantic, 11 a crucial region due to its position relative to the global northward heat transport system and its vulnerability to climate change. 12 The roles of orbitally driven summer cooling, volcanism, and human impact as triggers of local environmental changes in the 13 Holocene of Iceland, remain debated. While there are indications that human impact may have reduced environmental resilience during Late Holocene summer cooling, it is still difficult to resolve to what extent human and natural factors affected 14 15 Iceland's Late Holocene landscape instability. Here, we present a continuous Holocene fire record of northeastern Iceland 16 from proxies archived in Stóra Viðarvatn sediment. We use pyrogenic polycyclic aromatic hydrocarbons (pyroPAHs) to trace 17 shifts in fire regimes, paired with continuous biomarker and bulk geochemical records of soil erosion, lake productivity, and 18 human presence. The molecular composition of pyroPAHs and a wind pattern reconstruction indicate a naturally driven fire 19 signal that is mostly regional. Generally low fire frequency during most of the Holocene significantly increased at 3 ka and 20 again after 1.5 ka BP, before known human settlement in Iceland. We propose that shifts in vegetation type caused by cooling 21 summers over the past 3 kyr, in addition to changes in atmospheric circulation, such as shifts in North Atlantic Oscillation 22 (NAO) regime, led to increased aridity and biomass flammability. Our results show no evidence of faecal biomarkers associated with human activity during or after human colonisation in the 9th century CE. Instead, faecal biomarkers follow the 23 24 pattern described by erosional proxies, pointing toward a negligible human presence and/or a diluted signal in the lake's 25 catchment. However, low post-colonisation levels of pyroPAHs, in contrast to an increasing flux of erosional bulk proxies, 26 suggest that farming and animal husbandry may have suppressed fire frequency by reducing the spread and flammability of 27 fire-prone vegetation (e.g., heathlands).

Overall, our results describe a fire frequency heavily influenced by long term changes in climate through the Holocene. They also suggest that human colonisation had contrasting effects on the local environment by lowering its resilience to soil erosion

30 while increasing its resilience to fire.

31 **1 Introduction**

32 Iceland is highly sensitive to most mechanisms controlling the evolution of Holocene climate in the North Atlantic, from 33 millennial (e.g., shifts in deep water formation and ocean current positions) to sub-decadal timescales (e.g., variability of the 34 North Atlantic Oscillation) (Harning et al., 2021; Mjell et al., 2016; Moossen et al., 2015; Petit et al., 2020). Recent lake 35 sedimentary records in Iceland (Alsos et al., 2021; Geirsdóttir et al., 2009a, 2013, 2019, 2020; Harning et al., 2016, 2020; 36 Hiles et al., 2021; Larsen et al., 2011, 2012; Richter et al., 2021) draw a comprehensive picture of Icelandic environments 37 during the Holocene (last 11.7 kyr). These Holocene paleoclimate reconstructions derived from lake sediments in Iceland show 38 first-order millennial trends that reflect orbitally-driven changes in Northern Hemisphere summer insolation, and millennial to 39 sub-millennial changes that are primarily impacted by northern North Atlantic ocean circulation and to a part by local 40 volcanism (e.g., Flowers et al., 2008; Geirsdóttir et al., 2013, 2020; Harning et al., 2018b; Larsen et al., 2012). These Holocene 41 climate reconstructions further indicate a major shift from occasional to increasingly severe landscape instability and soil 42 erosion occurring at least 300 years before the acknowledged settlement of Iceland (ca 870 CE; The Book of Icelanders "Íslendigabók", by Ari Thorgilsson, 12th century CE, e.g., Smith, 1995), suggesting that human impact had a secondary role 43 44 to climate by lowering the resilience of the environment to an already ongoing naturally driven erosion (e.g., Bates et al., 2021; 45 Geirsdóttir et al., 2009b, 2020). The ability to generate high-resolution Holocene terrestrial climate records, along with 46 Iceland's relatively short settlement history, makes Iceland an ideal location to attempt disentangling the impact of natural 47 climate variability and human activities on the changes in the local landscape during the Late Holocene.

48 In this study, we use multiple organic proxies from a Holocene sediment core from the Stóra Viðarvatn lake in northeast 49 Iceland to investigate the effects of natural and anthropogenic drivers on the local Icelandic environment. First, we focus on 50 tracing the evolution of fire regimes using pyrogenic polycyclic aromatic hydrocarbons (pyroPAHs; Lima et al., 2005). Fires 51 can have a significant impact on ecosystems, affecting vegetation patterns, nutrient cycling, and wildlife habitat (e.g., 52 Goldammer and Furyaev, 1996). The frequency, intensity, and spatial extent of fires can provide insights into past climate and 53 environmental conditions (e.g., Marlon, 2009; Power et al., 2008) and, to our knowledge, there are no such records for the 54 Holocene in Iceland, while limited data is available for the surrounding regions (Chen et al., 2023; Marlon et al., 2013; Segato 55 et al., 2021; Zennaro et al., 2014). Second, as fire frequency can be influenced by human activities as well (e.g., Marlon et al., 56 2009, 2013; Zennaro et al., 2015), we also analyse faecal markers of human presence (Vázquez et al., 2021). By analysing 57 these biomarkers from deglaciation to present, we can define their natural, pre-settlement background levels and thus 58 potentially trace anthropogenic impact on the local environment, pinpointing human arrival in the lake catchment.

- 59 Finally, by coupling fire and human presence biomarker records with established proxies for environmental change (e.g., soil
- 60 erosion and primary productivity, e.g., Argiriadis et al., 2018; Geirsdóttir et al., 2013; Gross, 2017), we test what control
- 61 natural and/or human factors had on the evolution of the Holocene landscape in Iceland.

62 2 Methods

63 2.1 Study site

- Stóra Viðarvatn (SVID) is a lake (2.6 km² surface area) located in NE Iceland (Fig. 1A-B) at an elevation of 151 m asl. SVID has a maximum depth of 48 m, a catchment area of 17 km² (including the lake surface), and a volume of ca 3.6×10^7 m³ (this study, based on data from the National Land Survey of Iceland, Landmælingar Íslands, 2023; Axford et al., 2007). The lake is surrounded by Quaternary age basaltic lavas and glacial hyaloclastites formed by subglacial eruptions, as well as some Holocene soil with several mm-cm thick tephra layers (Hjartarson and Sæmundsson, 2014). The nearby Raufarhöfn station (Icelandic Meteorological Office, 2022) provides weather data for the 1961–1990 CE interval: mean annual temperature is
- 70 2 °C with a maximum in July-August (8 °C), while the lake surface is usually frozen between November and March; mean
- annual precipitation is 733 mm a⁻¹ with lowest values occurring in May (28 mm) and the highest in October (ca 86 mm),
- suggesting a lake-water residence time between five and nine years.
- 73 In February 2020, we recovered a 8.93 m long core 20-SVID-02 (66.236867° N; -15.837837° E; 1C) from 17.4 m water
- depth near the centre of the lake (Harning et al., 2023). The sediment was retrieved in seven drives of ~150 cm each, using
- ⁷⁵ lake ice as a coring platform. The core was subsequently stored in a cool room (4 °C) at the Institute of Arctic and Alpine
- Research, University of Colorado Boulder, where it was subsampled. Previously, two studies have analysed an 8 m long core
- 77 (04-SVID-03; 1C) retrieved in February 2004 to trace Holocene temperature (Axford et al., 2007) and δ^{18} O from chironomid
- remains, as well as the δD , $\delta^{13}C$, $\delta^{15}N$ of total organic matter (Wooller et al., 2008) at a 1–0.2 kyr resolution.



Figure 1: (A) Study area in NE Iceland; (B) Location of the Raufarhöfn climatological station and Ytra-Áland site (Karlsdóttir et al., 2014), which are 20 km NNW and 13 km ESE from Stóra Viðarvatn (SVID), respectively; (C) Location and catchment area of the Stóra Viðarvatn lake: 20-SVID-02 core is marked by a pink hexagon and an older 04-SVID-03 core by a red square (Axford et al., 2007); SVID bathymetry (10 m isolines) is reported by Axford et al. (2007); watershed catchment and contour lines (10 m) are calculated via ArcGIS (Esri, 2023) based on digital elevation models provided by the National Land Survey of Iceland; basemap sourced from Esri.

86 2.2 Tephrochronology

87 Our sediment core chronology takes advantage of the geochemical fingerprints of visible Icelandic tephra layers and their 88 correlation to marker tephra of known age. Thirteen tephra layers were sampled along the vertical axis, sieved to isolate glass 89 fragments between 125 and 500 µm, and embedded in epoxy plugs. At the University of Iceland, individual glass shards were 90 analysed on a JEOL JXA-8230 electron microprobe using an acceleration voltage of 15 kV, beam current of 10 nA, and a 91 beam diameter of 10 µm. The international A99 standard was used to monitor for instrumental drift and maintain consistency 92 between measurements. Tephra origin was then assessed using major oxide compositions, following the systematic procedures 93 outlined in Jennings et al., 2014 and Harning et al., 2018a. Briefly, based on SiO₂ wt% vs total alkali (Na₂O + K₂O) wt%, we 94 determined whether the tephra volcanic source is mafic (tholeiitic or alkalic), intermediate and/or rhyolitic. From here, we 95 objectively discriminate the source volcanic system through a detailed series of bi-elemental plots produced from available 96 compositional data on Icelandic tephra. Source eruption was then determined using the geochemical fingerprint and relevant 97 stratigraphic information. See supplemental information for complete major oxide compositions and bi-elemental plots.

Using the 13 marker tephra layers of known age (Table 1), we generated a Bayesian age model using the R package rbacon
2.5.7 (Blaauw and Christen, 2011; R Core Team, 2020) and default model functions (Fig. 2). We used the 'slumps' function
for the thicker tephra layers (e.g., Hekla 3 and Hekla 4) to reflect their instantaneous deposition on geologic timescales.

101 **2.3** Sample preparation and analysis

102 At the University of Colorado Boulder, we retrieved a total of 196 sediment core samples at an average spacing of 4.5 cm, 103 providing a temporal resolution of decadal to centennial time scales. We freeze-dried samples for 24-48 hours, and ground 104 and homogenised them (mean weight 1.5 g, range 0.6–6.6 g) using an agate mortar and pestle. Using 13–70 mg of sediment, 105 we measured total carbon (TC), total nitrogen (TN), and δ^{13} C (relative to VPDB) on an elemental analyser linked to a Thermo 106 Delta V isotope ratio mass spectrometer (EA-IRMS) in the Earth Systems Stable Isotope Laboratory at the University of 107 Colorado Boulder; samples were analysed against a suite of secondary laboratory standards that are extensively calibrated to 108 international standard reference materials to correct for size, blank-mixing, linearity and drift effects (Harning et al., 2018b). 109 We did not decalcify samples as the contribution of inorganic carbon to TC in Icelandic lake sediment is considered to be 110 negligible (see par. 5.1; Geirsdóttir et al., 2020). We analysed 9-11 mg of sediment for biogenic silica by Diffuse Reflectance 111 Fourier Transform Infrared Spectrometry (FTIRS) on a Bruker Vertex 70 with a Praying Mantis diffuse reflectivity accessory 112 (Harrick) and report values in FTIRS - Fourier Transform Infrared Spectroscopy absorbance units (e.g., Harning et al., 2018b). 113 We processed 86 selected samples for organic biomarker analyses. We extracted 0.4–2 g of dry sediment with an accelerated 114 solvent extractor (Dionex ASE350). Sample size, matrix, and richness in organic compounds greatly varied along the core and 115 initial tests occasionally showed coloration in the sample extract even after 4-5 cycles, likely due to the persistence of organics. 116 Thus, to maintain consistency in lipid yields throughout the core, we used dichloromethane (DCM):methanol (MeOH) 9:1 for 117 six cycles of five minutes (static time), 100 °C, and 1,500 psi. After extraction, we spiked the total lipid extract (TLE) with 118 1000 ng of 3-methyl-heneicosane (CAS#: 6418-47-9, Sigma-Aldrich), 20 ng of p-terphenyl (CAS#: 92-94-4, TCI), and 50 ng 119 of pregnanol (5 β -Pregnan-3 α -ol, CAS#: 4352-07-2, Steraloids) as internal standards for the quantification of *n*-alkanes, PAHs, 120 and sterols, respectively. We concentrated the TLE under a gentle flow of nitrogen and then mixed it with HCl-activated copper 121 shots to remove elemental sulphur as copper sulphide precipitates. We then filtered the samples through a NaSO₄-packed 122 Pasteur column to remove any residual water and copper sulphide and concentrated them under N₂. We subsequentially 123 separated the TLE into six chromatographic fractions using a Pasteur pipette packed with silica gel (60-200 µm - 60 A) and 124 solvents of increasing polarity. We calculated the column's dead volume (DV) with *n*-hexane, and then eluted samples with 125 1.5 DV of n-hexane (F1), 2 DV of n-hexane: DCM 4:1 (F2), 1.5 DV of DCM (F3), 2 DV of DCM: acetonitrile (F4), 1.5 DV of 126 acetonitrile (F5), and 3 DV of MeOH (F6). We derivatised fraction F4, containing the sterols/stanols, using TMS-BSTFA 127 (Supelco) and pyridine (50:50) at 70 °C for 15 minutes, then dried under N₂ and redissolved it in *n*-Hexane. We added 1 ng of

- 128 p-terphenyl D₁₄ (CAS: 1718-51-0, Sigma-Aldrich) and 50 ng of 5α-cholestane (CAS: 481-21-0, Sigma-Aldrich) to fractions
- 129 F2 (PAHs) and F4 (sterols), respectively, as injection standards to check the recovery and quantification consistency of
- 130 analyses.
- 131 We analysed the n-alkanes, PAHs, and sterols using a Thermo Scientific Trace 1310 gas chromatograph (GC) equipped with 132 a PTV inlet and a Restek glass liner interphase to a TSQ8000-Evo triple quadrupole mass spectrometer (MS). We used a 60 133 m DB1 column (DB-1MS, 0.25 mm, 0.25 µm film thickness, Agilent, USA) to separate n-alkanes and a DB-5 column (DB-134 5MS, 0.25 mm, 0.25 μ m film thickness, Agilent, USA) for PAHs and sterols, and He (1.2 ml min⁻¹) as a carrier gas. For *n*alkane analysis, we injected samples in splitless mode at 65 °C and the PTV was ramped to 400 °C at 3 °C s⁻¹ and held for 5 135 136 min. The GC oven temperature was programmed from 60 °C to 220 °C (25 °C min⁻¹) and then to 315 °C (2.5 °C min⁻¹, held 137 13 min). n-Alkanes were analysed in full scan (50-600 m/z) using the following MS conditions: 300 °C EI source at 70 eV 138 electron energy, 50 uA emission current, and 15 V electron lens voltage, with a transfer line at 315 °C. For PAH analysis, all 139 samples were injected in splitless mode at 45 °C and the PTV was ramped to 400 °C at 11.6 °C s⁻¹ and held for 2 min. The GC 140 oven temperature was programmed from 60 °C (held 1 min), to 150 °C (40°C min⁻¹), to 320 °C (3 °C min⁻¹, held 15 min). MS 141 conditions were as follows: 250 °C EI source at 70 eV electron energy, 50 uA emission current, and 15 V electron lens voltage, 142 with a transfer line at 320 °C. For sterol/stanol analysis, all samples were injected in splitless mode at 90 °C, evaporated at 143 100 °C (0.1 min), and the PTV was ramped to 400 °C at 8 °C s⁻¹ and held for 1 min). The GC oven temperature was programmed 144 from 80 °C (held 1 min), to 200 °C (20 °C min⁻¹), to 320 °C (5 °C min⁻¹, held 20 min). MS conditions were the same as for n-145 alkanes. PAHs and sterols/stanols were analysed in selected reaction monitoring (SRM) using the collision energies and mass
- 146 transitions reported in Table A1 and Table A2).

147 **2.4** Analysis of air parcel back-trajectory patterns

148 To define the potential regional extent of airborne PAHs arriving to SVID's catchment area, we traced the back-trajectory of 149 air parcels using HYSPLIT (hybrid single particle lagrangian integrated trajectory; Draxler et al., 1998; Stein et al., 2015). 150 Using a modified version of an R script originally developed to trace precipitation patterns (Caves Rugenstein and 151 Chamberlain, 2018), we analyse data from the NOAA Global Data Assimilation System (GDAS; resolution 1° by 1°) at a six 152 hours frequency tracing back trajectories for three days (72 h) and two weeks (336 h) during two years characterised by 153 opposite North Atlantic Oscillation (NAO; Hurrell et al., 2003) configuration (2009-2010, NAO-; 2013-2014, NAO+; NOAA, 154 2023). PAHs deposition, which is enhanced by low temperatures, occurs not only via precipitation but in dry conditions as 155 well (Arellano et al., 2018; Feng et al., 2017; Golomb et al., 2001; Halsall et al., 2001). Thus, we present data for air parcel 156 trajectories that did and did not produce precipitation within six hours from the endpoint (SVID), initialising the trajectories 157 at four different altitudes: 1000, 1500, 2000 m asl (water vapour usually advects within an altitude of 2 km; Bershaw et al., 158 2012; Lechler and Galewsky, 2013; Wallace and Hobbs, 2006), and 150 m asl (SVID surface elevation).

159 **3 Background on proxies**

160 **3.1 Polycyclic aromatic hydrocarbons (PAHs)**

We use pyrogenic PAHs (pyroPAHs) as tracers for the frequency/intensity of fire episodes, and the PAH perylene as a biogenic PAH related to terrestrial organic matter input. PAHs are semi-volatile compounds that can be of pyrogenic, petrogenic, or biogenic origin (Kozak et al., 2017; Lima et al., 2005). Low molecular weight (LMW; see Table A1 for group definition) PAHs in their non-alkylated form (Page et al., 1999; Yunker et al., 2002) constitute the majority of the PAHs produced by the combustion of plant biomass, while the relative amount of high molecular weight (HMW) PAHs increases along with higher fire temperatures (McGrath et al., 2003). LMW PAHs tend to be airborne and show high aqueous solubility and higher volatility, whereas HMW PAHs are usually in a solid phase (associated to either soot or char), show lower volatility, and are

- 168 likely sourced locally (Hoffmann and Wynder, 1977; Junk and Ford, 1980; Karp et al., 2020; Lammel et al., 2009; Lima et al.,
- 169 2005; Purushothama et al., 1998). Thus, low contributions of HMW PAHs in environmental samples are often considered
- 170 indicative of either low temperature fires (e.g., Denis et al., 2012) or a distal source, while high relative amounts generally
- 171 point toward a more local signal. Finally, perylene is a 5-hexa-ring PAH often detected in aquatic sediments and considered
- to be mostly of in situ biogenic origin, probably from precursor compounds present in saprophagous and mycorrhizal fungi
- 173 (e.g., Aizenshtat, 1973; Jiang et al., 2000; Slater et al., 2013; Wang and Huang, 2021), and thus likely linked to higher organic
- 174 matter content and terrigenous input (Guo and Liao, 2020; Hanke et al., 2019).

175 **3.2** Sterols/stanols as markers of plant sources and animal digestion

176 Stanols are saturated isomers of sterols (e.g., Patterson, 1971). When the bacterially mediated reduction of sterol double bounds 177 occurs in an open environment (e.g., soil), it leads almost exclusively to the production of 5α stanol isomers. When the 178 reduction of sterols occurs in the animal's digestive tract, their enteric bacterial flora maximises the production of 5ß stanols 179 (Hatcher and McGillivary, 1979; Murtaugh and Bunch, 1967). Humans (and partially other omnivores and carnivores) 180 maximise the production of coprostanol (5β-cholestan-3β-ol) through the saturation of animal derived cholesterol (5-en-181 cholest-3 β -ol). Ruminants such as sheep and cattle, on the other hand, maximise the production of 5 β -stigmastanol and 5 β -182 campestanol (Leeming et al., 1996, 1994) from plant derived sterols like stigmasterol, sitosterol, and campesterol (e.g., Goad, 183 1977; Goad and Goodwin, 1966; Pancost et al., 2002). Higher/lower ratios of coprostanol and its derived epimer epi-184 coprostanol (5 β -cholestan-3 α -ol; McCalley et al., 1981; Quirk et al., 1980; Wardroper et al., 1978) to 5 β -stigmastanol or 5 β -185 campestanol are considered to be a proxy for higher/lower faecal input from human sources relative to ruminant sources, and 186 have been widely applied to samples from modern/ancient sewage material and manured soil (e.g., Birk et al., 2012; Bull et 187 al., 2001, 2002; Cordeiro et al., 2008; Evershed et al., 1997; He et al., 2018; Lerch et al., 2021; Simpson et al., 1999; Tyagi et 188 al., 2009).

189 3.3 *n*-Alkanes

190 Plants synthesise *n*-alkanes and other *n*-alkyl lipids as part of their waxy coating with a characteristic strong odd-over-even 191 chain length predominance (Eglinton and Hamilton, 1967), which is summarised by their higher carbon preference index (CPI; 192 Bray and Evans, 1961; Marzi et al., 1993). In contrast, lower CPI values are usually indicative of petrogenic, algal, or bacterial 193 sources (Grimalt and Albaigés, 1987; Han and Calvin, 1969). Aquatic sources such as macrophytes and mosses (e.g., 194 Sphagnum) maximise their leaf wax *n*-alkane production at mid-length homologues (C_{21-25}), while terrestrial plants (e.g., 195 grasses, sedges, trees, shrubs) are generally skewed toward longer homologues (C₂₇₋₃₁), allowing for use of source 196 discriminating ratios and indices such as the aquatic plant index (Pag, Ficken et al., 2000) and average chain length (ACL, 197 Gagosian and Peltzer, 1986).

198 **3.4 Bulk geochemistry proxies**

199 Aquatic and terrestrial catchment productivity, flux of inorganic sediments, and organic matter preservation are the main 200 factors determining the level of total organic carbon content in lacustrine sediments (Meyers and Ishiwatari, 1993). The molar 201 carbon to nitrogen ratio (C/N) in plant tissue varies between aquatic plants and phytoplankton (<10) and terrestrial plants and 202 bryophytes (>10; Meyers, 1994). Thus, increases in C/N are usually interpreted as an increased catchment erosion and input 203 of terrestrial organic matter and/or as a relative decrease of aquatic plant productivity (Fernández-Martínez et al., 2021; 204 Kaushal and Binford, 1999; Meyers, 1997; Meyers and Teranes, 2001; Rieger et al., 1979). Shifts in the abundance of diatom 205 derived biogenic silica (BSi) can trace lake productivity (Colman et al., 1995; Conley, 1988; Conley and Schelske, 2002). The 206 conservation potential of diatom frustules is strongly related to sedimentation rate, with higher rates leading to better 207 preservation. When sedimentation rates are considered relatively constant, shifts in BSi can reflect qualitative changes in

- spring/summer temperature in high-latitude lakes, such as Iceland (Geirsdóttir et al., 2009a; McKay et al., 2008). The stable
- 209 isotopic composition of carbon (δ^{13} C) can trace shifts in the relative contribution of organic matter sources, with terrestrial
- 210 plants (but also bryophytes) and associated soils showing more ¹³C-depleted values (ca -32‰ to -25‰), aquatic plants
- 211 exhibiting more ¹³C-enriched values (ca -20% to -10%), and freshwater algae and phytoplankton showing a wider isotopic
- range (Meyers, 1994; Prokopenko et al., 1993; Rundel et al., 1979; Smith and Epstein, 1971; Geirsdóttir et al., 2020 and refs
- therein). The physical mixing or stratification of a lake water column can also influence the carbon isotopic signature of aquatic
- 214 sources (Hernández et al., 2011).

215 **4 Results**

216 **4.1 Age model**

- 217 Based on major oxide composition and stratigraphical information, we identified 13 marker tephra layers of known age (Table
- 1). Our Bayesian tephra age model shows nearly constant sediment accumulation rates throughout the Holocene (Fig. 2).
- 219 There is increased uncertainty in age control between the G10ka tephra series and Kverkfjöll/Hekla 6200 due to fewer marker
- tephra layers being present. However, the Late Holocene, particularly during the historical period of settlement, features numerous tephra layers that result in substantially lower age estimate uncertainty.
- 222

223 Table 1: Marker tephra layers of known age identified in 20-SVID-02 and used to develop the age model.

Composite depth (cm) Tephra layer ID		Layer age (a BP)	Reference	
31.5	Bárðarbunga-Veiðivötn 1717	233 ± 2	Thorarinsson (1974)	
42.0	Grímsvötn 1619?	352 ± 2	Thorarinsson (1974)	
56.0 - 57.0	Veiðivötn 1477	473 ± 2	Larsen et al. (2002)	
70.0	Hekla 1300	650 ± 10	Thorarinsson (1967)	
82.5	Hekla 1104	846 ± 10	Thorarinsson (1967)	
102.2	Kverkfjöll	1125 ± 50	Óladóttir et al. (2011)	
242.5 - 248.5	Hekla 3	3010 ± 54	Dugmore et al. (1995)	
344.0 - 345.0	Hekla 4	4200 ± 42	Dugmore et al. (1995)	
414.0 - 414.5	Kverkfjöll	5200 ± 100	Óladóttir et al. (2011)	
503.2	Kverkfjöll and Hekla	6200 ± 100	Óladóttir et al. (2011)	
815.0	G10ka Series (top)	9900	Óladóttir et al. (2020)	
853.5	G10ka Series (bottom)	10400	Óladóttir et al. (2020)	
886.5	Askja S	10830 ± 57	Bronk Ramsey et al. (2015)	



Figure 2: Stóra Viðarvatn age model generated in Bacon (Blaauw and Christeny, 2011). Green horizontal lines denote the age and uncertainty of marker tephra layers, red line reflects mean values of model iterations, the grey lines denote the 95% confidence envelope, and darker shading reflects more likely ages. Gray vertical bars mark the 'slumps' used for the Hekla 3 and Hekla 4 tephra layers.

To facilitate the interpretation of downcore records, we present and discuss data (1) divided into nine time intervals (I–IX; Table 2) representing the most distinguishable periods of variability with respect to background values, and (2) separately for the sections preceding and following the G10ka tephra series (Óladóttir et al., 2020). We tried analysing some material from the G10ka series (a very thick tephra unit; Óladóttir et al., 2020) but the yields of organics were too low to be reliable (as expected; its contents are almost 100% inorganic). Thus, here we only included one sample at the bottom limit of the G10 ka tephra series and one at the upper limit.

237	Table 2. Age intervals (approximate) and descriptions of the nine subdivisions of the 20-SVID-02 record used in this study
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Interval	Age (ka BP)	Description
Ι	10.85-10.63	Potential Preboreal cooling
II	10.63-10.40	Pre-Boreal warming
III	10.4-9.9	G10 ka tephra series
IV	9.9-8.8	Early Holocene warming (rebound after G10 ka event)
V	8.8-7.5	Early Holocene instability (8.2 ka event?)
VI	7.5-3.0	Middle Holocene plateau and trend inversion
VII	3.0-1.3	Late Holocene cooling
VIII	1.3-0.25	Medieval period and Little Ice Age
IX	0.25-present	End of LIA and contemporary warming

238 4.2 Bulk geochemistry

239 The C/N ratio (Figs. 3-4B) ranged from ~1 to ~9.5, showing lowest values at the beginning of the record (I). At ca 10.63 ka 240 BP, C/N increased sharply and reached mid-range values (5-6), remaining relatively stable throughout most of the remaining 241 Holocene (IV–VI), except for two drops, after the G10ka Series tephra and between 8.8 and 7.75 ka BP (V). In the last 2 kyr, 242 C/N values steadily increased, leading to the highest values in the most recent portion of the record. The two periods with decreased C/N values, as well as the initial increase (I to II), generally paralleled the behaviour of the total carbon (Figs. 3-243 244 4E), biogenic silica (Figs. 3-4H), and δ^{13} C (Figs. 3-4D) records. TC increased steadily throughout the Holocene from ~0% 245 while BSi rapidly increased at the beginning of the record (30 to 90) to then stabilise at 110–120 for more than 5 kyr (V–VI). 246 Both records peaked at ca 3.5 ka BP (2.8%, TC; ~160, BSi max) and temporarily dropped between ca 3 and 2.25 ka BP. Subsequently, TC increased to its maximum value (3.6%, modern) while BSi decreased in a stepwise manner, reaching its 247 lowest value of the last 10 kyr (~50) at ca 0.2 ka BP. The δ^{13} C record showed the most 13 C-enriched values (-20‰) in the 248 249 oldest interval (I); it then decreased (to -26‰, modern) steadily throughout the Holocene, except for two major drops to its 250 most depleted values (ca -27‰) during periods II and V.



Figure 3: Erosional and primary productivity proxies from the pre-G10ka Series tephra interval (pink vertical band) of 20-SVID-02 core; all concentrations are on g of dry samples. (A, black) sum of C_{19–35} *n*-alkanes concentration aquatic plant index derived from *n*-alkanes; (B, green) carbon to nitrogen ratio; (C, purple) perylene concentration; (D, black-yellow) stable isotopic composition of total carbon; (E, brick red) percentage of total carbon; (F, dark blue) *n*-alkanes carbon preference index (CPI_{19–31}); (G, red) aquatic plant index derived from *n*-alkanes; (H, light blue) biogenic silica.





Figure 4: Erosional and primary productivity proxies of 20-SVID-02 core. (A, black) sum of C₁₉₋₃₅ *n*-alkanes concentration aquatic
 plant index derived from *n*-alkanes; (B, green) carbon to nitrogen ratio; (C, purple) perylene concentration; (D, black-yellow) stable
 isotopic composition of carbon; (E, brick red) percentage of total carbon; (F, cyan) northern hemisphere summer insolation at 65°
 N (Berger and Loutre, 1999); (G, red) aquatic plant index derived from *n*-alkanes; (H, light blue) biogenic silica; (I, dark blue)
 carbon preference index (CPI₁₉₋₃₁). Red dotted line marks the conventional age (870 CE) of the settlement of Iceland (Landnám).
 Black vertical dashed lines mark the subdivision of the 20-SVID-02 record into nine intervals (Table 1).

264 4.3 *n*-Alkanes

We detected *n*-alkane homologues from C_{19} to C_{33-35} (Fig. A3) in most samples, with a total sum that ranged from 0.3 to 50 µg g⁻¹ (700–4000 µg g⁻¹ TC; Figs. 3-4A). The 10.8 to 4.2 ka BP interval showed relatively low and stable values (~5 µg g⁻¹); concentrations roughly doubled from 4 to 1.5 ka and then again after 0.5 ka BP, reaching its maximum value at the end of the record. The CPI showed a stable odd-over-even predominance (3 to 6.5) through the whole record, except for low values (1 to 3) seen in the interval preceding the G10ka Series tephra (Figs. 3F-4I). The most abundant homologues were C_{23-27} (45%) in the 10.8 to 3 ka BP interval and C_{29-31} (40%) in the last 3 kyr. This regime change was highlighted by a shift in P_{aq} from relatively high values (up to 0.8; avg. 0.6) through the early-mid Holocene to lower values (down to 0.2; avg. 0.3) after 3 ka

272 BP (Fig. 4G).

273 4.4 Faecal sterol/stanols

274 We detected the three main faecal stanols: the plant derived 5 β -stigmastanol was ~5–10 times more abundant (up to 3-500 ng g⁻¹; Figs. 5-6A) than coprostanol + epi-coprostanol (10-30 ng g⁻¹; Figs. 5-6B). The oldest interval (I) showed the lowest 275 276 concentrations for all three stanols, while the following interval (II) displayed high (highest for coprostanol and epi-277 coprostanol) concentrations. All three stanols showed low and stable concentrations between ~9.5 and 7.5 ka BP, gradually 278 increased from ~7.5 ka BP before reaching a relative maximum around 1.5 ka BP (VII), dropping again during interval VIII, 279 and peaking during the last 200 to 300 years (IX). Parent sterols (β -sitosterol, β -stigmasterol, except cholesterol) and the α -280 stanol isomers, follow patterns similar to the β -stanols throughout the Holocene, but with 10 to 100 times higher concentrations 281 (Fig. A2).









Figure 6: Sterols/stanols of the 20-SVID-02 core. (A, red) 5β-stigmastanol and (B, dark blue) sum of coprostanol and epi-coprostanol
 concentration on g of dry sample (full line) and on g of TC (dotted line); (C, black) β-sitosterol to 5α-stigmastanol ratio; (D, yellow)
 β-sitosterol concentration; (E, light blue) 5α-stigmastanol concentration. Red dotted line marks the conventional age (870 CE) of the
 settlement of Iceland (Landnám). Black vertical dashed lines mark an arbitrary subdivision of 20-SVID-02 record into nine intervals
 (Table 1).

292 **4.5** Polycyclic aromatic hydrocarbons (PAHs)

293 PAHs were present in all samples and generally in higher concentrations in more recent compared to older samples (Fig. A1). Perylene (Figs. 3-4C), which accounts for more than 97% of detected PAHs, maintains low concentrations (0-20 ng g⁻¹) from 294 10.5 to 4-3 ka BP. A first increase to 0.5–1.5 µg g⁻¹ occurred between 3.5 and 2.8 ka BP, followed by a decrease loosely coeval 295 296 to the decrease in TC and BSi (VII). After ca 1.5 ka BP, perylene increases to a maximum value (~6 µg g⁻¹, ca 0.15 ka BP), 297 which generally matches the pattern of TC. The second most abundant compound was phenanthrene $(0.01-31.1 \text{ ng g}^{-1})$, followed by pyrene (0.1–11.5 ng g^{-1}) and fluoranthene (0.01–17.6 ng g^{-1}). The least abundant PAHs were naphthalene, 298 299 acenaphthylene and acenaphthene. However, since the detection of these three low molecular weight compounds could have 300 been influenced by evaporation losses during sample preparation, their reported concentrations are likely to be underestimated. 301 Given the overwhelming dominance of perylene and its likely biogenic rather than pyrogenic origin, we removed it from total 302 PAHs abundance calculations to provide a record with features that were not apparent in perylene's trend. In terms of total 303 pyroPAHs abundance, we observe five distinguishable intervals (Fig. 7A). First, a 3 kyr-long interval (~80 years average 304 temporal resolution) starting at ca 10.5 ka BP displays a relatively stable low concentrations (< 5 ng g⁻¹, SD σ = 2.3). The only exception is a point taken within the G10ka Series tephra (up to 10 ng g⁻¹) at ca 10 ka BP. Second, from ~7.5 to 2.9 ka BP 305 306 (\sim 320 years average temporal resolution), the total PAHs concentration increases from \sim 5 to 10 ng g⁻¹, although with enhanced 307 variability ($\sigma = 3.7$). Third, led by the increase of low molecular weight PAHs (LMW, Fig. 7B), during the 2.9 to 0.7 ka BP interval, values fluctuate ($\sigma = 7.5$) between ~10 and 20 ng g⁻¹, with two major peaks reaching 45 and 35 ng g⁻¹ at ~2.7 to 2.3 308 309 ka BP and 1.5 to 1.3 ka BP, respectively. Fourth, between 0.7 and 0.3 ka BP there is a relatively brief although clear drop (~10 310 ng g⁻¹, $\sigma = 1.5$), led by both LMW and MMW PAHs. In the last 250 years we observe a sharp, 10-fold increase in PAHs 311 concentration leading to the highest recorded values (~200 ng g⁻¹). When normalised for TC (Fig. A1), the absolute values 312 increase 10 to 100-fold, but the patterns do not substantially change.



Figure 7: PAHs record of the 20-SVID-02 core. (A, black) sum of pyrogenic PAHs concentrations; (B, fuchsia) low, (orange) medium, and (blue) high molecular weight pyrogenic PAHs concentration. Red dotted line marks the conventional age (870 CE) of the settlement of Iceland (Landnám). Vertical pink band marks the G10ka Series Tephra. Black vertical dashed lines mark an arbitrary subdivision of 20-SVID-02 record into nine intervals (Table 1).

318 **4.6 HYSPLIT**

We calculated a total of 11,392 air trajectories, which we split by year, season, dry (i.e., not associated with precipitation), or precipitation bearing trajectories (Fig. A4). Most trajectories, even for two weeks intervals, show air parcels originating mostly from Iceland and surrounding areas of the North Atlantic, regardless of season or NAO configuration, while the contribution from nearby terrestrial regions (potential PAHs sources) such as Greenland, British Isles, or Scandinavia is negligible. As the marine environment is not conducive to combustion nor redeposition of particulates, this implies a dominantly Icelandic signal for PAH production. Since wildfires are concentrated in the relatively dryer, snow-free summer season (Mccarty et al., 2021), we focus particularly

on the JJA air trajectory data (Fig. 8). These results show that: (1) 95% of back-trajectories originate from Iceland and its

nearby waters; (2) 90% of back-trajectories originate within 100–150 km radius from SVID; (3) dry trajectories more likely

328 originate inland relative to precipitation carrying trajectories; (4) 95% of trajectories from a NAO- year tend to be confined to

329 northern Iceland while during a NAO+ year trajectories more commonly originate from inland; (5) these patterns are consistent

even when scaled from three days to two weeks intervals (Fig. A4).





Figure 8: HYSPLIT back trajectories of air parcels for 2010 (NAO-) and 2013 (NAO+), annual and summer (JJA). Trajectories are calculated on a two-weeks (336 h) interval at a 6-hour frequency; "precip" indicates trajectories that produced precipitation within 6 h from the SVID endpoint, while "dry" trajectories did not. Contour colours indicate the frequency at which air parcels part of a trajectory travel above a certain area.

336 5 Discussion

337 5.1 Primary aquatic production vs erosion/terrestrial input

338 In Icelandic lacustrine environments, $\delta^{13}C$ and C/N are generally considered proxies for the relative contribution of terrestrial 339 vs aquatic organic matter and shifts in primary productivity, as total carbon is virtually solely of organic origin (e.g., Geirsdóttir 340 et al., 2009a). Iceland's bedrock is dominantly comprised of basaltic bedrock, including the catchment of SVID (Hjartarson 341 and Sæmundsson, 2014), meaning there is negligible carbonate available. Although some dissolved inorganic carbon (DIC) 342 has been measured in Icelandic rivers, it is greatly outweighed by organic carbon (Kardjilov et al., 2006), whereas the amount 343 of inorganic carbon measured in soils is negligible (Mankasingh and Gísladóttir, 2019). This is important to consider as SVID 344 has no river inflow and water inflow is dominated by runoff from the catchment through soil. In addition, while there is some 345 evidence for the transport of Saharan dust to Iceland within the last decade (Varga et al., 2021), there is currently no evidence 346 of such transport during the Holocene. Additional pools of inorganic carbon from aquatic invertebrates, such as ostracods, are 347 also considered to be negligible (Alkalaj et al., 2019). Ostracods crystallise their shells in very close equilibrium to the carbon isotopes of DIC (Decrouy, 2012), which for Iceland is notably enriched relative to bulk organic matter carbon isotopes 348 349 (Sveinbjörnsdóttir et al., 2020). If ostracods were a substantial contributor to the total carbon pool, we would expect lake 350 sediment carbon isotopes to deviate significantly from modern terrestrial and aquatic plant carbon isotope values. As Icelandic 351 lake sediment bulk geochemistry is consistent with the fields of modern plants (see Geirsdóttir et al., 2020), inorganic carbon 352 from aquatic invertebrates are not considered a significant contributor to the total carbon pool.

In addition, in SVID, the similarity of the C/N record to the perylene curve reinforces its significance as a proxy for terrigenous
 input.

355 5.1.1 11-7.5 ka BP: Postglacial warming

Deglaciation in the NE of Iceland set in between 15 and 13 ka BP and proceeded in a stepwise fashion, with two main glacier re-advances at ca 12.7 (Younger Dryas) and 10.9 ka BP (Preboreal; Geirsdóttir et al., 2009b; Norðdahl and Pétursson, 2005). Our record captures sediment below the Askja S tephra layer (10.83 \pm 0.57 ka BP, Bronk Ramsey et al., 2015), showing icefree conditions and the start of organic sedimentation by 10.85 ka BP at SVID's location. Except for the oldest sample, high δ^{13} C and low TC values indicate a primarily aquatic source of carbon during the oldest interval (I) (Fig. 3). This suggests an absence of substantial terrestrial vegetation, consistent with a postglacial landscape and possibly a cooler climate associated with the Preboreal period.

- 363 TC, BSi, and C/N increase suddenly at ca 10.65 ka BP, maintaining higher values for two-three centuries (vice versa for δ^{13} C), 364 indicating an enhanced terrestrial input likely resulting from a retreating glacier, development of soil and vascular plants, and 365 generally warming conditions (Fig. 3). After the G10ka Series tephra, all proxy values decrease, likely due to the destructive 366 impact of substantial volcanic ash fallout on both terrestrial and aquatic vegetation and related water chemistry alteration (e.g., 367 δ^{13} C dropping due to acidification; Kilian et al., 2006) (Fig. 4). Following the volcanic event, all proxies increase at ca 9.75 ka 368 BP, whereas terrestrial-relative to aquatic-sourced carbon temporarily increase (ca 8.7–7.5 ka BP). The observed decrease in 369 δ^{13} C (and, partially, C/N) between 8.7 and 7.5 ka BP is identified in other Icelandic lake sediment records between 8.8 and 7.9 370 ka BP (e.g., Eddudóttir et al., 2018; Geirsdóttir et al., 2013; Harning et al., 2018b; Larsen et al., 2012) and has been attributed 371 to the likely impact of meltwater pulses into the northern North Atlantic due to the retreating Laurentide ice sheet and/or local
- 372 effusive volcanic eruptions (Geirsdóttir et al., 2013; Larsen et al., 2012).
- The total sum of *n*-alkanes (C_{19-35} ; Fig. 4A), which is heavily controlled by C_{29} and C_{31} (Fig. A3), increases throughout the
- 374 Holocene similar to the pattern described by C/N and perylene as a result of an increased terrigenous input. As inferred by
- 375 high CPI values, most *n*-alkanes in SVID originate from plants (terrestrial and possibly also aquatic) through the Holocene
- 376 record (Fig. 4I). The relatively low CPI values in the oldest interval (I) indicate a negligible contribution from plant sources

- 377 (relative to phytoplankton) to the carbon pool, which is consistent with a still cold, relatively barren, deglacial environment.
- 378 The CPI curve shows a similar but opposite pattern to the δ^{13} C record until ca 8 ka BP, reinforcing δ^{13} C as a proxy mostly
- 379 controlled by terrigenous input (*n*-alkanes from aquatics show lower CPI values than terrestrial plants; e.g., Bray and Evans,
- 380 1961; Duan et al., 2014; Eglinton and Hamilton, 1967; Li et al., 2020). The reason for the change in the relationship between
- 381 CPI and δ^{13} C (which become positively correlated after 8 ka BP) is unclear. While it matches the timing of increasing
- temperatures (Axford et al., 2007; Fig. 10D) and Betula expansion in the region (Karlsdóttir et al., 2014; Fig. 10E), its
- interpretation is complicated by the fact that CPI can also be influenced by factors such as changes in mean annual precipitation,
- seasonality, plant community, and algal productivity (Li et al., 2020).

385 5.1.2 7.5-4.2 ka BP: Mid-Holocene Plateau and trend inversion

- 386 Overall, all proxies suggest that the interval between ~7.5 and 4.2 ka BP, was characterised by relatively stable climatic 387 conditions, generally warmer (Axford et al., 2007) and wetter (Moossen et al., 2015) than both the preceding and following 388 periods. These conditions likely led to an enhanced primary productivity within the lake, as suggested by high values of BSi, 389 Pag, CPI, and pollen inferred vegetation communities (Eddudóttir et al., 2016; Karlsdóttir et al., 2014). In fact, modelled 390 reconstructions of Icelandic vegetation cover throughout the Holocene show the highest values during this interval (Ólafsdóttir 391 et al., 2001). In particular, partly due to the retreat/disappearance of most glaciers, between 50% and 60% of Iceland was likely 392 covered in vegetation, of which a quarter was birch forest, throughout the mid-Holocene, with two peaks, one at the Holocene 393 Thermal Maximum (8–7 ka BP) and one at ~3.5 ka BP (Ólafsdóttir et al., 2001). The warm and moist climate, paired with the 394 expansion of vegetation, may have stabilised the local environment, reducing erosion. Such stability is consistent with the low 395 C/N, Perylene, and *n*-alkanes values throughout this interval.
- 396 Though this four thousand year period is broadly categorised by stability, a more detailed view reveals important inflection 397 points in the long term trends of many proxies. For example, while some proxies keep increasing (e.g., TC, and summer 398 temperature, Axford et al., 2007), some stop rising and remain relatively flat throughout this interval (e.g., C/N, Perylene), 399 while others even invert their trends (δ^{13} C, P_{aq}, CPI). Another trend inversion occurred around 5 ka BP with the inception of 400 neo-glaciation in Iceland when glaciers started to expand again (Geirsdóttir et al., 2019). This could be interpreted as a slow 401 inertial response of the local environment to the decreasing NH summer insolation, likely reducing its resilience to short term 402 events such as volcanic eruptions and NAO shifts, until some kind of threshold was finally reached around 4 ka BP (Geirsdóttir 403 et al., 2013, 2019).

404 5.1.3 4.2 ka BP: Increased erosion in a cooling climate

405 Our SVID proxy datasets generally agree with previous work using bulk geochemistry proxies in Icelandic lakes, which 406 collectively point toward decreasing primary productivity and increasing landscape instability in response to declining 407 Northern Hemisphere summer insolation (e.g., Geirsdóttir et al., 2013, 2019; Harning et al., 2018b, 2020; Larsen et al., 2012). 408 The decreasing trend in δ^{13} C is generally anticorrelated with the TC curve, indicating that TC is increasingly controlled by 409 terrestrial input. BSi, TC, and C/N values drop or invert their trend after ca 4 ka BP, consistent with a general decrease in 410 productivity. This is possibly related to a combination of decreasing moisture and/or summer temperatures (Axford et al., 411 2007), and the effect of the Hekla 4 volcanic event, pushing the local environment beyond a threshold (Eddudóttir et al., 2017). 412 Absolute amounts of *n*-alkanes also increase starting at 4 ka (Fig. 4A), led by an increase in $n-C_{29}$ (Fig. A3), typical of terrestrial 413 plants. The ratio between the mid- and long-chain homologues (aquatic plant index or Paq; Ficken et al., 2000) is often 414 interpreted as a proxy for a wetter/dryer environment. However, this is likely an oversimplification as questions remain about 415 the relationship between n-alkane chain length and vegetation source, particularly for aquatic plants which are often a minor 416 component of the leaf wax pool in Arctic lakes (Dion-Kirschner et al., 2020; Hollister et al., 2022). Nevertheless, when coupled 417 to the concentration data of *n*-alkanes (Fig. 4A; Fig. A3), P_{ag} can here be more safely interpreted as indicative of lower/higher

- 418 terrigenous input. The SVID record (Fig. 4H; Sect. 3.3) shows higher P_{aq} values (mean 0.6) during the 10.8 to 4-3 ka BP
- interval, indicating an environment with significant aquatic plant production and likely limited erosion/in-wash. The record
 then switches abruptly to lower values (mean 0.3) at 3 ka BP, highlighting a shift toward greater in-wash of terrestrially derived
 material.
- 422 The massive Hekla 3 eruption (the most severe Hekla eruption of the Holocene; ~3,010 a BP; Larsen, 1977; Larsen and 423 Eiríksson, 2008), was likely the cause, or at least the trigger, of this abrupt shift at 3 ka BP in most SVID proxy records, 424 particularly the ones related to primary production and erosion (Larsen et al., 2011). In fact, the volcanic fallout likely killed 425 terrestrial plants by burning, root suffocation, and reduced photosynthesis (e.g., Ifkirne et al., 2022; Mack, 1981; De Schutter 426 et al., 2015), and had likely similar effects on aquatic flora as well, also through increased turbidity and acidity of lake waters 427 (e.g., Ayris and Delmelle, 2012). The subsequent reduced coverage of terrestrial plants likely exposed the soil to increased 428 erosion, resulting in more terrestrial in-wash (as reflected by a sudden perylene peak), skewed toward the inorganic components 429 of soil (as reflected by a sudden drop in TC and *n*-alkanes lasting roughly a century). The increased terrestrial in-wash would 430 have further reduced primary productivity within the lake, as suggested by the drop in BSi. At the same time, the short C/N 431 peak and the major drop in P_{ag} seem to indicate that the productivity and contribution of terrestrial plants remained higher than 432 aquatic sources (Larsen et al., 2011). The post-3 ka trend is temporarily interrupted by what appears to be a partial rebound in 433 primary productivity (BSi increases too) and diminished in-wash of terrestrial material until ca 1.5 ka BP, whereafter it 434 continues to decline.

435 **5.2** Is there geochemical evidence for human settlement in the SVID catchment area?

436 In paleoclimate studies, relative shifts to above natural background levels of β -stanols have been used as a proxy for human 437 settlement, marking the appearance of humans and domesticated animals in specific areas of the world (Shillito et al., 2020; 438 Sistiaga et al., 2014), often in lake catchments (Battistel et al., 2016; Callegaro et al., 2018; Raposeiro et al., 2021; Sear et al., 439 2020; Vachula et al., 2019, 2020). This method has detected the arrival of Viking settlers in other Nordic regions, such as in 440 the Lofoten islands in northern Norway (D'Anjou et al., 2012, Fig. 9B) and in the Faeroe Islands (Curtin et al., 2021, Fig. 9C). 441 However, SVID sterol/stanol records show no evident human signals at or around the time of colonisation (i.e., 9th century CE, 442 ca 1.1 ka BP). While a relative maximum of sterol/stanol concentrations found at ca 1.4 ka BP resembles the timing of an 443 earlier-than-colonisation stanol signal found in the Faeroes (Curtin et al., 2021), as well as an analogous signal in the Lofoten 444 Islands (D'Anjou et al., 2012), we cannot confidently interpret this peak as indication of human presence as its amplitude is 445 comparable in magnitude to the inherent variability in the record (i.e., low signal to noise ratio).



Figure 9: Holocene sub-Arctic records of faecal stanols in North Atlantic Islands. (A) 5β-stigmastanol (dark red) and sum of coprostanol and epi-coprostanol (dark blue) from core 20-SVID-02, NE Iceland (this study); (B) 5β-stigmastanol (pink) and coprostanol (light blue) from cores LILA09-LILC09 from Lilandsvatnet lake, Lofoten Islands (D'Anjou et al., 2012); (C) 5βstigmastanol (orange) and coprostanol + epi-coprostanol (green) from core EI-D-01-15 from Eiðisvatn lake, Faroe Islands (Curtin et al., 2021).

452 The lack of a clear anthropogenic faecal biomarker signal could be explained by either (1) a scarce/null incidence of human 453 activities in the catchment (unlikely, given the archaeological evidence in nearby areas; Gísladóttir et al., 2012; Lebrun et al., 454 2023) and/or by (2) dilution of the signal due to the relatively large size of the lake paired to a small catchment. The sterol/stanol 455 records show a general increase throughout the Holocene (Fig. A2) in a pattern that matches the C/N, n-alkanes and perylene 456 trends, suggesting that the primary driver of SVID's sterol signal is likely landscape instability and soil erosion rather than 457 human/ruminant presence. Furthermore, ratios of sterols to their derived 5β-5α stanols can trace redox conditions in various 458 environments (e.g. Andersson and Meyers, 2012; Canuel and Martens, 1993; Jaffé et al., 1996; Routh et al., 2014) and thus, 459 potentially, human presence in a lake catchment, as anthropogenic activities tend to mobilise more soil and increase in-wash 460 of organic material, fostering reducing conditions (Argiriadis et al., 2018). In SVID, the stanol values (5β - 5α) are consistently 461 lower than their respective sterol precursors, suggesting a generally oxidising environment throughout the Holocene (Fig. 6). The only exception to this trend is in the earliest part of the record (ca 10.8-10.6 ka BP; Fig. 5), indicative of a more reducing 462 463 environment, though not linked to an increased organic input (low TC values), but more likely to lake stratification with 464 deglacial water sinking at the bottom of the lake (Sugiyama et al., 2021).

465 **5.3 Holocene fire frequency**

Pyrogenic PAHs are considered a reliable proxy for fire frequency on a local scale, within and around a catchment (Denis et al., 2012). Although other factors can influence the PAH signal in sedimentary archives (e.g., accumulation rates, degradation;
Stogiannidis et al., 2015), we interpret SVID pyroPAHs data as a record of NE Iceland fire history through the Holocene.

- 469 The trend in pyroPAHs does not match the erosional signal described by bulk geochemical proxies and *n*-alkanes, suggesting
- 470 that soil erosion is not a mechanism for Holocene pyroPAHs variability. We exclude chemical degradation as a source of the
- 471 signal, as PAHs are relatively stable molecules on long time scales (e.g., Johnsen et al., 2005). In fact, the ratio between low
- 472 molecular weight (more prone to chemical degradation and leeching) and high molecular weight PAHs (as defined in Fig. A5)
- 473 remains above 1 in most samples. Similarly, the pyrene/coronene ratio shows high and stable values throughout the record,
- 474 indicating good preservation, with no significant degradation or preferential removal of less recalcitrant PAHs such as pyrene
- 475 (Fig. A5 and refs therein).
- 476 The pyroPAH record presents two peaks at ca 2.8 and 1.5 ka BP, both predating acknowledged human settlement. PyroPAH values subsequently drop (VIII) and increase again in the last two centuries reaching maximum values in the present (Fig. 7A). 477 478 Analysing PAH data subdivided in molecular weight classes (see 3.1; Table A1; Fig. 7B) can help explain these two features 479 as well as the general trend. HMW pyroPAHs show low and stable relative contributions (12±7% of total pyroPAHs) through 480 the whole record and rise to 70% in the last 200 years. This trend, paired with increased pyroPAH concentrations, is consistent 481 with the burning of coal/oil, the use of internal combustion engines, and increased human presence (Abas and Mohamad, 2011; 482 Kozak et al., 2017). MMW pyroPAHs show relatively stable concentrations through most of the record (until ca 0.2 ka BP). 483 Their relative abundance (~68±35%) slowly decreases through the Holocene, proportionally to the increase of LMW 484 pyroPAHs ($\sim 20\pm 16\%$). The latter, which are predominantly present in the gaseous phase (Karp et al., 2020), peak at 3–2.8 and 485 1.5 ka BP, and substantially control the shape of the pyroPAH record (Fig. 7A-B). Together, the (1) low and stable values of 486 HMW pyroPAHs, the (2) stable MMW values, and the (3) increasing/peaking values of LMW pyroPAHs are consistent with 487 a general increase in the frequency of low temperature fires (e.g., peat fires or crawling fires) at a regional level. While the 488 Hekla 3 event (3.01 ka BP, the largest rhyolitic eruption during the Holocene; Larsen, 1977; Larsen and Eiríksson, 2008) 489 occurs just before the first pyroPAH peak, it is unlikely to be its unique or even main cause. However, in an environment such 490 as Iceland, is legitimate to ask if the several and frequent eruptions might have influenced the PAHs' natural background and 491 if this is detectable in our records. As discussed below, we are confident that volcanic eruptions have had no significant direct 492 impact on the amount and distribution of PAHs in the SVID archive.
- 493 Volcanic activity does produce PAHs (both in the gaseous and in the particulate phases) but generally their long term
 494 contribution is considered negligible, particularly in the modern world where the main PAHs source is the burning of fossil
 495 fuels (Guiñez et al., 2020; Kozak et al., 2017).
- 496 Overall, due to the high temperatures involved, volcanic eruptions tend to produce medium-high molecular weight PAHs in 497 gases and particulates (Guiñez et al., 2020; Ilyinskaya et al., 2017). Volcanic layers can contain pyrogenic (unsubstituted but 498 also alkylated) PAHs with a molecular weight distribution resembling modern fossil fuel combustion, dominated by 499 unsubstituted forms (Murchison and Raymond, 1989) as well as traces of nitro- and oxy-PAHs (Guiñez et al., 2020). We do 500 not see this distribution in any of our samples, not even when they include parts of tephra layers. This suggests that (1) either 501 our sampling method does not capture volcanic layers (maybe due to its resolution) or that (2) there is no such signal in the 502 SVID archive.
- Regarding the first hypothesis (sampling method not capturing volcanic PAHs), Kozak et al. (2017) analysed the impact of 2010 and 2011 Icelandic eruptions in Svalbard (Arctic Norway), finding high abundances of 4-5 ring PAHs in volcanic mud, great variations in the contribution of different eruptions to the total PAHs detected in sampled surface water, significant increases in the total abundance of PAHs during eruption years, but also that this increase in PAHs abundance as well as shifts
- 507 in PAHs distribution do not seem to last beyond the eruption years. Considering the temporal resolution of our record (max

- 508 10-50 years), it is unlikely that any eruption would have impacted it significantly: from a geological point of view, eruptions
- 509 tend to be short-living, and unless they relate to a massive event sustained over a long period of time, they are unlikely to have
- 510 a strong impact on a sample that represents 10-50 years of sedimentation. No discernible correlation arose between the PAHs
- 511 curve and the detected tephra, the only exceptions being the Hekla 3 and, to a lesser extent, the Hekla 4 tephra layers. However,
- 512 these major volcanic events, besides marking the beginning of major shifts in most proxies, correlate to the initial phase of
- 513 major shifts in low (and not high nor medium) molecular weight PAHs. This suggests that, if a connection between these two
- 514 eruptions and PAH shifts exists, it must be indirect and, more likely, the two events acted as a general destabilising factor in
- an environment already subjected to increasing cooling and erosion.
- 516 The only possible example of an increased PAH concentration due to volcanic sources could come from the sample obtained
- from the lower limit of the G10ka series tephra (Fig. 7), which has a different composition (mostly inorganic) compared to the rest of the organic-rich samples in our record. In fact, HMW PAHs seem to spike here, even if they still exhibit an overall lower concentration than LMW PAHs; this could be due too its massive nature (Óladóttir et al., 2020).
- Regarding the second hypothesis (missing signal of volcanic PAHs), it is possible that no detectable volcanic PAHs were present in the SVID archive due to (1) its geographical location, relatively far from volcanic sources and formations (Hjartarson and Sæmundsson, 2014), and (2) the same nature of Icelandic volcanic eruptions, which are characterised by relatively lowviscosity basaltic lava rather than highly explosive pyroclastic flows (Thordarson and Höskuldsson, 2008), thus reducing the chance of ash production and deposition, particularly in distal locations such as SVID.
- 525 The effects of tephra fallout on vegetation and related PAH deposition, seem to be quite short lived, with fires events likely 526 coeval to the eruption and vegetation recovering within a few decades (Eddudóttir et al., 2017; Pickarski et al., 2023), while
- 527 SVID pyroPAH peaks are clearly led by increases in LMW PAHs on a longer timescale.
- Notably, this shift in fire regime at ca 3 ka BP in Iceland falls within a wider pattern of increasing fire frequency emerging from the analysis of several Holocene fire records throughout Europe (Marlon et al., 2013). This is linked to either an increase of cultivated land (fire was used to clear land for agriculture) and/or, particularly in Europe, to increasing aridity (Marlon et al., 2013). We hypothesize that the latter is the most probable explanation for the shifts in NE Icelandic fire regimes as discussed in Section 5.4.

533 5.4 Regional drivers of precipitation and their role on fire frequency

- 534 Fuel moisture content and, more generally, environmental moisture, are the main variables controlling flammability in 535 vegetational communities typical of temperate/sub-arctic regions (Marino et al., 2010; Plucinski et al., 2010; Santana and 536 Marrs, 2014). The North Atlantic Oscillation (NAO; Hurrell et al., 2003) modulates the intensity of the westerly storm track 537 and thus the amount and source of precipitation in Iceland. Its positive mode (NAO+) brings intervals of higher precipitation 538 resulting in a wetter (and often warmer) climate than NAO- intervals, which are characterised by weaker westerlies, stronger 539 northerly winds, and drier (and often colder) conditions (Hurrell, 1995; Trouet et al., 2009). Major changes in precipitation 540 regimes usually lead to changes in the hydrogen stable isotopic value of environmental water (δD ; Dansgaard, 1964) which 541 translate into shifts in the δD of plant waxes (e.g., *n*-alkanes; Sachse et al., 2012). This relationship has been calibrated in 542 various environments, including the Arctic (e.g., Berke et al., 2019; Bush et al., 2017; McFarlin et al., 2019; Thomas et al., 543 2016) and applied for paleo-precipitation reconstructions (e.g., Ardenghi et al., 2019; Niedermeyer et al., 2016; Tierney et al., 544 2017; Wilkie et al., 2013). A C₂₉ n-alkane δD record from a fjord core in NW Iceland (Fig. 10C; Moossen et al., 2015) describes 545 a relatively stable NAO+ configuration (wetter - more D-depleted) throughout the Holocene, and two major shifts toward 546 NAO- (dryer - less D-depleted) conditions at ca 3–2.5 and 1.5–1.0 ka BP, matching the timing of first SVID pyroPAH peak 547 and at least partially overlapping to the second one. A similar correlation of fire frequency shifts to NAO- configurations has 548 recently been suggested for other Arctic sites, particularly in Svalbard (Chen et al., 2023).
- 549 Biomass typology (i.e., the kind of vegetation on site) also influences fuel flammability (Chandler et al., 1983; Fernandes and

550 Cruz, 2012; Santana et al., 2011; Scarff and Westoby, 2006). In Iceland, many plant taxa appeared shortly after deglaciation 551 (e.g., Alsos et al., 2021; Harning et al., 2023). Increasing summer temperatures led to the expansion of thermophilic woody 552 plant taxa (e.g., birch) during/after the Holocene Thermal Maximum (e.g., Eddudóttir et al., 2016; Geirsdóttir et al., 2022; 553 Karlsdóttir et al., 2014). Since ~6 ka BP, the birch woodland in the NE region has evolved into more open heathland and 554 peatland, until the birch population decreased around 3 ka BP (Roy et al., 2018), along with a general temperature decrease 555 (Axford et al., 2007). In this context, at Ytra-Áland (Fig. 1B), two major drops in *Betula* pollen coeval to two increases in Ericales (heather's order) pollen closely follow the NAO- shifts at 3 and 1.5 ka BP (Fig. 10E; Karlsdóttir et al., 2014). 556 557 Heathlands, especially in low moisture conditions, are associated with higher flammability, particularly high sustainability 558 (i.e., how well the combustion proceeds). Thus, heathlands are prone to longer, more stable fires, and with an increased potential for igniting higher canopy elements and underlying peat layers (Plucinski et al., 2010; Rein et al., 2008; Santana and 559 560 Marrs, 2014). More frequent, more stable, slow crawling fires involving dense bushes and peat would increase the amount of 561 pyroPAHs produced and deposited in the region while skewing their distribution toward LMW components (George et al., 562 2016; Iinuma et al., 2007; Siao et al., 2007), as observed in the SVID pyroPAH signal (Fig. 7). Lastly, the observed shifts in 563 pyroPAHs are unlikely to be the result of a change in their source area. Our back-trajectory analysis reveals that more air 564 parcels originate over Iceland or in the surrounding North Atlantic (Fig. 8). The analysis further reveals that trajectories with 565 terrestrial origins are more likely in NAO+ than NAO- regimes. As such terrestrial trajectories would be the ones responsible 566 for bringing combustion products to SVID, we might therefore expect a stronger pyroPAH signal during NAO+ intervals. However, our results display the opposite trend, with the initial increase in pyroPAH abundances at ~3 ka occurring during an 567 568 NAO- mode and their subsequent drop occurring during a strong NAO+ interval. Thus, the late Holocene increase in pyroPAHs 569 in SVID is likely to record a substantial increase in local fires (driven by vegetation change and NAO-modulated aridity) rather 570 than a shift in the compounds' sources.

571 Overall, SVID pyroPAH signal describes a major shift from a relatively stable (i.e., low fire frequency) early and mid-Holocene 572 environment to a dryer late-Holocene environment at ca 3 ka BP, naturally more prone to long term persistence of low 573 temperature wildfires. This is likely the result of the combination of (1) recorded cooling and related shifts in vegetational 574 communities, (2) NAO- shifts and associated dryer conditions.



575

Figure 10: Regional comparison of climatic, fire, and vegetation records. (A, purple) levoglucosan concentrations from the RECAP ice core, south-eastern Greenland (Segato et al., 2021); (B, black) sum of pyrogenic PAHs concentrations – red line indicates a 3 points running average; (C, blue) stable isotopic composition of hydrogen of sedimentary *n*-alkanes from marine core MD99-2266 off the coast of NW Iceland (Moossen et al., 2015) – cyan line indicates a 3 points running average; (D, red) chironomid derived temperatures from core 04-SVID-03 (Axford et al., 2007); (E) pollen percentages of *Betula* (green), Poaceae (yellow), and Ericales (blue) in a peat section from the Ytra-Áland site, NE Iceland (Karlsdóttir et al., 2014).

582 **5.5** Evidence for a human influence on fire frequency?

583 Unlike other proxies, pyroPAHs return to background levels after reaching high values at 1.5 ka BP, and then remain low 584 through the Medieval Warm Period (ca 900-1200 CE) and most of the Little Ice Age (1300-1900 CE), before peaking again 585 in the last 150–200 years. A similar drop in fire markers (anhydrosugars) is also observed in eastern Greenland (Segato et al., 586 2021; Fig. 10A). Low pyroPAHs levels during an interval of known human presence in Iceland suggests that human activities 587 might have curbed regional fire frequency, thus modulating the natural signal, which would have otherwise remained relatively 588 high due to the persistence of colder conditions and more flammable plant communities (regardless of NAO shifts). In fact, 589 while increased pressure from grazing lowered environmental resilience to soil erosion (e.g., Bates et al., 2021; Eddudóttir et 590 al., 2016; McGovern et al., 2007), it likely also decreased fuel flammability (which is dependent on the amount of dead 591 biomass; Davies and Legg, 2011; Santana and Marrs, 2014) in the predominant heathland environment (Lake et al., 2001). 592 Additionally, the creation of farmland and pastureland at the expenses of areas with woody vegetation and heathland likely 593 reduced the extent of the biomes naturally prone to fires. This reduction of local wood availability is reflected in changes of

- 594 foraging habits, as settlers shifted to relying more heavily on more abundant fuel sources such as peat and turf, as well as other
- 595 marine derived substances (e.g., seal oil, seaweed; Bold, 2012), while driftwood and imported wood (from Europe or North
- 596 America, often with ad hoc expeditions) satisfied most of the need for timber (e.g., Bold, 2012; Edvardsson, 2010; Mooney,
- 2016; Pinta, 2021; Sveinbjarnardóttir et al., 2007). A general mechanism for fire suppression due to the expansion of cultivated
 land has already been proposed for global data (Marlon et al., 2013), but assumed to be likely asynchronous in different regions
- and strongly influenced by local climatic, environmental, and social conditions.
- 600 We speculate that the drop in fire markers in Iceland from the reduction in wildfire risk due to husbandry and farming exceeded
- 601 the production of fire markers due to human necessities (e.g., warming), resulting in an overall suppressed fire signal. This
- 602 would also be consistent with the low population density, which started to increase only in the 1800s CE (Iceland Statistical
- 603 Service, 2023; Jónsson and Magnússon, 1997), matching the coldest interval of the Little Ice Age and the sharp rise in HMW
- 604 pyroPAHs. From this perspective, the pressure of human activities would have fostered erosion through decreased
- 605 environmental resilience while, at the same time, suppressing natural fire frequency.

606 6 Conclusions

- Our multiproxy analysis of Holocene sediments from Stóra Viðarvatn provides new insight into the coupled vegetation, fire,
 erosion, and climate regimes of NE Iceland:
- Bulk geochemistry proxies show that the general climatic evolution of NE Iceland is primarily driven by summer
 insolation: an initial deglacial warming followed by a relatively warm and stable climate until ca 4–3 ka BP, after which
 declining summer temperatures result in accelerating catchment erosion.
- Faecal biomarkers, traditionally linked to human activities, do not show an elevated signal at or around colonisation (9th 613 century CE). Instead, faecal biomarkers roughly trace the erosional signal described by bulk geochemical proxies. This 614 may result from a combination of (1) low local anthropogenic pressure (although sparse settlements existed a few km 615 from the study area), and (2) signal dilution, due to the large lake size and its relatively small watershed. Therefore, we 616 urge caution when interpreting faecal biomarkers as unequivocal proxies of human presence, particularly when highly 617 sensitive analytical tools like the one used in this study are involved.
- 618 PyroPAHs carry a regional (mostly confined to northern and north-eastern Iceland) and predominantly natural signal 619 (i.e., controlled by parameters such as precipitation and moisture availability, vegetation typology and flammability). 620 After generally low fire frequency throughout most of the Holocene, we observe major regime changes at 3 ka and 1.5 621 ka BP, before known human colonisation in Iceland. During this interval, the distribution of pyroPAHs point toward a 622 regional increase in low temperature fire frequency. This can be linked to a change in vegetation typology driven by the 623 cooling of the last 4 to 3 kyr, coupled to major shifts in atmospheric circulation (i.e., NAO regimes) that led to increased 624 aridity and thus flammability. Finally, low levels of pyroPAHs characterise the time following known human colonisation, before rising again (but with a molecular composition more distinctive of fossil fuels) in the last ~200 years. 625 626 This suggests that human activities, particularly husbandry and farming, may have suppressed fire frequency by reducing 627 the range and flammability of environments more prone to fire, effectively modulating the natural signal while decreasing 628 the resilience of the local environment to soil erosion.



Figure A1: Concentration curves of all 20-SVID-02 pyrogenic PAHs recovered in this study. All concentrations are expressed as ng per g of dry sample, except for the last curve (red-dotted) which is in ng per g of TC. Note that several vertical axes have been adjusted to minimise the rise in the last 2–3 centuries. Compounds are listed in chromatographic order and grouped by molecular weight through colour shading (LMW in magenta; MMW in orange; HMW in blue).





Figure A2: Concentration curves of all 20-SVID-02 faecal sterol/stanols of interest recovered in this study. All concentrations are
 expressed as ng per g of dry sample. Note that several vertical axes have been adjusted to minimise the rise in the last 2–3 centuries.
 Compounds are grouped and colour shaded by structure (cholesterol and stanol derivatives in blue purple; sito-stigma- sterols and
 stanols in red orange; campesterol and campestanol in green).



Figure A3: Concentration curves of all 20-SVID odd-numbered *n*-alkane homologues from *n*-C₁₉ to *n*-C₃₅ recovered in this study.
 All concentrations are expressed as ng per g of dry sample.







647 Figure A4: HYSPLIT back trajectories of air parcels for December 2009 to November 2010 (NAO-) and May 2013 to April 2014 648 (NAO+, except for October 2014). Trajectories are calculated on a three days (72 h, A-C) and a two weeks (336 h, B-D) intervals at 649 a 6 hours frequency; "precipitation" (top) indicates trajectories that produced precipitation within 6 h from the SVID endpoint, 650 "dry" (middle) vice versa; bottom plots are the sum of "precipitation" and "dry" trajectories. Contour colours indicate the frequency 651 at which air parcels part of a trajectory travel above a certain area.



652

653 Figure A5: PAH indices used as an indication of PAH preservation.

654The L/H index (black) is a ratio between low and high molecular weight unsubstituted PAHs, defined as L/H = (Phenanthrene +
Anthracene + Fluoranthene + Pyrene) / (Benzo[a]anthracene + Chrysene + Benzo[k]fluoranthene + Benzo[a]pyrene +
Indeno[1,2,3,c,d]pyrene + Dibenzo[a,h]anthracene + Benzo[g,h,i]perylene) (Magi et al., 2002; Stogiannidis et al., 2015 and refs.
therein).

The Pyrene-Coronene index (orange) defined as pyrene / (pyrene + coronene) is based on the assumption of an higher preservation

659 potential of the HMW coronene over the lighter, more soluble pyrene (Denis, 2016; Denis et al., 2017; May et al., 1978); higher, more

660 stable values point toward good preservation for both HMW and LMW PAHs.

Table A1: Polycyclic aromatic hydrocarbons (PAHs) analysed in this study. Pyrogenic PAHs are grouped into low, medium, and high molecular weight. Elution order and SRM transitions are reported for each compound.

Elution order	Group	Compound name	Mass	Product mass a/b	Collision energy
1		Naphthalene*	128	128/102	8
2		Acenaphthylene*	152	152	8
3	M	Acenaphthene*	154	153/154	8
4	Ľ	Fluorene	166	166	8
5		Phenanthrene	178	178	8
6		Anthracene	178	178	8
7		Fluoranthene	202	202	8
8	~	Pyrene	202	202	8
11	M	Retene	234	234/219	8
12	Ą	Benzo[a]anthracene	228	228	8
13	F-	Triphenylene	228	228	8
14		Chrysene	228	228	8
15		Benzo[k]fluoranthene	252	252	8
16		Benzo[j]fluoranthene	252	252	8
17	8	Benzo[a]pyrene	252	252	8
19	ž	Indeno[1,2,3-C,D]pyrene	276	276	8
20	H	Dibenzo[a,h]anthracene	278	278	8
21		Benzo[g,h,i]perylene	276	276	8
22		Coronene	300	300	8
18		Perylene**	252	252	8
9		p-Terphenyl D14 (IS)	244	244	8
10		p-Terphenyl (IS)	230	230	8

* Compound(s) difficult to quantify correctly and thus excluded from final sums. ** Non-pyrogenic PAH.

Table A2: Faecal sterols and stanols analysed in this study. Elution order and SRM transitions are reported for each compound.

Elution order	Name	Detailed name	CAS #	Quantitative qualitative	Mass	Product Mass	Collision Energy
1	Pregnanol (IS)	5β-pregnan-3α-ol	4352-07-2	Q q	361 361	215 191	10 10
2	5α-Cholestane (IS)	5α-cholestane	481-21-0	·	372	217	10
3	Coprostanol	5β-cholestan-3β-ol	360-68-9	Q q	370 460	215 215	10 10
4	Epi-Coprostanol	5β-cholestan-3α-ol	516-92-7	Q q	370 460	215 215	10 10
5	Cholesterol	5-en-cholest-3β-ol	57-88-5	Q q	368 458	145 129	20 50
6	Cholestanol	5α-cholestan-3β-ol	80-97-7	Q q	370 460	215 215	10 10
7	5β-Campestanol	24R-methyl-5β-cholestan-3β-ol	33947-18-1	Q q	384 474	215 215	10 10
8	5β-Stigma(sito)stanol	24R-ethyl-5β-cholestan-3β-ol	4736-91-8	Q q	383 488	147 215	20 10
9	Campesterol	24R-methyl-5-en-cholest-3β-ol	474-62-4	Q	382 472	255 129	5 50
10	5α-Campestanol	24R-methyl-5α-cholestan-3β-ol	474-60-2	Q q	384 474	215 215	10 10
11	β-Stigmasterol	24S-ethyl-5,22E-dien-cholest-3β-ol	83-48-7	Q q	394 484	255 211	5 20
12	β-Sitosterol	24R-ethyl-5-en-cholest-3β-ol	83-46-5	Q q	396 484	255 394	10 10
13	5α-Stigma(sito)stanol	24R-ethyl-5 α -cholestan-3 β -ol	83-45-4	Q q	383 488	147 215	20 10

668 8 Data availability

- 669 All raw data will be available on the NOAA National Centers for Environmental Information
- 670 (https://www.ncei.noaa.gov/access/paleo-search/study/38503.).
- The data will also be made available upon request.

672 9 Author contributions

673 GHM, ÁG and JS conceptualized research and obtained financial support for the NSF project ILLUME (Iceland landscape 674 reconstruction using molecular proxies); GHM, ÁG, JHR, DJH, and NA participated in the field campaign to retrieve the 675 sediment core; JS and GHM provided laboratory and analytical infrastructure; NA, DJH, and BRH processed all sediment 676 samples; NA performed method development and sample analysis. TT and DJH performed the tephra analysis and developed 677 the age model; NA performed the HYSPLIT analysis; NA wrote the manuscript draft, except for the age model paragraph

678 (DJH); GHM, ÁG, JS, DJH, and JHR reviewed and edited the manuscript.

679 10 Competing interests

680 The authors declare that they have no conflict of interest.

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