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Proxy reconstruction of ultraviolet-B irradiance at the Earth's surface, and its relationship with solar activity and ozone thickness

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DOI: 10.1177/0959683619875798

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Document Version Peer reviewed version

Citation for published version (Harvard):

Jardine, PE, Fraser, WT, Gosling, WD, Roberts, CN, Eastwood, WJ & Lomax, BH 2019, 'Proxy reconstruction of ultraviolet-B irradiance at the Earth's surface, and its relationship with solar activity and ozone thickness', *The Holocene*. https://doi.org/10.1177/0959683619875798

Link to publication on Research at Birmingham portal

Publisher Rights Statement: Checked for eligibility: 18/07/2019

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1	Proxy reconstruction of ultraviolet-B irradiance at the Earth's surface, and
2	its relationship with solar activity and ozone thickness
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26 Abstract

27 Solar ultraviolet-B (UV-B) irradiance that reaches the Earth's surface acts as a biotic stressor and has the potential to modify ecological and environmental functioning. The challenges of 28 reconstructing UV irradiance prior to the satellite era mean that there is uncertainty over long-29 term surface UV-B patterns, especially in relation to variations in solar activity over 30 centennial and millennial timescales. Here, we reconstruct surface UV-B irradiance over the 31 32 last 650 years using a novel UV-B proxy based on the chemical signature of pollen grains. We demonstrate a statistically significant positive relationship between the abundance of UV-33 B absorbing compounds in Pinus pollen and modelled solar UV-B irradiance. These results 34 35 show that trends in surface UV-B follow the overall solar activity pattern over centennial 36 timescales, and that variations in solar output are the dominant control on surface level UV-B flux, rather than solar modulated changes in ozone thickness. The *Pinus* biochemical response 37 demonstrated here confirms the potential for solar activity driven surface UV-B variations to 38 impact upon terrestrial biotas and environments over long timescales. 39

40

Keywords: Ultraviolet-B irradiance; pollen; vegetation; palaeoclimate; solar activity; ozone
42

43 Introduction

The role of solar activity in influencing the Earth system has received an increase in 44 attention over recent years (Ermolli et al., 2013; Gray et al., 2010; Solanki et al., 2013). 45 Predominantly, the focus has been on total solar irradiance (TSI, defined as the amount of 46 solar energy reaching the upper atmosphere) and its contributions to climatic changes versus 47 anthropogenic inputs (Solanki et al., 2013). In addition to TSI, which affects temperature and 48 atmospheric circulation patterns through 'bottom up' warming of the Earth's surface, there is 49 a growing awareness of the importance of ultraviolet (UV) irradiance as a climate forcing 50 mechanism (Gray et al., 2010; Ineson et al., 2015). UV irradiance stimulates production and 51

destruction of ozone via absorption driven processes (the Chapman cycle), resulting in a
warming of the stratosphere and exerting a 'top down' influence on regional climatic and
oceanic patterns through dynamical coupling with the underlying troposphere (Ermolli et al.,
2013; Gray et al., 2010; Ineson et al., 2015; Solanki et al., 2013).

56

Solar UV-B (280-315 nm) radiation that reaches the Earth's surface (referred to 57 hereafter as surface UV-B) is an important stressor on biotic systems, and has the potential to 58 drive larger-scale ecosystem-level processes (Rozema et al., 1997). As well as directly 59 damaging DNA, enhanced UV-B levels can lead to morphological and phenological changes 60 61 in plants and possibly alter competitive relationships among species (Caldwell et al., 1998; Rozema et al., 1997). UV-B stimulates production of secondary metabolites that in part act as 62 UV protective compounds, which are both a metabolic cost to the plant and can influence 63 herbivory levels, plant decomposition and carbon cycling (Meijkamp et al., 1999; Rozema et 64 al., 1997). Increased surface UV-B can also directly enhance leaf litter decomposition through 65 photodegradation, and impact upon the activities of organisms such as animals, fungi and 66 bacteria that play a role in decomposition and nutrient cycling (Gehrke et al., 1995; Rozema 67 et al., 1997). While much attention has been focused on multidecadal increases in surface 68 69 UV-B flux due to anthropogenic reductions in ozone thickness (Caldwell et al., 1998; Lomax et al., 2008), considerably less is known about the surface UV-B changes that result from long 70 term (centuries to millennia) variations in solar activity, and what impact these have on 71 ecological and environmental functioning. 72

73

Temporal changes in surface UV-B flux are a result of variations in solar UV
irradiance, ozone thickness, and their interaction. Variations in solar activity have been
characterized by satellite measurements since 1978 (all dates given in calendar years CE), and
historical proxies such as sunspot counts from 1610 and cosmogenic radionuclides (primarily

¹⁰Be and ¹⁴C) through the Holocene (Solanki et al., 2013; Steinhilber et al., 2012; Svalgaard 78 79 and Schatten, 2016). These indicators reveal solar cycles ranging in length from ~27 days to several millennia, as well as irregularly spaced, sustained 'Grand' minima and maxima of 80 81 solar activity (e.g. the Maunder Minimum, ~1645 to 1710) (Solanki et al., 2013; Usoskin, 2017). Temporal variations in solar UV irradiance are still poorly understood, partly because 82 of the discontinuous nature of spectral solar irradiance (SSI) satellite measurements 83 84 (Haberreiter et al., 2017), but mostly because of the challenges of reconstructing UV irradiance beyond the satellite era. Nevertheless, SSI satellite measurements have revealed 85 UV cycles (in particular the 11-year solar cycle) in phase with those of TSI (Usoskin, 2017), 86 87 and the long term constancy of this relationship has been assumed in models of TSI and SSI such as the semi-empirical SATIRE-T model (Spectral And Total Irradiance REconstructions 88 for the Telescope era), which has provided UV reconstructions back to the Maunder 89 90 Minimum (Krivova et al., 2010), and the empirical NRLSSI2 model (Naval Research Laboratory Solar Spectral Irradiance), which has recently been extended back to 850 (Lean, 91 92 2018).

93

Increased UV-C (100-280 nm) flux during enhanced solar activity stimulates ozone 94 production, limiting the flow of UV-B to the Earth surface. It follows that while incoming 95 (top of atmosphere) UV-B and TSI may be correlated through time, surface level UV-B will 96 be anticorrelated with both (Rampelotto et al., 2009; Rozema et al., 2002), and this is 97 supported by ground based measurements of UV-B across the 11 year solar cycle 98 (Rampelotto et al., 2009). Rozema et al. (2002) hypothesized that this relationship will be 99 consistent across longer-term solar variations, with higher levels of surface UV-B flux during 100 solar activity lows such as the Maunder Minimum, even though overall UV and TSI are 101 reduced. Empirical evidence to test this hypothesis is currently lacking, however. Surface 102 UV-B proxy reconstructions based on the abundance of photoprotective pigments in fossil 103

cladocera (water flea) carapaces in arctic and subarctic lakes (Nevalainen et al., 2015, 2016, 104 105 2018) demonstrated a positive correlation between surface UV-B and solar activity over the last millennium, which is the opposite pattern to that predicted by Rozema et al. (2002). 106 107 However, UV-B proxies based on aquatic organisms such as cladocerans are influenced by water transparency as well as ambient UV-B, and so relate at least in part to local climatic and 108 vegetation conditions and anthropogenic land use changes (Nevalainen et al., 2015, 2018). 109 110 Although these impacts should have been limited in the arctic lake records used by Nevalainen et al. (2016), where water and UV transparency are high, it is still not clear what 111 the long-term relationship between solar activity and surface UV-B is, and therefore what 112 113 biotic and environmental impacts can be expected from solar variability in the future.

114

To address these uncertainties, we take advantage of a novel proxy for surface UV-B 115 irradiance based on the chemistry of pollen grains (Fraser et al., 2014; Rozema et al., 2001; 116 Seddon et al., 2019). Plants produce UV absorbing compounds (UACs) to protect their cells 117 from the harmful effects of UV-B, and up-regulate production in response to increased UV-B 118 doses (Fraser et al., 2014; Gao et al., 2004; Lomax et al., 2008; Rozema et al., 2001; Singh et 119 al., 2014). Pollen grains and spores preserve well in the fossil record because their outer wall, 120 121 or exine, is made of sporopollenin, a highly resistant biopolymer (Mackenzie et al., 2015). Critically, the UAC signal within the exine is also preserved (Jardine et al., 2016), and 122 remains stable over geological time (Fraser et al., 2012). Therefore, by measuring the 123 124 concentration of UACs in fossil and sub-fossil pollen grains, UV-B flux in the past can be reconstructed (Blokker et al., 2005, 2006; Fraser et al., 2014; Jardine et al., 2016; Lomax et 125 126 al., 2008; Rozema et al. 2001; Seddon et al., 2019; Willis et al., 2011).

127

128 The UV-B response mechanism is thought to be an ancient evolutionary adaptation to 129 terrestrial environments and occurs across the land plant phylogeny (Christie et al., 2012;

Jardine et al., 2016; Rizzini et al., 2011; Rozema et al., 1997), which means that a wide array 130 131 of taxa are available for sampling from the pollen and spore record. To date, a positive correlation between UV-B and sporopollenin UAC levels has been demonstrated for 132 Lycopodium (clubmoss) (Fraser et al., 2011; Jardine et al., 2016, 2017; Lomax et al., 2008, 133 134 2012; Watson et al., 2007), *Pinus* (pine) (Willis et al., 2011), *Cedrus atlantica* (Atlas cedar) (Bell et al., 2018), Vicia faba (broad or fava bean) (Rozema et al., 2001), and Poaceae 135 136 (grasses) (Jardine et al., 2016), confirming the broad phylogenetic applicability of the UAC proxy. Furthermore, because this proxy is based on terrestrial plants, it is less biased by 137 changes in the surrounding environment than those derived from aquatic organisms. UAC 138 139 concentrations in pollen and spores are determined by the UV dose experienced by the parent 140 plant, and are thought to represent the clear skies maximum UV level across the growing or pollen/spore production period (i.e. a timescale of several weeks prior to pollen/spore release) 141 (Jardine et al., 2016; Lomax et al., 2012). The impact of short-term variations in cloudiness 142 on UAC levels should therefore be limited, and inter-annual comparisons of UV-B flux can 143 144 be achieved. Since the UAC proxy detects surface UV-B flux it is sensitive to changes in ozone column thickness (Lomax et al., 2008), which means that variations in surface-level 145 146 UV-B caused by changes in ozone through time can be recovered.

147

Here, we use a maar lake sedimentary record from Nar Gölü in central Turkey, and 148 analyse UAC concentrations in Pinus pollen to reconstruct surface UV-B flux over the last 149 150 650 years. This record is then used to test for the correlation between surface UV-B and modelled solar UV-B irradiance, following the assumption that TSI and solar UV-B 151 152 irradiance will vary in phase though time. A negative correlation would support the hypothesis of Rozema et al. (2002), with solar activity highs leading to ozone production and 153 decreased UV-B flux to the surface. A positive correlation would support the cladoceran UV-154 B reconstructions of Nevalainen et al., (2015, 2016, 2018), and would suggest that the 155

Page 6

relationship between solar activity and ozone thickness observed on shorter timescales (e.g.
Rampelotto et al., 2009) cannot simply be scaled up across centuries and millennia.

158

159 Materials and Methods

Nar Gölü (38°20'24"N, 34°27'23"E; 1363 m a.s.l.) is a maar lake in central Turkey, ~0.7 km² in area and ~26 m deep, with a sediment record extending through the Holocene and into the last glacial (Dean et al., 2015). The upper 2500 years of the sedimentary sequence is continuously annually laminated (varved), which has allowed for a precise chronology to be developed (Dean et al., 2015; Jones et al., 2005). The Nar Gölü sediment record has been the focus of previous sedimentological, mineralogical, palynological and geochemical research (Dean et al., 2013, 2015; England et al., 2008; Jones et al., 2005, 2006).

167

The sediment core used in this study was collected in 2001, and the pollen samples 168 were initially documented in England et al. (2008). The age model for this core was based on 169 170 varve counting, which was carried out independently by two workers, who recounted until agreement to within 3 laminae for each 6 cm section of core was reached (Jones et al., 2005). 171 Replication of varve counts from additional cores has provided a maximum age uncertainty of 172 173 2.5% of the given age (Jones et al., 2006). The pollen samples were collected at \sim 20 year inter-sample resolution, with most samples representing 3 (sometimes 4 or 5) years of 174 sediment accumulation. The whole sequence covers 640 years and ~160 cm of sediment core. 175 176 The samples were processed according to standard palynological protocols, using 10% HCl, 10% NaOH, 60% HF, and acetolysis (England et al., 2008). We used the same pollen 177 preparations in this study to maintain stratigraphic consistency with the pollen count data, and 178 because standard processing protocols, including acetolysis (oxidation), do not impact upon 179 the recoverability of variations in UAC concentrations across samples (Jardine et al., 2015, 180

2016, 2017). Furthermore, Bell (2018) showed that UAC levels were similar in acetolysed
and untreated *Pinus* pollen.

183

We selected *Pinus* as the target taxon because it is abundant through most of the upper 184 part of the Nar Gölü record, with relative abundances of 13% to 45% of the pollen sum and 185 influx rates of 300 to 4400 cm²/year (England et al., 2008). Furthermore, compared to lower 186 187 stature vegetation the impact of localized shading on UV reconstructions (Fraser et al., 2011; Jardine et al., 2016) should be minimal. A positive association between UAC levels and UV-188 B has also previously been demonstrated for *Pinus* pollen, across both a modern latitudinal 189 190 gradient and over the last 9.5 kyr (Willis et al., 2011), suggesting that a measurable signal is recoverable from the Nar Gölü Pinus pollen record. Pinus pollen within the Nar Gölü 191 sediments represents mostly regional rather than local vegetation, and is mostly derived from 192 193 the Taurus Mountains >70 km south and southeast of Nar Gölü, although Pinus was also planted near the lake in the 1980s (England et al., 2008). In the modern day, the three main 194 195 Pinus species in this region are Pinus brutia, Pinus nigra and Pinus sylvestris (Woldring and Bottema, 2003), and these are anticipated to have been the major contributors to the Nar Gölü 196 197 Pinus pollen record during the study interval.

198

We used Fourier Transform Infrared (FTIR) microspectroscopy to generate the 199 chemical data, because previous analyses (Bell et al., 2018; Fraser et al., 2011; Jardine et al., 200 2016, 2017; Lomax et al., 2008, 2012; Watson et al., 2007) have shown that this can 201 successfully capture variations in UAC abundances at small sample sizes. To prepare the 202 samples for FTIR analysis, individual *Pinus* pollen grains were picked out from the processed 203 sediment samples and mounted on ZnSe windows. To pick the pollen grains we used an 204 inverted microscope with a micromanipulator attachment, the full set-up comprising 205 Narishige MMN-1 and MMO-202ND course and fine control micromanipulators, an IM-11-2 206

pneumatic microinjector, with a Microtec IM-2 inverted microscope. The picked pollen grains 207 208 were arranged in groups of 4 to 5 grains on the ZnSe windows, with 5 replicate groups per sample. This means that each FTIR spectrum represents 4 or 5 pollen grains, and each pollen 209 sample is represented by 5 replicate FTIR spectra. We generated the data using a Thermo 210 Scientific (Waltham, MA, USA) Nicolet Nexus FTIR bench unit connected to a Continuum 211 IR microscope fitted with an MCT-A liquid nitrogen-cooled detector and a Reflachromat 15x 212 213 objective lens. FTIR spectra were generated in transmission mode using a microscope aperture of 100 x 100 μ m recording the mean of 256 scans with a resolution of 1.928 cm⁻¹ 214 wavenumbers. Five of the 33 samples in the study interval had insufficient Pinus pollen for 215 216 FTIR analysis, resulting in a dataset of 28 samples.

217

Peak height measurement and data analysis were carried out in R v3.4.0 (R Core 218 Team, 2017). The package 'baseline' v1.2-1 (Liland and Mevik, 2015) was used to baseline 219 correct the IR spectra, by subtracting a 2nd order polynomial baseline from each spectrum 220 (Figure 1). We quantified UAC concentrations by measuring the height of the 1510 cm⁻¹ 221 aromatic (C=C) peak (Fig. 1), because this peak records changes in the abundance of the 222 phenolic compounds para-coumaric acid and ferulic acid that act as UACs in sporopollenin 223 (Fraser et al., 2014; Watson et al., 2007). Absorbance values in IR spectra relate to the 224 thickness of material being analysed, so following previous research (Fraser et al., 2011: 225 Jardine et al., 2015, 2016, 2017; Lomax et al., 2008, 2012) the 1510 cm⁻¹ aromatic peak 226 height was normalized against the hydroxyl (OH) band centred on 3300 cm⁻¹ (Figure 1). 227 Although the aromatic/OH ratio has not yet been calibrated to UAC concentrations or UV 228 levels, it does provide a successful proxy whereby higher aromatic/OH ratio values equate to 229 higher UV-B flux (Fraser et al., 2011; Jardine et al., 2016, 2017; Lomax et al., 2008, 2012). 230 Short-term variations in ambient UV-B flux experienced by the pollen-producing plants will 231

add noise to the UAC reconstruction; possible sources of additional variability are considered

233	in the Discussion. The raw data (sample ages and peak height measurements) are available for
234	download from figshare (Jardine et al., 2019). [NB For review please used this private link
235	to access the data: <u>https://figshare.com/s/45c1f29f1d76c1cbc01d</u>].
236	
237	[Insert Figure 1]

238

We used the historical SSI reconstruction of Lean (2018) to obtain solar UV-B irradiance estimates for the last 650 years. This SSI reconstruction covers the period 850 to 2016, and provides an annually resolved time series that incorporates information from spacebased irradiance observations, sunspots and cosmogenic radionuclides (full details in Lean 2018). Within the ultraviolet the SSI estimates are resolved to 1 nm wavebands. We therefore integrated the irradiance values within the range 280 to 315 nm, to obtain an irradiance reconstruction integrated across the UV-B part of the solar spectrum (shown in Figure 2b).

246

We used Spearman's rank order correlation to test the association between UACs and 247 solar UV-B. For the UACs we used the mean of the five replicates within each pollen sample, 248 249 and for solar UV-B we used the mean UV-B irradiance values within the calendar years 250 represented by each pollen sample. Spearman's rank order correlation is appropriate because it is a non-parametric test that does not assume normality of distributions or a linear 251 relationship among variables. To examine the influence of shared long-term temporal trends 252 253 among the variables, we detrended the data by taking the residuals from linear regressions of each variable against time. The residuals were then used as variables in the correlation test. 254 255

256 **Results**

The Nar Gölü *Pinus* UAC record (Figure 2a) shows that surface level UV-B irradiance
has varied over the last 650 years. Visual comparison with the solar UV-B reconstruction of

259	Lean (2018) (Figure 2b) reveals many of the same features, including an initial high value at
260	\sim 1350, minima at $\sim\!\!1460$ to 1550 (the Spörer Minimum), 1645 to 1710 (the Maunder
261	Minimum), ~1790 to 1820 (the Dalton Minimum), and 1880 to 1920, and the rise from the
262	Maunder Minimum to the late 20 th Century. The Spearman's rank order correlation between
263	the UAC data and solar UV-B reconstruction demonstrates statistically significant positive
264	relationships for both raw ($n = 28$, $r_s = 0.52$, $p = 0.005$) and detrended datasets ($n = 28$, $r_s =$
265	0.55, $p = 0.004$) (Figure 3). These correlations show that the visual similarities between the
266	surface and solar UV-B data are robust.
267	
268	[Insert Figure 2]
269	
270	[Insert Figure 3]
271	
272	Discussion
273	Our results show a positive correlation between the Pinus UAC data and the solar UV-
274	B reconstruction of Lean (2018), demonstrating that solar activity and surface UV-B trends
275	have been concordant over the last 650 years. These results are in agreement with the
276	cladoceran-based surface UV-B reconstructions of Nevalainen et al. (2015, 2016, 2018), but
277	are not consistent with the hypothesis of Rozema et al. (2002) that surface level UV-B should
278	be anti-correlated with solar activity across grand solar minima and maxima. Our results
279	therefore suggest that any variations in the thickness of the ozone layer were not sufficient to
280	alter the incoming UV-B flux.
281	
282	These results also demonstrate that the anti-correlation between solar activity and
283	ground-based measurements of UV across the 11 year solar cycle (Rampelotto et al., 2009)
284	cannot simply be scaled up to longer timescales (Rozema et al., 2002). Whether this implies a

different relationship between solar activity, UV and ozone thickness across the 11 year solar 285 cycle and longer-term cycles and trends is currently unclear. While solar UV irradiance at 286 wavelengths under 242 nm leads to ozone production, longer wavelength UV destroys it (Ball 287 et al., 2016), therefore given the right balance of change across the UV spectrum decreases in 288 ozone creation across solar minima could be cancelled out by decreases in ozone destruction 289 (and vice versa during solar maxima). Ozone concentrations are also modulated by hydrogen, 290 nitrogen, and chlorine catalytic cycles (Lary, 1997), and long-term variations in the 291 atmospheric concentrations of HOx, NOx and ClOx radicals may influence how ozone 292 thickness changes in response to solar UV. This question deserves further research, both with 293 294 more instrumental measurements of spectral UV irradiance and surface level UV-B, and 295 additional high-resolution UAC-based records over longer timescales.

296

While the relationships between our UAC data and solar UV-B are statistically 297 significant (Figure 3), the strength of the correlations are moderate ($r_s = 0.52$ for the raw data, 298 and $r_s = 0.55$ for the detrended data). In the Nar Gölü record we have identified three main 299 factors that, in addition to solar UV-B, may have been responsible for variation in the UAC 300 signal. First, the *Pinus* pollen does not represent local vegetation, but is thought to be largely 301 302 sourced from the Taurus Mountains (England et al., 2008). The pollen signal therefore likely represents a mix of altitudes and incoming UV-B levels (Lomax et al., 2012), which will 303 contribute to within-sample variance (Seddon et al., 2019). To the extent that the UV-B 304 305 response differs among the *Pinus* species contributing pollen to the Nar Gölü record, any variations in their relative abundances over time may have added further noise to the UAC 306 307 signal. The recently planted *Pinus* trees near Nar Gölü (England et al., 2008) may also have impacted upon the UAC measurement from the most recent sample in the record, which 308 covers the years 1998 to 2001. 309

310

Second, variations in cloud cover modify surface UV irradiance levels (Calbó et al.,
2005; Fraser et al., 2011). Although the UAC proxy is unlikely to be affected by short-term
variations in cloudiness (Lomax et al., 2012), interannual variability in cloud cover may add
additional noise to UAC time series where each sample represents several years of pollen
release.

316

Third, any errors in the sediment core chronology will add noise to the correlations. 317 Although dating precision in the Nar Gölü record is thought to be better than the maximum 318 age uncertainty of 2.5% would suggest (Dean et al., 2013), small errors are likely to be 319 320 inevitable even in continuously varved sediments, and the magnitude of change in solar UV-B irradiance across the 11 year solar cycle (Figure 2b; Lean, 2018) suggests that age model 321 deviations could add substantial noise to the UAC-UV relationship. While dating errors of a 322 few years could artificially impose a positive correlation between the solar UV-B 323 reconstruction and UAC data, the sustained, multi-decadal solar UV-B lows during both the 324 325 Spörer Minimum and Maunder Minimum (Figure 2b) coincide with intervals of low UAC concentrations (Figure 2a). This suggests that the positive correlation demonstrated here is not 326 an artefact of minor errors in the varve chronology, but rather represents a genuine signal. 327 328

Despite these sources of variability, the UAC proxy has successfully recovered the 329 major solar UV-B signal, demonstrating that it has much to offer as a means of examining 330 331 solar inputs to the Earth system and their contributions to biotic and climatic change. As a surface UV-B proxy UAC measurements allow us to test hypotheses relating to changes in 332 333 ozone thickness through time (Lomax et al., 2008), and can aid in deconvoluting the effects of solar activity and ozone-related variations on UV-B at the Earth's surface. The pollen UAC 334 proxy could also be used in conjunction with UV proxies based on aquatic organisms 335 (Nevalainen et al., 2016) to separate out solar UV from water transparency effects 336

(Nevalainen et al., 2015, 2018), and therefore constrain key factors impacting on aquatic
ecosystems. More generally, the UAC proxy can be used to test for the impacts of surface
UV-B changes on biotic systems, in relation to solar UV irradiance variations, longer term
cycles in the Earth's orbit around the Sun (Jardine et al., 2016), and periods of ozone layer
disruption (Lomax et al., 2008; Visscher et al., 2004).

342

Our Pinus UAC data, taken together with the cladocera data of Nevalainen et al. 343 (2015, 2016, 2018), demonstrate that variations in solar UV irradiance are sufficient to drive 344 biochemical responses in disparate environments and groups of organisms. Future changes in 345 346 surface UV-B flux, whether driven by variations in solar activity or atmospheric composition, 347 will alter not just organismal stress but also the metabolic costs of producing UV protective compounds, and may influence interspecific competitive relationships and ecosystem-level 348 processes such as decomposition and carbon cycling (Rozema et al., 1997). The relative 349 importance of these responses therefore needs to be assessed in future studies of ecological 350 change. 351

352

353 Conclusions

354 We have provided the first detailed proxy reconstruction of surface level UV-B flux on centennial timescales. By linking this with a published solar UV-B reconstruction, we have 355 shown that solar UV-B flux at the surface follows a similar long-term trend to top of 356 357 atmosphere UV-B. On the timescales considered here, incoming solar UV flux will therefore be the dominant control on surface-level UV, rather than UV modulated ozone thickness. In 358 359 addition to better understanding and modelling sources of variance in UAC reconstructions, future research in this area needs to focus on calibrating the UAC UV-B proxy to a specific 360 dose-response relationship, to quantify the magnitude of change across different timescales 361 (Rozema et al., 2001, 2002, 2009; Seddon et al., 2019). Developing an action spectrum 362

363	(Herman, 2010; Rozema et al., 2001; Seddon et al., 2019) for UAC production will also be
364	important for quantifying how UAC concentrations vary with changes in ozone levels.
365	Together, these measures will help towards understanding how variations in solar UV
366	irradiance impact on Earth's climate and biota through time, both in the past and in the future.
367	
368	Acknowledgements
369	The raw data for this study are available on figshare:
370	https://dx.doi.org/10.6084/m9.figshare.8075519 [NB For review please used this private
371	link to access the data: https://figshare.com/s/45c1f29f1d76c1cbc01d]. We thank NERC
372	(grant NE/K005294/1) for funding this research, and a previous NERC award
373	(NER/S/A/2002/10316) to University of Birmingham-based PhD student, Ann England. Andy
374	Moss is thanked for lab support. We also thank Benjamin Bell and Liisa Nevalainen for
375	insightful reviews, and the anonymous reviewers whose comments improved earlier versions
376	of this paper.
377	
378	References
379	Ball WT, Haigh JD, Rozanov EV, et al. (2016) High solar cycle spectral variations
380	inconsistent with stratospheric ozone observations. Nature Geoscience 9: 206-209.
381	Bell, BA. (2018) Advancing the Application of Analytical Techniques in the Biological
382	Chemistry of Sporopollenin: Towards Novel Plant Physiological Tracers in
383	Quaternary Palynology. PhD Thesis, University of Manchester, UK.
384	Bell BA, Fletcher WJ, Ryan P, et al. (2018) UV-B-absorbing compounds in modern Cedrus
385	atlantica pollen: The potential for a summer UV-B proxy for Northwest Africa. The
386	Holocene 28: 1382-1394.

387	Blokker, P, Boelen, P, Broekman, R and Rozema, J. (2006) The occurrence of <i>p</i> -coumaric
388	acid and ferulic acid in fossil plant materials and their use as UV-proxy. Plant Ecology
389	182: 197-207

Blokker, P, Yeloff, D, Boelen, P, Broekman, R and Rozema, J. (2005). Development of a

- 391 proxy for past surface UV-B irradiation: A thermally assisted hydrolysis and
- methylation py-GC/MS method for the analysis of pollen and spores. *Analytical Chemistry* 77: 6026-6031.
- Calbó J, Pagès D and González J-A. (2005) Empirical studies of cloud effects on UV
 radiation: A review. *Reviews of Geophysics* 43.
- Caldwell MM, Björn LO, Bornman JF, et al. (1998) Effects of increased solar ultraviolet
 radiation on terrestrial ecosystems. *Journal of Photochemistry and Photobiology B:*
- *Biology* 46: 40-52.
- Christie JM, Arvai AS, Baxter KJ, et al. (2012) Plant UVR8 photoreceptor senses UV-B by
 tryptophan-mediated disruption of cross-dimer salt bridges. *Science* 335: 1492-1496.
- 401 Dean JR, Jones MD, Leng MJ, et al. (2015) Eastern Mediterranean hydroclimate over the late

402 glacial and Holocene, reconstructed from the sediments of Nar lake, central Turkey,

- 403 using stable isotopes and carbonate mineralogy. *Quaternary Science Reviews* 124:
- 404 162-174.
- Dean JR, Jones MD, Leng MJ, et al. (2013) Palaeo-seasonality of the last two millennia
 reconstructed from the oxygen isotope composition of carbonates and diatom silica
 from Nar Gölü, central Turkey. *Quaternary Science Reviews* 66: 35-44.
- 408 England A, Eastwood WJ, Roberts CN, et al. (2008) Historical landscape change in
- 409 Cappadocia (central Turkey): a palaeoecological investigation of annually laminated
 410 sediments from Nar lake. *The Holocene* 18: 1229-1245.

411	Ermolli I, Matthes K, Dudok de Wit T, et al. (2013) Recent variability of the solar spectral
412	irradiance and its impact on climate modelling. Atmospheric Chemistry and Physics
413	13: 3945-3977.
414	Fraser WT, Lomax BH, Jardine PE, et al. (2014) Pollen and spores as a passive monitor of
415	ultraviolet radiation. Frontiers in Ecology and Evolution 2.
416	Fraser WT, Scott AC, Forbes AES, et al. (2012) Evolutionary stasis of sporopollenin
417	biochemistry revealed by unaltered Pennsylvanian spores. New Phytologist 196: 397-
418	401.
419	Fraser WT, Sephton MA, Watson JS, et al. (2011) UV-B absorbing pigments in spores:
420	biochemical responses to shade in a high-latitude birch forest and implications for
421	sporopollenin-based proxies of past environmental change. Polar Research 30: 8312.
422	Gao W, Zheng Y, Slusser JR, et al. (2004) Effects of Supplementary Ultraviolet-B lrradiance
423	on Maize Yield and Qualities: A Field Experiment. Photochemistry and Photobiology
424	80: 127-131.
425	Gehrke C, Johanson U, Callaghan TV, et al. (1995) The impact of enhanced ultraviolet-B
426	radiation on litter quality and decomposition processes in Vaccinium leaves from the
427	Subarctic. Oikos 72: 213-222.
428	Gray LJ, Beer J, Geller M, et al. (2010) Solar Influences on Climate. Reviews of Geophysics
429	48.
430	Haberreiter M, Schöll M, Dudok de Wit T, et al. (2017) A new observational solar irradiance
431	composite. Journal of Geophysical Research: Space Physics 122: 5910-5930.
432	Herman JR. (2010) Use of an improved radiation amplification factor to estimate the effect of
433	total ozone changes on action spectrum weighted irradiances and an instrument
434	response function. Journal of Geophysical Research 115.
435	Ineson S, Maycock AC, Gray LJ, et al. (2015) Regional climate impacts of a possible future
436	grand solar minimum. Nature Communications 6: 7535.

Page 17

437	Jardine PE, Abernethy FAJ, Lomax BH, et al. (2017) Shedding light on sporopollenin
438	chemistry, with reference to UV reconstructions. Review of Palaeobotany and
439	Palynology 238: 1-6.
440	Jardine PE, Fraser, WT, Gosling, WD, et al. (2019) Data from "Reconstruction of ultraviolet-
441	B irradiance at the Earth's surface, and its relationship with solar activity and ozone
442	thickness". figshare, https://dx.doi.org/10.6084/m9.figshare.8075519.
443	Jardine PE, Fraser WT, Lomax BH, et al. (2015) The impact of oxidation on spore and pollen
444	chemistry. Journal of Micropalaeontology 34: 139-149.
445	Jardine PE, Fraser WT, Lomax BH, et al. (2016) Pollen and spores as biological recorders of
446	past ultraviolet irradiance. Scientific Reports 6: 1-8.
447	Jones MD, Leng MJ, Roberts CN, et al. (2005) A Coupled Calibration and Modelling
448	Approach to the Understanding of Dry-Land Lake Oxygen Isotope Records. Journal
449	of Paleolimnology 34: 391-411.
450	Jones MD, Roberts CN, Leng MJ, et al. (2006) A high-resolution late Holocene lake isotope
451	record from Turkey and links to North Atlantic and monsoon climate. Geology 34:
452	361.
453	Krivova NA, Vieira LEA and Solanki SK. (2010) Reconstruction of solar spectral irradiance
454	since the Maunder minimum. Journal of Geophysical Research 115.
455	Lary DJ. (1997) Catalytic destruction of stratospheric ozone. Journal of Geophysical
456	Research: Atmospheres 102: 21515-21526.
457	Lean JL. (2018) Estimating Solar Irradiance Since 850 CE. Earth and Space Science 5: 133-
458	149.
459	Liland KH and Mevik B-H. (2015) baseline: Baseline Correction of Spectra. 1.2-1 ed.
460	Lomax BH, Fraser WT, Harrington G, et al. (2012) A novel palaeoaltimetry proxy based on
461	spore and pollen wall chemistry. Earth and Planetary Science Letters 353-354: 22-28.

462	Lomax BH, Fraser WT, Sephton MA, et al. (2008) Plant spore walls as a record of long-term
463	changes in ultraviolet-B radiation. Nature Geoscience 1: 592-596.
464	Mackenzie G, Boa AN, Diego-Taboada A, et al. (2015) Sporopollenin, The Least Known Yet
465	Toughest Natural Biopolymer. Frontiers in Materials 2: 1-5.
466	Meijkamp B, Aerts R, van de Staaij J, et al. (1999) Effects of UV-B on secondary metabolites
467	in plants. In: Rozema J (ed) Stratospheric ozone depletion: The effects of enhanced
468	UV-B on terrestrial ecosystems. Leiden: Backhuys, 71-99.
469	Nevalainen L, Luoto TP, Rantala MV, et al. (2015) Role of terrestrial carbon in aquatic UV
470	exposure and photoprotective pigmentation of meiofauna in subarctic lakes.
471	Freshwater Biology 60: 2435-2444.
472	Nevalainen L, Rantala MV, Luoto TP, et al. (2016) Long-term changes in pigmentation of
473	arctic Daphnia provide potential for reconstructing aquatic UV exposure. Quaternary
474	Science Reviews 144: 44-50.
475	Nevalainen L, Rantala MV, Rautio M, et al. (2018) Spatio-temporal cladoceran
476	(Branchiopoda) responses to climate change and solar radiation in subarctic ecotonal
477	lakes. Journal of Biogeography 45: 1954-1965.
478	Rampelotto PH, da Rosa MB and Schuch NJ. (2009) Solar cycle and UV - B comparison for
479	South America–South of Brazil (29° S, 53° W). In: Nakajima T and Yamasoe MA
480	(eds) AIP Conference Proceedings. AIP, 490-493.
481	R Core Team (2017) R: A language and environment for statistical computing. 3.4.2 ed.
482	Vienna, Austria: R Foundation for Statistical Computing.
483	Rizzini L, Favory JJ, Cloix C, et al. (2011) Perception of UV-B by the Arabidopsis UVR8
484	Protein. Science 332: 103-106.
485	Rozema J, Blokker P, Mayoral Fuertes MA, et al. (2009) UV-B absorbing compounds in
486	present-day and fossil pollen, spores, cuticles, seed coats and wood: evaluation of a

487 proxy for solar UV radiation. *Photochemical and Photobiological Sciences* 8: 1233-

488 1243.

- Rozema J, Broekman RA, Blokker P, et al. (2001) UV-B absorbance and UV-B absorbing
 compounds (*para*-coumaric acid) in pollen and sporopollenin: the perspective to track
 historic UV-B levels. *Journal of Photochemistry and Photobiology B: Biology* 62:
- 492 108-117.
- 493 Rozema J, van de Staij J, Björn L-O, et al. (1997) UV-B as an environmental factor in plant
 494 life: stress and regulation. *Trends in Ecology and Evolution* 12: 22-28.
- 495 Rozema J, van Geel B, Björn L-O, et al. (2002) Toward Solving the UV Puzzle. *Science* 296:
 496 1621-1622.
- 497 Seddon AWR, Festi D, Robson TM, et al. (2019) Fossil pollen and spores as a tool for
- reconstructing ancient solar-ultraviolet irradiance received by plants: an assessment of
 prospects and challenges using proxy-system modelling. *Photochemical and Photobiological Sciences* 18: 275-294.
- 501 Singh SK, Reddy KR, Reddy VR, et al. (2014) Maize growth and developmental responses to

502 temperature and ultraviolet-B radiation interaction. *Photosynthetica* 52: 262-271.

Solanki SK, Krivova NA and Haigh JD. (2013) Solar Irradiance Variability and Climate.
 Annual Review of Astronomy and Astrophysics 51: 311-351.

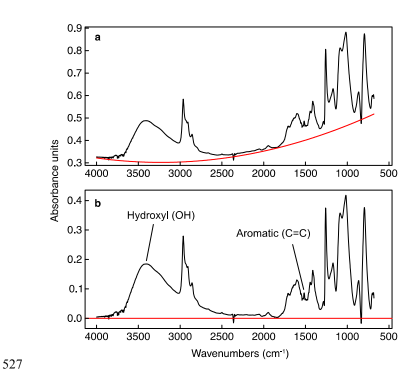
Steinhilber F, Abreu JA, Beer J, et al. (2012) 9,400 years of cosmic radiation and solar
activity from ice cores and tree rings. *Proceedings of the National Academy of*

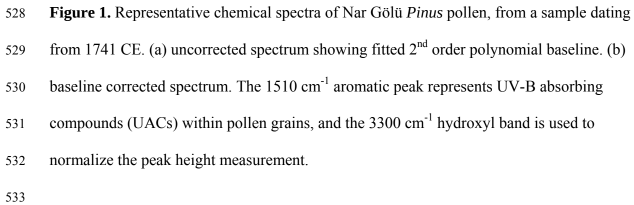
- *Sciences* 109: 5967-5971.
- Svalgaard L and Schatten KH. (2016) Reconstruction of the Sunspot Group Number: The
 Backbone Method. *Solar Physics* 291: 2653-2684.
- 510 Usoskin IG. (2017) A history of solar activity over millennia. *Living Reviews in Solar Physics*
- 511 14.

512	Visscher H, Looy CV, Collinson ME, et al. (2004) Environmental mutagenesis during the
513	end-Permian ecological crisis. Proceedings of the National Academy of Sciences of the
514	United States of America 101: 12952-12956.
515	Watson JS, Septhon MA, Sephton SV, et al. (2007) Rapid determination of spore chemistry
516	using thermochemolysis gas chromatography-mass spectrometry and micro-Fourier
517	transform infrared spectroscopy. Photochemical and Photobiological Sciences 6: 689-
518	694.
519	Willis KJ, Feurdean A, Birks HJB, et al. (2011) Quantification of UV-B flux through time
520	using UV-B-absorbing compounds contained in fossil Pinus sporopollenin. New
521	Phytologist 192: 553-560.
522	Woldring, H and Bottema, S. (2003) The vegetation history of east-central Anatolia in
523	relation to Archaeology: the Eski Acégöl pollen evidence compared with the Near
524	Eastern environment. Palaeohistoria 43/44: 1-31.

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Figures 526





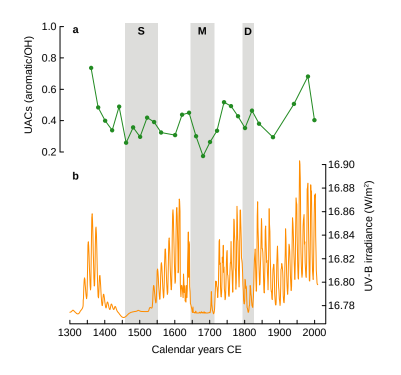
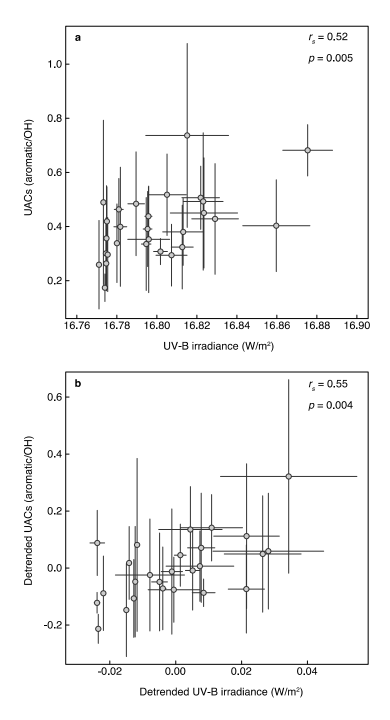




Figure 2. Surface and solar UV-B records. (a) Pinus UAC data from Nar Gölü, shown as the
mean of five replicates (solid green line with points showing samples) ±1 standard deviation
(shaded area). (b) Modeled solar UV-B irradiance, from Lean (2018). Grey shaded regions
show solar minima, D = Dalton Minimum, M = Maunder Minimum, S = Spörer Minimum.



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Figure 3. *Pinus* UAC data plotted against modeled solar UV-B irradiance (Lean 2018), for both raw (a) and detrended (b) data. For the UAC data, the points show the mean of five replicates, and the error bars are 1 standard deviation. For the UV-B irradiance reconstruction, the points are the mean values within the calendar years represented by each pollen sample, and the error bars are 1 standard deviation. r_s = Spearman's rank order correlation coefficient, p = p value of correlation.

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