

1 Prospects for dendroanatomy in paleoclimatology – a case study on *Picea* 2 *engelmannii* from the Canadian Rockies

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15

16 **Abstract**

17 The continuous development of new proxies as well as a refinement of existing tools are key
18 to advances in paleoclimate research and improvements in the accuracy of existing climate
19 reconstructions. Herein, we build on recent methodological progress in dendroanatomy – the
20 analyses of wood anatomical parameters in dated tree rings – and introduce the longest (1585
21 – 2014 CE) dendroanatomical dataset currently developed for North America. We explore the
22 potential of dendroanatomy of high-elevation Engelmann spruce (*Picea engelmannii*) as a
23 proxy of past temperatures by measuring anatomical cell dimensions of 15 living trees from
24 the Columbia Icefield area. X-ray maximum latewood density (MXD) and its blue intensity
25 counterpart (MXBI) have previously been measured, allowing comparison between the
26 different parameters. Our findings highlight anatomical MXD and maximum radial cell wall
27 thickness as the two most promising wood anatomical proxy parameters for past
28 temperatures, each explaining 46% and 49%, respectively, of detrended instrumental July-
29 August maximum temperatures over the 1901–1994 period. While both parameters display
30 comparable climatic imprinting at higher frequencies to X-ray derived MXD, the anatomical
31 dataset distinguishes itself from its predecessors by providing the most temporally stable
32 warm-season temperature signal. Further studies, including samples from more diverse age
33 cohorts and the adaptation of RCS-based standardization, are needed to disentangle the
34 ontogenetic and climatic components of long-term signals stored in the wood anatomical traits
35 and to more comprehensively evaluate the potential contribution of this new dataset to
36 paleoclimate research.

37 **Keywords:** Dendroanatomy, *Picea engelmannii*, Canadian Rockies, tree rings, latewood
38 density, temperature reconstruction, paleoclimatology

39

40 **1. Introduction**

41 Tree rings form the backbone of high-resolution palaeoclimatology of the Common Era by
42 providing precisely dated, annually resolved, spatially widespread and easily accessible
43 archives of climate proxy data. Tree-ring archives make up more than half of all publicly
44 available temperature proxy records and are greatly influential in multi-proxy hemispheric-
45 scale temperature reconstructions (PAGES 2k Consortium 2017). They are vital for spatially
46 explicit mapping of the Medieval Climate Anomaly, the Little Ice Age, and other important
47 climate periods (e.g., PAGES 2k Consortium 2013), and the study of temporally distinct
48 cooling events caused by volcanic eruptions (e.g., Schneider et al. 2015; Stoffel et al. 2015;
49 Wilson et al. 2016). Moreover, tree-ring based climate reconstructions play a key role in many
50 of the emerging proxy-model comparison efforts (e.g., Goosse 2017; Luterbacher et al. 2016;
51 Pages k-PMIP3 group 2015; Phipps et al. 2013; Seftigen et al. 2017).

52

53 The most frequently and successfully used tree-ring parameters for the study of past
54 temperature variations at high latitudes and altitudes are ring width and maximum latewood
55 density or simply maximum density (MXD) (e.g., Esper et al. 2018). While ring width is the
56 most easily acquired proxy of year-to-year variations in climate, the parameter often proves
57 difficult to interpret as it may represent distorted transformations of the underlying climate (e.g.,
58 Frank et al. 2010; Lücke et al. 2019). In particular, ring width may exhibit amplified low-
59 frequency signals (von Storch et al. 2004) resulting from lagged growth processes in response
60 to climate (Esper et al. 2015) or non-climatic processes (Rydval et al. 2015). Consequently,
61 the presence of prominent decadal variability should not be taken as evidence of
62 corresponding variability distribution in climate observations, and an overestimation of low-
63 frequency signals is often observed (e.g., Franke et al. 2013; Seftigen et al. 2017; Wilson et
64 al. 2016). The MXD parameter, in contrast, generally contains a stronger climate signal with
65 higher signal-to-noise ratios (e.g., Briffa et al. 2002; Ljungqvist et al. 2020), as well as less
66 biological persistence (Esper et al. 2015) and age-related signal-muting (Konter et al. 2016),
67 and is less influenced by stand disturbances (Rydval et al. 2018). However, a number of recent
68 studies (Björklund et al. 2020; Björklund et al. 2019; Edwards et al. 2022) have proposed the
69 accuracy of the MXD parameter to be sensitive to measurement resolution. Björklund et al.
70 (2019) showed that increasingly lower resolution of MXD data could result in an increased
71 artificial similarity to the climate response of ring width, and thus that several of the issues
72 facing ring width as a climate proxy may also represent non-negligible constraints on the MXD
73 parameter.

74

75 To reduce uncertainties, future reconstruction efforts could profit from the development of new
76 proxy types and parameters for paleoclimatology, as well as new and expanding

77 methodologies. Recently, dendroanatomy – the analysis of wood anatomical traits in dated
78 tree rings (Fonti et al. 2010; Pacheco et al. 2018) – has become more accessible through
79 semi-automated approaches to quantify wood cell anatomy (Prendin et al. 2017; von Arx and
80 Carrer 2014; von Arx et al. 2016). Analysis of anatomical cell dimensions is now possible at
81 the scale required for high-quality climate reconstruction over centuries to millennia (Björklund
82 et al. 2020). Unlike ring width, anatomical traits of temperature-limited conifers appear to be
83 less affected by biological memory effects and are imprinted with strong and mechanistically-
84 grounded temperature signals (Björklund et al. 2019; Cuny et al. 2019; Cuny et al. 2014).
85 Moreover, cell anatomical measurements have unprecedentedly high temporal resolution
86 relying on the base unit of the xylem – the tracheid cell, and their biological foundations and
87 functional links are comparably well understood (e.g., Bouche et al. 2014; Pittermann et al.
88 2011; Wilkinson et al. 2015).

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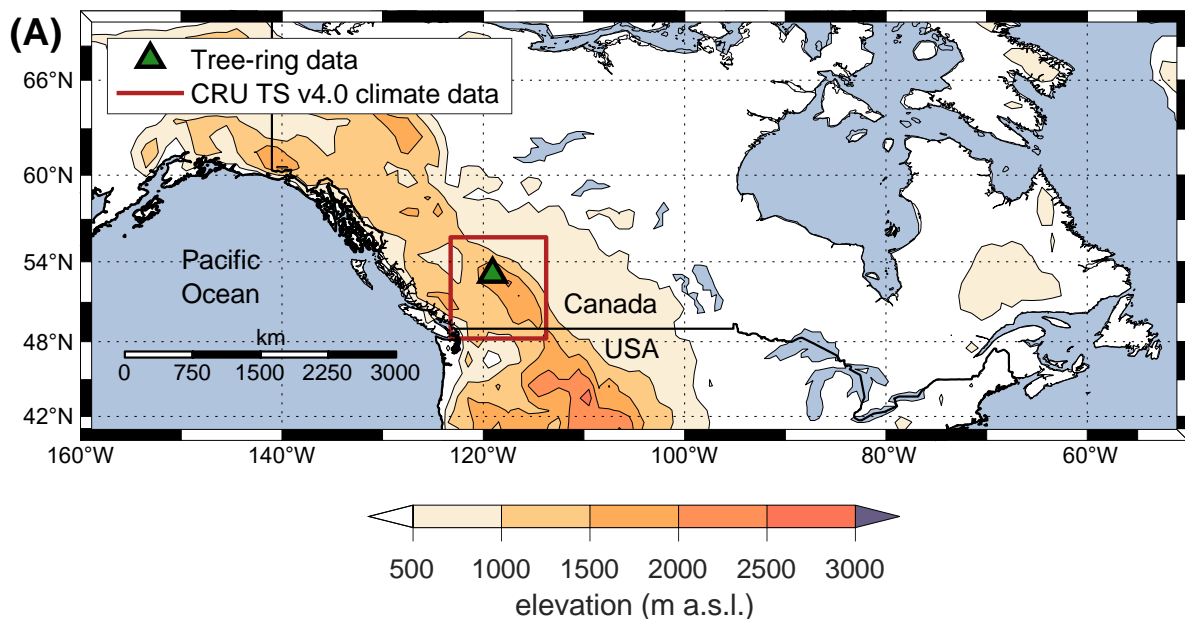
90 In this article, we aim to explore the value of dendroanatomy for high-elevation living
91 Engelmann spruce (*Picea engelmannii*) trees as a proxy of past temperatures. We make use
92 of tree samples from the Columbia Icefield area of the Canadian Rockies (Fig. 1) – a site
93 known for hosting the longest (950–1994 CE) available temperature-sensitive tree-ring
94 densitometric collections for boreal North America (Luckman et al. 1997; Luckman and Wilson
95 2005). The Icefield collection, originally comprising ring width and MXD measurements, have
96 previously been used in regional (Luckman 1997; Luckman 2000; St. George and Luckman
97 2001) and hemispheric-scale (Briffa et al. 2002; D'Arrigo et al. 2006; Esper et al. 2002; Mann
98 et al. 1999) temperature reconstructions, including the recent large-scale Northern
99 Hemisphere summer temperature reconstruction syntheses (Anchukaitis et al. 2017;
100 Schneider et al. 2015; Wilson et al. 2016). The analysis of the new dendroanatomical dataset
101 produced here includes an assessment of its signal strength and the imprint of temperature
102 within a number of wood anatomical traits spanning the period 1585–2014 CE. We detail
103 common variance amongst selected anatomical parameters and emphasize the
104 reconstruction potential of this dataset. The availability of MXD from the Columbia Icefield area
105 (Luckman et al. 1997; Luckman and Wilson 2005), produced with the Walesch Electronic
106 Dendro2003 technique (Eschbach et al. 1995) and its predecessor (Schweingruber et al.
107 1978) (hereafter referred to as X-ray MXD), and latewood blue-intensity (referred to as MXBI)
108 (McCarroll et al. 2002) measurements allow for an optimal opportunity for testing the skill and
109 potential advantages of dendroanatomical parameters as climate proxies. This work is part of
110 a larger ongoing collaborative effort dedicated to developing a network of long (~500–1000
111 years) wood dendroanatomical chronologies across the northern hemisphere. The ultimate
112 ambition of this initiative is to sharpen signal interpretations of the dendrochronological records

113 and to optimize seasonal and temporal fidelity of the proxy-based reconstructions in order to
114 revise (or reinforce) previous conclusions about pre-industrial climate variability and the
115 mechanisms causing this variability. This work also represents a first step towards a
116 millennium long anatomical *P. engelmannii* dataset for the Columbia Icefield area, Canada.
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118 2. Data and methods

119 2.1 Sample preparation and dendroanatomical measurements

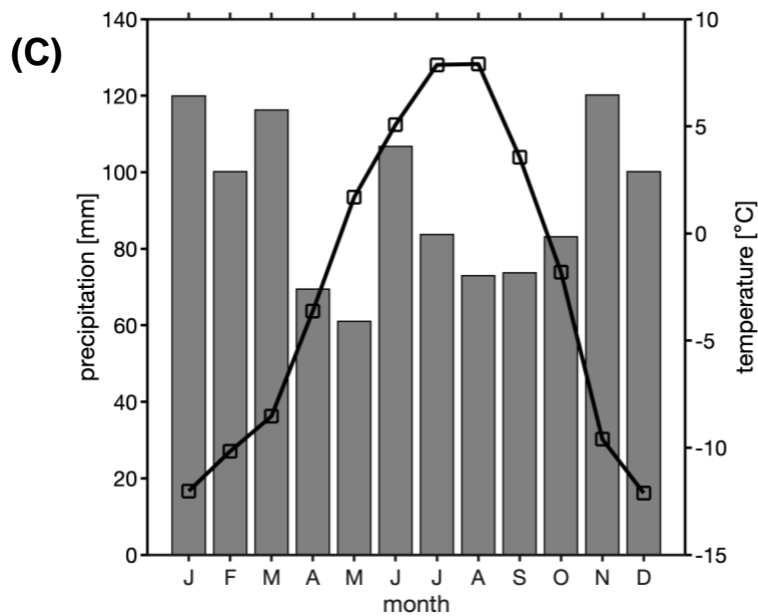
120 Fifteen living *P. engelmannii* trees (one core per tree) were selected for dendroanatomical
121 measurements from a collection sampled in 2015, from tree-line sites (2000–2100 m a.s.l.)
122 adjacent to the Athabasca Glacier in the Columbia Icefield area of the Canadian Rockies
123 (52.13 N, 117.14 W) (fig. 1). The selection of cores was based on 1) the visual appearance of
124 the material (cores with obvious defects were avoided), 2) the temporal coverage of the series
125 (we strived to have an even replication through time) and, 3) the common signal strength
126 based on RBAR statistics (Wigley et al. 1984) of the ring-width measurements (in general,
127 cores with higher than average RBARs were selected for wood anatomy). The selection was
128 primarily dictated by 1) and 2), and only secondarily by 3).



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131

132 **Figure 1:** A) Location of the Columbia Icefield site, Canadian Rockies where wood cores for
 133 dendroanatomical measurements were collected in 2015. B) The tree-ring sampling site at the
 134 Columbia Icefield. The view is to the north from the Athabasca Glacier forefield, September
 135 2018. The 2015 samples were obtained from sites east and west of the Icefield Centre
 136 (building located in the middle of the image). The Athabasca Glacier extended to the foot of
 137 the slope in the left of the photo in the 1840s. C) Monthly mean temperature and total
 138 precipitation (1970–2018 averages) for the CRU TS v4.03 grid point (52.25° N, 117.25° W)
 139 covering the Columbia Icefield area.

140

141 Wood cores were refluxed in alcohol for 24 hours using a Soxhlet apparatus to remove resin
 142 and other soluble substances, and subsequently embedded in paraffin using a Tissue
 143 Processor TP1020 and Histocore Arcadia Embedding Center (Leica, Germany). A rotary
 144 microtome RM2245 (Leica Biosystems, Germany), equipped with N35 disposable microtome
 145 blades (Feather, Japan), were used to cut 12 μm thick transverse sections from the wood

146 cores. The thin-sections were stained with a 1:1 safranin-astrablue solution and mounted on
147 slides with Euparal (Carl Roth, Germany), following standard procedures (von Arx et al. 2016).
148 Digital images from each section were taken with a Zeiss Axio Scan Z1 (Carl Zeiss, Germany)
149 at a resolution of 2.3 pixels μm^{-1} . Tree-ring borders and individual tracheid cells were then
150 semi-automatically identified, and ring width as well as the position and anatomical dimension
151 of each tracheid cell were measured in the digital images using the image analysis software
152 ROXAS (v3.1) (von Arx and Carrer 2014). The anatomical parameters included, for instance,
153 cell lumen area and cell wall thickness (CWT), where the latter was measured in four directions
154 to obtain the average cell wall thickness, i.e. two radial and two tangential cell walls per
155 tracheid cell (Prendin et al. 2017). Each tree ring was divided into 20 μm wide bands with a
156 10 μm overlap, parallel to the ring border (the tangential extension of each band encompasses
157 ~75–100 tracheids). In order to minimize the influence of outliers, for each anatomical
158 parameter the values of all cells corresponding to the 75th percentile within each 20 μm wide
159 band were retained for further analysis. That is, from the cells within the radial and tangential
160 extension of each band, the 75th percentile of each cell dimension was calculated, building up
161 intra-annual measurement profiles of CWT, Lumen area, and other anatomical traits. The
162 maximum parameters (e.g., Max. radial CWT) thus retain the highest value of the profile for
163 each year, the minimum parameters retain the lowest value of the profile for each year, and
164 the earlywood and latewood parameters retain the average value of the profile for the
165 earlywood and latewood portions of the ring, respectively. The anatomical density was derived
166 as the ratio of wall area to overall cell area (that is, including both wall and lumen area) in each
167 20 μm wide band. Mork's index (Denne 1989) was used to separate the earlywood and the
168 latewood portions of the ring. For further details regarding the dendroanatomical
169 measurements, see Björklund et al. (2020).

170

171 *2.2 Chronology development*

172 From the potentially large number of possible dendroanatomical parameters, we narrowed
173 down subsequent analyses to seven parameters of anatomical dimensions, and three wood
174 density parameters based on anatomical dimensions, which are directly comparable to X-ray
175 and blue intensity-based microdensitometric parameters. The parameters are listed in [table 1](#).
176 For comparative purposes, we also retained X-ray derived measurements of MXD (Luckman
177 and Wilson 2005), and the previously unpublished latewood blue intensity counterpart (MXBI)
178 measured on *P. engelmannii* from the Columbia Icefield area. The X-ray MXD was produced
179 using radiodensitometric techniques (Schweingruber et al. 1978) from 1.2-mm-thick laths, cut
180 using a twin-blade saw along the tree cores but perpendicular to the fiber direction (see
181 Luckman and Wilson 2005 for details). For the production of MXBI, the methodology outlined

182 in (Rydval et al. 2014) was adopted. The MXBI measurements were conducted using the
183 CooRecorder software (ver. 8.1) (<http://www.cybis.se/forfun/dendro/index.htm>).
184 Corresponding time series of ring-width were also obtained and hereafter referred to as
185 “original ring-width”, as opposed to “ROXAS ring-width”, which were measured in program
186 ROXAS on the fifteen cores used for the dendroanatomical measurements. The X-ray MXD
187 and MXBI datasets were originally developed from living trees and snag material, however, to
188 ensure consistency for the parameter comparison, we used X-ray MXD, MXBI and original
189 ring-width measurements from living trees only (X-ray MXD: N = 78 series, MXBI: N = 182,
190 and original ring width: N = 182, see [table 1](#)). The dendroanatomical analysis was performed
191 on tree cores for which original ring-width and MXBI measurements were available. Thus, an
192 additional subset based on the fifteen trees was retained for the latter two parameters to
193 ensure also a direct comparison with the dendroanatomical chronologies. For the full MXBI
194 dataset (N = 182), we additionally derived eight partly overlapping percentile chronologies
195 based on *absolute* ring-width (fig. S1), to assess whether a similar ring-width dependence as
196 previously reported by Björklund et al. (2019) from Northern Fennoscandia could also be
197 detected in the Icefield dataset, i.e. a ring-width related differences of MXBI measurements
198 taken in narrow versus wide rings. The following ring-width percentile intervals were used: 0
199 – 30th, 10th – 40th, 20th – 50th, 30th – 60th, 40th – 70th, 50th – 80th, 60th – 90th, and 70th – 100th to
200 derive the sub-sampled MXBI chronologies. Thus, for example, the 70th – 100th percentile
201 chronology is computed from MXBI-values measured in the 30% *widest* rings, while the 0 –
202 30th percentile chronology corresponds to MXBI-values from the 30% of the *narrowest* rings.
203 Unfortunately, a similar comparative analysis was not possible to conduct for the X-ray based
204 MXD, as the corresponding ring-width measurements originally developed, were unavailable
205 to us in the current study.

206

207 Since the analysis was performed on data derived from a cohort of same-aged living trees,
208 capturing low-frequency variability (i.e. decadal and longer) with RCS-type methods is a
209 challenge (e.g., Briffa et al. 1992). This is because living-trees-only, share potential climate
210 signal on lower frequencies even if they are aligned by cambial age (trend-in-signal) (Briffa
211 and Melvin 2011). Any attempt, even using signal-free approaches, will provide indices, that
212 likely would have to be revised when implementing the same technique on a large multi-
213 generational material, and ultimately reflect certainty where there is little. Thus, we primarily
214 focused here on the year-to-year signals in the tree-ring anatomical parameters. To
215 emphasize the interannual variations, the individual dendroanatomical series were detrended
216 in the program MATLAB (version R2021a), by 1) fitting a cubic smoothing spline function with
217 50% frequency response cutoff at 35 years to the raw tree-ring series (Cook and Peters 1981),
218 2) subtracting the fitted values from the observed values to obtain detrended series (division

219 was used to standardize the ring-width measurements), and finally 3) averaging the detrended
220 series by simple arithmetic mean to produce the final parameter-specific chronologies
221 (hereafter referred to as detrended data). The same detrending procedure was performed on
222 the MXBI, X-ray derived MXD and original ring-width series, in order to obtain data that are
223 comparable with the dendroanatomical datasets. All chronologies were truncated to the 1700–
224 1994 period in the subsequent analyses, to ensure a consistent overlap between datasets as
225 well as a sample depth ranging between 9 and 15 cores for the anatomical dataset.

226

227 *2.3 Statistical methods*

228 To evaluate the strength of the between-series common signal and establish the replication
229 needed to obtain mean chronologies meeting the commonly accepted standard, we used the
230 RBAR (defined as the mean Pearson’s correlation coefficient between all possible pairs of
231 individual tree-ring series) (Wigley et al. 1984) and Expressed Population Signal (EPS) (Briffa
232 et al. 1992) statistics. To assess the degree to which the various parameters co-vary, principal
233 component analysis (PCA) and pairwise correlations were computed over the 1700–1994
234 period.

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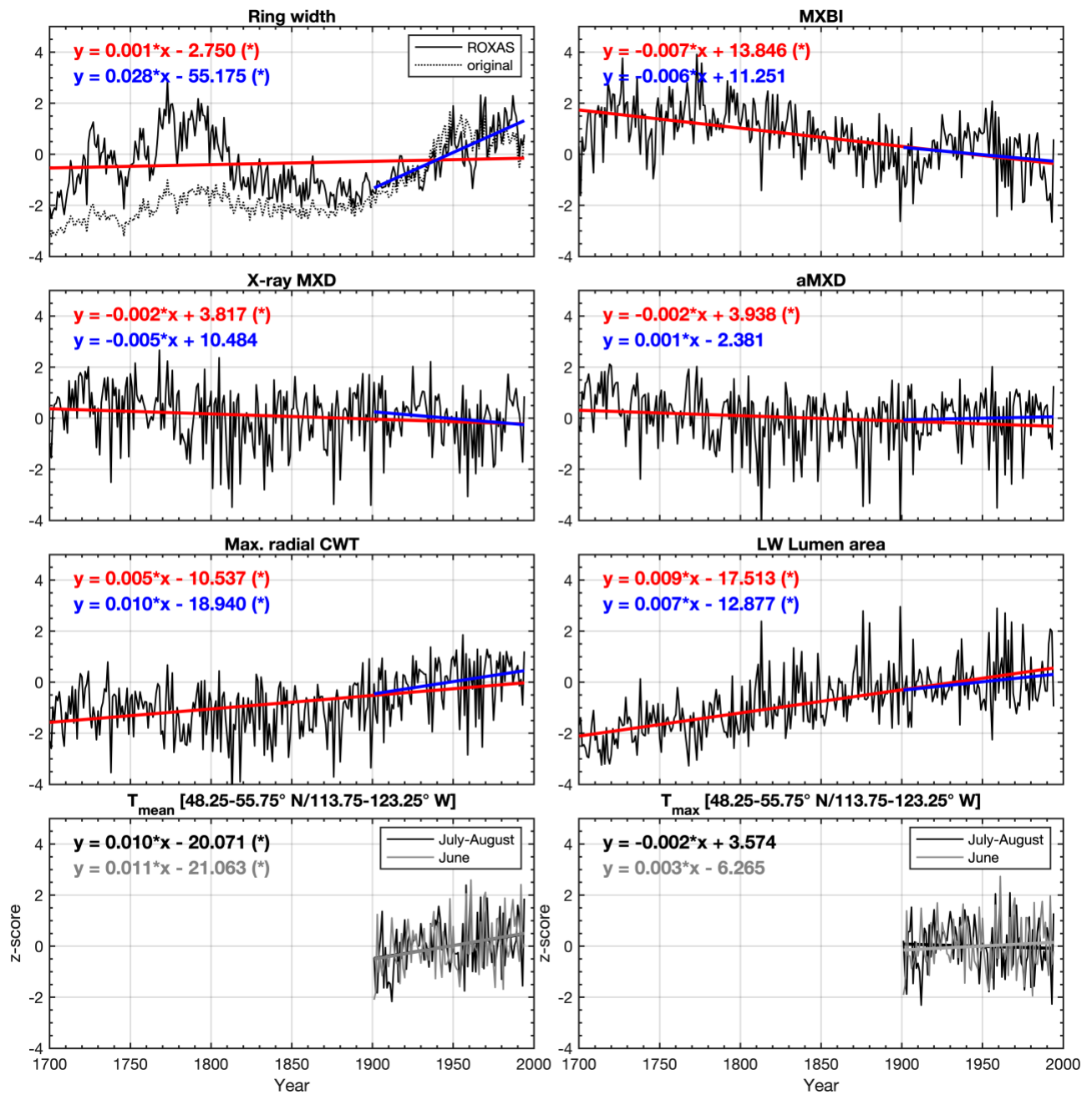
236 Detrended tree-ring parameter chronologies were assessed for their relationship to regional
237 monthly mean (Tmean) and maximum (Tmax) temperatures, by correlation against the
238 monthly 0.5° x 0.5° gridded CRU TS v4.03 dataset (Harris et al. 2020) for the grid point
239 average bounded by the latitude/longitude coordinates 48.25–55.75° N/113.75–123.25° W
240 (Fig. 1, 2). Tmax was included in the analysis because previous work has demonstrated
241 slightly stronger calibration statistics than for Tmean when using MXD and ring-width
242 chronologies for climate reconstruction in this region (e.g., Heeter et al. 2021; Wilson et al.
243 2019; Wilson et al. 2014; Wilson and Luckman 2003). The associations with monthly
244 precipitation totals and minimum temperatures were also tested, but not included here due to
245 weak significant empirical relationships. The lack of precipitation sensitivity of *P. engelmanni*
246 in the Icefield area was already noted in (St. George and Luckman 2001) which is not
247 surprising as the trees are growing in temperature limited upper tree-line environments.
248 Pearson’s correlations were calculated between parameter-specific chronologies and monthly
249 meteorological variables over the 1901–1994 period, and the 1901–1948 and 1949–1994
250 subperiod to evaluate temporal stabilities of the climate responses. A paired *t*-test was used
251 to test whether the calibration statistics differed between tree-ring parameters and sub-
252 periods. To make the climate sensitivity analysis comparable to previous studies from the
253 Columbia Icefield area, we also included the homogenized (1895–present) 50 x 50 km gridded
254 temperature data (Vincent and Gullett 1999; Zhang et al. 2000) originally developed by the
255 Meteorological Service of Canada and previously used in Luckman and Wilson (2005) to

256 reconstruct last-millennium summer temperatures for the Canadian Rockies. Similar to
257 Luckman and Wilson (2005), we used the mean of four grids closest to the Columbia Icefield
258 area. Calibration trials with these data are provided in the supplement (fig. S2 and S3). To
259 ensure the climate analysis was not affected by long-term trends, all temperature data were
260 filtered prior to analysis using the same 35-year filter as was used to detrend the tree-ring
261 parameters (henceforth referred to as detrended data).

262

263 Further, the dynamic nature of the temperature signal (i.e. optimal target season and its
264 temporal stability) was evaluated through moving window correlation analysis between
265 detrended tree-ring chronologies and detrended average daily temperature data (grid 52.5°
266 N, 118.5° W) from the Berkeley Earth dataset (<http://berkeleyearth.org/data/>) (Rohde and
267 Hausfather 2020) covering the 1880–recent period. Pearson’s correlations were computed for
268 30-year sliding windows with a 1-year offset. For each 30-year block, temperatures were
269 averaged in 30-day long windows which were shifted at daily time steps throughout the year
270 (sensu Jevsenak and Levanic 2018).

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Figure 2: Average non-detrended time series of selected tree-ring parameters, z-scored over the 1901–1994 reference period. The blue and red lines show the linear trends over the 1901–1994 and 1700–1994 periods, respectively. Seasonally averaged June–August (48.25–55.75° N/113.75–123.25° W CRU TS v4.03 subset average) mean (T_{mean}) and maximum (T_{max}) temperatures are provided for comparison. (*) indicates a significant trend ($\alpha = 0.05$) estimated by the Mann-Kendall trend detection test. Abbreviations used in the figure are LW (latewood), CWT (cell wall thickness), aLWD (anatomical latewood density) and aMXD (anatomical maximum latewood density). See supplement fig. S4 for full 1586–2015 CE period chronologies.

282 3. Results and discussion

283 3.1 *Picea engelmannii* dendroanatomy characteristics

284 Besides the conventional width parameters (i.e., ring width, earlywood- and latewood width,
285 referred to as “ROXAS” in [table 1](#)), seven anatomical parameters and three anatomically-
286 based density parameters were retained for analysis. Basic chronology assessments of
287 detrended data over the common 1700–1994 CE period are provided in [table 1](#), and non-
288 detrended mean chronologies for selected parameters are shown in [figure 2](#). In line with
289 previous work by Björklund et al. (2020) on temperature-sensitive conifers, we find that that
290 maximum radial cell wall thickness (Max. radial CWT) and anatomical MXD (aMXD) are the
291 two anatomical parameters with the highest mean inter-series correlation (R_{BAR} = 0.47 and
292 0.48, respectively). For both parameters, EPS reaches the 0.85 threshold (Wigley et al. 1984)
293 with 6 series ([table 1](#)). Notably, these values are of comparable strength to the R_{BAR} and
294 EPS of X-ray based MXD (R_{BAR} = 0.49, 6 trees required for EPS = 0.85). By comparison, the
295 R_{BAR} for MXBI is surprisingly low at 0.19 and the replication needed to attain the EPS of 0.85
296 is 24 series. These MXBI chronology statistics are lower than for ring width (R_{BAR} = 0.22 and
297 0.28 for original and ROXAS ring width, respectively) – an observation noted previously by
298 (Rydval et al. 2014; Wilson et al. 2019). The R_{BAR} and EPS values for MXBI slightly decrease
299 if computed only on the 15 trees that have been pre-selected for the dendroanatomical
300 analysis. This is surprising given that the selection of the cores for dendroanatomy was partly
301 based on its ring-width signal strength ([see sect. 2.1](#)), and that the R_{BAR} and EPS statistics
302 for ring width actually improve when narrowing the analyses down to these 15 trees ([see table](#)
303 [1](#)). Although the BI-based density parameters typically require a larger sample size than ring
304 width (e.g., Blake et al. 2020; Wilson et al. 2021) for a robust chronology, the MXBI chronology
305 statistics obtained for *P. engelmannii* from our site are still lower than the previously reported
306 MXBI findings for the same species across British Columbia, Canada (Wilson et al. 2014).

307
308 Notably, several anatomical and density parameters are found to exhibit a rather low common
309 signal, yet a reasonably strong temperature sensitivity ([see sect. 3.2](#)). These include, in
310 decreasing order of signal strength: earlywood (EW) cell wall area (R_{BAR} = 0.13), EW lumen
311 area (R_{BAR} = 0.12), EW density (R_{BAR} = 0.10), EW cell area (R_{BAR} = 0.09) and latewood
312 (LW) cell area (R_{BAR} = 0.09). The replication required to attain robust EPS statistics ranges
313 between 38 (EW cell wall area) to 57 trees (EW cell area and LW cell area).

314
315 **Table I:** Basic summary statistics for each detrended parameter chronology, based on the
316 common 1700–1994 period. Abbreviations used in the table are EW (earlywood), LW
317 (latewood), CWT (cell wall thickness), aLWD (anatomical latewood density) and aMXD
318 (anatomical maximum latewood density). Estimations of the number of trees needed to reach

319 the arbitrary EPS threshold level of 0.85 are derived from the EPS equation (see Wigley et al.
 320 1984) using the RBAR statistic for each tree-ring parameter. Parameters highlighted in grey
 321 are those requiring the lowest sample replication to reach the threshold level.

	# samples	RBAR	n for EPS (0.85)
Width parameters			
Original ring-width (from Luckman 1997; Luckman and Wilson 2005, and later unpublished updates)	182	0.22 (0.27 for N = 15)*	20 (15 for N = 15)*
ROXAS ring-width	15	0.28	15
ROXAS EW width	15	0.26	16
ROXAS LW width	15	0.19	24
Earlywood anatomy			
EW cell area	15	0.09	57
EW Lumen area	15	0.12	42
EW cell wall area	15	0.13	38
Latewood anatomy			
LW cell area	15	0.09	57
LW Lumen area	15	0.31	13
Max. radial CWT	15	0.47	6
Max. tangential CWT	15	0.34	11
Density parameters			
EW density	15	0.10	51
aLWD	15	0.28	15
aMXD	15	0.48	6
MXBI (unpublished)	182	0.19 (0.16 for N = 15)*	24 (30 for N = 15)*
X-ray MXD (from Luckman and Wilson (2005))	78	0.49	6

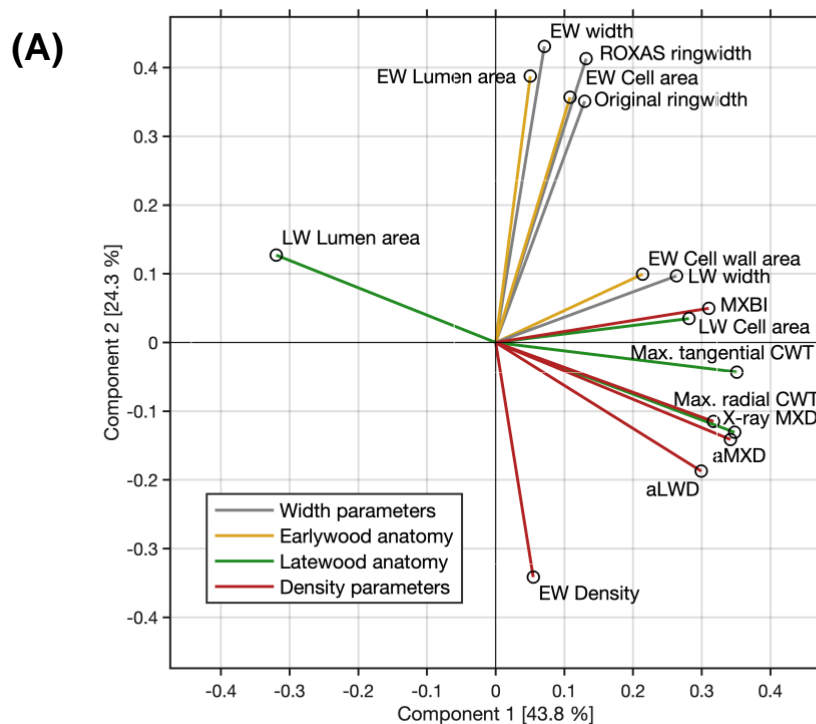
322 *The RBAR and EPS values in parentheses are for the original ring-width and MXBI time-series
 323 computed for exactly the same 15 trees that have been used to produce the wood anatomy datasets.

324
 325 The co-variability between the various parameters over their common 1700–1994 period was
 326 assessed through principal component analysis and pairwise correlations (fig. 3). The first two
 327 components together represent 68.1% of the total variation. The PC1 alone explains 43.8% of
 328 variance, and is dominated by latewood-related parameters, including both anatomy and
 329 density parameters. We found that aMXD, Max. radial CWT and X-ray MXD cluster together
 330 in the bivariate plot, showing that all three parameters express comparable signals (also
 331 corroborated by the correlation matrix in fig. 3b). The MXBI also loads strongly positively on
 332 PC1, but slightly separates from this cluster by being positively correlated to PC2. Among the
 333 LW density-related components, MXBI is the parameter best correlated with ring-width and
 334 latewood-width chronologies (fig. 3b), although these correlations are only moderate ($r_{\text{MXBI vs.}}$
 335 original ring width = 0.43, $r_{\text{MXBI vs. latewood width}} = 0.66$). The principal component analysis including the

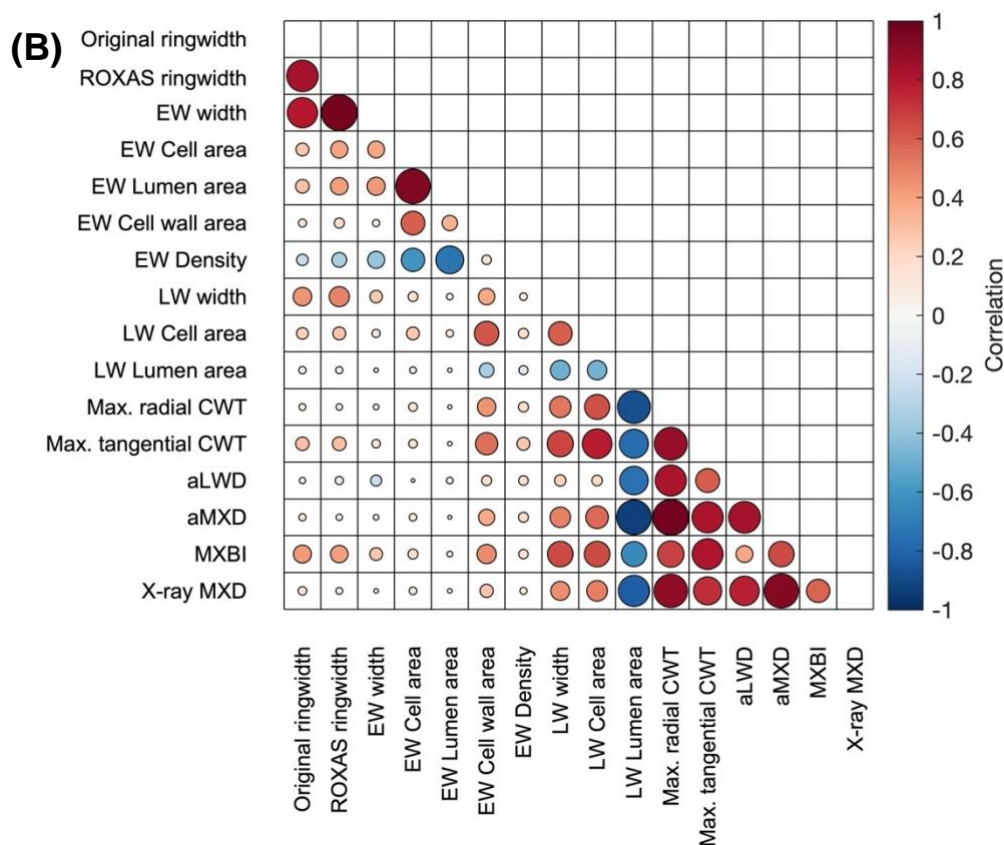
336 subsampled MXBI percentile chronologies based on the *absolute* corresponding ring widths
 337 reveal that the correlation coefficients against the latewood width, and to some degree also
 338 ring width, successively increase for the “narrow-ring MXBI chronologies” (fig. S5). The “wide-
 339 ring MXBI chronologies” (i.e., ~50th-100th percentiles) are, on the other hand, more similar to
 340 the aLWD, Max. radial CWT, aMXD and X-ray MXD chronologies. This observed ring-width
 341 inclination of MXBI suggest that the dataset might be subject to a resolution bias (Björklund et
 342 al. 2019). More detail on this issue in [sect. 3.4](#).

343

344 The variance of PC2 (24.3 % of total variability) is dominated by ring width and earlywood-
 345 related density and anatomy parameters. Amongst these, EW density stands out by loading
 346 strongly negatively on the PC2 axis (reflecting its negative association with early-summer
 347 temperatures, see [sect. 3.2](#)). Moreover, the EW cell wall area stands out by loading more
 348 strongly on the PC1 axis than on the PC2 axis, and by clustering more closely with the
 349 latewood than with the earlywood components (reflecting its late-summer temperature
 350 sensitivity, see [sect. 3.2](#)).



351



352

353 **Figure 3:** A) biplot of the first two principal components of the PCA performed 1700–1994 CE
 354 period on the width, anatomy and density parameters. The colors of the vectors correspond
 355 to the parameter grouping used in table 1. The first two components together represent 68.1%
 356 of the total variation. B) Pearson's correlation matrix between various anatomical and width
 357 parameters. X-ray MXD and MXBI are included for comparison. Correlations are computed
 358 over the common 1700–1994 period using detrended chronologies. The color and size of the
 359 markers denote the direction and strength of the relationships.

360

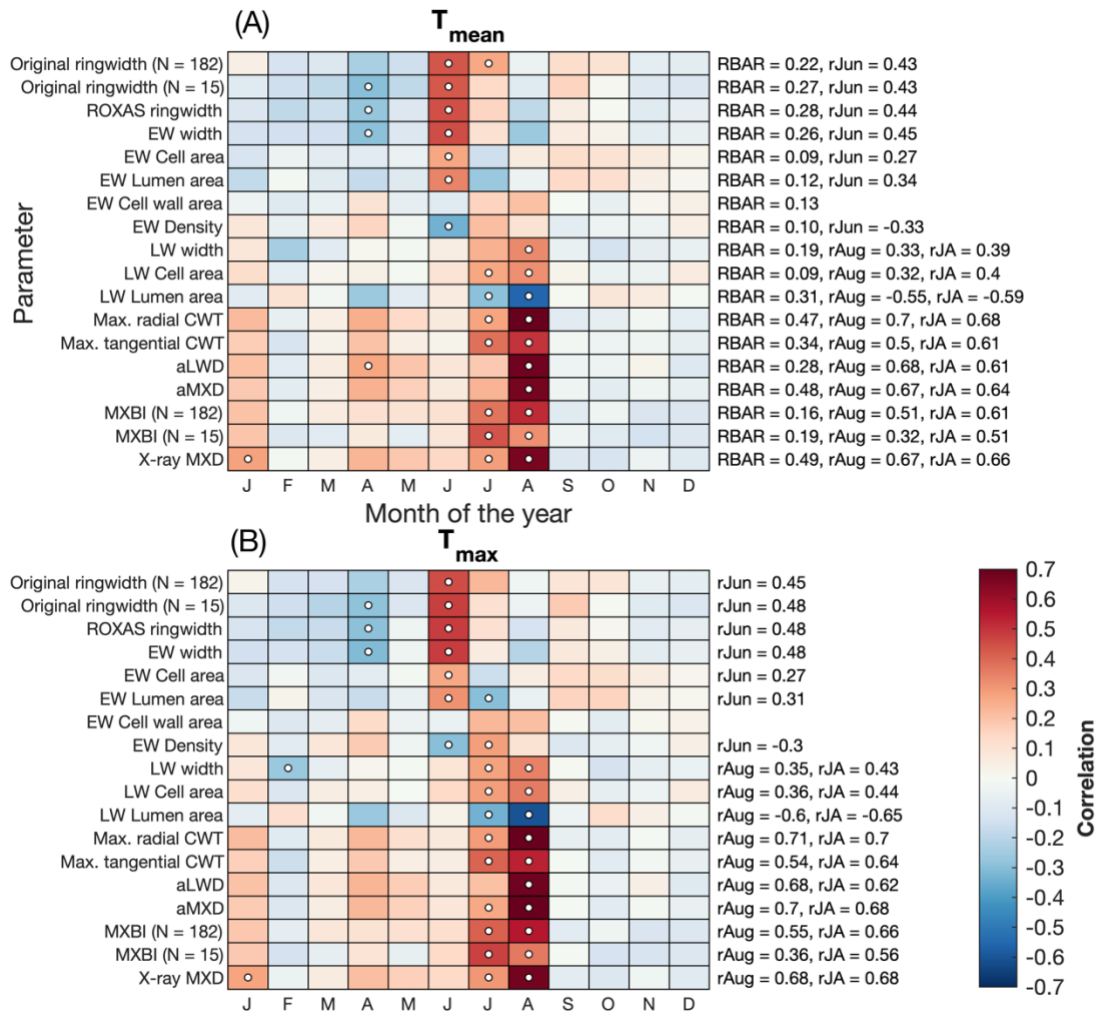
361 3.2 Climate response

362 Simple linear correlations between selected parameters and monthly CRU TS mean (Tmean)
 363 and maximum (Tmax) detrended temperatures are shown in fig. 4. In line with previous work
 364 from North America (Harley et al. 2021; Heeter et al. 2021; Luckman and Wilson 2005; Wilson
 365 et al. 2014; Wilson and Luckman 2003), our results reinforce the importance of Tmax
 366 temperatures for wood formation and growth of *P. engelmannii* in the region by providing, in
 367 general, slightly higher correlation values for Tmax than for Tmean. Interestingly, the pattern
 368 observed in North America contrasts to many other temperature-limited regions of the
 369 Northern Hemisphere, where conifers have generally been noted to correlate stronger to
 370 Tmean than to Tmax (observation made by the author team, results not published). Whether

371 this is actually grounded in a tree physiological mechanism is still an open question.
372 Furthermore, the general pattern revealed by the climate response analysis shows that the
373 various dendroanatomical traits respond to consecutive temporal windows within a short
374 seasonal window extending from June to August, in line with our understanding of the
375 successive physiological processes (i.e., cell expansion and cell wall thickening) behind wood
376 formation and growth (e.g., Fonti et al. 2013). These results support the climate-response
377 pattern that has generally been observed for conifers across the Canadian Rockies (Luckman
378 and Wilson 2005) and the adjacent Interior British Columbia (Wilson et al. 2014; Wilson and
379 Luckman 2003), yet contrasts to the seasonally wide temperature imprint (extending between
380 May–August and occasionally even between April–September) within latewood density of
381 *Picea mariana* in the eastern Canadian taiga (Wang et al. 2020). This is also the case when
382 comparing our results with the previous study of (Björklund et al. 2020) on latewood
383 anatomical traits of *Pinus sylvestris* in northern Scandinavia, where the temperature response
384 window extends from April to September. The narrow window of response patterns seen here
385 is most likely constrained by the distinct and short warm season characterizing the climatology
386 of the study site, where average monthly temperatures rise above 0 °C only in five months of
387 the year (fig. 1c).

388

389 We find that latewood-related parameters in general display a late-summer (July–August)
390 temperature sensitivity, while ring width and earlywood-related density and anatomy
391 parameters most strongly correlate with mid-summer (June–July) temperatures (fig. 4). The
392 strongest temperature signals are found in anatomical components of the latewood, which are
393 also the parameters displaying the highest RBAR statistics (table 1). In particular, aMXD and
394 Max. radial CWT stand out. The imprints of year-to-year temperature variability within these
395 two parameters are, over the 1901–1994 period, very similar, if not identical, to that of the
396 MXD derived from the X-ray technique. By comparison, the exceptionally weak inter-series
397 signal strength of the MXBI parameter (table 1) is compensated by high replication (N = 182),
398 and thus MXBI is also rather similar to aMXD, Max. radial CWT and X-ray MXD. However, the
399 temperature signal of MXBI is shifted earlier by expressing stronger correlation with July
400 temperatures but weaker with August compared to aMXD, Max. radial CWT and X-ray MXD.
401 The aggregated July–August temperature response of MXBI is thus in fact only marginally
402 weaker than that of X-ray MXD, aMXD and Max. radial CWT.



403

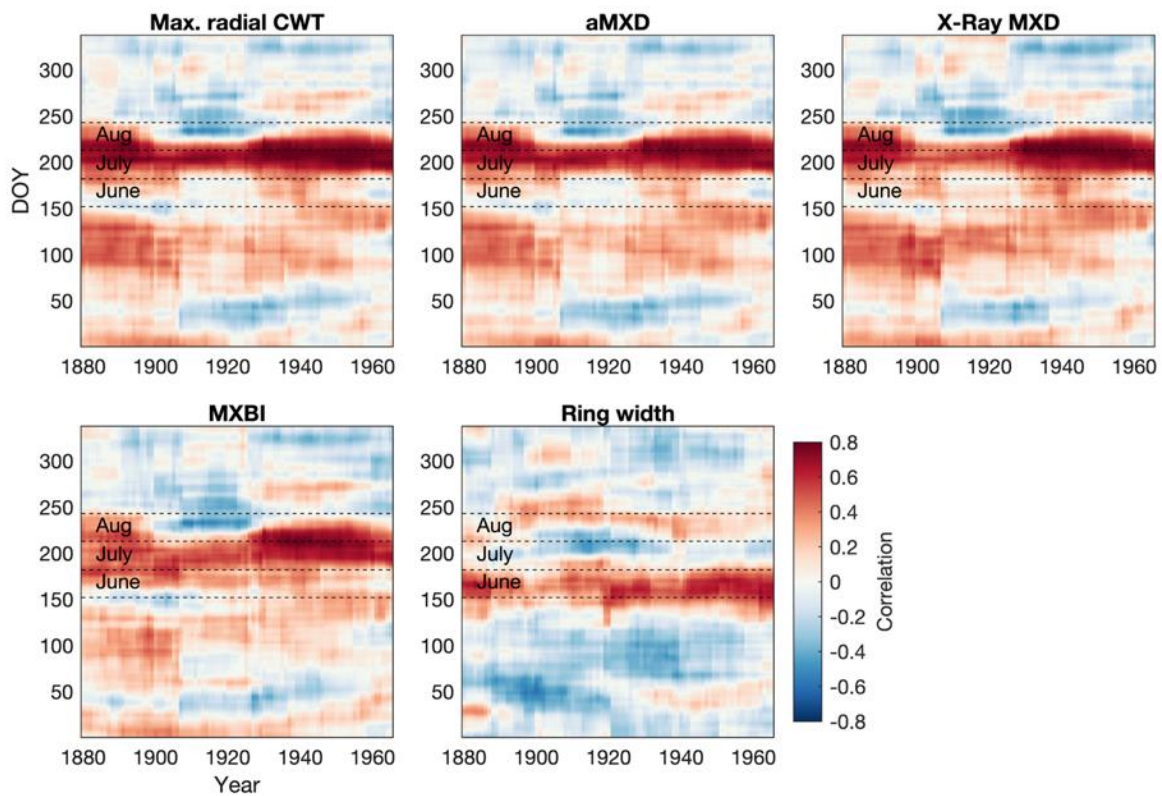
404 **Figure 4:** Correlations between tree-ring parameters and monthly (A) average (T_{mean}) and (B)
 405 maximum (T_{max}) temperatures from the CRU TS v4.03 product (48.25–55.75° N/113.75–
 406 123.25° W subset average). Pearson's correlation coefficients are computed over the 1901–
 407 1994 period using detrended data. The RBAR statistics for each parameter chronology, and
 408 correlation coefficients for the best temperature target season are provided on the right side
 409 of the plots (June for ring width and EW parameters, July-August and August for LW
 410 parameters). For original ring width and MXBI, results are also provided for chronologies
 411 (denoted as N = 15) built from the same 15 trees that are used to produce the dendroanatomy
 412 data. Significant correlations ($p < 0.01$) are marked with white circles. Correlations with
 413 temperature data produced by the Meteorological Service of Canada are given in the
 414 supplement (fig. S2).

415

416 3.3 Temporal signal stability

417 Focusing only on anatomical traits with the highest temperature sensitivity (aMXD and Max.
 418 radial CWT), comparison against daily temperatures (fig. 5) confirms a significant and strong
 419 mid/late summer signal over the 1880–1994 period. Breaking down the climate response in

420 daily increments reveals that the strongest signal ($r > 0.5$) occurs on average between day
 421 192 and day 251 of the year (i.e. July 11th until September 8th-9th, with a peak correlation of
 422 0.74 for Max. radial CWT and aMXD occurring between 21st of July-20th of Aug and 23rd of
 423 July-22nd of August), respectively. The temperature associations at the margins of the target
 424 season are, however, more unstable. We note, for example, that the September signal
 425 disappears around the first half of the 20th century for both anatomical parameters. A similar
 426 correlation structure holds for X-ray derived MXD and to a lesser degree MXBI (N = 182), but
 427 the two parameters exhibit enhanced correlation coefficients in the second half of the 20th
 428 century compared to the early period (also corroborated by the split-period calibration in [figure](#)
 429 [6](#)). Moreover, despite the high sample replication, MXBI shows slightly weaker correlations
 430 with daily data than the other density-related parameters, particularly in the early 1880–1930
 431 period, when ring widths coincidentally are the narrowest in the record (see [fig. 8](#)-and [sect.](#)
 432 [3.4](#)). For comparative purposes we also include anatomically derived ring width, which shows,
 433 on average, the strongest correlations ($r = 0.3$ to 0.5) with temperatures between day 146 and
 434 206 of the year (i.e. May 26th to July 25th).



435
 436 **Figure 5:** Moving correlation between the full tree-ring parameter datasets and Berkeley Earth
 437 gridded daily temperatures (grid 52.5°N 118.5°W, 1880–1994 period). A 30-year moving
 438 window, shifted by one year, was used in the analysis. Temperatures were averaged over a
 439 30-day window and shifted throughout the year at daily steps. The days on the x- and y-axis
 440 thus show the first day of the 30-year and 30-day windows, respectively. E.g., day 152 on the

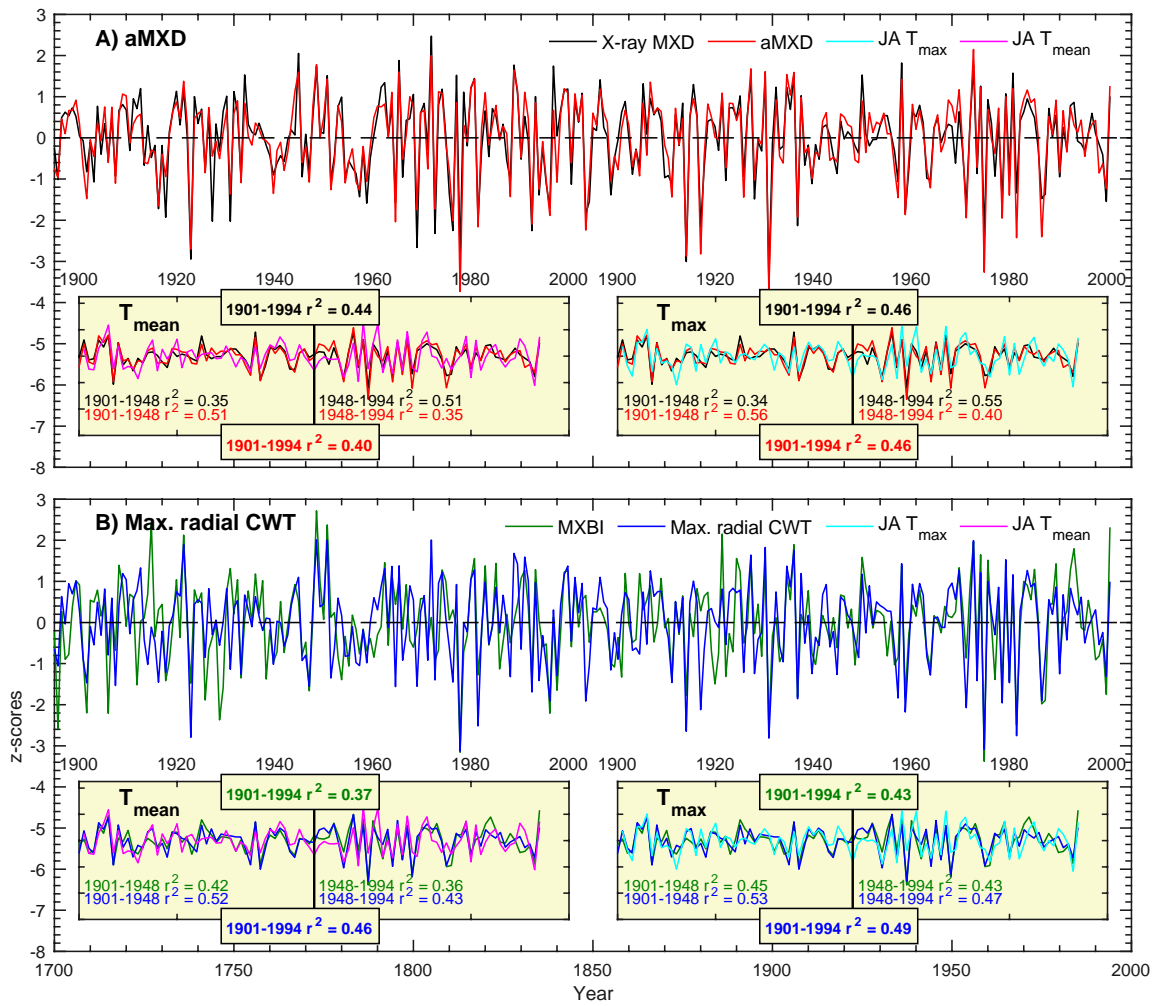
441 *y-axis represents the period from June 1 to June 30. Both tree-ring and temperature data have*
442 *been detrended prior to analysis. The June-August months are highlighted to aid*
443 *interpretation. Pearson's correlations between monthly aggregated (i.e. average daily data for*
444 *each calendar month) Berkeley Earth temperatures and tree-ring parameters are provided in*
445 *figure S6 for comparison.*

446

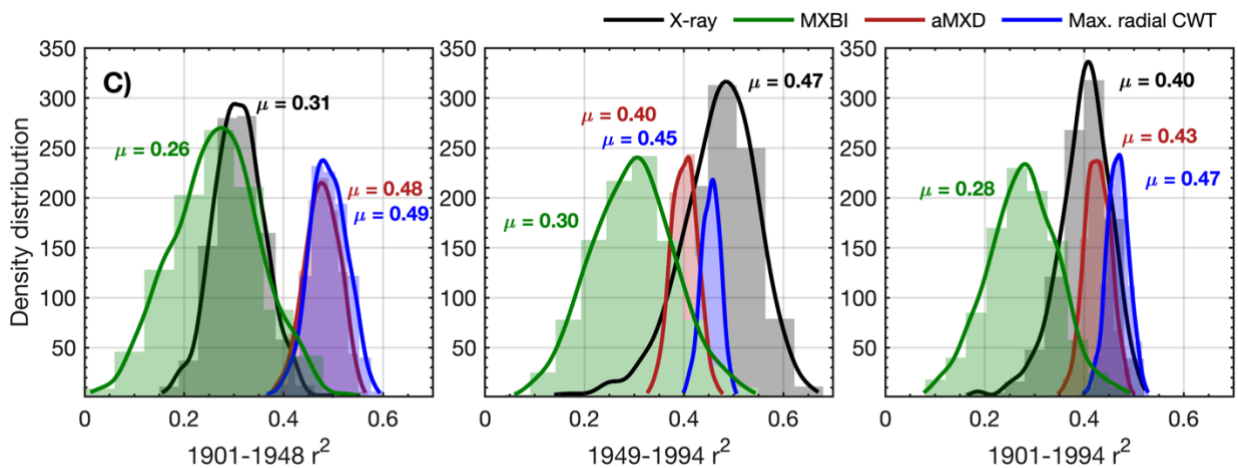
447 The stability of the July-August temperature signals of detrended aMXD and Max. radial CWT,
448 along with X-ray MXD and MXBI, were further assessed by a split-period calibration procedure
449 (1901–1948 and 1949–1994) (fig. 6). The two wood anatomical parameters calibrate more
450 strongly to the early period compared to the late, both when using Tmean and Tmax. However,
451 especially for Max. radial CWT, the calibration differences in the two periods are slight ($r^2 =$
452 53% and 47% against Tmax, respectively). By comparison, the X-ray MXD calibrate more
453 strongly in the latter half of the instrumental period and show more pronounced temporal
454 instabilities ($r^2 = 34\%$ and 55% against Tmax, respectively). This contrasts to the prior finding
455 (Luckman and Wilson 2005), where no such instabilities in the early 20th century were
456 detected. These contrasting results are most likely not related to using different climate data
457 products because similar results (fig. S3) were obtained when using the Luckman and Wilson
458 (2005) temperature data, originally produced by the Meteorological Service of Canada.
459 Instead we suspect that the discrepancy can be attributed to either using a larger network of
460 MXD data than used in this study, or that Luckman and Wilson (2005) used multivariate
461 regression models (including ring width and lagged growth responses) to explain a wider target
462 season than attempted here.

463

464 Calibration trials with detrended data over the full period 1901–1994 reveal that Max. radial
465 CWT performs overall best (Tmax $r^2 = 49\%$), closely followed by aMXD ($r^2 = 0.46\%$) and X-
466 ray MXD ($r^2 = 0.46\%$). The temporal instability of X-ray MXD and by comparison the robust
467 and strong signals of the aMXD and especially the Max. radial CWT parameters are further
468 confirmed by the resampling calibration trials presented in fig. 6c, where 10 random series are
469 drawn from the sample cohorts 1000 times without replacement, and the resulting parameter
470 chronologies are subsequently correlated against July-August Tmax. The reason for the X-ray
471 MXD loss in signal is difficult to disentangle, but it is unlikely related to having different samples
472 for the X-ray and anatomical datasets because the resampling scheme clearly show that the
473 r^2 -distributions are different (fig. 6c) (also corroborated by a two-sample t -test at a significance
474 threshold of 0.05, indicating that the r^2 -statistics comes from two populations with unequal
475 means).



476



477

478 **Figure 6:** A-B): Full (1901–1994) and split-period (1901–1948, 1949–1994) calibration
 479 statistics for the Max. radial CWT (blue line), aMXD (red line), X-ray MXD (black line) and
 480 MXBI (green line) chronologies against July-August mean and maximum CRU TS
 481 temperature. Time-series in the figures show non-detrended mean chronologies, z-scored

482 *over the instrumental 1901–1994 period. C): The density distribution and its mean (μ) of r^2 -*
483 *values obtained from 1000 calibration trials where parameter chronologies are built from 10*
484 *series randomly drawn without replacement from the sample cohort. The resampling trials are*
485 *based on detrended climate and tree-ring data. Calibrations are performed against July-*
486 *August maximum temperatures.*

487

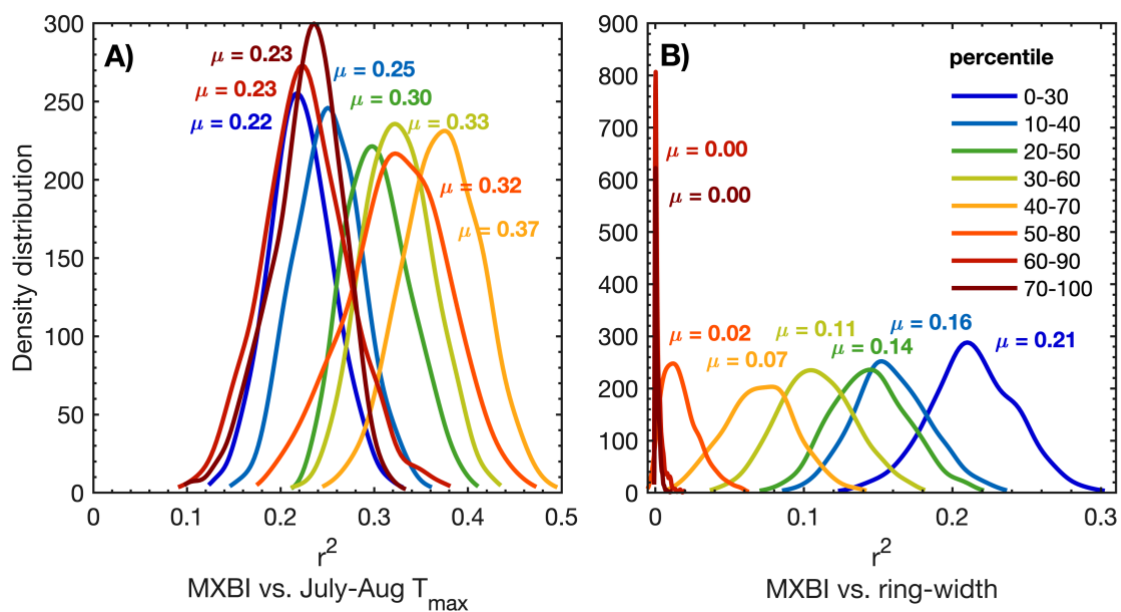
488 *3.4 Possible implications of measurement resolution on climate signal*

489 As shown in the previous sections, the climate imprint withing the anatomical LW density
490 components slightly differs from its X-ray and BI based counterparts, although all these
491 parameters essentially measure the same component in wood. As previously noted by
492 Björklund et al. (2020), the main difference between these metrics is the measurement
493 resolution, while factors such as the cell wall density is of marginal importance (Björklund et
494 al. 2021). Thus, as part of a multi-parameter approach, the higher resolution of dendroanatomy
495 may serve to evaluate the potential risk of a resolution bias (in X-ray MXD and MXBI) when
496 implementing these parameters both on shorter and longer timescales.

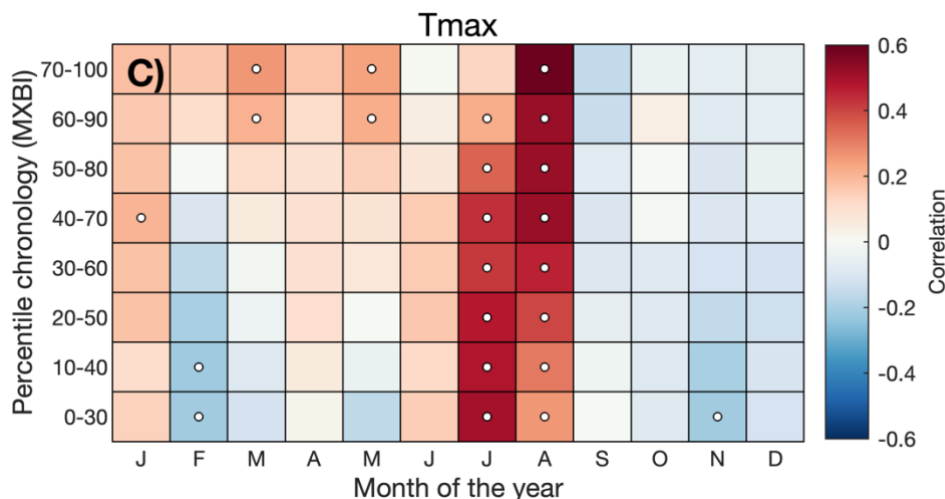
497

498 We have seen that the monthly correlations of the full MXBI dataset (N = 182) differ slightly
499 from the more physically direct density and anatomy parameters, which we hypothesize could
500 partially be related to the lower measurement resolution that artificially makes it more similar
501 to ring width and latewood width (Björklund et al., 2019). The pairwise correlation between
502 parameter chronologies (fig. 3) and the PCA biplot based on the percentile MXBI chronologies
503 (fig. S5) confirms this enhanced relationship with ring width/latewood width. To test this theory
504 further, we have correlated the percentile MXBI chronologies against the target July-August
505 Tmax (fig. 7a) and against the full (N = 182) detrended original ring-width chronology (fig. 7b),
506 using resampling of data. Unfortunately, corresponding latewood width measurements are not
507 available for MXBI, so this comparative analysis is restricted to ring width. Nevertheless, we
508 find that when using the full July-August season the poorest temperature imprint is found in
509 the MXBI values of the narrowest (~40%), and the widest (~40%) of the rings, while the
510 strongest July-August signal can be recovered from the MXBI-values in rings that are close to
511 average in width (40th – 70th percentile). Expanding the climate correlation analysis to monthly
512 Tmax data (fig. 7c) reveals, however, a gradual transition from predominantly an August
513 temperature signal in the wide ring MXBI chronologies to a more July dominated signal in the
514 narrow ring MXBI chronologies. MXBI-values in rings that are close to average in width
515 correlate equally strong to both July and August, which explains the overall better performance
516 of these data when comparing to the July-August target (fig. 7c). Importantly, we find no
517 correlation between the MXBI and ring-width in the widest rings (fig. S5). However, as we
518 move towards narrower rings, the MXBI-values become successively more like ring

519 width/latewood width (fig. 7b and fig. S5). All in all, these results suggest that an effect of low
 520 measurement resolution may be present for narrower ring widths/latewood widths. If so, this
 521 means that the MXBI parameter may become subject to greater target seasonal uncertainty,
 522 which may fluctuate between July and August signals through time, largely depending on the
 523 absolute ring width/latewood width of the analyzed tree-ring sample collection and the
 524 resolution of the captured image. Although posing a challenge for paleoclimate
 525 reconstructions, this resolution issue is likely to become a less relevant methodological
 526 problem in a near future, as more laboratories are currently investing in the development of
 527 high-resolution image capturing systems and other analytical techniques to enhance the
 528 precision of the BI data.



529



530

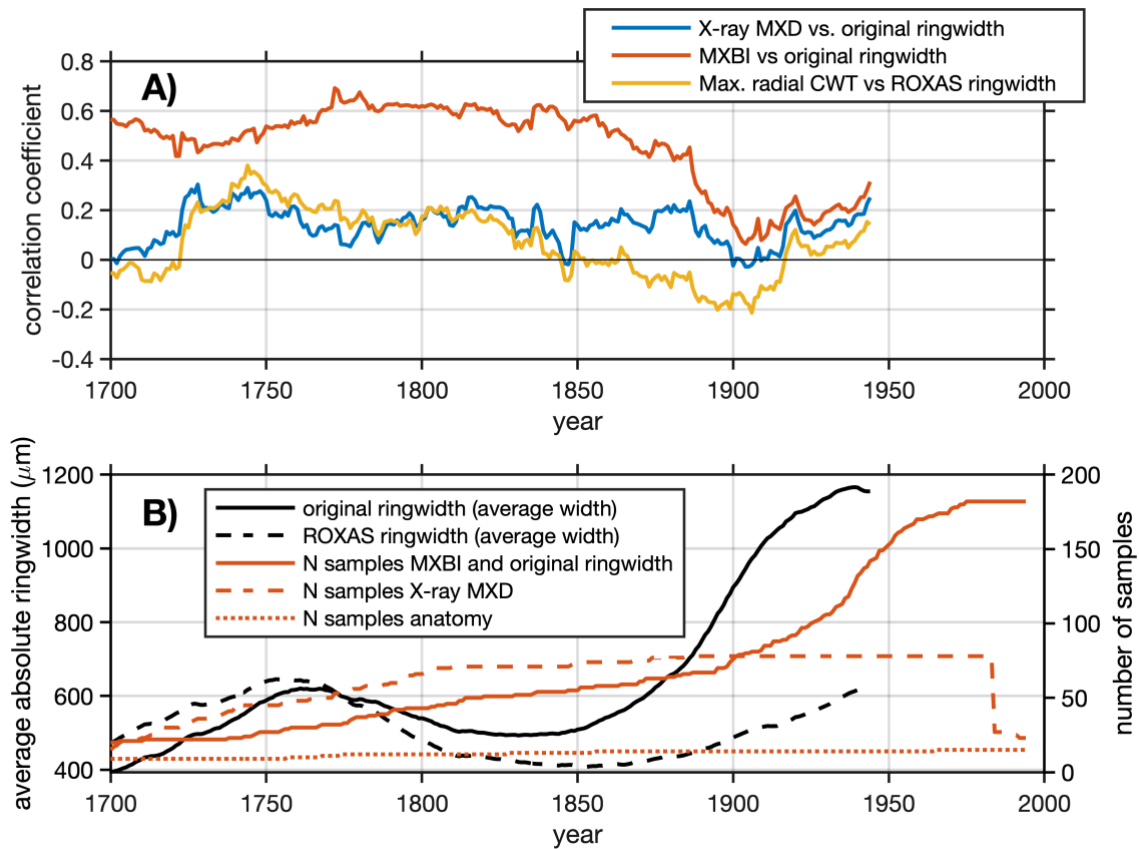
531 **Figure 7:** A)-B): The density distribution of r^2 -values obtained from 1000 calibration trials
 532 (1901–1994 period) where MXBI chronologies are built from 100 series randomly drawn from
 533 the total of 182 series without replacement. The detrended MXBI values are sorted into
 534 percentiles based on the absolute ring-width (e.g., the 0-30 percentile are the corresponding

535 *MXBI-values for the narrowest 30% of the rings), and then averaged into percentile*
536 *chronologies. A) the calibration r^2 -values between these chronologies and detrended July-*
537 *August CRU TS Tmax, B) same as A) but calibrated against the full ($N = 182$) detrended ring-*
538 *width chronology. C) Correlation between the MXBI percentile chronologies and monthly*
539 *maximum (Tmax) temperatures from the CRU TS v4.03 product (48.25–55.75° N/113.75–*
540 *123.25° W subset average). Correlation coefficients are computed over the 1901–1994 period*
541 *using detrended tree-ring and temperature data. Significant correlations ($p < 0.01$) are outlined*
542 *with white circles.*

543

544 Further, we note that the correlations between the various latewood parameters against ring
545 widths change from the early to late 20th century periods, *and* that the correlations slightly
546 differ in magnitude and sign (fig. 8). The MXBI is positively correlated with ring width, whereas
547 the correlations for X-ray MXD range from non-significant to weakly positive. The Max. radial
548 CWT, on the other hand, show a non-significant or weak negative correlation with ring width
549 during the 20th century. This gradual, and slightly larger shift in moving window correlation
550 against ring width during the early 20th century may thus be an indication that both MXBI and
551 to some degree X-ray MXD are challenged by comparatively low measurement resolution. If
552 this is the case, then the inter-annual climate signal may potentially become muted when ring
553 (latewood) widths are narrow. This dependence could, in fact, affect the lower frequencies,
554 and inflate multi-decadal variability (Esper et al. 2015). Moreover, the fidelity to the monthly
555 temperature targets may exhibit instability when rings (latewoods) are narrow, shifting back
556 and forth between August or July dominated signals (as seen in fig. 7c). It is at the moment
557 unclear how this phenomenon could affect the lower frequencies of our chronologies, as a
558 robust picture of long-term trends in dendroanatomical parameters can only emerge from
559 analysis of millennial length, multi-generation, composite chronologies suitable for RCS-type
560 detrending (Briffa and Melvin 2011). Moreover, periods with persistence in narrow ring widths
561 will force MXBI, and perhaps also X-ray MXD, to exhibit persistently low densitometric values
562 (Björklund et al. 2019). Exacerbating this issue is that persistently narrow ring width/latewood
563 width may not even be a product of the distinct and earlier temperature target (June-July, fig.
564 4), but could also be related to stand dynamics/disturbances (Rydval et al. 2018), and thus
565 pass down non-climatic distortions of decadal to centennial variations to X-ray MXD and MXBI.
566 This clearly needs further scrutiny because it may be important for the interpretation of inferred
567 climate signals back in time, particularly because the ring-width correlation converges for the
568 X-ray and anatomy data but dramatically diverges for MXBI (fig. 8). The lower late-period
569 (1949–1994) signal of the anatomical parameters compared to X-ray MXD requires a different
570 explanation (fig. 6). According to the distribution of the r^2 -values in the resampling scheme of

571 [figure 6c](#), the late period Tmax signals are not appreciably different, so perhaps this is simply
 572 by chance compounded by having five times higher X-ray MXD replication.



573
 574 **Figure 8:** A) running Pearson's correlation (a 50-year window shifted by one year) between
 575 selected density parameters and ring width. The years on the x-axis show the first year of the
 576 50-year correlation windows. Note that for X-ray MXD, the ring-width data are not obtained
 577 from the same tree cores as have been used for the density measurements, which is otherwise
 578 the case for both MXBI and anatomy. B) running average of absolute ring widths (original and
 579 ROXAS datasets) computed using a 50-year window shifted by one year, together with the
 580 chronology sample depths of the X-ray MXD, MXBI and dendroanatomical datasets.

581

582 Concluding remarks

583 Tree-ring based reconstructions of pre-industrial climate provide a key insight into Earth's
 584 present and future changing climate, yet their full potential will remain unexploited without a
 585 concerted effort to overcome several critical challenges. This study is part of a larger ongoing
 586 synergetic effort (e.g., Björklund et al. 2020, and other work currently in preparation) directed
 587 at exploring the efficacy of highly temperature sensitive tree-ring data frequently used in large
 588 scale temperature reconstructions (e.g., Wilson et al. 2016), with the ambition to improve upon
 589 these existing records using dendroanatomical techniques. This is because dendroanatomy
 590 represents the direct morphological refinement of current microdensitometric techniques

591 where it is possible to control within-ring specific location of the measurements down to the
592 cellular level (von Arx and Carrer 2014).

593

594 In summary, based on the collective comparison between the new wood anatomical dataset
595 of *P. engelmannii* from the Columbia Icefield and the two predecessors X-ray MXD and MXBI,
596 we are able to draw the following conclusions:

597 1. Maximum radial cell wall thickness and anatomical MXD are the two most promising
598 wood anatomical proxy parameters for estimating past temperatures, each explaining
599 >45% in instrumental detrended July-August maximum temperatures. Both
600 parameters display a comparable climatic imprint and strength of signal to the X-ray
601 derived MXD. It does, however, appear that the stability of the temperature signal over
602 time is more robust for the maximum radial cell wall thickness than for X-ray MXD.

603 2. For these anatomical parameters, the number of trees needed to reach the commonly
604 accepted quality threshold for chronologies used in dendroclimatic analyses is, for our
605 experimental site and species, exemplary with just six trees. However, this high
606 common signal strength is matched by the X-ray MXD parameter and thus does not
607 constitute an obvious advantage by itself. Nevertheless, if the temperature signal is
608 more stable in maximum radial cell wall thickness, it is advantageous to know that very
609 few trees are needed to reach chronology confidence. This is especially true given that
610 the problem of fading records, i.e. the general decrease in sample replication and
611 between tree correlations back in time (Esper and Büntgen 2021), poses a severe
612 constraint to almost all chronologies extending up to or beyond the last millennium.

613 3. The higher resolution of dendroanatomy appears to positively influence the high-
614 frequency temperature signal stability. Using anatomical parameters as opposed to
615 density parameters, be it from X-ray or anatomy, may also be beneficial for data quality
616 and the mechanistic interpretation of the proxy record. However, further research is
617 needed to consolidate this and other important potential effects regarding the low
618 frequency fidelity of long-term temperature reconstructions based on X-ray
619 densitometry.

620

621 Justification of the cost and time constraints currently associated to the production of long
622 dendroanatomical datasets requires that there must be an information gain not obtainable
623 from conventional techniques. In fact, high-resolution, cell-based, measurements already offer
624 an advantage when it comes to the understanding of the structure – function relationships
625 (e.g., Bouche et al. 2014; Pittermann et al. 2011; Wilkinson et al. 2015), the complex
626 mechanisms behind tree-ring formation (Rathgeber et al. 2016), with relative timestamps
627 (Ziaco 2020) of brief intra-seasonal climate extremes, such as late growing season cold spells

628 or initiation of volcanic cooling episodes (Edwards et al. 2022; Piermattei et al. 2020). The
629 question remains, however, whether dendroanatomy can also provide additional paleoclimate
630 information. Despite the encouraging results detailed herein, it is necessary to continue to
631 extend this dataset by adding more series from multiple age classes across the last millennium
632 to more thoroughly evaluate the multi-centennial to millennial scale variations of this key
633 temperature proxy site. The work detailed here is the first piece of a puzzle to explore
634 dendroanatomy of the *P. engelmannii* sample set for the Columbia Icefield area in Canada,
635 formerly analyzed with X-ray and BI techniques (Luckman and Wilson 2005). As such, it also
636 represents the longest (1585–2014 CE) dendroanatomical dataset currently developed for
637 North America.

638

639 **Author contributions**

640 KS and JB conceptualized the research and obtained the funding to support it. MF performed
641 the dendroanatomical measurements, using wood material collected by BL and RW. GvA
642 aided the interpretation of the dendroanatomical data, and MR of the BI-measurements. KS
643 carried out the analysis and drafted the paper. All authors contributed to the planning and
644 structuring of the paper.

645

646 **Data availability**

647 The dendroanatomical chronologies from the Icefield area, Canada, will be available on
648 request.

649

650 **Competing interests**

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

652

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660

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