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Article

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 gradient

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43 ABSTRACT

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45 Mature berries of Pinot Noir grapevines were sampled across a latitudinal gradient in Europe, from southern Spain to central Germany. Our aim was to study the influence of 46 latitude-dependent environmental factors on the metabolite composition (mainly 47 phenolic compounds) of berry skins. Solar radiation variables were positively correlated 48 with flavonols and flavanonols and, to a lesser extent, with stilbenes and cinnamic 49 acids. The daily means of global and erythematic UV solar radiation over long periods 50 (bud break-veraison, bud break-harvest and veraison-harvest), and the doses and daily 51 52 means in shorter development periods (5-10 days before veraison and harvest) were the variables best correlated with the phenolic profile. The ratio between trihydroxylated 53 and monohydroxylated flavonols, which was positively correlated with antioxidant 54 55 capacity, was the berry skin variable best correlated with those radiation variables. Total 56 flavanols and total anthocyanins did not show any correlation with radiation variables. Air temperature, degree days, rainfall and aridity indices showed fewer correlations 57 with metabolite contents than radiation. Moreover, the latter correlations were restricted 58 to the period veraison-harvest, where radiation, temperature and water availability 59 60 variables were correlated, making it difficult to separate the possible individual effects 61 of each type of variable. The data show that managing environmental factors, in particular global and UV radiation, through cultural practices during specific 62 63 development periods, can be useful to promote the synthesis of valuable nutraceuticals and metabolites that influence wine quality. 64

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66 Keywords: Vitis vinifera cv. Pinot Noir, latitudinal gradient, phenolic composition,

- 67 *berry skins, solar radiation, ultraviolet radiation, hydroxylation ratios, Europe*
- 68

69 INTRODUCTION

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Environmental factors, such as air temperature, ambient solar radiation (including UV) 71 72 and photoperiod, vary with latitude. In turn, variations in these environmental factors 73 may cause changes in physiological and/or biochemical characteristics of plants. Yet, this is not always the case as plant responses to latitudinal climatic conditions may be 74 75 masked by, for example, local climatic factors, cultivational measures, or pest and diseases. Thus, there is a need for latitudinal studies that help to identify the 76 77 environmental factors that impact most on plants, as well as the traits most affected. Such studies are important in terms of understanding ecological processes (especially in 78 79 the context of climate change), but also have a direct relevance for the agricultural 80 industry. A number of plant traits have been studied in relation to latitude, including 81 plant height, seed production, growth, biomass production, photosynthesis rates, 82 chlorophyll fluorescence, photosynthetic pigment composition, mineral nutrient contents and ratios, water relations and secondary metabolite contents.¹⁻⁸ Most of these 83 traits have been measured in leaves, whereas only a few studies have used fruits. 84 Latitude-related environmental variables that have been hypothesized to explain 85 86 changes in plant traits include air temperature, degree days, rainfall, aridity indices, soil moisture, total solar radiation doses, and UV radiation doses. Most latitudinal studies 87 have been carried out using wild species, while only a few studies have dealt with 88 commercially interesting species, such as juniper,³ ryegrass⁷ and currant.⁸ To our 89 knowledge, no study has dissected the effects of latitudinal gradients, and the associated 90 environmental parameters, on grapevine, although latitude is a recognized factor used, 91 for example, to predict the suitability of territories for grapevine culture.⁹ 92

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94 Remarkably, the effects of latitude and associated environmental parameters on the phenolic composition of grapevine berries have not been studied, in spite of the fact that 95 similar studies have been conducted on other species with less commercial impact.^{3-5,7,8} 96 97 This omission is even more remarkable, given that the phenolic compounds synthesized 98 in grapevine berries decisively determine wine characteristics and quality, including the presence of important nutraceuticals and nutritionally-desirable antioxidants.^{10,11} Berry 99 skin is the main source of many of these phenolic compounds, including anthocyanins, 100 flavonols and stilbenes.12-14 101

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The present study was conducted on Pinot Noir grapevines. This variety is the tenth 103 104 most cultivated grapevine worldwide, and the seventh fastest-expanding winegrape variety in the period 2000-2010.¹⁵ Pinot Noir grapevines occupy more than 86,000 ha in 105 106 the world (1.88% of the total grapevine acreage), especially in Europe, where it 107 occupies 3% of the total acreage. Pinot Noir is especially adapted to cold climates, thus ascending to higher latitudes than other varieties. In fact, the European distribution of 108 this cultivar ranges from southern Spain to central Germany. Given this wide ranging 109 distribution, our aim was to identify the influence of latitude and associated 110 environmental parameters (air temperature, global and UV radiation, rainfall and 111 aridity) on the metabolite composition of berry skins of Vitis vinifera cv. Pinot Noir in 112 Europe. This study will inform management of those environmental parameters that 113 affect berry skin composition. In turn, a better understanding of the influence of these 114 115 parameters can help improve wine quality.

116 MATERIALS AND METHODS

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118 Collection sites and environmental variables

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Berries of Pinot Noir grapevines (Vitis vinifera L.) were collected in 2013 from 11 120 121 localities in Spain, France, Italy, Hungary, Austria, Slovenia, the Czech Republic and Germany (Figure 1, Table 1). This represented a latitudinal gradient of almost 14° (36.7-122 50.0 °N) and a linear distance of around 1,500 km, covering most of the commercial 123 Pinot Noir growing latitudes in the Northern Hemisphere (35-55°).¹⁶ Vinevard age 124 varied between 6 and 30 years, and vineyard soils were mostly calcareous and neutral-125 alkaline (pH between 7.0 and 8.5). No fertilization or irrigation had been applied to the 126 vineyards. 127

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129 In each locality, berry samples were collected from three separate plants (replicates) at commercial maturity, always around noon-time, and on a sunny day. Collection dates 130 varied from 31 July to 22 October, depending on the location. Three clusters were 131 collected for each replicate. As row orientation varied between vineyards, clusters were 132 133 always picked from a SE-orientated shoot. In situ, every berry was separated from its cluster by cutting the pedicel. Subsequently, berry density was determined as 134 floatability in a NaCl solution series, which allowed for harvesting berries of a similar 135 ripeness using a non-destructive method.^{17,18} To reduce the variability that is normally 136 found within a cluster, berries with a density between 140-160 g NaCl 1⁻¹ were selected, 137 rinsed in distilled H₂O and immediately transported to the laboratory in a portable 138 139 icebox. In the laboratory, berries were frozen in liquid nitrogen and kept at -80°C until 140 further analyses.

141

142 Relevant environmental data were obtained for each locality. Daily values of mean 143 temperature, rainfall and ground-station global radiation (GGR) were obtained for the 144 period bud break-harvest from the nearest meteorological observatory to each vineyard. 145 For most vineyards, meteorological stations were located less than 200 m from the actual vineyards. Remaining stations were located less than 20 km away, except in the 146 case of Lednice (Czech Republic) where the station for GGR measurement was located 147 50 km from the vineyard. In the latter cases, it was ascertained that meteorological 148 stations were located at a similar latitude and altitude as the respective vineyards, which 149

makes the assumption that data were homogeneous. Based on these data, two aridity 150 indices were calculated: the ratio Rainfall/ETP, where ETP is the potential 151 152 evapotranspiration computed according to Hargraves formula (based on solar global 153 radiation and mean air temperature), and the Gaussen Index (the ratio between rainfall and twice the mean daily temperature). In addition, daily values of DSSF (Downward 154 155 Surface Shortwave Flux) global radiation and TEMIS-derived erythematic UV radiation (T UVery) were obtained for the period bud break-harvest. Daily DSSF was calculated 156 by integrating the 30 minutes of data downloaded from the LandSaf web page 157 (http://landsaf.meteo.pt). The data in this archive take into account the differences in the 158 159 day-length of the various locations. T UVery was downloaded from the ESA-TEMIS web page (http://www.temis.nl) and estimated on the basis of Meteosat data (to assess 160 cloud cover), SCIAMACHY data (to assess O_3 column) and a radiative transfer 161 model.¹⁹ The degree days (using 10°C as base temperature) and the daily doses of GGR, 162 163 DSSF and T UVery were integrated over three different periods: bud break-veraison, bud break-harvest, and veraison-harvest. Additionally, DSSF and T UVery doses were 164 integrated for 5 and 10 days before veraison, and for 5 and 10 days before harvest, 165 because the periods around veraison and prior to harvest are important for the synthesis 166 of phenolic compounds in grapevine berries and, thus, for their commercial quality.²⁰⁻²² 167

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169 Analysis of berries

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Frozen berries were allowed to partially thaw and the skin was carefully removed from the flesh using a scalpel, and without rupturing the hypodermal cells. The content of total soluble solids (TSS) was measured in ^oBrix in the flesh, using a digital refractometer. The skins were immediately submerged in liquid nitrogen, weighed and lyophilized. Lyophilized berry skins were weighed and ground to obtain a homogeneous powder for each replicate. Then, all the samples were shipped to one laboratory for detailed analysis of metabolites.

178

For each analytical sample used for the analysis of phenolic compounds, 50 mg of skin
powder was frozen in liquid nitrogen and ground again in a TissueLyser (Qiagen,
Hilden, Germany). The total content of methanol-soluble phenolic compounds
(MSPCs), mainly located in the vacuoles,¹⁸ was measured by spectrophotometry. For
this analysis, 2 ml of a mixture of methanol: water: 7M HCl (70:29:1 v:v:v) was added

for extraction (24 h at 4°C in the dark). The extract was centrifuged at 6000 g for 15 min 184 185 and the supernatant was selected for spectrophotometry. The level of MSPCs was 186 measured as the area under the absorbance curve in the wavelength intervals between 280-315 and 280-400 nm (AUC₂₈₀₋₃₁₅ and AUC₂₈₀₋₄₀₀ respectively) and normalised per 187 unit of dry weight (DW),²³ using a λ 35 spectrophotometer (Perkin-Elmer, Wilton, CT, 188 USA). Individual phenolic compounds were analysed by ultra-performance liquid 189 chromatography (UPLC) using a Waters Acquity UPLC system (Waters Corporation, 190 Milford, MA, USA).²³ Solvents were: A, water/formic acid (0.1%), and B, acetonitrile 191 with 0.1% formic acid. The gradient program employed was: 0-7 min, 99.5-80% A; 7-9 192 min, 80-50% A; 9-11.7 min, 50-0% A; 11.7-15 min, 0-99.5% A. The UPLC system was 193 coupled to a micrOTOF II high-resolution mass spectrometer (Bruker Daltonics, 194 Bremen, Germany) equipped with an Apollo II ESI/APCI multimode source and 195 196 controlled by the Bruker Daltonics DataAnalysis software. The electrospray source was operated in positive or negative mode. The capillary potential was set to 4 kV; the 197 drying gas temperature was 200 °C and its flow 9 1 min⁻¹; the nebulizer gas was set to 198 3.5 bar and 25 °C. Spectra were acquired between m/z 120 and 1505 in positive mode 199 for anthocyanins and in negative mode for the remaining phenolic compounds. The 200 201 different phenolic compounds analysed were identified according to their order of 202 elution and the retention times of the following pure compounds: myricetin, quercetin, catechin, epicatechin, astilbin, trans-resveratrol, p-coumaric acid, caffeic acid and 203 204 ferulic acid (Sigma, St. Louis, MO, USA); kaempferol-3-O-glucoside, isorhamnetin-3-O-glucoside, syringetin-3-O-glucoside, procyanidin B1 and malvidin-3-O-glucoside 205 (Extrasynthese, Genay, France); isorhamnetin, quercetin-3-O-glucoside, quercetin-3-O-206 207 galactoside, quercetin-3-O-glucopyranoside, quercetin-3-O-glucuronide and quercetin-208 3-rutinoside (Fluka, Buchs, Germany). Quantification of compounds that were not 209 commercially available was carried out using the calibration curves belonging to the most similar compound: myricetin for its glucosides; isorhamnetin for isorhamnetin-3-210 211 O-glucuronide; quercetin for quercetin-3-O-arabinoside; astilbin for taxifolin-3-Oglucoside; trans-resveratrol for its glucoside; p-coumaric acid for p-coumaroyl-tartaric 212 213 acid; caffeic acid for p-caffeoyl-tartaric acid; ferulic acid for feruloyl-tartaric acid; and malvidin-3-O-glucoside for anthocyanins. Total contents of the different phenolic 214 groups were obtained as the sum of the individual compounds. The ratios between 215 trihydroxylated and dihydroxylated (3',4',5'-OH/3',4'-OH) anthocyanins, and between 216

trihydroxylated and monohydroxylated (3',4',5'-OH/4'-OH) and trihydroxylated and
dihydroxylated (3',4',5'-OH/3',4'-OH) flavonols, were also calculated.

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For carotenoid and chlorophyll extraction,²⁴ 6 ml of a mixture of methanol, acetone, and 220 hexane (1:1:1 v:v:v) was added to a glass tube containing 50 mg of lyophilized skin 221 powder. The mixture was vortexed for 30 s and then stirred for 30 min at 4°C in the 222 dark. After the addition of 2 ml of MilliQ water the tube was vigorously shaken for 1 223 min and then centrifuged for 1 min at 1500 g. The non-polar phase containing 224 225 carotenoids and chlorophylls was recovered. The extraction was repeated by adding 2 ml of hexane to the remaining mixture. The two extracts were pooled and the volume 226 reduced to 1 ml by vacuum evaporation. The extract was filtered through 0.2-µm filters 227 and immediately subjected to high-performance liquid chromatography (HPLC) 228 229 analysis as follows. Separation was performed at room temperature by a Spectra System 230 P4000 HPLC, equipped with a UV 6000 LP photodiode array detector (Thermo Fisher Scientific, Waltham, MA, USA) using a Zorbax ODS column (5 µm particle size, 250 x 231 4.6 mm, Agilent Technologies, Santa Clara, CA, USA). HPLC separation was carried 232 out at a flow rate of 0.8 ml min⁻¹ using the following linear gradient: 0 min, 82% A 233 (CH₃CN), 18% B (methanol/hexane/CH₂Cl₂ 1:1:1 v:v:v); 20 min, 76% A, 24% B; 30 234 min, 58% A, 42% B; 40 min, 39% A, 61% B. The column was allowed to re-equilibrate 235 in the starting solution (82% A, 18% B) for 5 min before the next injection. Different 236 237 individual chlorophylls and carotenoids were detected by their absorbance at 445 nm.

238

The antioxidant capacity of berry skins was measured by generating the radical cation 239 2.2'-azinobis(3-ethylbenzothiazoline-6-sulfonic acid) (ABTS⁺⁺).²⁵ The radical solution 240 was diluted in ethanol to obtain an absorbance of 0.700 ± 0.020 at 734 nm (Perkin-241 Elmer λ 35 spectrophotometer). After addition of 1 ml of diluted ABTS⁺⁺ solution to 100 242 μl of skin extract (250 μg of skin powder in 1 ml of a mixture of methanol: water: 7M 243 244 HCl 70:29:1 v:v:v), the decrease in absorbance was monitored and compared to that of 245 the Trolox standard (Sigma) exactly 4 min after initial mixing. Antioxidant capacity was 246 expressed in terms of Trolox equivalent antioxidant capacity (TEAC) per g DW of skin. 247

DNA isolation from lyophilized berry skins was carried out using the ZenoGene40
Plant DNA Purifying Kit (Zenon Bio Kft., Szeged, Hungary). Concentration of the
samples was measured with a Genova Nano Spectrophotometer (Jenway, Staffordshire,

UK). DNA content per DW of berry skin (ng mg⁻¹ DW) was calculated using the formula: mean of DNA concentration (ng μ l⁻¹) multiplied by the volume of extraction (μ l) and divided by the DW of the lyophilized sample (mg). This analysis served to calculate the metabolite concentrations on a DNA basis.

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256 Statistical analysis

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Pearson correlation coefficients (r) were used to examine the relationships between all 258 259 the variables studied, both the environmental-geographical parameters and the traits analyzed in berry skins, including the total contents of the different groups of phenolic 260 compounds. Correlations were considered significant when p < 0.05. The sampling 261 localities were ordinated by Principal Components Analysis (PCA), taking into account 262 MSPCs and the total contents of the different groups of phenolic compounds. All the 263 statistical procedures were performed with SPSS 19.0 for Windows (SPSS Inc., 264 Chicago, IL, USA). 265

266 **RESULTS**

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268 Variation in environmental variables

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270 The latitudinal gradient used in this study was associated with substantial differences in several meteorological variables (Table 2). For the period from bud break to harvest, 271 these differences were, amongst others, around 5°C in mean daily temperature, 500 272 degree days, almost 300 mm in rainfall, almost 900 MJ m⁻² in DSSF dose, and 241 kJ 273 m^{-2} in T UVery dose. Interestingly, the parameters displaying the greatest differences 274 were the DSSF and T UVery doses accumulated during the 10 days before harvest. For 275 276 these variables, the differences between the maximum and the minimum values along the gradient were more than 80% of the maximum value. The highest and lowest values 277 278 of temperature variables were usually recorded in Pécs and Rioja, respectively, except 279 for the veraison-harvest period, in which they were recorded in Spanish localities (Jerez or Girona) and Lednice, respectively. The highest mean values of solar radiation (GGR, 280 DSSF, T UVery) were always recorded in Jerez, and this included also the highest 281 accumulated doses in the 5 or 10 days before veraison and before harvest. The highest 282 283 accumulated doses over longer periods were recorded in Spanish localities (either Rioja, Girona or more rarely Jerez) or in Lednice, depending on the length of the period 284 considered, because those periods were longer in Rioja, Girona or Lednice than in Jerez 285 286 (see Table 1 for the length of the period bud break-harvest). The lowest values of radiation variables were generally recorded in Geisenheim or Lednice. 287

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289 Variation in berries variables

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291 Metabolite contents were obtained and normalized against both berry skin DW (Table 3) and DNA amount. The correlations between metabolites and environmental 292 293 parameters were similar irrespective of the normalization approach, given that DNA amount and berry skin DW were significantly correlated (r = 0.79, p < 0.01, n = 11). 294 Therefore, results are only described on a per berry skin DW basis. MSPC values varied 295 between 9.7 and 40.3 (as AUC₂₈₀₋₃₁₅ mg⁻¹ DW) and between 17.1 and 74.3 (as AUC₂₈₀₋ 296 ₄₀₀ mg⁻¹ DW). Absorption levels in the two wavelength regions were strongly and 297 positively correlated (Table S1). The highest and lowest MSPC values were found in 298 Girona and Lednice, respectively. We quantified 29 phenolic compounds: 24 flavonoids 299

(14 flavonols, 5 anthocyanins, 3 flavanols –monomeric or dimeric tannins-, and 2 300 301 flavanonols) and 5 non-flavonoids (3 cinnamic acids and 2 stilbenes). Great differences 302 in the concentrations of most groups of phenolic compounds were found between 303 localities. Anthocyanins were the most abundant group, showing values between 18.9 (Bilje) and 110.1 (Girona) mg g⁻¹ DW. In every locality, malvidin-3-O-glucoside was 304 the major anthocyanin. Flavonols were the second most abundant group of flavonoids, 305 ranging between 1.76 (Bilje) and 7.7 (Girona) mg g⁻¹ DW. The major flavonol was 306 quercetin 3-O-glucuronide. Flavanonols (between 0.18 and 1.14 mg g⁻¹ DW, in Bilje 307 and Jerez, respectively) and flavanols (between 0.21 and 0.99 mg g⁻¹ DW, in Lednice 308 and Bilje, respectively) were less abundant. Among non-flavonoids, cinnamic acids 309 were the most abundant group, and also the group showing the greatest variability 310 between localities, with values between 0.16 (Lednice) and 7.2 (Firenze) mg g^{-1} DW. 311 Finally, the least abundant compounds were stilbenes, which also showed a great 312 variability (between 14 and 928 μ g g⁻¹ DW, in Lednice and Girona, respectively). 313

314

The antioxidant capacity of berry skin extracts varied between 3592 (Lednice) and 9104 (Firenze) μ M TE g⁻¹ DW. Chlorophylls and all carotenoids showed the highest values in Rioja and the lowest in Pécs. β -Carotene was the most abundant carotenoid. The berry fresh weight varied between 1.1 (Girona and Bordeaux) and 2.1 g (Geisenheim), although most localities showed values between 1.1 and 1.3 g. TSS varied between 19.1 (Bilje) and 23.7 °Brix (Jerez).

321

322 Correlations between variables

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The correlations between all the environmental and plant response variables were 324 325 determined (Table S1). Unless otherwise stated, the correlations mentioned in this text were significant (p < 0.05) and positive. With respect to the correlations between berry 326 327 skin variables, MSPCs were correlated with the contents of most phenolic compounds 328 (except flavanols) and carotenoids. The total contents of flavonols, flavanonols, 329 stilbenes and anthocyanins were correlated with one another, whereas the total content of cinnamic acids was only correlated with that of flavanonols. Total flavanol content 330 was not correlated with the total content of any other phenolic group. The antioxidant 331 capacity of berry skin extracts was correlated with anthocyanins, MSPCs, flavonols, the 332 ratio 3',4',5'-OH/3',4'-OH flavonols and, less significantly, with flavanonols, cinnamic 333

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acids, the ratio 3',4',5'-OH/4'-OH flavonols, and carotenoids. There was no correlation between the antioxidant capacity and contents of stilbenes or flavanols. Carotenoid and chlorophyll contents were correlated with each other, and carotenoid levels were also correlated with those of stilbenes.

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Possible correlations between environmental-geographical parameters and berry skin variables were also explored. It was found that latitude was negatively correlated with MSPCs and the total contents of flavonols, flavanonols and stilbenes, but not flavanols, cinnamic acids, anthocyanins and carotenoids (Figure 2).

343

Correlations between temperature variables and berry variables were few for the periods 344 bud break-veraison and bud break-harvest. The mean daily temperature and degree days 345 346 in the period bud break-veraison (but not bud break-harvest) were correlated 347 (negatively) with carotenoids, chlorophylls and TSS, only. Degree days in the period bud break-veraison were also correlated with flavanonols. No temperature variable in 348 these two periods was correlated with the total content of any other phenolic group, 349 although there were some correlations between temperature variables and individual 350 351 compounds. For the period veraison-harvest, the mean daily temperature and degree 352 days were correlated with MSPCs and the total contents of flavonols and flavanonols. In addition, the mean daily temperature was correlated with the ratios 3',4',5'-OH/4'-OH 353 and 3',4',5'-OH/3',4'-OH flavonols, and the degree days with the total content of 354 anthocyanins. 355

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Rainfall and aridity indices were hardly correlated with berry skin variables for the periods bud break-veraison and bud break-harvest. Only quercetin showed somewhat consistent (positive) correlations with rainfall, the Rainfall/ETP ratio and Gaussen Index (but only in the period bud break-harvest). For the period veraison-harvest, rainfall and aridity indices were negatively correlated with the total content of flavonols and flavanonols. In addition, Gaussen index was negatively correlated with MSPCs and the ratios 3',4',5'-OH/4'-OH and 3',4',5'-OH/3',4'-OH flavonols.

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Radiation variables, particularly DSSF and T UVery variables, correlated well with berry skin variables for the three periods considered. The daily means of DSSF and T UVery in the periods bud break-harvest and veraison-harvest, the DSSF doses in the 10

days before harvest, the daily mean of T UVery in the 5 and 10 days before veraison, 368 369 and the T UVery doses in the 5 and 10 days before veraison were all correlated with 370 MSPCs. The same variables, together with the T UVery doses in the 10 days before 371 harvest and in the period bud break-harvest (in this last case, with a lower significance 372 level), were correlated with the total contents of flavonols and flavanonols. Total stilbene content was only correlated with the DSSF and T UVery doses in the period 373 bud break-harvest, and total cinnamic acid content only with the daily mean and the 374 dose of T UVery in the 10 days before veraison. Total flavanol and anthocyanin 375 contents were not correlated with any radiation variable. Regarding individual 376 377 compounds, the strongest correlations were found between contents of several flavonols and flavanonols and the daily means of DSSF and T UVery in the periods bud break-378 379 harvest and veraison-harvest, as well as with the DSSF and T UVery doses in the 380 periods of 5 or 10 days before veraison or harvest. Levels of two flavanols, one 381 anthocyanin and the three cinnamic acids analyzed were also correlated with some of those T UVery expressions. 382

383

The ratio 3',4',5'-OH/3',4'-OH anthocyanins was not correlated with any radiation or 384 temperature variable. Yet, the ratios 3',4',5'-OH/4'-OH and 3',4',5'-OH/3',4'-OH 385 386 flavonols were the berry skin variables that displayed the strongest correlations with 387 specific radiation variables, such as the daily means of DSSF and T UVery in the 388 periods bud break-harvest and veraison-harvest, and the accumulated doses in the 10 days before veraison and harvest. This correlation did, however, not extend to the 389 390 accumulated doses in longer periods, as Figure 3 shows for the period bud break-391 harvest. Finally, the number of days from bud break to harvest and from veraison to 392 harvest were negatively correlated with total and several individual flavanols.

393

394 Principal Components Analysis

395

The localities studied were ordinated by PCA using MSPCs and the different groups of phenolic compounds. The accumulated variance by the first three axes was 94.0% (67.3% for axis I, 17.3% for axis II and 9.4% for axis III). The plot using the first two axes, together with the loading factors and their significance, is shown in Figure 4. The total contents of all the phenolic groups, except flavanols, were significant loading factors for the positive part of axis I, which broadly ordinated the localities on the basis of their latitude, with southernmost localities situated towards the positive part of the
axis and the northernmost ones towards the negative part. Total flavanols and total
cinnamic acids were the only significant loading factors for the positive part of axis II,
which separated localities 4, 6, 9, 7 and 1 from the remaining ones. No significant
loading factor was found for the negative part of axes I and II.

407 **DISCUSSION**

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Environmental-geographical gradients, such as those related to latitude, can be exploited 409 410 to explore and predict the physiological and/or biochemical responses of plants by using a space-for-time substitution.⁶ This type of study cannot necessarily pinpoint the 411 influence of one particular environmental parameter on a plant response, as can be done 412 in controlled studies. However, the strength of latitudinal studies is that plant responses 413 are studied under realistic conditions (i.e. commercial vineyards), where plants are 414 exposed to a natural combination of ambient, environmental parameters. In this study a 415 range of metabolites were measured in skins of Pinot Noir berries, originating from 11 416 vineyards along a latitudinal gradient of nearly 14°. The levels of the various 417 metabolites measured in Pinot Noir berry skins were broadly in agreement with levels 418 measured in other studies using this, or other cultivars.^{12,18,23} 419

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421 Radiation is an important determinant of berry skin metabolite profile

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A key finding of this study is that the contents of MSPCs, flavonols, flavonols and 423 stilbenes in Pinot Noir berry skins increased with decreasing latitudes. Previously, 424 similar results were found for MSPC contents in leaves of *Lolium perenne*⁷, but no 425 comparative results existed for specific phenolic compounds nor for grapevine. It might 426 427 be argued that negative correlations between latitude and the abovementioned phenolic groups are due to the longer berry maturation period at lower latitudes. However, we 428 429 consider this unlikely because (1) latitude was not significantly correlated with the 430 number of days from veraison to harvest, and (2) the latter variable was not correlated with the contents of those phenolic compounds. Rather, the correlations between 431 432 latitude and contents of phenolic compounds were probably due to the negative correlation between latitude and radiation (both global and UV) variables. Radiation 433 434 variables were strongly and positively correlated with the total contents of most 435 phenolic groups, mainly flavonols and flavanonols, and to a lesser extent with stilbenes 436 and cinnamic acids, together with MSPCs. The relationship between radiation levels and the content of these phenolic compounds had previously been reported for berry 437 skins of several red grapevine varieties, such as Pinot Noir, Merlot, Malbec and 438 Cabernet Sauvignon,²⁶⁻²⁹ although not in relation with latitudinal gradients. 439

440

Rather than radiation in general, the means of DSSF and T UVery over long periods 441 (bud break-veraison, bud break-harvest and veraison-harvest) and the means or doses in 442 important development periods (5-10 days before veraison and harvest) were the 443 variables best correlated with phenolic compounds, particularly flavonols, flavanonols 444 and cinnamic acids. This is related to the fact that the periods around veraison and prior 445 to harvest are important for the synthesis of phenolic compounds.²⁰⁻²² The stimulation of 446 flavonol acumulation was expected because these compounds are radiation-reactive and 447 concentrations are well known to increase with increasing levels of solar radiation 448 (particularly UV-B) in grapevine berry skins.^{13,18,27,29-33} 449

450

It is not simply total flavonol levels that correlate with radiation parameters, the ratios 451 3',4',5'-OH/4'-OH and 3',4',5'-OH/3',4'-OH flavonols were the berry skin variables 452 best correlated with specific radiation variables, such as the mean values or doses of 453 454 DSSF and T UVery radiation in critical periods (5-10 days before veraison and harvest), but not with the accumulated doses over long periods (Figure 3). Thus, higher solar 455 radiation values (both total and UV) in those critical periods might increase the B-ring 456 hydroxylation level of flavonols in Pinot Noir berry skins. Previously, it was shown that 457 the hydroxylation level depends on both the grape variety¹² and environmental factors, 458 such as the radiation level. The effect of radiation, in turn, may depend again on the 459 variety considered: the hydroxylation ratios increased with increasing total or UV 460 radiation in Pinot Noir (this study), but decreased with increasing total or UV radiation 461 in Sangiovese²² and Tempranillo.^{18,34} This complexity may be caused by the intricate 462 regulation mechanism of the genes and enzymes involved in the synthesis of flavonols 463 with different hydroxylation levels.^{21,30,31} In petunia, the highest level of B-ring 464 hydroxylation was caused by the specific effect of increased UV-B radiation.³⁵ The 465 antioxidant activity of flavonoids strongly depends on the number of hydroxyl groups 466 bound to the aromatic B-ring.³⁶ Given that the hydroxylation ratios were positively 467 468 correlated with the antioxidant capacity in our study, flavonols may be important as both sunscreens and antioxidants in Pinot Noir berry skins, and their role as antioxidants 469 470 would increase in those localities with higher radiation levels.

471

Flavanonols (dihydroflavonols) are bioactive compounds that contribute to tolerance to
fungal infections and colour expression in some red wines.³⁷ Given that flavanonols
comprise a relatively small fraction of total wine flavonoids, their regulation by, and

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475 responses to, radiation were not clear. However, the results in this paper show that 476 flavanonol levels were positively correlated with radiation. This observation is 477 consistent with a previous study that reported increases in flavanonols in Malbec berry 478 skins following exposure to higher solar radiation levels due to cluster thinning.³⁷ 479 Similarly, flavanonol levels were found to be elevated in berries exposed to ambient 480 UV-B, in comparison with berries receiving no UV-B.¹³

481

The reported data indicate positive correlations of cinnamic acid levels with radiation. 482 Consistently, higher values of caffeoyl-tartaric acid were found in skins of Pinot Noir 483 berries exposed to solar radiation when compared with shaded berries.²⁶ However, not 484 all studies show increases in cinnamic acids with increasing radiation. Coumaroyl-485 tartaric acid levels showed no response to solar UV-B radiation exposure in Malbec 486 berry skins.²⁸ Probably, the synthesis of cinnamic acids in berries is more influenced by 487 the radiation received prior to veraison, because contents are highest before berry 488 maturation.¹⁴ Besides, there is some debate on whether cinnamic acids are 489 predominantly present in pulp, rather than skin. Furthermore, the response of cinnamic 490 acid levels to variations in radiation appears to be influenced by the specific year,³⁸ and 491 each specific cinnamic acid seems to react in a different way.¹⁸ 492

493

In contrast to flavonol and flavonool content, the levels of total stilbenes were only 494 correlated with the global and UV radiation doses over long periods (bud break-495 496 harvest). Both stilbenes and flavonoids derive from coumaroyl-coenzyme A in the 497 general phenylpropanoid metabolism, but stilbenes are synthesized by stilbene synthase instead of chalcone synthase. Stilbene synthase is found in berry skins during all stages 498 of fruit development,³⁹ which could explain the correlation of total stilbene contents 499 with global and UV doses over long periods. Yet, similar to flavonols, stilbenes 500 (resveratrol) were also found to be UV-induced, as was demonstrated in studies using 501 Malbec berry skins.²⁸ 502

503

It was found in this study that the total content of anthocyanins was not correlated with any radiation variable. This finding is congruent with previous findings on Pinot Noir berry skins, which showed that anthocyanin content was not affected by sun exposure.²⁶ The finding is also consistent with the fact that anthocyanin biosynthesis is controlled by a different system than that controlling flavonol biosynthesis.⁴⁰ In general,

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anthocyanins are accumulated under conditions of low temperature and high radiation
levels,^{8,41} but contradictory data have been reported in grape berries as a consequence of
differences in cultivar, site, season, sampling and analytical techniques.⁴² In addition, it
has often been difficult to separate the effects of light and temperature.

513

The ratio 3',4',5'-OH/3',4'-OH anthocyanins was also not correlated with any radiation 514 variable (unlike the hydroxylation ratio of flavonols). Previous studies had shown that 515 the hydroxylation ratio of anthocyanins may increase⁴³ or decrease^{31,44} with increasing 516 (total or UV) radiation in different grapevine varieties, and even the responses may vary 517 depending on the year of study.^{27,30} These diverse responses to radiation may be due not 518 only to a complex regulation of the synthesis of differently hydroxylated anthocyanins 519 520 in the different varieties (as occurred with respect to the hydroxylation ratios of 521 flavonols), but also to the specific responses of each individual anthocyanin. For 522 example, in our study the trisubstituted malvidin-3-O-glucoside was the only anthocyanin (positively) correlating with radiation variables, thus affecting the response 523 of the ratio to radiation. 524

525

526 Total flavanol levels were not correlated with any radiation variable nor with levels of 527 any other phenolic group. A likely explanation for this observation is that flavanols are synthesized during the early stages of berry development and that their levels remain 528 529 fairly stable during subsequent berry growth. Several authors have reported that flavanol levels are stable, and show little responsiveness to changes in radiation or other 530 environmental parameters.^{14,44,45} Nevertheless, there is no consensus on this point, as 531 solar UV exclusion has been reported to decrease flavanol content,²⁹ and responses to 532 temperature and water availability have also been reported.⁴⁵⁻⁴⁷ 533

534

Thus, it is concluded that radiation is strongly correlated with Pinot Noir berry skin phenolic profile. Radiation-related changes in phenolic profile are highly specific. Radiation appears to affect one class of metabolites, while other compounds are not affected. Such specific regulatory interactions offer scope to precision manipulation of berry skin metabolite profiles, in order to increase berry and wine quality.

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543 Effects of temperature and water supply on berry skin metabolic profile

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Along the latitudinal gradient studied, the effect of temperature on overall phenolic 545 546 composition of Pinot Noir berry skins was weaker than the effect of radiation, because temperature variables were correlated with phenolic composition only when they were 547 calculated for the period veraison-harvest. In this case, MSPCs, flavonols, flavanonols, 548 anthocyanins, and the ratios 3',4',5'-OH/4'-OH and 3',4',5'-OH/3',4'-OH flavonols, 549 were positively correlated with the mean daily temperature and/or degree days. These 550 correlations might be due to the fact that temperature and radiation variables were also 551 correlated for that period (Table S1), and it may be difficult to differentiate radiation 552 and temperature effects.⁴² It may not be surprising that the effects of temperature were 553 more clear in the most important period for berry maturity (veraison-harvest).²⁰ 554 particularly in the case of anthocyanins, which increase strongly in that period.²⁰⁻²² 555 Anthocyanins are known to be influenced by specific temperature conditions, such as 556 ambient temperatures recorded after veraison.^{27,41,47,48} Results are also congruent for 557 flavonols because, although more influenced by radiation, these compounds can also 558 respond to temperature.⁴ Flavanols are known to be influenced by specific temperature 559 conditions, but in this study effects of a limited range of temperatures were tested, and it 560 561 is possible that more extreme temperatures are required to impact on these phenolics. With respect to cinnamic acids, their synthesis in the first stages of berry development 562 and the strong decrease in concentrations after veraison²⁰ may mask the influence of 563 temperature on their content at harvest, thus concealing any correlation between 564 565 temperature parameters and cinnamic acid concentrations.

566

Rainfall and aridity indices showed a similar behavior as temperature variables, and 567 568 were correlated with some phenolic compounds only when the period veraison-harvest was considered. In this period, water availability variables were correlated with 569 570 temperature and radiation variables, and thus the individual effect of each variable could not be differentiated. Water availability typically shows strong relationships with 571 different plant traits,⁴⁹ but direct effects on the contents of grape skin phenolic 572 compounds are considered to be relatively minor.^{50,51} This could be due to the fact that 573 the effects of water availability on berry skin composition are mainly mediated by 574 changes in berry size which subsequently affect the proportion of skin in relation to 575 total berry, or by changes in photosynthesis rates modifying source-sink relationships.⁴² 576

Nevertheless, changes in anthocyanins, flavonols and stilbenes caused by water deficit or excess have been described, sometimes in contradictory ways,^{42,52} and drought conditions have been reported to increase the expression of different genes involved in the biosynthesis of phenolic compounds.^{31,52} Overall, correlations between water availability and phenolic composition were not conclusive in our study.

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584 In summary

585

PCA was used to summarize the results described above. Axis I mostly represented a 586 latitude gradient, and was determined by nearly all different groups of phenolic 587 compounds that are present in berry skins (flavonols, flavanonols, anthocyanins, 588 589 stilbenes and cinnamic acids, together with MSPCs). Thus, Pinot Noir berry skins from 590 southern localities were more enriched in most phenolics than those from northern latitudes. This is congruent with the general variation in phenolic compounds (except 591 anthocyanins) with latitude.⁴ Changes in phenolic composition can influence wine 592 quality and will contribute to wine genuineness in each locality. Given that, in our 593 594 study, latitude was more often correlated with radiation variables than with temperature 595 or water availability variables, radiation appeared to be the most important factor contributing to the differentiation of berry skin composition at the localities studied. 596 597 Nevertheless, in the most important period for phenolic ripeness (veraison-harvest), 598 latitude and radiation, temperature and water availability variables were correlated with 599 one another, and the effect of each type of variable was difficult to separate. Thus, apart 600 from the effect of radiation in every period considered, the interaction of radiation, 601 temperature and water availability in the period veraison-harvest was strongly correlated 602 with the phenolic composition of berry skins along the latitudinal gradient considered. Flavanols and cinnamic acids were the only phenolic compounds that define axis II of 603 604 the PCA, thus contributing to the differentiation of berry skins from some localities, in 605 particular those situated to the positive part of the axis II, such as Bilje, Firenze, Retz, 606 Potoče and Jerez.

607

608 Genetic and environmental factors (other than radiation, temperature and water 609 availability) have not been considered in our study, but may also affect the metabolite 610 composition of berry skins. In particular, a clone effect cannot be excluded. However,

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this effect has been demonstrated to be relatively minor and/or non-significant in 611 previous studies using both Pinot Noir^{48,53} and other grapevine cultivars.⁵⁴ On the other 612 hand, additional environmental factors related to the so-called "terroir" and not analyzed 613 in detail in our study, such as soil type or mineral nutrition, could have influenced 614 metabolites composition,^{54,55} although it is doubtful whether the impacts of such 615 variables would have been correlated with latitude. Overall, in spite of having used 616 different clones, plant ages and soils, a significant relationship between metabolites 617 composition and the latitude-dependent environmental changes in radiation, temperature 618 and water availability was found. It is likely that this environmental influence masked 619 620 the possible effects of genetic factors and other non-considered environmental variables.

Particularly relevant is the finding that skin phenolic composition was correlated with 622 623 the DSSF and T UVery means and doses in relatively short development periods (5-10 624 days before veraison and harvest). Thus, increasing the total and/or UV radiation received by the clusters in those periods through management practices, such as leaf 625 removal or supplemental UV exposure, could promote the synthesis of valuable 626 phenolic metabolites. This may eventually contribute to improved wine quality because 627 628 of the notable contribution of phenolic compounds to wine flavor and also by increasing the amount of nutraceuticals and healthy antioxidants, such as flavonols, flavanonols, 629 stilbenes and cinnamic acids.^{10,11} Among others, UV radiation has been demonstrated to 630 be an important factor correlated with berry skin composition in our study. Although 631 some of the effects observed, such as the increase in MSPCs, flavonols and cinnamic 632 acids, have been repeatedly attributed to UV (particularly UV-B) radiation, 13,18,29,31 633 634 more specific manipulative experiments are needed to prove the specific effects of this fraction of solar radiation across the latitudinal gradient considered. 635

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It is concluded that radiation in several development periods, and an interaction between radiation, temperature and water availability in the period veraison-harvest, were the environmental factors most correlated with the phenolic composition of Pinot Noir berry skins along a latitudinal gradient in Europe. In addition, it was demonstrated that effects of environmental variables may be different for different compounds and that some compounds were more responsive (for example, flavonols) than others (flavanols).

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645	ASSOCIATED CONTENT									
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647	Supporting information									
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649	Table S1. Correlation coefficients among environmental-geographic and berry									
650	variables. Significant correlations are indicated in different colours depending on the									
651	significance level: purple, $p < 0.001$; fuchsia, $p < 0.01$; pink, $p < 0.05$. Bb, bud break; v,									
652	veraison; h, harvest; see the remaining abbreviations in Table 2 and 3 legends.									
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871	FIGURE AND TABLE LEGENDS
872	
873	Figure 1. Geographic location of the 11 European sampling localities used in this study.
874	1, Jerez de la Frontera (Spain); 2, Girona (Spain); 3, La Rioja (Spain); 4, Firenze (Italy);
875	5, Bordeaux (France); 6, Bilje (Slovenia); 7, Potoče (Slovenia); 8, Pécs (Hungary); 9,
876	Retz (Austria); 10, Lednice (Czech Republic); 11, Geisenheim (Germany).
877	
878	Figure 2. Regressions between selected berry variables, including carotenoids and the
879	different groups of phenolic compounds, and latitude. Determination coefficients (R^2)
880	and <i>p</i> values are shown.
881	
882	Figure 3. Regressions between the ratio trihydroxylated / monohydroxylated flavonols
883	and selected radiation variables. DSSF, Downward Surface Shortwave Flux. T UVery,
884	TEMIS-derived erythematic UV. For both variables, the daily mean in the period bud
885	break-harvest, and the accumulated dose in the same period and in the 10 days before
886	harvest, were used for calculations. Determination coefficients (R^2) and p values are
887	shown.
888	
889	Figure 4. Ordination, through Principal Components Analysis (PCA), of the 11
890	sampling localities used in this study, taking into account the total content of methanol-
891	soluble phenolic compounds (MSPC) and the total contents of the different groups of
892	phenolic compounds. Significant loading factors for the positive and negative parts of
893	each axis, together with their corresponding significance levels, are shown (***,
894	p < 0.001; **, $p < 0.01$; *, $p < 0.05$). Axis 1 is the horizontal one, and axis 2 is the vertical
895	one. Each mark on the axes represents 0.5 units.
896	
897	Table 1. Geographic location (latitude, longitude and altitude) of the 11 European
898	sampling localities used in this study, together with the number of days from bud break
899	to harvest.
900	
901	Table 2. Ranges of the environmental variables in the 11 European sampling localities
902	used in this study, together with the localities in which each extreme value was recorded
903	(between brackets). ETP, potential evapotranspiration. GGR, Ground-station Global
904	Radiation. DSSF, Downward Surface Shortwave Flux. T UVery, TEMIS-derived

- 905 erythematic UV. The different variables were calculated along three periods: bud break-
- veraison (white background), bud break-harvest (light grey background) and veraison-
- 907 harvest (dark grey background). In addition, DSSF doses were calculated in the 10 days
- 908 before harvest, and T UVery (mean values and total doses) in different periods.
- 909
- **Table 3.** Values (means \pm SE) of the variables analyzed in Pinot Noir berries in the 11
- 911 European sampling localities used in this study. MSPC, methanol-soluble phenolic
- 912 compounds. AUC, area under curve. TSS, total soluble solids.

	sampling site	country	latitude (°N)	longitude (°E)	altitude (m)	days from bud break to harvest
1	Jerez de la Frontera	Spain	36.7	-6.2	40	141
2	Girona	Spain	41.8	2.6	150	174
3	La Rioja	Spain	42.5	-2.3	342	175
4	Firenze	Italy	43.9	11.2	280	131
5	Bordeaux	France	44.8	-0.6	22	176
6	Bilje	Slovenia	45.9	13.6	70	143
7	Potoče	Slovenia	45.9	13.8	120	140
8	Pécs	Hungary	46.1	18.1	200	152
9	Retz	Austria	48.8	15.9	324	172
10	Lednice	Czech Republic	48.8	16.8	176	183
11	Geisenheim	Germany	50.0	8.0	95	170

Table 1. Geographic location (latitude, longitude and altitude) of the 11 European sampling localities used in this study, together with the number of days from bud break to harvest.

	min	max
mean daily temperature (°C)	16.4 (3)	21.2 (8)
mean daily temperature (°C)	16.6 (10)	21.1 (8)
mean daily temperature (°C)	13.1 (10)	24.4 (1)
degree days (°C)	936 (3)	1367 (8)
degree days (°C)	1197 (3)	1703 (8)
degree days (°C)	113 (10)	381 (2)
rainfall (mm)	155 (4)	439 (5)
rainfall (mm)	196 (4)	481 (5)
rainfall (mm)	0 (1)	103 (10)
rainfall/ETP	0.31 (4)	0.80 (5)
rainfall/ETP	0.28 (1)	0.82 (9)
rainfall/ETP	0 (1)	0.9 (9,10)
Gaussen Index	4.0 (4)	12.8 (5)
Gaussen Index	4.9 (4)	13.7 (5)
Gaussen Index	0 (1)	4.7 (10)
GGR (mean) (MJ m-2 d-1)	12.7 (9)	24.2 (1)
GGR (mean) (MJ m-2 d-1)	11.2 (9)	24.9 (1)
GGR (mean) (MJ m-2 d-1)	8.1 (9)	28.6 (1)
GGR (dose) (MJ m ⁻²)	1487 (9)	3035 (3)
GGR (dose) (MJ m ⁻²)	1939 (9)	3718 (2)
GGR (dose) (MJ m ⁻²)	370 (4)	759 (10)
DSSF (mean) (MJ $m^{-2} d^{-1}$)	18.3 (11)	23.8 (1)
DSSF mean (MJ $m^{-2} d^{-1}$)	15.9 (11)	24.5 (1)
DSSF mean (MJ $m^{-2} d^{-1}$)	10.1 (11)	28.4 (1)
DSSF (dose) (MJ m^{-2})	2201 (11)	2908 (2)
DSSF (dose) (MJ m ⁻²)	2684 (11)	3542 (2)
DSSF (dose) (MJ m ⁻²)	384 (4)	695 (10)
T UVery (mean) (kJ $m^{-2} d^{-1}$)	3.0(11)	3.8(1)
T UVery (mean) (kJ $m^{-2} d^{-1}$)	2.4 (11)	4.0 (1)
T UVery (mean) (kJ $m^{-2} d^{-1}$)	1.5 (11)	4.8 (1)
T UVery (dose) (kJ m ⁻²)	254 (11)	483 (3)
T Uvery (dose) (kJ m ⁻²)	329 (11)	570 (3)
T Uvery (dose) (kJ m ⁻²)	49 (4)	114(1)
DSSF (10-days-before-harvest dose) (MJ m ⁻²)	56.6 (11)	284(1)
T Uvery (5-days-before-veraison mean (kJ m ⁻² d ⁻¹)	2.0 (10,11)	5.1(1)
T Uvery (10-days-before-veraison mean (kJ m ⁻² d ⁻¹)	2.4 (10,11)	5.0 (1)
T Uvery (5-days-before-veraison dose) (kJ m ⁻²)	9.9 (10)	25.3 (1)
T Uvery (10-days-before-veraison dose) (kJ m ⁻²)	23.8 (10)	50.2 (1)
T Uvery (10-days-before-harvest dose) (kJ m ⁻²)	6.9 (11)	47.4 (1)

Table 2. Ranges of the environmental variables in the 11 European sampling localities used in this study, together with the localities in which each extreme value was recorded (between brackets). ETP, potential evapotranspiration. GGR, Ground-station Global Radiation. DSSF, Downward Surface Shortwave Flux. T UVery, TEMIS-derived erythematic UV. The different variables were calculated along three periods: bud break-veraison (white background), bud break-harvest (light grey background) and veraison-harvest (dark grey background). In addition, DSSF doses were calculated in the 10 days before harvest, and T UVery (mean values and total doses) in different periods.

	Jerez	Girona	La Rioja	Firenze	Bordeaux	Bilje	Potoče	Pécs	Retz	Lednice	Geisenheim
total content of MSPC											
AUC ₂₈₀₋₃₁₅ mg ⁻¹ DW	39.1 ± 1.5	40.3 ± 1.2	31.0 ± 3.0	32.3 ± 0.7	22.2 ± 1.3	14.7 ± 0.2	13.2 ± 0.4	32.3 ± 0.2	32.1 ± 5.3	9.7 ± 0.1	24.3 ± 1.2
AUC ₂₈₀₋₄₀₀ mg ⁻¹ DW	71.2 ± 3.5	74.3 ± 3.0	54.5 ± 5.4	58.4 ± 1.7	41.0 ± 2.4	24.5 ± 0.1	22.7 ± 0.4	56.9 ± 0.6	56.1 ± 9.9	17.1 ± 0.4	40.7 ± 2.2
flavonols (µg g ⁻¹ DW)											
myricetin	139± 20	153± 8	112± 24	234± 27	38.7 ± 5.6	7.3± 2.8	13.2 ± 3.4	74.1± 9.2	164± 31	2.5 ± 0.8	15.3± 1.8
myricetin-3-O-glucoside	1066 ± 137	1041 ± 62	864 ± 86	918 ± 112	487 ± 37	157 ± 17	277 ± 45	473 ± 38	535 ± 92	61.2 ± 16.1	272 ± 30
myricetin-3-O-glucuronide	391 ± 50	355 ± 54	183 ± 32	368 ± 21	117 ± 11	62.5 ± 6.8	86.1 ± 7.5	267 ± 23	68.5 ± 9.1	22.2 ± 6.0	47.4 ± 8.6
kaempferol-3-O-glucoside	177 ± 37	273 ± 61	78.5 ± 9.9	109 ± 30	106 ± 7	21.6 ± 5.0	43.9 ± 8.2	40.7 ± 5.2	145 ± 36	48.1 ± 20.9	106 ± 36
isorhamnetin 3-O-glucoside	319 ± 31	433 ± 49	324 ± 33	274 ± 25	252 ± 16	84.4 ± 8.2	109 ± 11	234 ± 6	252 ± 27	138 ± 39	283 ± 21
isorhamnetin 3-O-glucuronide	72.9 ± 8.1	92.2 ± 6.5	41.8 ± 3.3	79.5 ± 6.3	50.4 ± 3.4	22.3 ± 4.6	28.2 ± 1.3	66.5 ± 1.8	27.3 ± 5.2	77.0 ± 15.2	51.6 ± 5.3
syringetin 3-O-glucoside	171 ± 26	130 ± 15	139 ± 16	87.8 ± 12.2	132 ± 8	62.1 ± 3.7	68.5 ± 4.9	156 ± 5	66.3 ± 8.3	57.2 ± 10.7	106 ± 7
quercetin	4.3 ± 0.4	5.6 ± 0.7	3.9 ± 0.7	2.8 ± 0.3	7.3 ± 3.2	1.3 ± 0.2	1.3 ± 0.1	3.5 ± 0.3	5.8 ± 2.4	2.3 ± 0.3	3.4 ± 0.5
quercetin 3-O-glucoside	105 ± 12	160 ± 21	159 ± 26	133 ± 13	50.9 ± 2.7	17.7 ± 2.2	22.9 ± 3.6	92.9 ± 9.0	181 ± 26	27.7 ± 5.0	94.3 ± 10.8
quercetin 3-O-galactoside	240 ± 33	400 ± 68	174 ± 11	228 ± 32	187 ± 14	39.5 ± 9.0	51.2 ± 3.1	106 ± 3	133 ± 30	50.8 ± 9.3	120 ± 24
quercetin-3-O-glucopyranoside	1075 ± 100	1361 ± 122	849 ± 47	973 ± 90	825 ± 45	260 ± 47	447 ± 41	629 ± 19	599 ± 107	300 ± 51	622 ± 100
quercetin-3-O-arabinoside	24.9 ± 3.0	22.1 ± 2.3	16.6 ± 1.6	15.3 ± 2.0	17.8 ± 2.1	3.6 ± 1.1	10.9 ± 1.7	8.6 ± 1.4	10.7 ± 2.0	5.7 ± 1.0	13.0 ± 2.2
quercetin 3-O-glucuronide	2726 ± 177	3121 ± 128	1951 ± 103	3014 ± 108	2119 ± 89	995 ± 132	1211 ± 19	2900 ± 44	1430 ± 253	1454 ± 259	1656 ± 156
quercetin-3-O-rutinoside	272 ± 35	170 ± 23	76.4 ± 9.8	279 ± 22	114 ± 10	28.3 ± 5.3	51.4 ± 3.2	144 ± 3	107 ± 38	48.7 ± 13.5	57.1 ± 5.9
flavanols (µg g ⁻¹ DW)											
catechin	126 ± 9	110 ± 8.7	111 ± 14	224 ± 19	81.9 ± 7.4	355 ± 25	188 ± 48	66.4 ± 1.8	162 ± 23	77.9 ± 5.7	102 ± 5
epicatechin	8.8 ± 1.3	5.1 ± 0.6	8.4 ± 0.7	13.3 ± 1.3	5.9 ± 0.7	7.2 ± 1.2	4.5 ± 0.6	3.3 ± 0.3	9.2 ± 1.0	1.8 ± 0.2	2.7 ± 0.1
procyanidin B1	331 ± 27	324 ± 35	266 ± 23	467 ± 40	208 ± 18	633 ± 40	384 ± 59	173 ± 7	323 ± 40	130 ± 6	168 ± 10
flavanonols (μg g ⁻¹ DW)											
astilbin	715 ± 61	591 ± 68	629 ± 59	511 ± 40	568 ± 45	163 ± 12	265 ± 35	476 ± 17	493 ± 43	299 ± 58	257 ± 43
taxifolin-3-O-glucoside	429 ± 64	114 ± 14	194 ± 37	250 ± 19	168 ± 38	21.8 ± 8.4	75.0 ± 19.1	138 ± 11	141 ± 21	10.7 ± 2.2	27.2 ± 6.0
stilbenes (μg g ⁻¹ DW)											
resveratrol	54.7 ± 6.7	123 ± 28	105 ± 29	34.1 ± 12.1	31.4 ± 5.1	21.7 ± 8.5	6.4 ± 1.4	41.4 ± 4.3	57.1 ± 19.2	11.8 ± 6.5	15.4 ± 0.9
resveratrol-3-O-glucoside	395 ± 62	805 ± 77	385 ± 52	117 ± 32	120 ± 27	53.9 ± 27.6	17.7 ± 5.5	243 ± 32	303 ± 19	2.2 ± 0.6	29.2 ± 8.3
cinnamic Acids (μg g ⁻¹ DW)											
coumaroyl-tartaric acid	876 ± 142	221 ± 14	215 ± 37	1016 ± 143	208 ± 54	72.6 ± 32.0	89.4 ± 24.9	72.0 ± 50.1	824 ± 114	14.8 ± 9.5	48.7 ± 14.9
caffeoyl-tartaric acid	4943 ± 716	2101 ± 427	1763 ± 214	6195 ± 809	1870 ± 497	894 ± 282	1047 ± 244	1597 ± 296	5855 ± 967	144 ± 108	947 ± 315
feruloyl-tartaric acid	5.7 ± 0.4	5.1 ± 0.6	2.3 ± 0.2	5.9 ± 0.7	5.8 ± 0.8	3.6 ± 0.4	5.0 ± 0.8	5.7 ± 2.4	4.0 ± 0.3	1.8 ± 0.3	2.1 ± 0.4
anthocyanins (mg g ⁻¹ DW)											
delphinidin-3-O-glucoside	1.7 ± 0.2	2.9 ± 0.3	3.0 ± 0.5	2.9 ± 0.2	0.8 ± 0.1	0.2 ± 0.0	0.2 ± 0.0	1.4 ± 0.0	3.7 ± 0.5	0.3 ± 0.0	2.6 ± 0.3
cyanidin-3-O-glucoside	0.9 ± 0.1	4.4 ± 0.1	1.6 ± 0.2	1.0 ± 0.3	0.9 ± 0.0	0.2 ± 0.0	0.1 ± 0.0	1.8 ± 0.3	1.7 ± 0.2	0.3 ± 0.0	1.5 ± 0.1
petunidin-3-O-glucoside	5.0 ± 0.9	6.4 ± 0.8	4.3 ± 0.0	5.7 ± 0.7	1.8 ± 0.3	0.7 ± 0.1	0.9 ± 0.1	2.7 ± 0.0	4.8 ± 2.0	1.0 ± 0.1	2.8 ± 0.2
peonidin-3-O-glucoside	14.9 ± 1.6	34.9 ± 1.0	20.9 ± 1.3	13.8 ± 1.4	11.7 ± 1.2	5.7 ± 0.8	3.2 ± 0.2	25.9 ± 0.3	16.7 ± 2.7	5.8 ± 0.8	8.1 ± 0.8
malvidin-3-O-glucoside	54.6 ± 1.1	61.5 ± 0.7	39.8 ± 3.6	44.2 ± 0.3	26.4 ± 3.6	12.3 ± 0.2	17.4 ± 0.1	39.8 ± 1.4	36.2 ± 3.9	13.3 ± 0.6	27.1 ± 0.1
other variables											
antioxidant capacity ($\mu M TE g^{-1} DW$)	8013 ± 942	8639 ± 408	8637 ± 216	9104 ± 212	5576 ± 654	4134 ± 308	5111 ± 600	6330 ± 730	8212 ± 902	3592 ± 685	8424 ± 595
lutein (µg g ⁻¹ DW)	66.2 ± 0.8	55.5 ± 5.2	67.7 ± 1.2	32.9 ± 1.6	32.3 ± 1.2	24.1 ± 1.0	31.8 ± 1.3	16.1 ± 1.3	48.4 ± 0.6	20.2 ± 1.6	52.0 ± 10.1
zeaxanthin ($\mu g g^{-1}$ DW)	8.6 ± 0.4	8.4 ± 0.0	9.2 ± 0.7	3.7 ± 0.3	5.5 ± 0.4	3.7 ± 0.5	4.9 ± 0.3	2.1 ± 0.0	6.7 ± 0.4	2.6 ± 0.1	9.2 ± 0.4
β -carotene ($\mu g g^{-1} DW$)	171 ± 7	165 ± 6	195 ± 2	96.4 ± 3.8	112 ± 11	83.1 ± 3.5	68.8 ± 7.6	56.7 ± 5.0	129 ± 9	66.8 ± 4.8	148 ± 19
				36							

chlorophylls ($a+b$) (µg g ⁻¹ DW)	438 ± 22	424 ± 44	525 ± 14	227 ± 6	290 ± 32	188 ± 16	182 ± 9	117 ± 10	360 ± 16	135 ± 5	480 ± 51
fresh weight per berry (g)	1.4 ± 0.2	1.1 ± 0.1	1.3 ± 0.0	1.3 ± 0.1	1.1 ± 0.1	1.2 ± 0.2	1.2 ± 0.0	1.4 ± 0.1	1.7 ± 0.1	1.5 ± 0.1	2.1 ± 0.0
TSS (^o Brix)	23.7 ± 0.3	20.4 ± 0.4	22.3 ± 0.3	21.3 ± 0.0	21.1 ± 0.4	19.1 ± 0.1	20.1 ± 0.5	19.5 ± 2.0	23.2 ± 0.4	20.9 ± 0.1	22.0 ± 0.2

Table 3. Values (means \pm SE) of the variables analyzed in Pinot Noir berries in the 11 European sampling localities used in this study. MSPC, methanol-soluble phenolic compounds. AUC, area under curve. TSS, total soluble solids.



Figure 1. Geographic location of the 11 European sampling localities used in this study. 1, Jerez de la Frontera (Spain); 2, Girona (Spain); 3, La Rioja (Spain); 4, Firenze (Italy); 5, Bordeaux (France); 6, Bilje (Slovenia); 7, Potoče (Slovenia); 8, Pécs (Hungary); 9, Retz (Austria); 10, Lednice (Czech Republic); 11, Geisenheim (Germany).



Figure 2. Regressions between selected berry variables, including carotenoids and the different groups of phenolic compounds, and latitude. Determination coefficients (R^2) and p values are shown.



Figure 3. Regressions between the ratio trihydroxylated / monohydroxylated flavonols and selected radiation variables. DSSF, Downward Surface Shortwave Flux. T UVery, TEMIS-derived erythematic UV. For both variables, the daily mean in the period budbreak-harvest, and the accumulated dose in the same period and in the 10 days before harvest, were used for calculations. Determination coefficients (R^2) and p values are shown.



Figure 4. Ordination, through Principal Components Analysis (PCA), of the 11 sampling localities used in this study, taking into account the total content of methanol-soluble phenolic compounds (MSPC) and the total concentrations of the different groups of phenolic compounds. Significant loading factors for the positive and negative parts of each axis, together with their corresponding significance levels, are shown (***, p<0.001; **, p<0.01; *, p<0.05). Axis 1 is the horizontal one, and axis 2 is the vertical one. Each mark on the axes represents 0.5 units.

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