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Polyphenolic diversity in Vitis sp. leaves

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28 ABSTRACT

We identified and quantified the main constitutive polyphenolic compounds of the leaves of 29 seven *Vitis* species and of one interspecific cross, analysing leaf blades and veins separately, to spread 30 31 light on the strategic localization of polyphenols in leaf tissues. To the best to our knowledge, the main leaf constitutive polyphenols of V. candicans, V. coignetiae, V. vinifera sylvestris and Börner were never 32 33 described. V. riparia and V. rupestris (belonging to the same botanical series of Ripariae) displayed 34 similar flavan-3-ol and dimeric proanthocyanidin concentration and similar percentage incidence of 35 caftaric acid over total phenolic acids. V. riparia distinguished respect to the other genotypes for its high flavonol content, the highest percentage incidence of myricetin derivatives and an important 36 diversification in the type of accumulated flavonol. V. berlandieri (series Cinereae) and Börner (hybrid 37 of V. riparia x V. cinerea) accumulated low amounts of flavonol-glucosides comparing to the other 38 species, but they showed a wide profile diversification, as well. However, it was V. v. sylvestris, the wild 39 ancestor of Vitis vinifera subsp. sativa that displayed the widest flavonol profile diversification. The 40 41 differences in the flavonol profile could be related to the genus Vitis evolution: in fact, with domestication, the flavonoid biosynthetic pathway underwent a progressive simplification; for this, the 42 highest flavonol diversity found in *Vitis v. sylvestris* is probably a demonstration of its reduced or nil 43 level of domestication. V. amurensis, known for its cold tolerance and resistance to downy mildew, 44 anthracnose and white rot, markedly differentiated respect to the other genotypes, for its high 45 46 concentration of polyphenols, particularly of vein flavonols and flavanonols. Moreover, V. amurensis 47 leaves generally presented a constantly high concentration of constitutive polyphenols throughout the season that probably contributes to protect against adverse environmental condition. The abundance of 48 polyphenols in V. amurensis leaves emphasizes that this species is a source of natural bioactive 49 50 compounds that could find application for nutraceutical and pharmacological uses. V. berlandieri and Börner markedly distinguished respect to the other studied species for their exclusive capability to 51 accumulate flavones (mainly orientin, isoorientin, vitexin and isovitexin) in blades and in veins, in 52 considerable amounts. 53

54	Knowledge about these subjects could contribute to shed light on the identification of species-
55	related molecules involved in the plant-defense mechanisms, to the chemotaxonomy of the genus Vitis,
56	to the possibility of identifying specific natural bioactive compounds to use in plant-based preparation
57	for nutraceutical, cosmetic, feed/food-additive purposes.
58	
59	KEYWORDS: V. amurensis, V. candicans, V, riparia, V. rupestris, V. berlandieri, V. coignetiae, V.
60	vinifera sylvestris, Börner.
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63 1. INTRODUCTION

The genus *Vitis* comprises more than 60 species, mostly inhabiting temperate regions. The most known and economically important one is the western Eurasian species *Vitis vinifera* subsp. *vinifera*, which ancestor is the wild *Vitis vinifera* subsp. *sylvestris*, naturally occurring in Europe, Middle East and Northern Africa. Up to thirty grapevine species are native of East Asia and North America, areas that host a wide diversity of *Vitis* species (Wan et al., 2013).

69 Vitis vinifera is susceptible to many pests. Phylloxera (Daktulosphaira vitifoliae), downy and 70 powdery mildew (caused by *Plasmopara viticola* and *Erysiphe necator*, respectively), imported almost 71 simultaneously from North America to Europe in the XIX century, are among those causing important 72 economic loss in viticulture. Viticulturists, technicians and researchers overcame the problem tied to vinifera sensitiveness to Phylloxera by using American species or their hybrids as rootstocks. 73 Furthermore, the introduction of downy and powdery mildew pushed to develop breeding programs 74 based on the constitution of interspecific crosses vinifera-American species that, together with Asian 75 species, were disease-tolerant or resistant due to their coevolution with the pathogen. Vitis species such 76 77 as V. riparia, V. rupestris, V. californica and V. amurensis are known as highly tolerant to downy mildew (Gómez-Zeledón et al., 2013; Jürges et al., 2009); V. riparia, V. munsoniana, V. candicans, V. 78 rotundifolia as highly resistant to powdery mildew (Staudt, 1997; Wan et al., 2007) and V. rotundifolia 79 (subspecies Muscadinia) and V. arizonica as resistant to Pierce's disease (Ruel and Walker, 2006). 80 Besides, some species distinguish for their tolerance to environmental stressors thanks to their origin: it 81 is the case of V. amurensis, native of cold areas in northeastern China and Russian Siberia, that is highly 82 83 frost tolerant (Zhang et al., 2012).

To prevent disease or damage caused by biotic or abiotic stresses, plants employ a complex defense system, which involves a broad spectrum of physical and biochemical changes. Biochemical resistance, such as preformed defenses, has evolved to face environmental stressors and is also related, among others, to secondary metabolites, including polyphenols (reviewed in Llorens et al., 2017 and in Dixon, 2001). *Vitis vinifera* is known for its abundance and richness in polyphenols that have been and

89 currently are widely studied in the berries due to their implications in the technology of winemaking and in the quality of derived wines. Thousands of scientific studies were devoted to define the polyphenolic 90 composition of Vitis vinifera berries, and, to a much lesser extent, to vegetative organs; oppositely, less 91 92 is known about other Vitis species polyphenolic composition, particularly as to vegetative organs. Moore and Giannasi (1994) described the qualitative composition of flavonols and flavones in leaves of some 93 North American Vitis species. Flavonol profiles were described also in V. amurensis (Hmamouchi et al., 94 95 1996) and V. rotundifolia (Pastrana-Bonilla et al., 2003) leaves. Main phenolic compounds were also analyzed in V. labrusca leaves (Dani et al., 2010; Dresch et al., 2014). Chen et al. (2018) reviewed V. 96 97 amurensis polyphenolic composition and related pharmacological properties. At present, the scientific interest about Vitis sp. polyphenolic composition is at least dual: to investigate the health beneficial 98 effects of grapevine leaves as a possible source of natural bioactive compounds, that can be used in 99 100 nutraceutical and pharmaceutical applications, and to deepen knowledge about their involvement in 101 grapevine defense mechanisms. Different Vitis species might display different metabolic pathways and/or different gene regulations, resulting in the production of specific classes of bioactive polyphenols, 102 or of individual molecules. Knowledge about these subjects could contribute to shed light on the 103 identification of species-related molecules involved in the plant-defense mechanisms, to the 104 chemotaxonomy of the genus Vitis and to the possible identification of specific molecules to use in 105 specific plant-based preparation for nutraceutical, cosmetical, nutritional purposes. 106

107 The aim of the present study was to identify the main constitutive polyphenolic compounds in 108 the leaves of seven *Vitis* species and of one interspecific cross (the rootstock Börner, *V. riparia x V.* 109 *cinerea;* Figure 1). To the best to our knowledge *V. candicans, V. coignetiae, V. v. sylvestris* and Börner 110 main leaf constitutive polyphenols were never described. Leaf blades and veins were analysed separately 111 to spread light on polyphenol strategic localization in leaf tissues, to contribute to explain the grapevine-112 pathogen interaction in specific biological systems.

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115 2. MATERIALS AND METHODS

2.1 Plant Material. Four American (V. candicans, V. riparia, V. berlandieri and Börner), two 116 Asian (V. amurensis, V. coignetiae) and one Eurasian (V. v. sylvestris) grapevine genotypes were studied 117 (Table 1; Figure 1). All the studied accessions were previously checked for their trueness to type by 118 morphology and molecular markers. The studied *Vitis* genotype leaves were collected in the collection 119 vineyard of DISAFA, University of Turin, Grugliasco (Piedmont, Italy), sampled five times during the 120 season: $1 = 28^{\text{th}}$ of May (148 day of the year - DOY), $2 = 22^{\text{nd}}$ of June (DOY 173), $3 = 14^{\text{th}}$ of July (DOY 121 195), $4 = 3^{rd}$ of August (DOY 215), $5 = 28^{th}$ of August (DOY 214), in 2015. The collection vineyard 122 was managed routinely during spring and summer as already described (Kedrina-Okutan et al., 2018) 123 (Kedrina-Okutan et al., 2018). One field parcel was generally constituted by four consecutive plants 124 used to collect the leaves; leaves were detached between the fourth and seventh node of the main shoots 125 from the west side of the row. Pools of 15 adult and healthy leaves (visual evaluation) were collected at 126 each sampling time and divided into three replicates. After sampling, leaves were immediately 127 transported to the laboratory for further analyses. 128

2.2 Reference Compounds and Reagents. Orientin, isoorientin, astilbin and caftaric acid were 129 purchased from Sigma-Aldrich S.r.l. (Milan, Italy). Vitexin, trans-fertaric acid and trans-coutaric acid 130 were purchased from Phytolab (Vestenbergsgreuth, Germany). (+)-catechin, (-)-epicatechin, (-)-131 epicatechin gallate, (-)-epigallocatechin gallate, proanthocyanidin B₁, proanthocyanidin B₂, quercetin 3-132 O-glucoside, quercetin 3-O-glucuronide, kaempferol 3-O-glucoside, kaempferol 3-O-glucuronide, 133 myricetin 3-O-glucoside and isorhamnetin 3-O-glucoside from Extrasynthèse (Genay, France). 134 Folin-Ciocalteu reagent and tartaric acid were purchased from Merck (Darmstadt, Germany); sodium 135 sulfate and sodium metabisulfite were purchased from BDH Laboratory Supplies (Poole, England). 136

137 2.3 Measurement of Dry Matter. Leaf tissue dry matter was measured three times during the 138 season at DOY 186, DOY 200 and DOY 241. Leaf veins and blades were separated and dried inside an 139 oven for 72 hours at constant temperature of 110 °C. The weight differences after exsiccation were 140 measured, and results were expressed as percentage of dry weight over fresh weight.

2.4 Sample Extraction for Polyphenol Analyses. The 15 collected leaves were divided into 141 sub-samples of five, and veins and blades were immediately separated. Two grams of tissue were 142 randomly chosen form the three sub-samples and extracted in a hydroalcoholic buffer (pH = 3.9, 40%143 ethanol, 22 mL L⁻¹ of 1 N NaOH, 5 g L⁻¹ of tartaric acid, 2 g L⁻¹ of sodium metabisulfite). For sample 144 homogenization and polyphenol extraction, an Ultraturrax dispersing machine (IKA, Staufen, Germany) 145 was used for around 1 min setting the speed at 10 000 rpm followed by 10 min of centrifugation at 4000 146 rpm. The supernatant was separated, and the pellet was resuspended with the same buffer; the 147 resuspension was kept in the dark for 30 min and then centrifuged again. The two supernatants were 148 149 combined and brought to a final volume of 50 mL. Obtained leaf extracts were stored at -20 °C until further analysis. 150

2.5 Measurement of Total Polyphenols. The method of Singleton et al. (1999) was used and results were expressed as gram of (+)-catechin equivalents (CE) per kg of blades/veins dry weight (DW). Briefly, 100 μ L of leaf extract were mixed with 5 mL ultrapure water and 1 mL of Folin-Ciocalteu reagent was added, mixed and incubated at room temperature for 5 min. After incubation, 4 mL of 10% Na₂CO₃ and ultrapure water were added until final volume of 20 mL. The mixture was incubated in the dark for 90 min and the absorbance was read at 760 nm by UV/Vis spectrophotometer (PerkinElmer, Lambda 25, Beaconsfield, Bucks, UK).

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2.6 Analysis of individual polyphenols.

2.6.1 Sample preparation. Prior to the chromatographic analysis, the extracts were diluted with
1 M phosphoric acid 90/10 v/v and passed through 0.20 µm membrane filter GHP Acrodisc (PALL
Italia, Buccinasco, Milan, Italy) (Di Stefano and Cravero, 1992).

Individual phenolic compound analysis by HPLC-ESI-MS/MS. Chromatographic separation was performed by HPLC-DAD system (HPLC 1100, Agilent Technologies, U.S.) equipped with column Luna C-18 150 x 2 mm (Phenomex Aschaffenburg, Germany). Mobile phase solvent A was water with formic acid 0.1% and solvent B was methanol with formic acid 0.1%; the gradient program for solvent B was the following: 0-30 min 0-50 % B; 30-35 min 50-100 % B; 35-50 min 100 % B and back to initial 167 conditions from 50 to 55 min, followed by 10 min of isocratic flow. Injection volume was 5 µL, flow rate 200 µL min⁻¹ and individual phenolic compounds were detected at 280, 320, and 370 nm. For the 168 mass spectrometry analyses, a Bruker Daltonics esquire 3000^{plus} ion trap spectrometer (Bruker 169 170 Daltonics, HB, Germany) equipped with electrospray (ESI) was used, operating at positive and negative modes. The scan mode was between 100-800 m/z, with a scan resolution of 13 000 m/z/s until the ICC 171 target reached either 20 000 or maximum accumulation time of 200 ms. The MS instrument operated 172 with nitrogen as drying gas at a temperature of 330 °C (flow rate of 9 L min⁻¹). Ionization voltage of the 173 electrospray capillary source was 4000 V and tandem MS was carried out using helium as the collision 174 gas (4.21 x 10⁻⁶ mbar) with 1 V collision voltage. MS identification of metabolites was according to 175 mass spectra, product ion spectra, retention time and confirmed with authentic standards and published 176 data. 177

178 2.6.2 Individual phenolic compound quantification by HPLC-DAD. The quantification of phenolic compounds was performed by HPLC-DAD according to previously published methods (Di 179 Stefano and Cravero, 1992; Ferrandino and Guidoni, 2010). Stationary phase column was Licrosphere 180 100 RP5 µm) packed into LiChroCART 250-4 (25 × 0.4 cm ID) HPLC-Cartridge (Merck KGaA, 181 Germany) with guard column (LiChroCART 4-4). Mobile phase solvent A was phosphoric acid 10^{-3} M 182 and solvent B was pure methanol. Run time was 50 min, temperature 25 °C and DAD peaks were 183 detected at 280 nm, 320 nm and 360 nm. Compounds were identified based on compliance with data 184 185 obtained from available pure standards and quantified based on standard curve constructed per each 186 individual molecule. Among flavanonol, as exclusively dihydroquercetin-rhamnoside (astilbin) was available as commercial standard, semi-quantification of individual compounds was carried out using 187 the astilbin standard curve. The average flavonol profile of individual genotype was calculated averaging 188 189 results of five sampling dates.

190 **2.7 Statistical analysis.**

All data are averages of three biological replicates and standard errors. The analysis of variance was
 performed by one-way ANOVA with IBM SPSS Statistics software program version 24.0 for Windows

193	(SPSS Inc., Chicago, IL). In case of significant differences ($P \le 0.05$), means were compared by Tukey-
194	b post-hoc test. Results related to leaf blade and vein polyphenolic composition taken separately were
195	used to run a series of principal component analysis (PCA); correlated variables were progressively
196	excluded, when this correlation had a biological meaning. Species and dates of sampling were separated
197	on the basis of specific classes of polyphenols (i.e. variables). PCA analysis was performed by SAS 9.4
198	for Windows (SAS Institute Inc., Cary, NC, US).

Table 1. Origin and botanical sub-generic series of the studied genotypes.

Plant name	Abbreviation	Origin	Series ^{<i>a</i>}
Vitis candicans	CAN	American	Candicansae
Vitis riparia cv Gloire de Montpellier	RIP	American	Ripariae
Vitis rupestris cv du Lot	RUP	American	Ripariae
Vitis berlandieri	BER	American	Cinereae
Börner (V. riparia x V. cinerea)	BOR	American	Ripariae X Cinereae
Vitis coignetiae	COI	Asian	Lambruscae
Vitis amurensis	AMU	Asian	Flexuosae
Vitis vinifera subsp. sylvestris	SYL	Eurasian	Viniferae
"Sorias were defined according to Calat	(1099)		-

201 ^{*a*}Series were defined according to Galet (1988).



203 204

Figure 1. Leaves of eight *Vitis* genotypes.

206 3. RESULTS

Significant differences of water content occurred among *Vitis* species at the same picking time 207 (Table 2); however, differences did not exceed 7% in blades and 8% in veins. Within the species, water 208 content evolution during the vegetative season was not higher than 6%; in blades of V. candicans, V. v. 209 sylvestris, Börner and V. coignetiae dry matter (DM) content changes during the season were never 210 significant. The comparison between averages of all analyzed grapevine blades and veins showed that 211 dry matter content in veins was around 10% lower than in blades. Concentrations of compounds were 212 calculated and expressed as amounts per dry matter, even though leaf extracts were prepared from fresh 213 214 leaves: this choice was done to minimize the well detailed polyphenol analysis perturbations due to losses or to chemical alterations during preparation (Abascal et al., 2005). 215

	DOY 186		DOY 200		DOY 241		date
		#	Blades	#		#	Sampling dates
AMU	$32.09\pm0.03a$	e	$34.43\pm0.58b$	c	$35.29\pm0.23b$	bc	*
CAN	29.71 ± 0.30	cde	29.68 ± 0.18	ab	28.94 ± 1.63	а	ns
RIP	$27.78\pm0.69a$	ab	$28.05\pm0.79a$	a	$31.95 \pm 1.21b$	ab	*
RUP	$27.35\pm0.75a$	a	$27.16\pm0.80a$	a	$33.51 \pm 1.01b$	b	**
BER	$31.45\pm0.18a$	de	$31.86 \pm 1.07a$	bc	$37.57\pm0.61b$	c	**
COI	$29.59\pm0.43a$	bc	$32.09\pm0.70b$	bc	$32.93 \pm 0.53b$	ab	*
SYL	31.39 ± 0.62	de	29.70 ± 0.83	ab	31.76 ± 0.61	ab	ns
BOR	30.57 ± 0.66	de	31.49 ± 0.74	bc	32.88 ± 0.35	ab	ns
average	29.99 ± 0.37		30.56 ± 0.51		33.10 ± 0.56		
species		**		**		**	
			Veins				
AMU	$19.01\pm0.42a$	ab	$22.61\pm0.47b$	bc	$22.32\pm0.33b$	b	**
CAN	$23.14\pm0.70b$	b	$23.61 \pm 0.99b$	c	$19.32 \pm 0.60a$	а	*
RIP	21.21 ± 0.71ab	b	$20.52\pm0.52a$	ab	$22.88 \pm 0.21b$	bc	*
RUP	19.23 ± 1.43a	ab	$20.02\pm0.52a$	ab	$25.13\pm0.34b$	cd	**
BER	$21.37\pm0.40a$	ab	23.08 ± 1.12a	bc	$26.30\pm0.58b$	d	*
COI	17.82 ± 0.53	a	18.89 ± 0.72	a	19.27 ± 0.66	а	ns
SYL	$21.22\pm0.47a$	ab	$23.74\pm0.15b$	c	$27.28\pm0.48c$	d	**
BOR	$20.59\pm0.45a$	ab	$20.26 \pm 0.33a$	ab	$23.88 \pm 0.81b$	bc	**
average	20.45 ± 0.38		21.59 ± 0.42		23.30 ± 0.60		
species		*		**		**	

Table 2. Dry matter (%) in leaves of Vitis genotypes during the vegetative season a

^{*a*} Results are expressed as means \pm standard errors (SE; n=3). Different small letters in each individual row after averages \pm SE represent statistical differences among sampling dates, within the same species. In columns individuated with #, different letters represent statistical differences among genotypes, within the same tissue and date. General statistical differences among sampling dates and species were assessed by a post-hoc Tukey-b test for P \leq 0.05 (*), P \leq 0.01 (**); ns – not significant. Day of the year (DOY) refers to 5th of July (DOY 186); 19th of July (DOY 200); 29th of August (DOY 241).

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The total polyphenol (TP) content ranged from 60.2 to 165.9 g kg⁻¹ DW in blades and from 28.6 to 130.1 g kg⁻¹ DW in veins. *V. amurensis* blades (Figure 2) displayed higher TP concentration during the entire vegetative season compared to the other genotypes. TP was on average about 1.5 times lower in veins than in blades. TP accumulation trend was different among genotypes: in *V. riparia*, the accumulation trends in veins and blades were similar, whereas in *V. candicans*, TP concentration increased in blades and decreased in veins during the season.



Figure 2. Total polyphenol (TP) accumulation in leaves of eight *Vitis* genotypes during the vegetative season; A = blades and B = veins. Means of three replicates \pm SE.

3.1 Phenolic Compound Accumulation in *Vitis sp.* Leaves. Twenty phenolic compounds were identified in the two Asian species, 27 in the four American species and in the hybrid, and 18 in *V. v. sylvestris.* The detected compounds belonged to the groups of flavonols, phenolic acids, flavan-3-ols, proanthocyanidins, flavanonols and flavones. Anthocyanins were exclusively detected in trace amounts or in extremely low concentrations (data not shown) and, in line with what we previously found in varieties of *V. vinifera sativa* (Kedrina-Okutan et al., 2018), in healthy leaves they were found exclusively in veins.

242 *3.1.1 Flavonols.*

The concentration of flavonol glycosides ranged from 3.6 to 20.6 g kg⁻¹ DW in blades and from 0.8 to 7.7 g kg⁻¹ DW in veins (Figure 3); they were the most abundant phenolic compounds in leaves. We detected a net separation among genotypes based on their ability to accumulate flavonol glycosides during the season. A first group included *V. amurensis* and *V. riparia*, both species displaying the highest 247 total flavonol concentration, with a peak at DOY 196 in V. riparia and at DOY 215 in V. amurensis (Figure 3A). A second group with medium flavonol concentration included V. v. sylvestris and V. 248 rupestris with a stable concentration trend during the vegetative season. A third group included the 249 250 remaining species, displaying relatively low total flavonol concentration and slightly decreasing trends during the examined period. Similar results were found in veins: genotypes of the first two groups 251 displayed higher total flavonol concentration and genotypes of the third group lower concentration with 252 253 decreasing trend during the season (Figure 3B). The comparison between the two compartments within each genotype showed that total flavonol concentration was always higher in blades than in veins. The 254 255 prevalent flavonol was quercetin 3-O-glucuronide, ranging between 60% and 92% of all the detected flavonols (Figure 4; Supplementary Table 1A,B). The most complex flavonol profile was found in V. 256 v. sylvestris leaves accumulating seven out of the eight identified flavonol glycosides and trace amount 257 258 of kaempferol 3-O-glucoside (Figure 4). Kaempferol 3-O-rhamnoside was exclusively found in V. v. sylvestris that also accumulated significant amounts of quercetin 3-O-rhamnoside. The simplest flavonol 259 profile was found in V. amurensis and V. candicans that exclusively accumulated quercetin and 260 kaempferol glucuronides and glucosides. In V. berlandieri and in Börner, no kaempferol glycosides 261 262 were detected.

263 *3.1*.

3.1.2 Phenolic acids.

Total phenolic acid content in blades ranged from 3.2 to 10.2 g kg⁻¹ dry weight and showed 264 265 different accumulation trends among genotypes. In Asian species, V. amurensis and V. coignetiae, total phenolic acid concentration fell during the vegetative season. In American species, concentration firstly 266 increased and then it started to fall between July and beginning of August (DOY 173-215), whereas in 267 V. v. sylvestris, concentration decreased between the first and the second sampling date and then rose 268 from the end of June (DOY 173) to the beginning of August (DOY 215) (Figure 5A). In leaf veins, 269 phenolic acid content showed a significant drop between the first and the second sampling date in all 270 genotypes, followed by a relatively steady period (Figure 5B). Total phenolic acid concentration in veins 271 was lower than in blades in each individual species, except at the first sampling date in V. candicans, V. 272

273 *riparia* and *V. rupestris* when concentration in veins was higher than in blades. Hydroxycinnamoyl tartaric acids were the prevalent phenolic acids in leaves, and, based on their characteristic UV maximum 274 absorbance around 320 nm and mass spectra, they were identified as cis- and trans-forms of caftaric, 275 276 coutaric and fertaric acid (Table 3). Trans-forms were always prevalent over cis-forms The main phenolic acid in leaves was trans-caftaric acid, which comprised up to 90% followed by trans-coutaric 277 and *trans*-fertaric acid (Figure 5C, Supplementary Table 2A, B). On the contrary, in V. *rupestris* and V. 278 riparia trans-fertaric acid displayed higher concentration respect to trans-coutaric acid. Moreover, these 279 two species accumulated an additional molecule, a coutaric acid isomer, that was tentatively identified 280 through its pseudomolecular ion $[M-H]^-$ at m/z 295 and two product ions at m/z 163, 131 in MS². Low 281 amounts of protocatechuic acid-glucoside were detected in V. rupestris, V. riparia, V. amurensis and 282 283 Börner blades, with a slight increasing trend during the season (Supplementary Table 2A).



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Figure 4. Flavonol profile of leaf blades of eight *Vitis* genotypes calculated as average of results

289 obtained at five sampling times performed during the vegetative season.



Figure 5. Accumulation of phenolic acids in *Vitis sp.* leaves during the vegetative season. Total phenolic acid concentration in blades (A) and veins (B); results are means of three replicates \pm SE. Phenolic acid profile of leaf blades (C) and veins (D) during the season. For species acronyms see Table 1; numbers after the species acronyms refer to 1 = DOY 148 (28th of May); 2 = DOY 173 (22nd of June); 3 = DOY 195 (14th of July); 4 = DOY 215 (3rd of August); and 5 = DOY 240 (28th of August).

297	Table 3. Phenolic compound identification in leaf blades and veins of eight Vitis genotypes by HPLC-ESI-
298	MS/MS

ID	Rt (±0.2 min)	[M] ⁻ (m/z)	[MS ²] ⁻ (m/z)	[M] ⁺ (m/z)	[MS ²] ⁺ (m/z)	compound identification
1	14.5	593	425, 407, 289	595	291	(epi)gallocatechin-(epi)catechin
2	16.5	315	153, 123			protocatechuic acid-glucoside
3	18.3	311	179			cis-caftaric acid
4	18.8	577	451, 407, 289			procyanidin B ₃
5	19.9	311	179			trans-caftaric acid
6	21.8	289	245, 205, 179	291	273, 165, 123	catechin
7	22.5	295	163			cis-coutaric acid
8	23.0	577	451, 425, 407, 289	579	427, 409, 289	procyanidin B ₁
9	23.5	295	163			trans-coutaric acid
10	23.8	729	577, 441, 289	731	579, 409	procyanidin dimer gallate
11	24.9	595	576, 385, 355			unknown proanthocyanidin
12	25.1	325	193			trans-fertaric acid
13	25.7	289	245, 205, 179	291	273, 165, 123	epicatechin
14	27.7	465	303, 285, 151			dihydroquercetin-hexoside
15	27.7	295	163, 131			coutaric acid isomer
16	29.2	615	481, 407, 359			unknown
17	30.8	447	393, 357, 327	449	413, 383, 329	luteolin 8-C-glucoside (orientin)
18	31.1	449	287, 269, 151			dihydrokaempferol hexoside
19	31.4	447	429, 357, 327, 285	449	431, 413, 383	luteolin 6-C-glucoside (isoorientin)
20	31.8	479	317	481	319	myricetin 3-O-glucoside
21	32.3	431	341, 311, 283	433	415, 367, 283	apigenin 8-C-glucoside (vitexin)
22	32.7	493	317	495	319	myricetin 3-O-glucuronide
23	33.7	449	303, 285, 151			dihydroquercetin-rhamnoside (astilbin)
24	33.8	431	413, 341, 311, 283	433	415, 367, 337	apigenin 6-C-glucoside (isovitexin)
25	34.2	447	285	449	287	luteolin-7-O-glucoside (luteoloside)
26	35.1	463	301	465	303	quercetin 3-O-glucoside
27	35.6	477	301	479	303	quercetin 3-O-glucuronide
28	37.1	447	301	449	303	quercetin 3-O-rhamnoside
29	37.2	447	285	449	287	kaempferol 3-O-glucoside
30	37.5	461	285	463	287	kaempferol 3-O-glucuronide
31	38.3	431	285	433	287	kaempferol 3-O-rhamnoside

^aID numbers correspond to peaks reported in Supplement Figure 1, 2.

300

301 *3.1.3 Flavan-3-ols.*

Important quantities of total flavan-3-ols were detected in grapevine leaves. The concentration of flavan-3-ols was rather similar between blades and veins, oppositely to flavonols and phenolic acids that prevalently accumulated in blades. Specifically, the concentration of flavan-3-ols ranged from 0.44 to 3.39 g kg^{-1} in blades and from 0.04 to 1.55 g kg^{-1} in veins (Figure 6A and B). In *V. amurensis* blades, the concentration of total flavan-3-ols was significantly higher respect to the other genotypes, and it 307 displayed a decreasing trend, so that at the end of the examined period the differences respect to the other genotypes were less marked. In V. rupestris flavan-3-ol concentration increased almost 308 continuously during the entire vegetative season, oppositely to the other analyzed genotypes. The flavan-309 310 3-ol profile analysis showed that (+)-catechin was generally the prevalent flavan-3-ol, corresponding up to 73 % of total flavan-3-ols in blades (Figure 5C; Supplementary Table 3A) or the exclusive one in V. 311 coignetiae veins (Figure 5D; Supplementary Table 3B). In the Asian species V. amurensis, (-)-312 epicatechin was the prevalent flavan-3-ol in blades, similarly to V.v. sylvestris and V. rupestris veins 313 from the second sampling date onwards. Concentration of (-)-epigallocatechin showed an increasing 314 315 trend during the examined season and in blades it accounted up to 38% by the end of the season. Interestingly, in veins of V. coignetiae and Börner, (-)-epicatechin was absent during the whole season, 316 317 and in V. candicans it was exclusively detected at the first sampling date. The relative abundance of (-)-epicatechin gallate ranged between 4.7% and 14.9% and it was exclusively found in blades of V. 318 319 rupestris, V. berlandieri and Börner.

320 3.1.4 Proanthocyanidins.

Proanthocyanidin concentration ranged from 0.21 to 3.09 g kg⁻¹ in blades. Particularly high 321 concentration was detected in V. amurensis at first four sampling dates and in V. berlandieri at first two 322 sampling dates, followed by a sharp concentration fall (Figure 7A). In veins, PA exclusively 323 324 accumulated in V. amurensis and V. candicans during the whole season and in some dates in four species, also in V. coignetiae and V. v. sylvestris. PA content was generally higher in blades than in 325 326 veins, with the only exception of V. amurensis and V. candicans at DOY 148 (end of May) when vein PA content was higher respect to that of blades (Figure 7; Supplementary Table 4A,B). Procyanidin B₁ 327 accumulated both in blades and veins. Additionally, in V. amurensis, V. berlandieri and in Börner blades 328 and in V. amurensis veins, a procyanidin dimer gallate was tentatively identified by pseudomolecular 329 ion [M-H]⁻ at m/z 729; product ions at m/z 577, 441, 289 in MS² and characteristic UV maximum 330 absorbance at 280 nm, in line with other report (Flamini, 2013). In V. berlandieri and Börner blades, 331

- procyanidin dimer gallate accounted for around 50% of total PA, whereas in *V. amurensis*, it accounted
 for 20-25 % of total PA during the vegetative season.
- 334
 Furthermore, by HPLC-ESI-MS/MS (epi)gallocatechin-(epi)catechin was tentatively identified
- in blades (pseudomolecular ion [M-H]⁻ at m/z 593; product ions at m/z 425, 407, 289 in MS²; Table 3),
- 336 In addition, in V. amurensis blades, another B-type of procyanidin was identified, most likely
- B_3 procyanidin B_3 and one further proanthocyanidin that remains unknown (Table 3).



339 □ (-)-epigallocatechin □ (+)-catechin □ (-)-epicatechin □ (-)-epigallocatechin gallate □ (-)-epicatechin gallate

Figure 6. Accumulation of flavan-3-ols in *Vitis sp.* leaves during the vegetative season. Total flavan-3-ol concentration in blades (A) and veins (B); results are means of three replicates ± SE. Flavan-3-ol profile of leaf blades (C) and veins (D) during the season; see Figure 5 for genotype acronym specification.



Figure 7. Accumulation of proanthocyanidin in *Vitis sp.* leaves (A = blades; B = veins) during the vegetative season. Means of three replicates \pm SE; see Figure 5 for genotype acronym specification.

344

348 *3.1.5 Flavanonols.*

In blades, flavanonols were detected exclusively in V. amurensis and V. coignetiae, whereas in 349 350 veins, they were also found in V. v. sylvestris. (Figure 8; Supplementary Table 4A, B). In blades, concentration was much lower than in veins. Particularly high concentration was found in V. amurensis 351 veins with values ranging between 0.95 and 2.12 g kg⁻¹ (Figure 8B). In both Asian species, vein total 352 353 flavanonol concentration showed two peaks, the first at DOY 148 and the second at DOY 215 whereas in V. v. sylvestris no differences among sampling times were noticed. Three flavanonol glycosides were 354 quantified and identified in leaves by the characteristic absorbance maximum at 290 nm and 355 correspondent mass spectra (Table 3): dihydroquercetin-hexoside was the main flavanonol found in V. 356

amurensis and *V. coignetiae* where it was the exclusive flavanonol in blades and it accounted for up to
90% of total flavanonols in veins (Figure 8A,B). In addition, dihydrokaempferol-hexoside was detected
in *V. amurensis* and in *V. coignetiae* veins with no differences among sampling dates. Astilbin
(dihydroquercetin-rhamnoside) was exclusively detected in the Eurasian species *V. v. sylvestris* veins.

361



362

Figure 8. Accumulation of flavanonol content in *Vitis sp.* leaves (A = blades; B = veins) during the vegetative season. Means of three replicates \pm SE; DHQuer - dihydroquercetin; DHKaemp – dihydrokaempferol; hex – hexoside; rham – rhamnoside; see Figure 5 for genotype acronym specification.

367

368 *3.1.6 Flavones*

Flavones have rarely been detected in leaves and other vegetative organs of grapevine. In fact, 369 370 also in our analyzed genotypes, flavones accumulated exclusively in V. berlandieri and Börner. Total flavone concentration ranged between 2.5 and 4.9 g kg⁻¹ DW in blades (Figure 9A). In comparison with 371 V. berlandieri, Börner blades accumulated significantly higher concentration of total flavones from 372 373 DOY 148 (end of May) until DOY 195 (mid-July). Similar to the other classes of measured polyphenols, flavone concentration was lower in veins respect to blades, ranging from 0.7 to 1.2 g kg⁻¹ DW (Figure 374 9B). Five flavones accumulated in both tissues, two were apigenin glycosides and three luteolin 375 glycosides. Isomers of apigenin glucosides were differentiated by pseudomolecular ion $[M - H]^{-}$ at m/z376 431 and absorbance maximum at 270 nm and 337 nm. Moreover, the product ions of luteolin glycoside 377

378 and their intensities provided insight into individual compound structure. Apigenin 8-C-glucoside (isovitexin) product ions were detected at m/z 283, 311, 341 and apigenin 6-C-glucoside (vitexin) 379 product ions were detected at m/z 283, 311, 341, 413 (Table 3). The absence of product ion m/z 413 and 380 the lower intensity of product ion at m/z 341 $[^{0,3}X_0 - H]^-$ helped to differentiate apigenin 8-C-glucoside 381 from apigenin 6-C-glucoside, accordingly to Kim et al. (2018). Isomers of luteolin glucoside had 382 pseudomolecular ion $[M - H]^-$ at m/z 447 and absorbance maximum at 269, 350 nm. Furthermore, 383 comparison of product ion pattern of three luteolin glucoside isomers let us tentatively identify luteolin 384 8-C-glucoside (isoorientin), luteolin 6-C-glucoside (orientin) and luteolin 7-O-glucoside (luteolin) 385 386 (Table 3). The compound with absence of product ion at m/z 429 and high intensity of product ion at m/z 357 $[^{0,3}X_0 - H]^-$ was tentatively identified as luteolin 6-C-glucoside. Luteolin glucoside isomer with 387 product ion pattern of two high abundance fragment at m/z 285 [M - H - 162]⁻ and m/z 284 [M - H -388 389 162]^{-•} was tentatively identified as luteolin 7-*O*-glucoside, according to Li et al. (2016). Leaf flavone profile analysis showed that isoorientin together with isovitexin comprised up to 70% of all detected 390 flavones, thus 8-C was the predominant glycosylation site of flavone aglycones in Vitis sp. leaves 391 (Figure 9; Supplementary Table 5). 392





394

Figure 9. Accumulation of flavones in *Vitis sp.* leaves (A = blades; B = veins) during the vegetative season. Means of three replicates \pm SE; see Figure 5 for genotype acronym specification.

398 3.2 Overall polyphenolic patterns and genotype associations

In blades, the PCA model that explained the highest variance with three PRINs (72%), allowing 399 a net separation of genotypes, based on eleven variables (Table 4) that were gradually chosen to exclude 400 401 redundancy. Variables associated to the first PRIN were phenolic acids (the sum of fertaric acid, coutaric acid isomer and the cis and trans forms of caftaric acid) and the total concentration of dimeric 402 proanthocyanidins. Genotypes separated on the second PRIN on the basis of myricetin glycoside 403 (glucoside + glucuronide) concentration and of flavonol total amount. The third PRIN separated 404 genotypes based on total flavan-3-ol concentration, evidencing in particular, V. amurensis. Considering 405 406 that the eigenvalue on PRIN3 was almost 2.0 (1.79, Table 4), the flavone total concentration could be considered as the second variable associated (negatively) to the third PRIN. On this basis, Börner and 407 408 V. berlandieri, highly associated to the third PRIN negative values, evidenced their peculiar capability to accumulate high amounts of flavones in leaf blades (Figure 9; Figure 10). 409

In veins, the proposed PCA model (Table 5; Figure 11) explained the 75 % of total variance with 410 the first three PRINs. Variables associated to the first PRIN were total polyphenols, total flavan-3-ols 411 and flavonol glucuronic derivatives; these variables did not markedly identify varieties or specific time-412 413 points of the leaf vegetative cycle, with the only exception of V. amurensis, generally associated to positive values of the first PRIN. Flavanonols and glucosides of flavonols were negatively associated to 414 the second PRIN. Dimeric proanthocyanidins associated to the third PRIN, allowed to evidence that in 415 416 genotypes accumulating this class of compounds, concentration was higher at the first sampling (particularly in V. amurensis, V. candicans, V. berlandieri and in Börner). 417

418 **3.3 Seasonal polyphenol-related traits**

The flavonol concentration varied during the vegetative season, but the profile, strictly relatedto a specific genotype, did not change (Figure 4).

The phenolic acid profile was species-specific, as well, with *V. rupestris* showing the highest diversification both in blades and in veins. *Trans*-caftaric acid (the main phenolic acid) percentage incidence was generally stable during the season or it decreased in a few genotypes (Figure 5).

- 424 The proanthocyanidin profile evolved during the season in a genotype-specific manner in blades;
- 425 oppositely, it did not change in veins (Figure 7).

426 No major differences were noticed in relation to the flavone profile evolution in *Vitis berlandieri*

427 and in Börner during the season.

Table 4. Eigenvectors of eleven variables (polyphenols in leaf blades) on the first three principal
components (PRIN1, PRIN2 and PRIN3).

430

431		PRIN1	PRIN2	PRIN3
422	Myricetin glycosides (glucoside +	0.25	0.51	0.05
432	glucuronide)	-0.25	0.31	-0.05
433	Protocatechuic acid-glucoside	-0.24	-0.04	0.20
121	Dihydroquercetin-hexoside	-0.14	-0.32	0.36
454	Fertaric acid + coutaric acid isomer	-0.40	0.34	-0.02
435	Caftaric acids (cis + trans)	0.41	0.33	-014
436	Coutaric acids (cis + trans)	0.30	0.02	-0.02
	Flavones	0.26	-0.03	-0.43
437	Total phenolic acids	0.33	0.44	-0.15
438	Total flavonols	-0.14	0.46	0.43
120	Total flavan-3-ols	0.30	0.08	0.55
435	Dimeric proanthocyanidins	0.43	0.03	0.34
440	Eigenvalues	3.56	2.57	1.79
441	Total variance	0.32	0.23	0.16

442 Eigenvalues of the three PRINs and their contribution to total variance. In bold letters, the variables associated to the 443 appropriate PRIN.



Figure 10. Tridimensional distribution of *Vitis* species during the vegetative season (1 to 5) on the first
three principal components, obtained by polyphenol quantifications and profiles of leaf blade extracts
(see Table 4 for the list of used variables).

Table 5. Eigenvectors of nine variables (polyphenols in leaf veins) on the first three principal
components (PRIN1, PRIN2 and PRIN3).

433				
454		PRIN1	PRIN2	PRIN3
455	Total polyphenols	0.50	0.11	-0.14
456	Total phenolic acids	0.36	0.39	0.25
457	Total flavan-3-ols	0.38	0.32	0.38
458	Dimeric proanthocyanidins	0.30	-0.30	0.52
459	Total flavanonols	0.18	-0.56	0.21
460	Total flavones	-0.22	0.35	0.28
461	Sum of glucuronide flavonols	0.38	0.14	-0.33
463	Sum of glucoside flavonols	0.36	-0.38	-0.29
	Sum of rhamnoside flavonols	0.15	0.21	-0.42
464	Eigenvalues	3.27	2.17	1.34
465	Total variance	0.36	0.24	0.15

466 Eigenvalues of the three PRINs and their contribution to total variance. In bold letters, the variables associated to the467 appropriate PRIN.



468

Figure 11. Tridimensional distribution of *Vitis* species during the vegetative season (1 to 5) on the first
three principal components, obtained by polyphenol quantifications and profiles of leaf vein extracts
(see Table 5 for the list of variables used).

473 4. DISCUSSION

There is increasing interest in grapevine leaf polyphenols to deepen knowledge about their biological role in plant-defense mechanisms and to answer to the growing demand of natural bioactive compounds for the feed/food, pharmaceutical and cosmetic sectors. Grapevine leaves are already employed in the production of food ingredients, dietary supplements, pharmaceutical products and medicated cosmetics (Dani et al., 2010). The evolution of the polyphenolic composition of eight *Vitis* genotypes during the vegetative season allowed pointing out significant qualitative and quantitative differences.

481 **4.1 Genotype-related specific traits of leaf polyphenols**

The two studied American species, *V. riparia* and *V. rupestris*, belonging to the same subgeneric group of *Ripariae* (Table 1), involved in crossbreed of Regent, a cultivar displaying low susceptibility to *Plasmopara viticola* and to *Erysiphe necator*, shared common phenolic composition and accumulation trends. In particular, they showed similarities in the accumulation trend of total phenolic acids and total flavan-3-ols. Additionally, they displayed identical flavonol profiles (Figure 4). However, *V. riparia* accumulated higher concentration of total polyphenols, total flavonols and total phenolic acids
respect to *V. rupestris*, which could justify its specifically high resistance to downy mildew, as both
flavonols and phenolic acids have been associated with the ability to limit this pathogen diffusion (Ali
et al., 2012). Moreover, *V. riparia* total flavonol concentration both in blades and in veins was the
highest comparing to all the other analyzed genotypes, almost during the entire vegetative season (Figure
3).

The American genotypes Börner and V. berlandieri, known as low-susceptible to many diseases, 493 494 were classified by Galet (1988) into the same subgeneric serie of Cinereae (Table 1). Their main leaf polyphenolic traits showed many similarities, such as the concentration and the accumulation trend of 495 total polyphenols, total flavonols and total flavones, together with identical profiles of flavonols, 496 497 proanthocyanidins and flavones. Therefore, Galet's classification based on plant morphology matched 498 with phenol profile patterns of these two genotypes, suggesting a phylogenetic link between Vitis cinerea (Börner parent) and Vitis berlandieri. Flavones, bioactive compounds often extracted from 499 500 medical and herbaceous plants and involved in plant signaling and defense responses (Jiang et al., 2016), were exclusively detected in Börner and in V. berlandieri, among the studied genotypes. In grapevine, 501 they are generally considered as minor compounds mostly represented by luteolin, but in Börner and V. 502 *berlandieri* leaves their concentration reached 5 g kg⁻¹ DW (Figure 9) and showed a much wider profile, 503 504 consisting of five apigenin and luteolin derivatives: specifically, four of them were C-glycosides and 505 one O-glycoside (Table 3; Figure 9). C-glycoside flavonoids generally display higher antioxidant potential and better therapeutic properties compared to aglycons or to O-glycosylated flavonoids (Xiao 506 et al., 2016). However, as most studies focused on biological activities of the more common plant O-507 508 glycosylated-flavonoids, further research is required on grapevine C-glycosylated derivative health beneficial effect. McNally et al. (2003) demonstrated that in cucumber C-flavones acted as phytoalexins 509 in response to powdery mildew fungus Podosphaera xanthii and concentration increases of orientin and 510 isoorientin were reported to be induced by soil salinity in buckwheat (F. esculentum) (Yang et al., 2018). 511

V. *berlandieri* and Börner were also similar in the fact that they did not accumulate kaempferol
glycosides, in line with what was previously demonstrated in the grapevine series *Cinerea* (Moore and
Giannasi, 1987).

The Asian wild species V. coignetiae has been cultivated during the last two decades in Japan 515 for wine, juice and jam making (Kamiya et al., 2018). V. coignetiae leaf extract showed a strong radical-516 scavenging activity and potential hepato-protective effect on nonalcoholic steatohepatitis in liver 517 (Takayama et al., 2009). However, in V. coignetiae leaves of the present study, the concentration of leaf 518 519 polyphenols was generally lower comparing to the other genotypes (Figure 2). V. coignetiae along with V. amurensis and V. v. sylvestris was one of the few species which accumulated flavanonols; two 520 flavanonol compounds were detected in V. coignetiae, dihydrokaempferol-hexoside in veins and 521 522 dihydroquercetin-hexoside in both blades and veins (Figure 8A, B). Taxifolin (dihydroquercetin) glycoside is known to have anti-inflammatory activity (Kim et al., 2008) and potential positive effect 523 on atopic dermatitis treatment (Ahn et al., 2010). 524

The most peculiar species among the analyzed genotypes was V. amurensis, which is known for 525 its cold tolerance and disease resistance to downy mildew (Jürges et al., 2009), anthracnose and white 526 527 rot (Li et al., 2008). The leaves of V. amurensis reached the highest concentration of total polyphenols 528 and flavan-3-ols in blades, important amounts of flavonols, and the highest concentration of flavanonols in veins. V. amurensis leaves are used in conventional Chinese medicine and included in Korean herbal 529 530 Pharmacopoeia (Chen et al., 2018). Moreover, leaf extracts showed antimicrobial activity against Streptococcus mutans and Streptococcus sanguis (Yim et al., 2010) and neuroprotective effect (Jeong 531 et al., 2010). Bak et al. (2012, 2016) demonstrated that V. amurensis seed proanthocyanidins have 532 hepato-protective and antioxidative properties and a possible chemopreventive role in humans 533 534 hepatocarcinoma cells. In V. amurensis blades, concentration of total flavan-3-ols was significantly higher during the entire vegetative season comparing to the other genotypes (Figure 6A). Moreover, also 535 the concentration of (-)-epicatechin was particularly high. This compound was previously individuated 536 as a reaction of pear leaves against Erwinia amylovora and of apple leaves against Venturia inaequalis 537

(as discussed in Kedrina-Okutan et al., 2018). In non-Vitis vinifera genotypes, (-)-epigallocatechin was 538 additionally detected with increasing concentration during the vegetative season (Figure 6). 539 (Epi)gallocatechin-(epi)catechin was tentatively identified in V. amurensis, consistently with previously 540 published dimeric PA identification (Flamini, 2013). (Vagiri et al., 2017) reported a negative correlation 541 between (-)-epigallocatechin and the presence of septoria leaf spot caused by Mycosphaerella ribis in 542 black currant leaves. The high concentration of constitutive polyphenolic compounds throughout the 543 544 season in V. amurensis leaves contributes to explain this biotype excellent ability to protect against adverse environmental condition (frost and diseases). Furthermore, the abundance of polyphenols in V. 545 546 amurensis leaves emphasizes that this species has the potential for further use as a source of natural bioactive compounds for nutraceutical and pharmacological uses. 547

The polyphenolic composition of V. candicans leaves was studied here for the first time, to the 548 549 best of our knowledge. This American species has been described as highly vigorous, highly resistant 550 to downy mildew and tolerant to drought and salt (Ollat et al., 2016). V. candicans leaves displayed a low concentration of total polyphenols, total flavonols, but a relatively high concentration of 551 proanthocyanidins and this genotype was among the few able to accumulate procyanidin B_1 in veins 552 (Figure 7). Procyanidin B₁, the dimeric proanthocyanidin consisting in units of (-)-epicatechin and (+)-553 catechin, has an ecological significance in protecting plants against pathogens, insect pests and larger 554 herbivores (Dixon et al., 2005). For instance, previous studies demonstrated that B. cinerea remained 555 556 quiescent in immature strawberry until B proanthocyanidins are present at an essential high 557 concentration: this effect is related to the proanthocyanidin inhibitory potential of the fungal polygalacturonase (Jersch et al., 1989). 558

The Eurasian wild species *V. v. sylvestris*, the ancestor of the worldwide cultivated *V. vinifera sativa*, accumulated relatively high concentration of total polyphenols and total phenolic acids (Figure 2; Figure 5). Among phenolic acids, the predominant non-flavonoid polyphenols, caftaric acid was the prevalent form, representing up to 90% of total phenolic acids (Figure 5C), similarly to what was previously assessed in *V. vinifera sativa* leaves (Kedrina-Okutan et al., 2018). Caftaric acid is an 564 important bioactive component found in chicory, in the medical plant Echinacea purpurea (Bel-Rhlid et al., 2012) and, in V. coignetiae juice, where it exhibited anti-mutagenic and anti-inflammatory 565 properties (Kamiya et al., 2018). In V. v. sylvestris leaves, we found astilbin (Figure 8B), in 566 concentrations that were in line with those found in Cabernet Sauvignon, Grenache, Shiraz and Barbera, 567 that were much lower respect to those found in Nebbiolo and Pinot noir (Kedrina-Okutan et al., 2018). 568 Additionally, V. v. sylvestris displayed the most complex flavonol profile (Figure 4). In V. v. sylvestris 569 veins we identified and quantified seven flavonols out of the eight flavonol compounds 570 chromatographically separated; V. v. sylvestris was the exclusive genotype that accumulated kaempferol 571 572 3-O-rhamnoside (Figure 4, Table 3) and, compared to V. vinifera sativa, it accumulated additional flavonol rhamnosides (kaempferol- and quercetin-3-O-rhamnoside), showing a wider flavonol-profile 573 574 diversification.

575 The peculiar traits of individual Vitis species leaf polyphenolic compositions were highlighted by the proposed PCA models. In particular, V. riparia and V. rupestris displayed similar eigenvectors 576 on PRIN1 and PRIN3, highlighting their similar concentrations of dimeric proanthocyanidins, similar 577 percentage incidence of caftaric acid over total phenolic acids (on PRIN1) and similar concentration of 578 flavan-3-ols on PRIN3 (Table 4; Figure 10). V. riparia sharply distinguished on PRIN2 due its high 579 580 flavonol content (Figure 3A) and the highest percentage incidence of myricetin derivatives respect to the other analyzed genotypes (Figure 4). Flavonols allowed the species separation, also when 581 582 considering their concentration: in fact, flavonol glucosides, together with flavanonols, allowed a sharp 583 separation of V. amurensis samples and were negatively associated to the second PRIN of the vein PCA model (Figure 11; Table 5). Oppositely, V. berlandieri and Börner associated to positive values of 584 PRIN2, with reduced amounts of flavonol-glucosides comparing to the other species. Quercetin 3-O-585 glucoronide was the prevalent flavonol in the analyzed genotypes, consistently with other studies on V. 586 vinifera and V. labrusca leaves (Dresch et al., 2014; Kedrina-Okutan et al., 2018). The differences in 587 the flavonol profile might be related to the genus *Vitis* evolution: in fact, evidences about the reduction 588 in structural complexity and in molecule diversification were reported (Moore and Giannasi, 1994), and 589

590 they were also ascribed to the flavonoid biosynthetic pathway, that was shown to have undergone a simplification during evolution, particularly in domesticated genotypes. Interestingly, V. v. sylvestris, 591 which, being the wild form of V. v. sativa, underwent no or low selection pressure, showed the widest 592 593 flavonol profile diversification and comprised high concentration of guercetin 3-O-rhamnoside (22% of total flavonols); besides, V v. sylvestris also displayed a higher flavonol profile diversity respect to 594 cultivated varieties of V. vinifera. Quercetin 3-O-rhamnoside was also found in V. berlandieri and in 595 Börner. Hilbert et al. (2015) identified flavonol rhamnosides in berry skin extracts of V. cinerea, which 596 is native to the United States and belongs to the subgeneric group of *Cinereae* (Galet, 1988), the same 597 598 of V. berlandieri and Börner. Thus, flavonol rhamnosides seem to be genotype-specific compounds, therefore they have the potential to be exploited as specific markers of some species and/or subspecies. 599

600 Vein flavanonol concentration, associated to PRIN2 (Figure 11; Table 5) allowed the sharp 601 separation of V. amurensis, and, secondarily the separation of V. coignetiae and V. v. sylvestris. Since 602 flavanonols accumulate prevalently in leaf veins, their concentration on a per whole leaf basis is supposed to be low, considering the limited percentage incidence of veins on the total leaf area, but the 603 604 role of these tissue-specific compounds arouses the interest about their interaction with vein-located pathogens. In some V. vinifera sativa varieties, we previously identified these molecules, particularly in 605 veins (Kedrina-Okutan et al., 2018) where astilbin was the predominant compound. We also associated 606 astilbin accumulation to a possible defense-mechanism in cv Nebbiolo (Ferrandino et al., 2019). In fact, 607 in Nebbiolo leaves of plants affected by Flavescence dorée (FD), we found a significantly higher 608 609 concentration of flavanonols compared to healthy vines, hypothesizing a possible capability to limit the pathogen development. Interestingly, together with V. v. sylvestris, also V. amurensis, whose capability 610 to defend against pathogens is detailed and V. coignetiae, known for its strong antiradical role, 611 612 accumulate flavanonols.

613 As to the ecological role of leaf polyphenols, further studies would be of extreme interest to 614 understand: i) if genotype-related molecules such as flavanonols and flavones have specific capacities to limit diseases, starting from *in vitro* trials, ii) if grapevine plants can be induced to produce specific
molecules, once demonstrated that they can efficaciously serve to limit pathogen development.

As to the nutraceutical, pharmaceutical and feed/food ingredients, grapevine leaf are a source of
bioactive compounds that in future could be economically exploited, also exploiting their wide and still
little investigated, biodiversity.

620

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- 624 REFERENCES
- Abascal, K., Ganora, L., Yarnell, E., 2005. The effect of freeze-drying and its implications for botanical
 medicine: a review. Phyther. Res. 19, 655–660. https://doi.org/10.1002/ptr.1651
- Ahn, J.Y., Choi, S.E., Jeong, M.S., Park, K.H., Moon, N.J., Joo, S.S., Lee, C.S., Choi, Y.W., Kapsok,
- L., Lee, M.-K., Lee, M.W., Seo, J.S., 2010. Effect of taxifolin glycoside on atopic dermatitis-like
 skin lesions in NC/Nga mice. Phyther. Res. 24, 1071–1077. https://doi.org/10.1002/ptr
- Ali, K., Maltese, F., Figueiredo, A., Rex, M., Margarida, A., Zyprian, E., Salomé, M., Verpoorte, R.,
- 631 Choi, Y.H., 2012. Alterations in grapevine leaf metabolism upon inoculation with *Plasmopara*
- 632 *viticola* in different time-points. Plant Sci. 191–192, 100–107.
 633 https://doi.org/10.1016/j.plantsci.2012.04.014
- Bak, M., Jun, M., Jeong, W., 2012. Procyanidins from wild grape (Vitis amurensis) seeds regulate ARE-
- mediated enzyme expression via Nrf2 coupled with p38 and PI3K/Akt pathway in HepG2 cells
 801–818. https://doi.org/10.3390/ijms13010801
- Bak, M.J., Truong, V.-L., Ko, S.-Y., Nguyen, X.N.G., Ingkasupart, P., Jun, M., Shin, J.Y., Jeong, W.-
- 638 S., 2016. Antioxidant and hepatoprotective effects of procyanidins from wild grape (Vitis

- 639 *amurensis*) seeds in ethanol-induced cells and rats. Int. J. Mol. 17, 758.
 640 https://doi.org/10.3390/ijms17050758
- Bel-Rhlid, R., Page-Zoerkler, N., Fumeaux, R., Ho-Dac, T., Chuat, J.-Y., Sauvageat, J.L., Raab, T., 641 2012. Hydrolysis of chicoric and caftaric acids with esterases and Lactobacillus johnsonii in vitro 642 gastrointestinal 9236-9241. and in a model. J. Agric. Food Chem. 60, 643 644 https://doi.org/10.1021/jf301317h
- Chen, Q., Diao, L., Song, H., Zhu, X., 2018. *Vitis amurensis* Rupr: A review of chemistry and
 pharmacology. Phytomedicine 49, 111–122. https://doi.org/10.1016/j.phymed.2017.08.013
- Dani, C., Oliboni, L.S., Agostini, F., Funchal, C., Serafini, L., Henriques, J.A., Salvador, M., 2010.
- Phenolic content of grapevine leaves (*Vitis labrusca* var. Bordo) and its neuroprotective effect
 against peroxide damage. Toxicol. Vitr. 24, 148–153. https://doi.org/10.1016/j.tiv.2009.08.006
- Di Stefano, R., Cravero, M.C., 1992. The separation of hydroxycinnamates in wine. Sci. des Aliment.
 139–144.
- Dixon, R., Xie, D.-Y., Sharma, S.B., 2005. Proanthocyanidins a final frontier in flavonoid research?
 New Phytol. 165, 9–28.
- Dixon, R.A., 2001. Natural products and plant disease resistance. Nature 411, 843–847.
 https://doi.org/10.1038/35081178
- 656 Dresch, R.R., Dresch, M.K., Guerreiro, A.F., Biegelmeyer, R., Holzschuh, M.H., Rambo, D.F.,
- 657 Henriques, A.T., 2014. Phenolic compounds from the leaves of *Vitis labrusca* and *Vitis vinifera* L.
- as a source of waste byproducts: development and validation of LC method and antichemotactic
 activity. Food Anal. Methods 7, 527–539. https://doi.org/10.1007/s12161-013-9650-4
- Ferrandino, A., Guidoni, S., 2010. Anthocyanins, flavonols and hydroxycinnamates: An attempt to use
 them to discriminate *Vitis vinifera* L. cv "Barbera" clones. Eur. Food Res. Technol. 230, 417–427.

662 https://doi.org/10.1007/s00217-009-1180-3

- Ferrandino, A., Pagliarani, C., Kedrina-Okutan, O., Icardi, S., Bove, M., Lovisolo, C., Novello, V., 663 Schubert, A., 2019. Non-anthocyanin polyphenols in healthy and Flavescence dorée infected 664 Barbera and Nebbiolo leaves. BIO Web Conf. 03003. 1-5. 665 https://doi.org/https://doi.org/10.1051/bioconf/20191303003 666
- Flamini, R., 2013. Recent applications of mass spectrometry in the study of grape and wine polyphenols.
 ISRN Spectrosc. 2013, 45. https://doi.org/10.1155/2013/813563
- Galet, P., 1988. In Cepages Et Vignobles De France [Grapes and vineyards of France], 2nd ed.
 Imprimerie Charles Dehan, Montpellier.
- Gómez-Zeledón, J., Zipper, R., Spring, O., 2013. Assessment of phenotypic diversity of *Plasmopara viticola* on *Vitis* genotypes with different resistance. Crop Prot. 54, 221–228.
 https://doi.org/10.1016/j.cropro.2013.08.015
- Hilbert, G., Temsamani, H., Bordenave, L., Pedrot, E., Chaher, N., Cluzet, S., Delaunay, J.C., Ollat, N.,
- 675 Delrot, S., Mérillon, J.M., Gomès, E., Richard, T., 2015. Flavonol profiles in berries of wild *Vitis*
- accessions using liquid chromatography coupled to mass spectrometry and nuclear magnetic
 resonance spectrometry. Food Chem. 169, 49–58. https://doi.org/10.1016/j.foodchem.2014.07.079
- Hmamouchi, M., Es-Safi, N., Lahrichi, M., Fruchier, A., Essassi, E.M., 1996. Flavones and flavonols in
 leaves of some Moroccan *Vitis vinifera* cultivars. Am. J. Enol. Vitic. 47, 186–192.
 https://doi.org/10.1071/PP9960115
- Jeong, H.Y., Kim, J.Y., Lee, H.K., Ha, D.T., Song, K.S., Bae, K., Seong, Y.H., 2010. Leaf and stem of
 Vitis amurensis and its active components protect against amyloid β protein (25-35)-induced
 neurotoxicity. Arch. Pharm. Res. 33, 1655–1664. https://doi.org/10.1007/s12272-010-1015-6
- Jersch, S., Scherer, C., Huth, G., Schlosser, E., 1989. Proanthocyanidins as basis for quiescence of

- *Botrytis cinerea* in immature strawberry fruits. J. Plant Dis. Prot. 96, 365–378.
- Jiang, N., Doseff, A., Grotewold, E., 2016. Flavones: from biosynthesis to health benefits. Plants 5, 2752. https://doi.org/10.3390/plants5020027
- Jürges, G., Kassemeyer, H.H., Dürrenberger, M., Düggelin, M., Nick, P., 2009. The mode of interaction
- 689 between *Vitis* and *Plasmopara viticola* Berk. & Curt. Ex de Bary depends on the host species. Plant
- 690 Biol. 11, 886–898. https://doi.org/10.1111/j.1438-8677.2008.00182.x
- 691 Kamiya, T., Tanimoto, Y., Fujii, N., Negishi, T., Suzuki, T., Hatano, T., Arimoto-Kobayashi, S., 2018.
- 692 2,6-Dimethoxy-1,4-benzoquinone, isolation and identification of anti-carcinogenic, anti-mutagenic
- and anti-inflammatory component from the juice of *Vitis coignetiae*. Food Chem. Toxicol. 122,
- 694 172–180. https://doi.org/10.1016/j.fct.2018.10.028
- Kedrina-Okutan, O., Novello, V., Hoffmann, T., Hadersdorfer, J., Occhipinti, A., Schwab, W.G.,
 Ferrandino, A., 2018. Constitutive polyphenols in blades and veins of grapevine (*Vitis vinifera* L.)
- 697 healthy leaves. J. Agric. Food Chem. 66, 10977–10990. https://doi.org/10.1021/acs.jafc.8b03418
- Kim, B., Woo, S., Kim, M.J., Kwon, S.W., Lee, J., Sung, S.H., Koh, H.J., 2018. Identification and
- 699 quantification of flavonoids in yellow grain mutant of rice (*Oryza sativa* L.). Food Chem. 241,
- 700 154–162. https://doi.org/10.1016/j.foodchem.2017.08.089
- Kim, Y.J., Choi, S.E., Lee, M.W., Lee, C.S., 2008. Taxifolin glycoside inhibits dendritic cell responses
 stimulated by lipopolysaccharide and lipoteichoic acid. J. Pharm. Pharmacol. 60, 1465–1472.
 https://doi.org/10.1211/jpp/60.11.0007
- Li, D., Wan, Y., Wang, Y., He, P., 2008. Relatedness of resistance to anthracnose and to white rot in
 Chinese wild grapes. Vitis 47, 213–215. https://doi.org/10.1073/pnas.96.3.1146
- Li, Z.H., Guo, H., Xu, W. Bin, Ge, J., Li, X., Alimu, M., He, D.J., 2016. Rapid identification of flavonoid
- constituents directly from PTP1B inhibitive extract of raspberry (*Rubus idaeus* L.) leaves by

- 708
 HPLC-ESI-QTOF-MS-MS.
 J.
 Chromatogr.
 Sci.
 54,
 805–810.

 709
 https://doi.org/10.1093/chromsci/bmw016

 </t
- Llorens, E., García-Agustín, P., Lapeña, L., 2017. Advances in induced resistance by natural
 compounds: towards new options for woody crop protection. Sci. Agric. 74, 90–100.
 https://doi.org/10.1590/1678-992x-2016-0012
- McNally, D.J., Wurms, K. V., Labbé, C., Bélanger, R.R., 2003. Synthesis of *C*-glycosyl flavonoid
 phytoalexins as a site-specific response to fungal penetration in cucumber. Physiol. Mol. Plant
 Pathol. 63, 293–303. https://doi.org/10.1016/j.pmpp.2004.03.005
- Moore, M.O., Giannasi, D.E., 1994. Foliar flavonoids of eastern North American *Vitis (Vitaceae)* north
 of Mexico. Plant Syst. Evol. 193, 21–36.
- Moore, M.O., Giannasi, D.E., 1987. Foliar flavonoids of selected Vitis taxa in the Southeastern United
 States. Biochem Syst Ecol 15, 79–83.
- Ollat, N., Bordenave, L., Tandonnet, J.P., Boursiquot, J.M., Marguerit, E., 2016. Grapevine rootstocks:
 origins and perspectives. Acta Hortic. 1136, 11–22.
 https://doi.org/10.17660/ActaHortic.2016.1136.2
- Pastrana-Bonilla, E., Akoh, C.C., Sellappan, S., Krewer, G., 2003. Phenolic content and antioxidant
 capacity of muscadine grapes. J. Agric. Food Chem. 51, 5497–5503.
 https://doi.org/10.1021/jf030113c
- Ruel, J.R., Walker, M.A., 2006. Resistance to Pierce's disease in *Muscadinia rotundifolia* and other
 native grape species. Am. J. Enol. Vitic. 32, 155–158.
- 728 Singleton, V.L., Orthofer, R., Lamuela-Raventós, R.M., 1999. Analysis of total phenols and other
- oxidation substrates and antioxidants by means of Folin-Ciocalteu reagent. Methods Enzymol. 299,
- 730 152–178. https://doi.org/10.1016/S0076-6879(99)99017-1

- Staudt, G., 1997. Evaluation of resistance to grapevine powdery mildew (*Uncinula necator* [Scxw.]
 Burr., anamorph *Oidium tuckeri* BERK.) in accessions of *Vitis* species. Vitis 36, 151–154.
- Takayama, F., Nakamoto, K., Kawasaki, H., Mankura, M., Egashira, T., Ueki, K., Hasegawa, A., Okada, 733 S., Mori, A., 2009. Beneficial effects of Vitis coignetiae Pulliat leaves on nonalcoholic 734 steatohepatitis Med. Okayama 63. 105–111. 735 in model. Acta а rat 736 https://doi.org/10.18926/AMO/31835
- Vagiri, M., Johansson, E., Rumpunen, K., 2017. Phenolic compounds in black currant leaves an
 interaction between the plant and foliar diseases? J. Plant Interact. 12, 193–199.
 https://doi.org/10.1080/17429145.2017.1316524
- Wan, Y., Schwaninger, H., He, P., Wang, Y., 2007. Comparison of resistance to powdery mildew and
 downy mildew in Chinese wild grapes. Vitis 46, 132–136.
- Wan, Y., Schwaninger, H.R., Baldo, A.M., Labate, J.A., Zhong, G.Y., Simon, C.J., 2013. A
 phylogenetic analysis of the grape genus (*Vitis* L.) reveals broad reticulation and concurrent
 diversification during neogene and quaternary climate change. BMC Evol. Biol. 13, 141–161.
 https://doi.org/10.1186/1471-2148-13-141
- Xiao, J., Capanoglu, E., Jassbi, A.R., Miron, A., 2016. Advance on the flavonoid C-glycosides and
 health benefits. Crit. Rev. Food Sci. Nutr. 56, 29–45.
 https://doi.org/10.1080/10408398.2015.1067595
- Yang, L., Wen, K.S., Ruan, X., Zhao, Y.X., Wei, F., Wang, Q., 2018. Response of plant secondary
 metabolites to environmental factors. Molecules 23, 762-788.
 https://doi.org/10.3390/molecules23040762
- Yim, N., Thi, D., Nam, T., Pyo, J., Lee, S., Na, M., Jung, H., Su, H., Ho, Y., Bae, K., 2010. Bioorganic
 & Medicinal Chemistry Letters The antimicrobial activity of compounds from the leaf and stem of *Vitis amurensis* against two oral pathogens. Bioorg. Med. Chem. Lett. 20, 1165–1168.

- 755 https://doi.org/10.1016/j.bmcl.2009.12.020
- Zhang, J., Wu, X., Niu, R., Liu, Y., Liu, N., Xu, W., Wang, Y., 2012. Cold-resistance evaluation in 25
 wild grape species. Vitis J. Grapevine Res. 4, 153–160.
 https://doi.org/10.1017/S0020743810001558