

Contribution of soil elemental contents and Cu and Sr isotope ratios to the understanding of pedogenetic processes and mechanisms involved in the soil-to-grape transfer (Soave vineyard, Italy)

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1	Contribution of soil elemental contents and Cu and Sr isotope ratios to the understanding of
2	pedogenetic processes and mechanisms involved in the soil-to-grape transfer (Soave vineyard, Italy)
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11	
12	Abstract
13	The effect of soil on wine composition and flavor is controversially discussed in viticulture. Mineral
14	nutrition of the grapevine is one possibility for an influence of soil chemistry on winemaking. However,
15	effects of soil chemistry are difficult to isolate from other physical and biological factors.
16	Here two winegrowing plots in the Soave region (Italy) were investigated, lying side by side on
17	contrasted soil types. Factors that influenced soil formation and characterize soil chemistry and
18	mineralogy were determined. The influence of soil type on the elemental composition (Mg, Al, P, S, K,

Ca, Mn, Fe, Cu, Zn, Sr, Mo and Ba) of leaves and grapes was studied. Carbon isotope ratios were
 measured to evaluate water stress. Radiogenic Sr and stable Cu isotope ratios were studied to identify

21 the influence of different bedrocks and better understand the mechanisms involved in the soil-to-grape

22 continuum.

23	Even though the morphology of soils is different, chemical characteristics are similar in both vineyard
24	plots. Nevertheless, Sr isotope ratios of horizons of both soils show influence of different bedrocks on
25	their genesis. The composition of grapevine plants is similar between both plots even though there is a
26	tendency for higher elemental contents on more calcareous soil. Finally, isotope ratios show that
27	different mechanisms control Cu and Sr in plant: Cu seems to be controlled by regulation mechanisms of
28	the plants whereas Sr is absorbed in ratios similar to the soil, reflecting the different geological origins.
29	
30	Keywords: vineyard soil; mineral nutrition; soil formation; element transfers; Sr and Cu isotopes
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45 One way of possible influence of soil chemistry on wine quality is plant nutrition and translocation of 46 elements through the plant (Maltman, 2013). Some 17 elements are essential nutrients for plant 47 functioning and are taken up from the soil (Marschner and Marschner, 2012). Concentrations of most 48 nutrients are closely regulated in plant tissues in a process called homeostasis (Marschner and 49 Marschner, 2012). By this self-regulation process (including particularly mineral nutrition, temperature 50 adjustment, transpiration function, etc.), a plant can maintain biological conditions and stability of its 51 dynamic equilibrium facing changes occurring in the environment (Castro et al., 2018). Mineral nutrients 52 play important roles in biochemical plant functioning as catalyzers, electron transporters or regulators of osmotic pressure (Marschner and Marschner, 2012). However almost the whole periodic table of 53 54 elements can be found in plants and element concentrations vary systematically between different 55 wines (Almeida and Vasconcelos, 2001; Coetzee et al., 2014; Day et al., 1995; Greenough et al., 2005; 56 Kwan et al., 1979). For example, soil carbonate content is an important factor influencing plant 57 functioning, as certain nutrients are scarcely available (especially P and Fe) in carbonated environments and plants have to deploy specific mechanisms to satisfy their needs (Marschner and Marschner, 2012; 58 59 Strom, 1997). Moreover, interaction among plant nutrients can involve antagonistic or synergistic effects 60 that potentially influence element efficiency in the plant system (Rietra et al., 2017). So, mineral nutrition is a complex process and soil-to grape transfer mechanisms in vineyards are the result of 61 62 several interactions and processes, still poorly described (Likar et al., 2015).

Although differences in elemental contents of wines or grapes, likely do not have a proper taste, smaller variations in element availability may influence the synthesis of compounds essential for grape maturity and wine taste (Epke and Lawless, 2007; Sipos et al., 2012). For example the K/Ca equilibrium is important for acidity in wine but K was also found to be important in polyphenol synthesis (Brunetto et al., 2015; Daudt and Fogaça, 2008). Furthermore, many elements act as nutrients for fermenting yeasts or catalyzers for synthesis of aromatic compounds (Pohl, 2007). Variations in metal ion contents have

been shown to influence grape properties as sugar and essential amino acid content in grapes, mustsand wines (Pereira, 1988).

71 Mobility and phytoavailability of elements do not only depend on the soil content but are controlled by 72 various soil properties such as pH, Eh, cation exchange capacity or organic matter content (Kabata-73 Pendias, 2004; Tyler and Olsson, 2001). Therefore, a thorough pedologic investigation is needed to 74 identify soil properties controlling plant nutrition. Soil hydrological parameters have also been reported 75 to widely influence mineral nutrition and fruit composition, as a resultant of soil-plant transfer (Ramos 76 and Martínez-Casasnovas, 2006; Tournebize et al., 2012; van Leeuwen et al., 2009). Elemental contents 77 of grapevine organs and their link with soil properties have been subject to several studies, some of 78 them succeeded in identifying a direct link between soil concentrations or extractions whereas other do 79 not (Angel Amoros et al., 2013; Cugnetto et al., 2014; Likar et al., 2015; Mercurio et al., 2014; Vazquez 80 Vazquez et al., 2016). Only few studies have investigated grapevine nutrition on contrasted soils 81 (Mackenzie and Christy, 2005; Peuke, 2000). In these studies differences in elemental contents and 82 winemaking parameters have been identified (Mackenzie and Christy, 2005; Peuke, 2000). However, the 83 vineyards investigated in these studies lie several kilometers apart, thus meteorological conditions were 84 likely different, and no information is given on slope or exposition to sunlight. Thus, as it has been 85 demonstrated that meteorological factors influence elemental compositions of grapevine (Boselli et al., 86 1998), a new perspective could be to work on the same climate parameters to better isolate the impact of elemental soil-to-grape transfers. 87

Besides elemental techniques, isotope ratios, and especially Sr isotopes, have been successfully used to determine geographical origin of wines, firstly studied by Horn et al. (1993) and more recently by various ltalian, Portuguese or Canadian research teams (Martins et al., 2014; Petrini et al., 2015; Marchionni et al., 2013; 2016; Vinciguerra et al., 2016; Braschi et al., 2018). Actually, radiogenic Sr isotope ratios between ⁸⁷Sr and ⁸⁶Sr have been used to trace sources in geology and soil sciences for many years. Also

in vineyard environments Sr-isotope ratios have been used to support wine quality and guarantee good
traceability (Durante et al., 2018, 2015, 2013; Tescione et al., 2018). This is possible because stable Sr
isotope fractionation related to common low temperature reactions appears to be negligible compared
to source signature, also in biological systems (Blum et al., 2000; Capo et al., 1998). This makes Sr isotope
ratios a valuable tracer of the source of Ca nutrition, as the chemical behavior of the two elements is
extremely similar (Blum et al., 2000; Poszwa et al., 2000; Schmitt et al., 2017). This is especially
interesting when carbonates are involved in mineral nutrition of grapevine plants.

100 Moreover, recent studies have underlined the potential use of Cu isotope analysis in the context of 101 polluted agricultural soils (Babcsányi et al., 2016; Fekiacova et al., 2015) and their help in determining 102 the mechanisms of Cu mobility in soils (Blotevogel et al., 2018) as well as the mechanisms of elemental 103 transport towards plants or trees (Mihaljevič et al., 2018). The discussion on Cu isotope fractionation 104 induced by plants is still ongoing, but first results reported that Cu isotopes fractionate considerably 105 during uptake and translocation: they have been recently used to identify Cu uptake mechanisms (Jouvin 106 et al., 2012), including reduction at root surfaces and preferential translocation of light isotopes to 107 higher foliar levels (Weinstein et al., 2011). Diverging results from different growth settings also suggest 108 that fractionation of Cu isotopes depends on environmental conditions (Jouvin et al., 2012; Li et al., 109 2016; Ryan et al., 2013; Weinstein et al., 2011). Cu isotope analysis is thus an interesting tool to investigate Cu mobility in vineyard environments. 110

The present study aims at investigating soil formation and mineralogical as well as geochemical differences occurring between two adjacent plots in the Soave area (Veneto, Italy), using complementary techniques. Because different bedrock types underlie the soils of the two plots, the study of the pedological characteristics and of the elemental distribution along the soil-to-grape continuum will allow us to establish a link between soil properties and plant composition under the same metereological and

agronomic conditions. In particular, the use of Sr and Cu isotopes will help to better identify thebiogeochemical processes and elemental transfers occurring in this vineyard soil-plant system.

118

119 Materials and Methods

120 Area of study

121 The study was performed in the Cantina Filippo Filippi domain, in the Castelcerino wine growing area, in 122 the Soave appellation (45.464911 N, 11.236905 E). The Soave vineyard is located in the southern 123 foothills of the Alps, in Northern Italy, in the Veneto region. Geologically, the studied site lies between 124 different bedrocks: basalts and volcanic tuff mainly in the Alpone valley and limestone in the western 125 parts of the appellation area (Benciolini et al., 2006). Volcanic rocks come from tertiary volcanic activity 126 during the alpine orogenesis with the closest eruptive centers in Alpone valley and north of Cazzano di Tramigna village (De Vecchi et al., 1976). Limestones are from paleogenic and lower Miocene but 127 128 siliclastic marine sediments and marls are also found in the area as some parts of the Lessini shelf 129 became emerged in early Oligocene whilst others remained on the slope of Thetys ocean (Bassi and 130 Nebelsick, 2010; De Vecchi et al., 1976). Basalt outcrops are mainly found to the northeast, tufa to the 131 north and west and limestones in the south and west of the vineyard (Figure 1).

The plots are cultivated with the typical Garganega cultivar in organic viticulture by the same vine grower. Treatment, pruning and harvesting intervals differ only by days and treatment amounts are the same on both plots. Both plots are in viticulture for 65 years. Plant material is still the first generation planted thus no deep ploughing occurred since the beginning of cultivation. Vineyard management has been organic for the last 12 years and grapevine plants are irrigated in case of severe droughts. The soil between grapevine rows is grass covered.

138

139 Field pedological description

140 Soil profiles were studied along two catenas within the two studied parcels (Figure 1), following the main 141 slope. Soil profiles descriptions and pedological investigations were performed both through soil pits 142 until a depth of 50-60 cm for a more complete surface description and an Edelman soil corer was used 143 down to the depth of 120 cm maximum for subsoil study. Structure, texture, roots abundance, biological 144 activity and pedological features (e.g., mottling, coatings, secondary phases precipitations, root and 145 pores distributions) were described directly on the field (i.e., at field moisture). Soil structures below 50 146 cm depth are indicative structures as structure observation is biased during soil corer sampling. Colors 147 were determined using a Munsell color chart in the year 2000 revised version (Munsell, 2000). The 148 occurrence of carbonate phases and their types (primary versus secondary carbonates phases) was 149 established using 1 M HCI (effervescence method). Samples for bulk densities of surface horizons of B60, 150 B150, C60 and C155 were taken by the cylinder method. Soils were classified according to the World 151 Reference Base for soil resources from FAO/ISRIC/ISSS, 2014 (2015 update).

152

153 Hydrological and geomorphological properties

As reported by van Leeuwen et al. (2009), vine water status and especially water deficit stress are key factors of mineral nutrition and grape composition. Among the various methods largely employed to assess the hydric stress, stable isotope ratio in carbon (δ^{13} C) are reported to be highly significant (Ramos, 2017; Ramos and Martínez-Casasnovas, 2006; Tournebize et al., 2012; van Leeuwen et al., 2009).

158 δ^{13} C, the 13 C/ 12 C ratio expressed relative to the Pee Dee Belimnite standard (PDB), has been measured as 159 an integrative indicator of water status of grapevine plants. As explained by Farquhar et al., 1989 , 12 C is

160 preferentially taken up by the enzymes involved in photosynthesis and « isotope discrimination » 161 between air and plant occurs. In water stress conditions, this discrimination is less severe and plants contain more ¹³C compared to those produced when plant water status is not limiting. Van Leeuwen et 162 163 al. (2009) linked δ^{13} C ratios in grapevine plants to stem water potential, sugar content and acidity. That article also defines threshold values of no water stress (δ^{13} C < -27 ‰) and severe water stress (δ^{13} C > -20 164 ‰). However, it was reported that δ^{13} C values are dependent on grapevine variety (Gómez-Alonso and 165 García-Romero, 2009). In this study, δ^{13} C have been calculated with 13 C/ 12 C values measured on ground 166 leaf and grape samples, on an Isoprime 100 IRMS by Elementar[®] according to the Equation (1): 167

168 Equation (1)
$$\delta^{13}C = \left(\frac{\binom{1^{3}C}{^{12}C}_{sample}}{\binom{1^{3}C}{^{12}C}_{PDB}} - 1\right) * 1000$$

Hydric stress occurs when the soil is dry enough, and thus the rate of flow of water out of the soil is a key determinant on the occurrence of water deficit stress. The first order controls on the rate of drainage of soils are their hydraulic conductivity and the topographical slope: the higher they are, the faster will be the drainage. Here, hydraulic conductivity at saturation was estimated by infiltration tests at constant hydraulic head (Porchet and Laferre, 1935) in 45 cm deep standardized boreholes for the different sampling points from C and B catenas, as already reported by Blotevogel et al. (2018).

Moreover, in order to estimates slope distributions, the studied vineyard was mapped using a kinetic differential GPS and an elevation map grid was calculated by kriging between 47968 data points using Surfer 13 software package to propose a vector map indicating slope direction.

All these parameters allow a characterization of the two vineyard plots in terms of their hydrological andgeomorphological behaviors.

180

181

182 Sampling of soils, plants and rocks

183 Soil sampling was performed following the two catenas identified in Figure 1: "B" (basalt) catena for soils 184 over basaltic bedrock (in blue) and "C" (carbonate) catena for soils over calcareous bedrock (in red). The 185 numbers identifying each sampling spot correspond to the distances in meters from the gravel road on 186 top of the parcels. Rock samples of limestone, tufa and basalt were collected from the respective 187 outcrops denoted in geological maps of the Soave region and described by Benciolini et al. (2006), and 188 marked in Figure 1. Limestone outcrops are situated within the C catena, basalt outcrops are found uphill 189 from the B catena and tufa outcrops uphill from the C one. This sampling strategy aimed at determining 190 an endmember of each parent material to study pedogenetic mechanisms occurring in this area.

Leaves and grapes were sampled at the end of September (2015), just before grape harvest, as composites from 5 plants surrounding the soil sampling site. Only tissues that visually appeared healthy were sampled at chest height (~ 150 cm from soil) using a ceramic knife. Leaves were washed three times in ultrapure water and thereafter frozen and freeze dried. Grapes were sampled including the pedicel and rinsed with ultrapure water and subsequently freeze dried. Plant samples were ground under liquid nitrogen using an agate mortar.

197

198 Soil physico-chemical properties

Soil samples were air-dried for several days under a laminar flow hood and sieved to 2 mm. Soil pH was measured on 1 g in ultrapure water (18.2 M Ω) following ISO 11464 protocol. The cation exchange capacity (CEC) was measured according to the cobalthexamin method, described by Orsini and Remy (1976), Ciesielski and Sterckeman (1997) and Arrouays et al. (2011): 1 g of soil was shaken in 20 mL of a 0.017 mol L⁻¹ cobalthexamine solution during 1 h, solutions were subsequently centrifuged and supernatant filtered at 0.22 µm. Then cobalthexamine loss from solution was determined by absorbance

205 loss at 475 nm with a Varian Cary 50 spectrophotometer. Soil Organic Carbon and Soil Inorganic Carbon 206 (respectively SOC and SIC) were calculated after subsequent measures of raw and calcined samples on a 207 Horiba EMIA – 320V CS automate, according to the method reported in Bao et al. (2017), Blotevogel et 208 al. (2018) and Kania et al. (2019). For granulometric analysis, 2 g of soil samples were suspended in 209 distilled water and sieved to 500 µm. The fine fraction was subsequently treated with 10 mL hydrogen 210 peroxide to remove organic matter. Treatment was repeated until no foam or gas formation was 211 observed when adding H₂O₂. Samples were treated with an acetic acid/ammonium acetate buffer at pH 212 ~4.5 to eliminate carbonates phases. Treatment was then repeated until no effervescence was visible. 213 Finally, samples were washed in ultrapure water and grain size distribution was measured on a Horiba 214 LA-950 laser diffraction granulometer, as reported by Wanogho et al. (1987). Soil bulk densities were 215 determined after soil drying at 105°C during 48h (until constant mass is reached).

216

217 Mineralogy of rock and soil samples

218 Rock samples were washed with ultrapure water. Approximately 200 g of sieved soil samples (<2 mm 219 fraction, i.e., fine earth) and rock samples were crushed using a planetary mill made from agate. X-Ray 220 Diffraction (XRD) spectra were acquired directly using an INEL diffractometer equipped with a CPS 120 221 detector at 40 kV and 25 mA applied to a Co anticathode ($\lambda K\alpha = 0.179$ nm; $\lambda K\beta = 0.162$ nm) for crushed soil 222 samples and a Bruker D8 advance at 40 kV and 40 mA applied to a Cu anticathode ($\lambda K\alpha$ =0.154 nm; 223 λKβ=0.139 nm) for crushed rock samples. The crystallized compounds were identified through the 224 comparison with the COD 2013 database. Clay type was identified on oriented thin sections using the <2 225 µm fraction of decarbonated soil samples. Oriented thin sections were measured as natural sample, 226 heated to 550°C for two hours and treated with ethylene glycol. Diffraction spectra were then measured 227 using the Brucker D8 device with the setting described above.

228

229

230 Elemental contents in rock, soil and plant samples

For total elemental content analyses, 100 mg of each crushed soil or rock sample were digested in a CEM MARS 5 microwave oven using ultrapure acids (9 mL HNO₃ : 2mL HCl : 3mL HF). For each digestion run, an experimental blank and a standard (SRM 2709a or BCR-2) were included. Ground plant samples (200 mg) were digested in a CEM MARS 5 microwave oven using suprapure acids (10 mL HNO₃ : 3mL HCl : 0.2mL HF). For each digestion run, an experimental blank and a standard (SRM 1515) were included.

After acid digestion, element contents were measured with an Agilent® 7500ce ICP-MS at the GET laboratory in Toulouse and leaves with an iCAPQ Thermo Scientific at Hydrosciences in Montpellier using an In-Re internal standard to correct data for instrumental drift and plasma fluctuations. Ca, Mn and Fe contents in soil samples were measured using an ICP-OES Horiba ultima 2. Element concentrations were expressed in dry weight (mg kg⁻¹, DW). Quality and measurement traceability were ensured by measuring replicates of SLRS-5 river water standard. N content of leaves was measured during C-isotope analysis Isoprime 100 IRMS, as described above for C.

243

244 Sr isotope analysis

From the above described sample digests, aliquots were taken to contain 300 ng of Sr. They were dissolved again in 0.5 mL of 2M HNO₃ and added to a column containing 150 µl of Sr-Specific resin. The matrix was eluted using 0.4 mL of 2M HNO₃ followed by 1.5 mL of 7 M HNO₃ and another 0.3 mL of 2 M HNO₃. Strontium was then recovered in 1 mL of 0.05 M HNO₃. All acids used for purification were double sub-boiled prior dilution. Samples were evaporated to dryness and deposited on W-filaments. 250 Measurements of Sr isotope composition were carried out on a Thermo® Triton thermal ionization mass 251 spectrometer (TIMS). NBS 987 standards were run to verify the precision of the method. As differences 252 in Sr isotope ratios are mostly radiogenic (from ⁸⁷Rb decay), Sr-isotope ratios are expressed as ratios 253 between ⁸⁷Sr and ⁸⁶Sr.

254

255 Cu isotope analysis

256 For Cu isotope analyses, aliquots of digested samples containing 500 ng of Cu were purified using anionic 257 AG MP-1 resin. Separations were carried out at least twice per soil sample (elution with HCl and H_2O_2) 258 using AG MP-1 resin with Biorad® column, according to the protocol adapted from (Maréchal et al., 259 1999) and detailed in Blotevogel et al. (2018). Recovery was checked to be $100 \pm 5\%$ and BCR-2 standards were run to assure result quality. Our measured values (0.21 ± 0.03 ‰) are in line with formerly 260 261 published values of 0.20 ± 0.10 ‰ (Babcsányi et al., 2014) and 0.22 ± 0.05 ‰ (Bigalke et al., 2010a). 262 Purified solutions were spiked with IRMM Zn standard for measurements on MC-ICP-MS (Nu Plasma 263 500, Nu Instruments[®] at ENS in Lyon; Neptune plus, Thermo Finnigan[®] at GET in Toulouse). Exponential 264 laws were used to correct for mass bias using 66/64, 68/64 and 68/66 Zn isotope ratios (see Maréchal et 265 al. (1999) and Blotevogel et al. (2018) for more details). Cu isotope ratios are expressed in ‰ relative to 266 NIST 976 Cu standard, according to the Equation (2):

267 Equation (2)
$$\delta^{65}Cu = \left(\frac{\left({}^{65}Cu/{}^{63}Cu\right)_{sample}}{\left({}^{65}Cu/{}^{63}Cu\right)_{NIST976}} - 1\right) * 1000$$

268

269 Results

270 Field morpho-pedological description and soil identification

271 Studied soil profiles from the C catena feature granular to fine subangular blocky soil structure with 272 worm casts on top of the soil profile (i.e., A horizon, 0-10 cm). The structure evolves to blocky angular in 273 intermediate calcic horizons, whereas the deepest calcic soil horizons seems to display single grained 274 structure for C220 and powder-like structure in C155 (soil structure estimate using soil corer). Only C60 275 displays a more massive/prismatic structure in subsoil horizons. The texture is loamy clayey on topsoils 276 and gets coarser towards the bottom. Soil colors are mostly 10 YR 4/3 but the lowest horizon of C220 277 displays 10 YR 5/3 colors. All studied soil profiles from the C catena present an increase in carbonate 278 content with depth, as indicated by their reaction with 1 M HCl. The increase in carbonate phase content 279 with depth is due to two main processes: i) primary carbonate dissolution in surface horizons; and ii) 280 secondary carbonate formation in subsoil. The soils from the C catena were identified as Cambic Calcisols 281 (according to the World Reference Base for soil resources from FAO/ISRIC/ISSS, 2014), as described in 282 Figure 2.

283 Soil profiles from the B catena also show granular to subangular blocky structure within the first 10 cm. 284 The intermediate and deepest horizons investigated generally display angular blocky structures or quasi 285 massive soil structure. Indeed, cambic horizons located beneath 30 cm depth present massive structure 286 when humid and show wedge-shaped aggregates and shrink-swell cracks when dry suggesting vertic 287 properties. However, the deepest horizon of B60 seems to features a single grained structure and the 288 two deepest horizons of B150 have powder-like to more aggregated structure (supposed blocky angular). 289 Texture varies from loamy clayey in topsoil horizons to clayey in protovertic horizons. Again, the bottom 290 layer of B60 differs from the rest of the studied soil profiles being clayey-loamy-sand and thus much 291 coarser. Colors are darker than calcisols from the C catena (mainly 10 YR 3/2 and 10YR 2/1). Only B60 292 bottom layer is brighter with 10 YR 5/6. The occurrence of shrink swell cracks and wedge shaped 293 aggregates was established for most of the subsoil horizons after 30 to 40 cm depth. However, those 294 structures resulting from shrinking and swelling properties of clays are not considered as currently active

because of a quasi-constant humidity all year long (climate with no dry season and irrigation) so that vertic properties are poorly expressed at the soil profile scale. A secondary precipitation of carbonate phases was observed at various depth along the soil profile (see Figure 2) suggesting protocalcic properties within these soils. Colluvic material was only established for B150. B20 and B60 were identified as Vertic Cambisols protocalcic whereas B150 was identified as a Vertic Cambisol colluvic (WRB soil classification), as described in Figure 2.

301

302 Vineyard hydrological and geomorphological properties

The results of soil hydraulic conductivity are reported in Table 1. The soil permeability K is slightly lower in vertic soils (8×10^{-7} to 10^{-5} m s⁻¹) than in calcareous ones (1 to 7×10^{-5} m s⁻¹). The lowest hydraulic conductivity is measured where the infiltration test depth includes the vertic horizon. However, these values remain close, suggesting that the soil hydraulic properties are quite the same between the two vineyard plots.

Moreover, results of δ^{13} C measurements (Table 3) show no significant difference between the two plots (B or C catena). The measured δ^{13} C values ranging between -26.30 and -27.64‰ and -24.27 and -25.82 for respectively the leaves and the grapes from the 2 plots are in the upper range of the thresholds defined by van Leeuwen et al. (2009) for water deficit. This suggests that the plants did not suffer severe water stress and that there is no significant difference in water stress between the two plots (Table 3; non-parametric Mann-Whitney U-test: p-value=0.5 and 0.35 for leaves and grapes, respectively).

Concerning geomorphological properties, the vector map indicating slope direction is presented in Figure 315 3. Elevation of the vineyard plots is the highest in the north east and the lowest in the south west. There 316 is no strong variability of slope between the two vineyards, since the variation of elevation are mainly 317 related to localized, steep banks rather than through distributed and strongly variable sloping.

319 Mineralogical and physico-chemical properties of soils

320 The mineralogical and physico-chemical characteristics of the soil samples are summarized in Table 1.

321 The mineralogy of parental rocks and soils is reported in the XRD spectra of Figure 4. The XRD spectra of 322 limestones taken from the outcrops between the parcels exclusively show calcite peaks. The tufa 323 samples also contain the peaks typical of calcite, but also exhibit peaks of clay minerals identified as 324 smectites (Figure 4b). Only the basalt samples have more complex mineralogy (Figure 4a) containing 325 albite, titanite, chabazite and diopside. In most soils of the C catena calcite remains the main mineral. 326 Only in the topmost horizon of C60, quartz is the major phase (examples of diffraction spectra are shown 327 in Figure 4). In all soils, peaks belonging to smectite-type minerals are detected and clay mineralogy 328 appears to be the same (i.e., montmorillonite type smectite) in both catenas according to orientated thin 329 section analysis (data not shown). Furthermore, albite and titanite peaks are present in C60 and C155. 330 Diffraction patterns for soils of the B catena show the same minerals phases as basaltic rock, except 331 chabazite. Smectite is detected in all of those soils and some horizons show calcite. The presence of 332 calcite was expected as the vineyard was irrigated in case of need. In the lowest horizon of B60, calcite is 333 more abundant than the two other minerals. In B60 50-70 cm, quartz is found to be the most abundant 334 mineral.

All studied soil horizons have pH values higher than 7 (Table 1). The lowest pH values within a soil profile are systematically found in topsoils and ranges from 7.2 in C60 and C220 to 7.5 in B20. The pH values of subsoils vary between 7.6 in B150 (40-50 cm), C60 (30-50 cm) and C155 (10-50 cm). The highest pH value is 8.1 and was measured in the deepest horizon of C220. Cation exchange capacity is around 80 cmol kg⁻¹ in most horizons. However, the deepest horizons of B150 have slightly lower CEC with values of 70.0 and 72.2 cmol kg⁻¹, whereas the soil C220 has the lowest CEC with values between 66.0 cmol kg⁻¹ in the

topsoil and 55.8 cmol kg⁻¹ in the lowest soil horizon. The soil C220 is also the soil with the highest 341 342 inorganic carbon contents comprised between 3.7% in the topsoil and 5.6% in the lowest soil horizon. 343 Inorganic carbon contents of around 2% (wt) are measured in the deepest horizon of C60 and the two 344 deepest horizons of C155 alongside with the two topmost horizons of B20 (Table 1). Values between 1.4 345 and 1.6% are measured in the two upper horizons of C155 and the lowest horizon of B60. All other 346 horizons have inorganic carbon contents of less than 1%. Organic carbon contents decline in all soil 347 columns with depth. Organic carbon contents in topsoils ranges from 7.1% in B60 (0-10 cm) to 2.0% in 348 C155 and systematically declines to < 0.1% in the lowest soil layers. Only B20 (110-120 cm) and C220 (50-349 60 cm) contain measurable amounts of organic carbon (0.8 and 0.2% respectively).

350 Granulometric analyses after carbonate removal show that most of the soil horizons have similar 351 granulometric distribution, characteristic of silty clay to silt-clay-loam textures (Figure 5): between 30 352 and 40% of clay and very little sand (generally <10%). Higher clay contents are found in the soil B150 as 353 well as the topsoil of C60 that (about 50%), which can be classified as having a heavy clay texture. The 354 lowest clay contents are found in lowest horizons of B60 and C155 (15 and 10 %, respectively) and 355 horizons of C220 have slightly lower clay contents (24-28 %). The soils from the C catena have slightly 356 higher sand contents than the soils from the B catena. The highest sand contents of 11 to 19% are found 357 in B60 (70-90 cm), C155 (50-70 cm) and C220 (20-50 cm).

Surface A horizon from both catena soils display very low bulk density values from 0.7 to 1, in accordance with the intense biological activity (abundant worm casts) observed in the first ten centimeters of the soils. Bulk densities for the B_{ik} horizons of the B catena soil profiles are relatively low with values from 1 to 1.2 in accordance with the vertic properties of these soils, implying low bulk density during humid periods. Indeed, the volumetric water contents of the samples were estimated at approximately 40 to 50% during the sampling campaign. Subsurface horizons from the C catena show higher bulk densities values from 1.2 to 1.7.

366 Elemental contents in rocks and soils

367 The results of elemental contents in rocks and soils are reported in Table 2. The elemental concentrations of the major elements Mg, K, Ca and Mn in the basaltic rock (44.2, 9.9, 55.3 and 1.25 g kg⁻ 368 369 ¹ respectively) are within the range of values found in other studies in the area (De Vecchi et al., 1976). 370 Conversely, we find contents of Fe, Al and P that are 1.5 to 2 times higher than reported in the literature 371 (De Vecchi et al., 1976). Concentrations of all measured elements except Ca (394 g kg⁻¹) are relatively low in the limestone. Especially, the content of the transition metals Mn (0.67 g kg⁻¹), Fe (3.32 g kg⁻¹), Cu 372 373 (2.57 mg kg⁻¹) and Zn (3.81 mg kg⁻¹) is low compared to the basalt (1.25 and 81.9 g kg⁻¹ respectively for 374 Mn and Fe, and 49.1 and 120.4 mg kg⁻¹ for Cu and Zn). Molybdenum is the only element besides Ca that 375 is more concentrated in the limestone (0.25 mg kg⁻¹) than in the tufa (0.17 mg kg⁻¹), but still less 376 concentrated than in the basalt $(1.10 \text{ mg kg}^{-1})$.

377 The tufa rock concentrations of elements are intermediate between those of basalt and limestone for 378 most elements. Ba, Cu, P and Al have almost identical concentrations in the tufa as in the basalt, whereas 379 all other elements are more concentrated in the basalt. In basaltic rocks, the most abundant element 380 measured is Fe, in contrast to tufa and limestone where Ca is far more abundant. In soil profiles, Ca and 381 Fe are the most abundant elements in the order of hundreds of grams per kg of bulk soil. Some of the 382 elements show distinct variations with depth. This is the case for Cu and S which are much more 383 concentrated in the top of the soil columns. However, also K, Zn and, to some extent P, show increasing 384 concentrations in the top of the soil profiles. For Ca the trend is inversed, with higher concentrations in 385 the bottom horizons of the soils, especially on catena C. Other elements are more or less equally 386 distributed through the soil profile. Magnesium is depleted in the top soil horizons of B150 and C60. 387 There are few significant differences in elemental content between catena B and catena C. One sided

non-parametric Mann-Whitney U-test performed over all soil horizons of the respective plot, that Ca and
Ba contents are significantly higher in C catena soils. On the other hand significantly higher
concentrations in the B catena were detected for Mn, Fe, Sr and Mo.

391

392 Elemental contents in leaves and grapes

393 Elemental contents in leaves of both catenas are of the same order of magnitude (Table 3). However, 394 some significant differences identified using a Mann-Whitney U-test. K and Zn concentrations in leaves 395 of the C catena are higher than those in the B catena, whereas Sr concentrations are higher in leaves 396 from the B catena. No significant differences were detected for other elements contents in leaves. The 397 values measured are around average contents reported in the literature for Mg, P, Mn and Zn (Marschner and Marschner, 2012). Cu and Ca contents (75 mg kg⁻¹ and 32 g kg⁻¹, respectively) are higher 398 399 compared to the corresponding values of 6 mg kg⁻¹ and 5 g kg⁻¹ reported in literature. Nitrogen, K and Fe concentrations are low, with respectively, 14.3 g kg⁻¹, 7.2 g kg⁻¹ and 43.7 mg kg⁻¹, relative to literature 400 401 values of 23-43 g kg⁻¹, 10 g kg⁻¹ and 100 mg kg⁻¹ (Grechi et al., 2007; Marschner and Marschner, 2012).

For grapes, the elemental contents are consistent with the literature (Cugnetto et al., 2014; Scherz and Kirchhoff, 2006). However, comparison to most sources is difficult due to different sampling strategies (including pedicels, juice only etc.) and differences in notation (dry matter vs. fresh weight) (i.e. Almeida and Vasconcelos, 2003; Bertoldi et al., 2011). Significantly higher concentrations of K, Al and Mo are found in the C-catena whereas Ca and Sr concentrations are higher in grapes from the B-catena.

407

408 Sr and Cu isotopes in the soil-plant continuum

Isotope ratios for Sr and Cu are reported in Table 2 for soils and rocks, and in Table 3 for plant leaves. Cuisotope ratios could not be measured in grape samples because of the low concentrations of Cu in this
plant compartment and the interference of residual organic matter with the separation protocol.

Strontium isotope ratios are the highest in limestone with ⁸⁷Sr/⁸⁶Sr of 0.7070. Sr isotope ratios are equal
to 0.7032 in basaltic rock and 0.7067 in tufa rock, closer to limestone. Moreover, in soils as well in leaves,
Sr isotope ratios are always higher in samples from the C catena than in those of the B catena. This
difference is significant using a Mann-Whitney U-test.

416 Copper isotope ratios in soils were already published in Blotevogel et al. (2018), see there for more 417 details. Within the same soils, the values vary slightly but significantly between different soil types 418 though 2SD interval overlap. If we discard the B60 70-90 horizon for being obviously calcareous, the p-419 value (Wilcoxon Rank Sum test) is <0.002 comparing mean isotope ratios of 0.18‰ for calcaric cambisols 420 and 0.28‰ for vertic soils. The B60 surface horizon contains the heaviest measured Cu signature for soil 421 samples present vineyard (i.e., 0.37 ± 0.05‰). This is also the horizon containing the most Cu and 422 organic carbon, suggesting an organo-complexation of Cu with humic acid, as already suggested by 423 Bigalke et al. (2010b) and Blotevogel et al. (2018). The lowest isotope ratio is found in the deepest 424 horizon of B60 ($0.12 \pm 0.04\%$). That horizon contains a considerable amount of inorganic carbon (1.6%), 425 and calcite is the most abundant C-bearing phase. In plant leaves, Cu isotope ratio varies more than in 426 soil samples, varying from -0.86 to -1.83 for leaves of vine plants taken respectively in C155 and C60 427 soils.

428

429 Discussion

430 Influence of geology versus pedogenetic processes on soil properties in the Soave vineyard

In former studies of the Soave vineyard, only the presence of limestone and volcanic rocks was highlighted (Benciolini et al., 2006). However the presence of tufa rocks fits into a greater geological context with variations in land and ocean levels that left behind a variety of calcareous and siliclastic rocks in the Lessini mountains (Bassi and Nebelsick, 2010; De Vecchi et al., 1976).

As explained above, hydrological and geomorphological properties appear to be similar between the two vineyards plots, as suggested by the values of soil hydraulic conductivity (Table 1 and δ^{13} C measurements in leaves and grapes in Table 3). Even though there are slight differences in hydraulic conductivity, plants do not suffer severe water stress and especially there are no significant differences in δ^{13} C measurements between plants from the two plots. Thus hydraulic parameters were considered of no influence in our study, and their impact was not investigated in terms of soil-to-grape transfer of nutrients and pollutants.

442 According to our dataset, the soils of the two catenas are clearly dissimilar after field pedological 443 description in terms of morpho-pedological features. Color, structure and inorganic carbon content show 444 distinct features of the two soil types. There is a tendency for more massive or prismatic structures 445 associated with shrink swell-crack features in the B catena (depending on water content) and more 446 polyhedral to single grained structures in the C catena. Furthermore, soils of the C catena show 447 important calcic properties with calcic horizons lighter in color and in secondary carbonates contents, 448 and all cambic calcisol horizons show traces to large amounts of calcite in XRD spectra. Even if secondary 449 calcite phases are found in soils from the B catena, the studied Vertic Cambisols clearly have minerals 450 inherited from basalt, have greater amount of clay fraction and display vertic properties (shrink swell 451 cracking, wedge shapes aggregates, slickensides type features). Besides, soils from the C catena tend to 452 have higher sand contents even after elimination of carbonates. However, a more detailed look also 453 shows similarities. There are several calcic horizons in the upslope soils of the B catena and CEC values 454 are similar in all soils; only C220 shows lower CEC values than the other soils and pH values are >7 in all

455 horizons. This is attributable to a control of carbonate phases on soil pH in all soils. The chemical 456 composition of the soils is also surprisingly similar. The only difference is the higher Ca contents in 457 carbonated horizons whereas the contents in metals such as Al, Mn and Fe are relatively high in all soil 458 samples. The C220 soil contains more Ca and less transition metals than the other soil profiles, indicating 459 that this soil is less influenced by basaltic material than all other soils. All soils seem to have formed 460 under mixed influence of the three bedrocks present in the study area.

461 In the C catena a distinct increase in Ca content with depth as well as the increase in relative importance 462 of the calcite peaks in XRD spectra indicate that carbonate weathering is one of the soil forming 463 processes involved in this catena (Figure 4). However, the relatively high metal and smectite contents as 464 well as the increase in Al/Fe ratios in soils compared to limestone indicate that limestone weathering is not the only soil forming process. The influence of the tufa rock weathering would possibly explain these 465 observations and is supported by the ⁸⁷Sr/⁸⁶Sr values close to tufa and limestone rocks in the soils of the 466 467 C catena. Besides, soils of the C catena mainly in more upslope positions contain trace of feldspars and 468 titanite typically inherited from basaltic rock (Figure 4). The presence of minerals inherited from the 469 basalt indicates that there is at least some contribution of basalt to the soils from the C catena.

The soil forming processes along the B catena are also complex. The finer texture of soils downslope (B150) indicates that erosive transport (colluvic processes) occurs. The presence of calcic horizons together with non-calcic horizons in upslope soils (B20 and B60) indicates an influence of limestone through erosion processes (particulate transport) or secondary carbonate precipitation implying the circulation of water charged with Ca and dissolved carbonate species. The presence of quartz also confirms an erosive contribution from sedimentary rocks.

476 Copper and S contents decrease with depth within the soil profiles, which is likely due to fungicide477 treatment. XRD analyses on oriented thin sections indicate no difference between smectite types from

the B and the C catena (data not shown). This suggests that all soil smectites formed under the same conditions, possibly through basalt weathering. Some contribution of inherited smectites from tufa rock weathering during pedogenesis is expected in catena C. Therefore, such inherited smectites entrapped in the tufa rocks might have formed from past basalt weathering, which is still occurring in this area. Tufa rocks seem to have an important role in pedogenesis of the studied soils.

483 A mixing diagram (Figure 6) was drawn using the contents of Ca and Sr and the ⁸⁷Sr/⁸⁶Sr ratios of the 484 three bedrock types to illustrate their influence on the composition of the corresponding soil horizons 485 and grape leaves (see Langmuir et al., 1978, for details on mixing diagram interpretation). Almost all soil 486 horizons lie outside of the mixing triangle as a decrease in Ca/Sr ratios is expected during weathering 487 (e.g. Pett-Ridge et al., 2009). The C220 horizons lie on the mixing line between the limestone and the tufa with decreasing Ca/Sr ratios towards the surface. The ⁸⁷Sr/⁸⁶Sr ratios are unchanged compared to 488 489 the limestone bedrock, as already observed for various soils from typical bedrock endmembers (Braschi 490 et al., 2018). In our case, this observation confirms that weathering of limestone is a major soil forming process for the C220 soil. In addition, for the two topmost horizons of C60, the ⁸⁷Sr/⁸⁶Sr ratios are in the 491 492 range of values measured in tufa and limestone. Their shift towards lower Ca/Sr values indicates that 493 weathering of one or both of these rocks played a role in their formation. For the other horizons, the 494 ⁸⁷Sr/⁸⁶Sr values differ from all bedrocks indicating a mixed influence of bedrocks. The deeper horizons of C60 lie together with the C155 horizons and rather close to the ⁸⁷Sr/⁸⁶Sr values of tufa and limestone, 495 496 indicating a major influence in their formation by those rocks. If they were derived mainly from the tufa 497 bedrock, weathering played a minor role in their formation as Ca/Sr values are not far from those of tufa 498 rocks. If the main source material was limestone, weathering in those horizons was even stronger than in 499 C220 horizons. A basalt influence is also a possibility to explain the observed soil signatures. It would be 500 coherent with the presence of mineral phases from basalt such as titanite in the soils from the C catena.

501 Isotope ratio ⁸⁷Sr/⁸⁶Sr values of all B catena horizons align between the basalt pole and a pole with lower Ca/Sr ratio and higher ⁸⁷Sr signature than tufa and limestone (Figure 6). We suggest this pole 502 503 corresponds to secondary carbonate formation in basaltic soil. B150 soils are closest to the basalt pole. 504 Whereas B20 and B60 that were more strongly influenced by carbonate rocks and secondary carbonates have higher ⁸⁷Sr/⁸⁶Sr values. This illustrates the complex interplay between at least three parental rock 505 506 sources and pedogenetic processes (weathering, secondary phase precipitation, erosive transport) in the 507 area. Erosive transfer, pedogenesis and complex bedrock geometry make the vineyard zones less clearly 508 distinct in their biogeochemical properties than expected from macroscopic pedological observations.

509

510 Influence of soil characteristics on plant elemental content

511 The complex pedogenetic processes in the Soave soils have, to a large extent, homogenized elemental 512 contents of bulk soils even though structure and color are clearly different according to field observations. A dominant influence of bedrock chemistry on soil chemistry is visible only for few 513 514 elements. One remaining difference is the Ca content as marker of the degree of carbonate content. The 515 increase of Ca contents with depth in the C catena reveals a weathering gradient within the soil column, 516 underlining the fact that those soils have not been deeply ploughed recently. The presence of carbonates 517 is an important control on pH and thus on the mobility of elements (Kabata-Pendias, 2004). Cu and S 518 contents decrease with depth within the soil profiles. Such decrease is due to progressive migration of 519 those elements, which are frequently used in plant protection especially in organic viticulture (Blotevogel 520 et al., 2018). The gradients of nutrients K, Zn and P come more likely from the uptake at depth by the 521 plants and deposition through litter fall (Jobbagy and Jackson, 2001). The fact that Mn, Fe, Sr, Mo and Ba 522 are more concentrated in soils from the B catena is related to their higher abundance in the basaltic 523 bedrock.

On the mixing diagram (Figure 6), leaves from different catenas display different ⁸⁷Sr/⁸⁶Sr values. Leaves 524 525 from the B catena have values that are significantly lower than tufa and limestone rock indicating an 526 influence of basalts on their Sr-uptake and thus on Ca-nutrition as Sr is considered as an analog of Ca, 527 interchangeable in plant growth (Walsh, 1945). Plants from C catena all have Sr isotope ratios in the 528 range of tufa or limestone. Leaves from C155 and C220 also have higher Ca/Sr ratios than the leaves 529 from the B catena. This shows that, even in a context where plants are influenced to different degrees by 530 the same rock formation, the geological origin of their nutrition can be traced using alkaline earth 531 elements.

532 Among the vine leaf samples, Cu contents are high (from 34 to 125 mg kg⁻¹) in comparison with the 533 literature (Ko et al., 2007; Chopin et al., 2008) and large variations are observed. These variations cannot 534 be explained by variable soil Cu contents. The highest leaf Cu contents are measured in the upslope 535 sampling sites. Nevertheless, a clear link between Cu concentration and Cu isotope ratios is observed: 536 the higher the leaf content of Cu, the lighter the Cu isotopic composition (Figure 7). Isotopic ratios are 537 much lighter in leaves (-0.9 to -1.8‰) than in soils or pesticides (-0.49 to 0.89‰) reported in Blotevogel 538 et al. (2018). As a consequence, incomplete washing can be excluded as a source of the high Cu contents. 539 This points towards a control of Cu content and isotopic ratios by physiological processes. Weinstein et 540 al. (2011) suggested that the translocation of Cu to higher plant organs induce light isotope fractionation. In this study, an equation linking the leaf height (from the soil) to the isotope ratios was proposed. This 541 542 equation predicts surprisingly well (calculated δ^{65} Cu = -1.57‰) the isotope ratios measured in the field-543 grown grapevines (-0.9 to -1.8%). However, isotope fractionation within plants is still under discussion. 544 Jouvin et al., (2012) reported that light Cu was taken up from a nutrient solution and light Cu isotopes 545 become further enriched in aerial parts of the plants. On the contrary, another study reported uptake of 546 light Cu to roots and subsequent heavy Cu enrichment in aerial parts (Ryan et al., 2013). The last study 547 used extreme conditions with high doses of ionic Cu likely not occurring in nature. Light isotope

548 enrichment with height has also been reported for the chemically similar element Zn (see Caldelas and 549 Weiss, 2017 for a review and discussion of potential mechanisms). For Zn it was also reported that 550 transpiration flow controls root to shoot Zn fractionation on contaminated soils (Couder et al., 2015). 551 However similar water supply and exposition rule out such influence in our setting. The gradient of Cu 552 isotope ratios together with high Cu concentration points towards an immobilization mechanism of Cu in 553 leaves (Blotevogel et al., 2016). The immobilization of Cu as Cu(I)-S groups favors light isotope 554 enrichment in leaves and could thus explain our observations (Collin et al., 2014; Ehrlich et al., 2004; 555 Jouvin et al., 2012; Mathur et al., 2005; Zhu et al., 2002).

556 Finally, small but significant differences in plant nutrition have been detected for various elements 557 including some essential nutrients. It is difficult to directly link these differences to differences in soil 558 chemistry. Yet it has been established that differences in bioavailability in elements can induce different 559 elemental contents in plant tissues (Kuppusamy et al., 2018; Tyler and Olsson, 2001). Note that, these 560 differences do not necessarily include the element whose availability changed. That effect is on one hand 561 due to plant homeostasis mechanisms that can keep tissue concentration especially of essential 562 nutrients constant for a wide range of availability (Marschner and Marschner, 2012); on the other hand 563 regulation mechanisms of plants are not totally independent and exclusive for one element, so that 564 chemically similar elements can be taken up "passively along the way" (Ehlken and Kirchner, 2002; Tyler, 565 2004). However, as factors other than soil chemistry are comparable between the two plots, we 566 attribute detected differences to differences in soil chemistry.

In leaves, significant differences between the two plots were detected for the essential nutrients S, K and Zn. Note that higher concentrations were measured in plants grown on the C catena with more calcareous soils. No clear link can be drawn from leaf content to wine properties, still different tissue content of essential nutrients can have an impact on plant biosynthesis (Marschner and Marschner, 2012). In grapes, significant differences include concentrations and ratios of K and Ca, elements known

572 for their role in wine acidity and polyphenol synthesis (Brunetto et al., 2015; Daudt and Fogaça, 2008). 573 Potassium was more concentrated in grapes from the C catena whereas Ca was more concentrated in 574 the B catena, highlighting the difference in the ratio between both elements (Table 3), which is 575 particularly important for wine acidity (Brunetto et al., 2015; Daudt and Fogaça, 2008). It is interesting to 576 note that lower Ca concentrations were detected in grapes grown on soils with higher Ca content. This 577 observation highlights the regulation mechanisms carried out by plant during mineral nutrition, and 578 especially during element uptake from soil. But it also puts in evidence the distinct role of sink and 579 source organs (Joubert, 2013) during element translocation from the roots to the different shoots 580 compartments (leaves or grapes). Finally, via the uptake of elements and then their translocation 581 through the plant, this study highlights possible pathways of the influence of soil chemistry on wine 582 properties.

583

584 Conclusion

In the two examined Soave plots, pedologic differences are obvious from field observations. However, complex formation processes and interlayering of the different bedrocks resulted in rather homogenous chemical composition of soils. This shows that soils that appear to be morphologically different can have similar chemical functioning in terms of element mobility, retention and bioavailability.

The pedogenetic processes and the mechanisms involved in mineral nutrition in the Soave vineyard have then been investigated through the elemental contents in the soil-to-grape continuum and the isotope ratios for Cu and Sr. In the Soave vineyard, similar soil chemistry is reflected by similar elemental concentrations in grapevine leaves, although there appears to be a trend for higher elemental concentration in plants grown on more calcareous soil. Moreover, elements are finally not identically distributed in the plants form the two catena and along the different plant tissues (especially leaves or

grapes) implying different regulation mechanisms during the soil-to-grape continuum, through uptake
and translocation processes. Most importantly K and Ca contents and their ratio, known for regulating
wine acidity, vary significantly between the two plots.

Isotope ratios measurements show that radiogenic Sr isotopes compositions can be used as a tracer of the geological origin of grapevine nutrition, whereas Cu isotope ratios rather depend on plant regulation mechanisms.

Finally, as methods presented in this paper give first indications on the role of soil chemistry in the potential "terroir effect", it would be possible to go further by improving the sampling strategy (more contrasted soils with a higher number of sampling points, sampling tissues at different vine growth stages, and over several years). However, attention must be given to the other "terroir" components as meteorological parameters, agricultural practices management in the field, winemaking process, etc. in order to more isolate the soil influence.

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Figures



Figure 1 – The experimental site in the Soave vineyard including the two catenas.
 C stands for calcareous (point C60 to C220) and B for basaltic (point B20 to B150). Rock outcrops (basalt, limestone and tufa rocks) are represented by colored areas.
 The aerial image is taken from Google Earth[®].



B catena soil profiles

Figure 2 – Schematic illustration of the field observations for the soils of the two catenas investigated in the present study (B & C). Roots are particularly thick in the first 10-20 cm for all the sampling points. Soil profile description was stopped according to the soil depth, using an Edelman soil corer down to the depth of 120 cm maximum. Soil structures above 50 cm are indicative structures as structure observation is biased during soil corer sampling.



Figure 3 – Elevation and slope direction map of the studied vineyard plots. The aerial image is taken from Google Earth[®].



Figure 4 – XRD spectra of rocks and soil horizons (a- Basalt rock and B catena; b-Calcareous rock and C catena). For clarity, only the main peaks of the identified phases are reported for silicates (a).



Figure 5- Textural triangle of the studied soils after acid treatment to eliminate carbonates.



Figure 6- Mixing diagram of different source rocks, soils and vine leaves. The arrow on the left represents a hypothetic secondary soil carbonate pool.



Figure 7 – Cu isotope ratios in vine leaves as a function of their Cu-content.

Soil ID and depths (cm)	Horizon (WRB	Mineralogical phases	рН (Н₂0)	CEC	SIC	SOC	Clay	Silt	Sand	
	classification)		/							К
				cmol/kg	wt %	wt %	wt %	wt %	wt %	m.s ⁻¹
B20										1*10 ⁻⁵
0-10	Mollic, calcic	calc., felds., smec.	7.51	72.12	1.8	4.3	30	67	3	
10-30	Calcic	calc., felds., smec.	7.67	66.13	2.2	1.3	39	59	2	
50-70	Vertic	felds., smec.	7.80	71.91	0.5	0.7	25	70	5	
70-120	Vertic	felds., smec.	7.74	75.23	<0.1	0.8	34	64	2	
B60										2*10 ⁻⁶
0-10	Mollic	felds., smec.	7.17	82.94	0.5	7.1	38	61	1	
10-30	Cambic	felds., smec.	7.90	NA	0.4	1.3	42	57	1	
30-60	Vertic	felds., smec.	7.83	77.72	0.3	0.9	33	66	1	
60-70	Vertic	qtz. , felds., smec.	7.82	78.31	0.3	0.6	36	61	3	
70-90	Calcic	calc., felds., smec.	7.87	82.75	1.6	<0.1	15	65	20	
B150										8*10 ⁻⁷
0-10	Mollic	felds., smec.	7.29	80.16	0.2	3.9	55	45	0	
10-40	Cambic	felds., smec.	NA	NA	0.3	NA	50	50	0	
40-50	Vertic	felds., smec.	7.60	78.87	0.1	0.9	52	48	0	
50-70	Vertic	felds., smec.	NA	NA	0.1	NA	55	45	0	
70-90	Vertic	felds., smec.	7.82	69.97	0.2	1.3	54	46	0	
90-100	Vertic	felds., smec.	7.76	72.25	0.6	0.0	55	45	0	
C60										1*10 ⁻⁵
0-10	Mollic, calcic	qtz., calc., felds.	7.24	80.36	0.5	4.9	63	37	0	
10-30	Calcic	qtz., calc., felds.	7.67	77.87	0.6	1.3	32	64	4	
30-50	Cambic	qtz., calc., felds.	7.62	84.55	0.8	0.9	31	69	0	
50-70	Cambic	calc., qtz., felds.	7.79	79.61	1.9	<0.1	40	60	0	
C155										3*10 ⁻⁵
0-10	Mollic,	calc., felds., smec.	7.32	83.01	1.4	2.0	32	63	5	
10-50	Cambic	calc., felds., smec.	7.59	79.49	1.5	0.9	31	61	8	
50-70	Calcic	calc., felds., smec.	7.84	80.99	2.1	0.2	33	56	11	
70-100	calcic	calc., felds., smec.	7.91	82.63	1.9	<0.1	10	82	8	
C220										7 *10 ⁻⁵
0-10	Mollic, calcic	calc., felds smec.	7.24	65.97	3.7	4.9	24	65	11	
20-50	Calcic	calc., felds., smec.	7.90	55.12	4.9	0.5	28	52	20	
50-60	Calcic	calc., felds., smec.	8.14	50.80	5.6	0.2	NA	NA	NA	

Table 1- Mineralogical and physico-chemical properties of Soave soils

Table 2 – Elemental contents (expressed in g kg⁻¹ or mg kg⁻¹) and Sr isotope ratios in soil and rock samples. P-values of one-sided Mann Whitney U-test (MWU-test) between all horizons of the two catenas were performed to evidence significant differences. Differences are considered significant if $p \le 0.05$ and are annotated with a*.

	Туре	Depth	Mg	Al	Р	К	Са	Mn	Fe	S	Cu	Zn	Sr	Мо	Ва	⁸⁷ Sr/ ⁸⁶ Sr	2SD
			g kg ⁻¹	g kg ⁻¹	g kg⁻¹	g kg⁻¹	g kg ⁻¹	g kg⁻¹	g kg ⁻¹	mg kg⁻¹	mg kg⁻¹	mg kg⁻¹	mg kg ⁻	mg kg ⁻	mg kg⁻¹		
B20	Bulksoil	0-10	19.36	56.28	2.85	7.10	86.35	1.00	62.08	679	356.6	162	257	1.02	282	0.7056	0.00011
B20	Bulksoil	10-50	20.37	56.12	2.24	5.90	107.34	1.02	66.83	317	209.6	138	295	1.03	299	-	-
B20	Bulksoil	50-70	12.52	36.96	1.75	4.05	40.82	1.08	75.40	-	91.0	125	256	0.94	257	0.7047	0.0001
B20	Bulksoil	110- 120	10.90	57.53	1.03	4.23	23.34	1.35	84.36	76	63.6	118	179	0.57	258	0.7051	0.00008
B60	Bulksoil	0-10	15.83	49.21	3.36	7.17	42.37	1.18	66.74	965	563.7	196	221	1.36	306	0.7054	0.00011
B60	Bulksoil	10-30	17.71	61.77	1.78	4.12	39.62	1.16	72.27	-	135.9	115	193	1.06	265	-	-
B60	Bulksoil	30-60	11.70	54.29	1.57	4.45	32.68	1.33	77.28	177	79.9	112	188	0.90	301	0.7057	0.00013
B60	Bulksoil	60-70	15.55	51.80	1.79	6.25	31.78	1.15	82.37	-	56.7	112	176	2.69	342	-	-
B60	Bulksoil	70-90	33.04	50.89	4.17	2.39	95.64	0.97	77.08	-	40.3	101	406	1.17	373	0.7053	0.00013
B150	Bulksoil	0-10	7.93	36.57	3.34	8.76	31.16	1.08	71.77	547	277.0	163	250	1.40	343	0.7043	0.00011
B150	Bulksoil	40-50	8.62	46.07	2.42	6.09	27.82	1.27	83.35	155	72.8	140	217	1.25	296	0.7043	0.00011
B150	Bulksoil	70-90	18.22	61.35	2.88	7.48	45.06	1.20	93.37	-	61.0	139	420	1.34	394	-	-
B150	Bulksoil	90- 100	17.95	70.39	3.05	7.92	52.64	1.32	98.88	-	59.1	133	441	1.46	424	0.7037	0.00009

C60	Bulksoil	0-10	9.94	47.98	3.29	9.02	35.16	1.13	72.18	707	351.9	177	137	0.91	394	0.7067	0.00009
C60	Bulksoil	10-30	5.39	39.17	2.64	6.82	32.46	0.95	60.33	255	186.5	131	128	0.70	344	0.7066	0.00011
C60	Bulksoil	30-50	14.54	56.04	2.44	6.11	44.11	1.08	71.93	-	160.5	134	152	0.82	413	-	-
C60	Bulksoil	50-70	23.01	55.10	3.25	4.65	92.3	1.07	74.01	-	83.9	119	178	0.60	386	0.7064	0.00008
C155	Bulksoil	0-10	23.08	57.24	3.55	6.92	71.22	0.96	70.68	400	164.7	132	152	0.68	497	0.7061	0.00011
C155	Bulksoil	10-50	22.7	53.08	2.92	4.61	73.03	0.92	68.88	254	124.9	119	153	0.69	475	0.7061	0.00009
C155	Bulksoil	50-70	23.25	52.57	3.03	4.05	89.64	0.99	73.49	-	111.2	123	171	0.70	504	-	-
C155	Bulksoil	70- 100	27.50	60.79	3.57	2.77	92.84	0.92	73.07	-	37.3	113	147	0.48	563	-	-
C220	Bulksoil	0-10	12.87	33.72	3.17	5.21	135.1	0.70	42.68	784	261.9	157	187	0.90	286	0.7069	0.0001
C220	Bulksoil	20-50	14.23	40.23	2.24	2.80	194.42	0.66	46.29	240	69.8	83	217	0.98	300	0.7068	0.0001
C220	Bulksoil	50-60	11.31	36.49	1.98	1.37	232.26	0.42	35.76	168	31.4	66	196	0.69	137	0.7071	0.0001
MWU- test	p-value		0.32	0.14	0.08	0.17	0.02*	<0.01*	0.01*	0.40	0.41	0.24	<0.01*	<0.01*	0.02*	<0.01*	
Basalt	Rock		44.22	55.91	2.96	9.87	55.34	1.25	81.86	-	49.1	120	1516	1.10	930	0.7032	0.00001
Tufa	Rock		17.28	46.15	2.72	2.16	105.42	0.83	52.95	-	44.0	68	173	0.17	903	0.7067	0.00009
Limestone	Rock		3.07	0.86	0.33	0.10	394.17	0.67	3.32	-	2.6	4	169	0.25	10	0.707	0.00002

NB: Cu isotope ratios are presented in Blotevogel et al. (2018)

Table 3 – Elemental contents (expressed in g kg⁻¹ or mg kg⁻¹) in leaves and grapes, and C, Sr and Cu isotope ratios in leaves. P-values of onesided Mann Whitney U-test (MWU-test) between the two catenas were performed to evidence significant differences between the two plots. Differences are considered significant if $p \le 0.05$ and are annotated with a*.

Sampl e	Туре	N	Mg	Ρ	S	к	Са	AI	Mn	Fe	Cu	Zn	Sr	Мо	Ва	K/C a	δ ¹³ C	2S D	87Sr/8 6Sr	2SD	δ ⁶⁵ C u	2S D
		g kg ⁻¹	g kg⁻¹	g kg⁻¹	g kg⁻¹	g kg⁻¹	g kg⁻¹	mg kg⁻¹	mol mol -1	‰	‰			‰	‰							
B20	Leaf	13. 04	2.8 1	2.2 4	1.6 3	6.8 5	29. 7	24.4	35.7	33	99.3	18.4	49.9	0.11 8	8.79	0.2 4	- 26.5 6	0. 22	0.7065	0.00 008	- 1.73 5	0.0 98
B60	Leaf	12. 77	2.9 5	2.5 3	1.4 1	7.0 5	31. 7	25.3	23	34.4	70.5	24.1	66.3	0.03 6	8.34	0.2 3	- 27.6 4	0. 12	0.7061	0.00 012	- 1.28	0.1
B150	Leaf	15. 14	4.2 4	2.3 9	1.4 4	4.6 7	32. 1	27.1	51.9	44.3	43.6	23.2	54.2	0.04 2	18.4 1	0.1 5	- 26.3 0	0. 24	0.7062	0.00 01	- 1.06 7	0.1
C60	Leaf	14. 03	3.1 5	2.1 4	1.7 8	8.4	26. 9	29.9	49.6	64.2	125. 4	25.6	43.1	0.27 1	36.1 4	0.3 2	- 26.5 4	0. 33	0.7072	0.00 009	- 1.83 3	0.0 55
C155	Leaf	15. 00	5.4 9	4.0 8	1.6 7	7.3 2	37. 7	25.3	58.8	40.6	33.9	35.2	48.7	0.16 6	22.6 5	0.2 0	- 26.9 6	0. 05	0.7068	0.00 012	- 0.86 3	0.1 02
C220	Leaf	15. 80	2.7 4	2.7 7	1.6	9.0 4	32. 6	40.7	47.9	45.7	74.3	28.8	38.6	0.11 4	20.4 2	0.2 9	- 27.0 2	0. 05	0.707	0.00 013	- 1.13 4	0.0 09
MWU- test	p- valu e	0.2 0	0.5 0	0.3 5	0.1 0	0.0 5*	0.3 5	0.13	0.20	0.10	0.50	0.05 *	0.05 *	0.10	0.05 *	0.2 0	0.50		0.05*		0.50	
B20	Grap e	3.3 3	0.7 2	1.3 7		9.2 1	2.3 7	4.80	2.90	8.11	5.83	5.05	5.38	0.02 8	1.15	3.9 9	- 24.5	0. 83				

B60	Grap e	1.0 0	0.5 2	1.2 5	10 40	. 1.6 9	4.55	1.87	8.95	5.73	3.15	3.18	0.02 7	0.50	6.3 3	- 24.6 5	0. 19
B150	Grap e	3.3 8	0.7 0	1.2 5	8.4 1	2.0 8	5.03	6.04	10.8 7	4.92	4.11	3.47	0.03 0	1.28	4.1 6	- 24.2 7	0. 52
C60	Grap e	2.2 8	0.5 4	1.3 1	10 53	. 1.4 6	5.05	4.40	9.64	6.88	3.97	2.43	0.04 6	2.17	7.4 1	- 24.2 9	0. 58
C155	Grap e	2.5 1	0.5 6	1.4 4	10 83	. 1.4 5	5.10	2.62	9.40	4.82	4.12	1.57	0.04 9	0.90	7.6 8	- 25.8 2	0. 48
C220	Grap e	2.8 1	0.5 1	1.3 3	11 16	. 1.4 5	7.08	2.94	10.0 7	5.79	3.19	1.93	0.05 3	1.01	7.9 1	- 24.5 6	0. 01
MWU- test		0.3 5	0.2 0	0.1 9	0.(5	0 0.0 4*	0.05 *	0.50	0.35	0.50	0.50	0.05 *	0.05 *	0.50	0.0 5*	0.35	