

Understanding and managing nitrogen nutrition in grapevine: a review

Thibaut Verdenal, Ágnes Dienes-Nagy, Jorge E Spangenberg, Vivian Zufferey, Jean-Laurent Spring, Olivier Viret, Johanna Marin-Carbonne, Cornelis van

Leeuwen

► To cite this version:

Thibaut Verdenal, Ágnes Dienes-Nagy, Jorge E Spangenberg, Vivian Zufferey, Jean-Laurent Spring, et al.. Understanding and managing nitrogen nutrition in grapevine: a review. OENO One, 2021, 55 (1), pp.1-43. 10.20870/oeno-one.2021.55.1.3866 . hal-04123332

HAL Id: hal-04123332 https://hal.inrae.fr/hal-04123332

Submitted on 9 Jun 2023

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution | 4.0 International License



Understanding and managing nitrogen nutrition in grapevine: a review

Thibaut Verdenal¹, Ágnès Dienes-Nagy¹, Jorge E. Spangenberg², Vivian Zufferey¹, Jean-Laurent Spring¹, Olivier Viret³, Johanna Marin-Carbonne⁴ and Cornelis van Leeuwen⁵

¹Agroscope Institute, Avenue Rochettaz 21, 1009 Pully, Switzerland

²Institute of Earth Surface Dynamics, Faculty of Geosciences and Environment, University of Lausanne, Switzerland ³Direction générale de l'agriculture, de la viticulture et des affaires vétérinaires (DGAV), 1110 Morges, Switzerland ⁴Earth Science Institute, Faculty of Geosciences and Environment, University of Lausanne, Switzerland ⁵EGFV, Univ. Bordeaux, Bordeaux Sciences Agro, Inrae, ISVV, F-33882 Villenave d'Ornon, France

*corresponding author: thibaut.verdenal@agroscope.admin.ch

ABSTRACT

This review addresses the role of nitrogen (N) in vine balance and grape composition. It offers an integrative approach to managing grapevine N nutrition. Keeping in mind that N excess is just as detrimental to wine quality as N depletion, the control of grapevine N status, and ultimately must N composition, is critical for high-quality grape production. N fertilisation has been intensively used in the past century, despite plants absorbing only 30 to 40 % of applied N. By adapting plant material, soil management and vine balance to environmental conditions, it would be possible for grape growers to improve plant N use efficiency and minimise N input in the vineyard. Vineyard N management is a complex exercise involving a search for a balance between controlling vigour, optimising grape composition, regulating production costs and limiting pollution. The first part of this review describes grapevine N metabolism from root N uptake to vine development and grape ripening, including the formation of grape aroma compounds. The advantages and limits of methods available for measuring plant N status are addressed. The second part focuses on the parameters that influence grapevine N metabolism, distinguishing the impacts of environmental factors from those of vineyard management practices. Areas for further research are also identified.

KEYWORDS

nitrogen use efficiency, agronomic practices, physiology, partitioning, balance, leaf-to-fruit ratio, amino N, yeast assimilable nitrogen, vine, wine

ABBREVIATIONS

2-AAP: 2-aminoacetophenone AA(s): amino acid(s) AF: alcoholic fermentation Atom % : atomic percentage B: boron C: carbon CO₂: carbon dioxide Cu: copper DAP: diammonium phosphate DMS: dimethyl sulphide DW: dry weight H₂S: hydrogen sulfide MLF: malolactic fermentation N: nitrogen N₂: dinitrogen NH₃: ammonia (gas) NH_4^+ : ammonium NO₂⁻: nitrite NO_3 : nitrate NUE: nitrogen use efficiency S: sulphur YAN: yeast assimilable nitrogen

INTRODUCTION

Nitrogen (N) is a major nutrient for plants involved in many vital physiological processes. It is required in larger amounts than the other mineral nutrients and regulates plant vigour and development in the absence of water restriction. N was intensively applied to crops, mainly in the form of nitrate, during the twentieth century to increase production, regardless of the pollution resulting from crops using only 30-40 % of the fertiliser. In viticulture, optimum yield for high-quality grape is not the maximum allowed by the conditions of the vineyard. N fertilisation has consequently become a complex exercise in the search for a balance between optimising vigour and grape composition, controlling production costs and limiting pollution. Over the past decades, the application of N in vineyards has been reduced with the aim of adjusting vigour and yield. Moreover, the development of cover cropping has led to vines competing for N resources, which can be detrimental to the crop in some cases. This evolution of management practices has created situations with high grape N deficiencies, which can affect fermentation kinetics and wine flavours. White wines are particularly sensitive to grape N deficiency, as they can express a typical 'stress taste' often associated with strong bitterness, despite corrective winemaking techniques. Although several reviews about grapevine N metabolism have been published (Haynes, 1986; Wermelinger, 1991; Mengel and Pilbeam, 1992; Roubelakis-Angelakis and Kliewer, 1992; Loulakakis et al., 2009; Masclaux-Daubresse et al., 2010), the relationship between plant N status and grape composition is still not fully understood. The management of grapevine N status and, ultimately, grape N composition at harvest should be a prerequisite for grape production with a high-quality potential. The scope of this review is to compile state-of-the-art knowledge about grapevine N nutrition, ranging from plant biology to factors linked to N regulation. It will contribute to the implementation of sustainable practices in the vineyard. The first section focusses on N metabolism, with an emphasis on grapevine N requirement and monitoring. The mechanisms of N uptake, assimilation and efflux are addressed. The role of grape N in the formation of wine aroma is described. The second section gives a comprehensive description of the factors influencing grapevine N status. The agronomic parameters useful for growers to enhance N use efficiency and optimise grape composition, while minimising the use of fertilisers, are discussed. Perspectives for further research are also considered.

NITROGEN REQUIREMENTS AND MONITORING

1. Grape growing

N plays a key role in plant metabolism. As a macronutrient, it represents approximately 1.5 % of dry weight (% DW) of grapevine and enters the composition of key metabolites, such as proteins, amino acids (AAs), enzymes, DNA, RNA and chlorophyll.

1.1. Grapevine N requirements

nutrition The positive impact of N on biomass development well known is (Holzapfel and Treeby, 2007; Gatti et al., 2018). The production of 1 kg of biomass requires from 20 to 50 g of N (Xu et al., 2012). Grapevine N requirements are rather modest in comparison to nonperennial crops, even with high production objectives (Metay et al., 2014), and have already been studied under different environmental conditions (Löhnertz, 1988; Porro et al., 2007; Schreiner et al., 2018). In the context of the sustainable production of 12 tons/ha of grape in cool climate, Löhnertz (1988) estimated the average grapevine N requirement to be 50 kg/ha per year (Table 1). This estimation ensures optimal vegetative growth, taking into account that only the grapes are exported from the vineyard; leaves are restored to the soil, as is the pruned wood in most vineyards.

TABLE 1. N allocation for Riesling at harvest (Löhnertz, 1988). Estimations for a yield of 12 tons/ha of grapes.

N allocation at harvest	Nitrogen kg/ha per year
Wood and roots	27
Grapes	23
Total exported and immobilised	50
Shoots	5
Leaves	37
Total	92

1.2. Symptoms of N deficiency and excess

N metabolism largely controls plant vigour and vegetative development (Metay *et al.*, 2014), and it also influences plant productivity and fruit composition. Both N deficiency and N excess have negative impacts on grapevine development and grape composition.

N deficiency results in weak vine growth, short inter-nodes, small and light-green to yellow leaves, low berry set, reduced long-term bud fruitfulness and yield (Guilpart *et al.*, 2014), reduced grape N content and possible delayed maturation (Schreiner *et al.*, 2018).

N excess leads to high vigour, dense canopy, large dark-green leaves, extended vegetative growth period (competing with and delaying grape ripening) and increased grape sensitivity to fungal diseases (Thomidis *et al.*, 2016).

N status alters both vine production variables and grape composition to different degrees (Schreiner *et al.*, 2018). Vegetative growth is more constrained than reproductive growth as N status decreases, as illustrated in Figure 1.

1.3. Nitrogen seasonal cycle

Forecasting plant N status in perennial fruit crops requires an understanding of the seasonal plant N cycle. The N assimilation rate fluctuates depending on both the physiological stage (biotic parameters) and environmental conditions (abiotic parameters). Several reports have described grapevine seasonal N uptake and detailed N partitioning within the vine (Conradie, 1980; Conradie, 1991; Löhnertz, 1988; Wermelinger, 1991; Bates et al., 2002; Zapata et al., 2004a; Zapata et al., 2004b; Treeby and Wheatley, 2006; Weyand and Schultz, 2006; Williams. 2015: Zufferey al., 2015; Schreiner, 2016: et Holzapfel et al., 2019). A model of seasonal changes in N content of grapevine tissues is shown in Figure 2.

Except in vineyards close to the equator where vines grow continuously, annual grapevine N requirement is usually concentrated in the vegetative period. Before the onset of winter - under the influence of seasonal changes in light and temperature - grapevines enter a phase in which metabolic activity is minimal and growth stops (Cookson *et al.*, 2013). Growth resumes at bud break, which is induced by increasing temperatures. Growth after bud break mainly depends on the vine's reserves in its storage organs (roots and wood), which have accumulated during the previous summer and autumn. During winter, the grapevine N reserves are mainly stored in the roots (about 75 % in dormant vines), in the form of AAs and proteins (Zapata *et al.*, 2004a; Zapata *et al.*, 2004b).



FIGURE 1. Hypothetical model of vegetative versus reproductive development rates as a function of grapevine N status.



FIGURE 2. Changes in N content of plant parts in grapevines over two growing seasons. Four-year-old potted Chasselas cv. (Verdenal *et al.*, unpublished data, 2017-2018). Letters designate major phenological stages: BB, budbreak; FL, flowering; VR, veraison; HA, fruit harvest; PR, pruning (*hypothetic values).



FIGURE 3. Annual evolution of the N uptake rate of grapevine (adapted from Löhnertz, 1988).

From bud break (phenological stage 07 on the BBCH scale) to the stage of 5-6 leaves (BBCH 53), N uptake remains low. N reserves from the roots and, to a lesser extent, from the wood are mobilised to support initial growth until root N uptake becomes sufficient around flowering (BBCH 65) (Zapata *et al.*, 2004a; Zufferey *et al.*, 2015). Soluble N in the storage organs reaches a maximum just before budbreak, and it decreases thereafter until the beginning of fruit growth (Wermelinger, 1991; Williams, 2015). After harvest, approximately 85 % of the increase in root and wood N reserves is due to N translocation from the leaves before leaf fall (Williams, 2015).

N uptake and AAs synthesis are necessary for the synthesis of proteins and enzymes, which are in turn required for the photosynthetic activity and other biochemical pathways related to plant development. Young leaves first behave as a sink for N compounds to ensure their own development; during the reproductive stage, leaves behave as a source of AAs for grape development and the refilling of reserves (Kant et al., 2011). Substantial refilling of reserves can occur after harvest due to N relocation from the leaves prior to leaf fall. In warmer countries, the post-harvest period (from harvest to complete leaf fall) may last for up to four months, and N uptake during that period may contribute up to 30 % of the annual refilling of the N reserve (Conradie, 1992; Conradie, 2005). An increased supply of nitrogenous compounds is necessary for optimum flowering and berry development; grapes start accumulating N during the first growth stage, with major N uptake occurring from two weeks before flowering until four weeks after flowering (BBCH 65) (Figure 3) (Linsenmeier et al., 2008; Holzapfel et al., 2019). A lag phase is observed at the onset of grape ripening (veraison, BBCH 85), and then a second uptake peak occurs at the beginning of grape ripening (Löhnertz, 1988; Ribéreau-Gayon *et al.*, 2017). During ripening, NH_4^+ content decreases and organic N content increases in grape berries. Most of the berry N is imported in the form of glutamine (Keller, 2015), which is then converted in the berry into other AAs via transamination.

At the end of the vegetative period, some of the N migrates from the leaves to the roots. The refilling of root N reserves usually starts before grape maturity and continues until leaf fall (Holzapfel and Treeby, 2007; Rossouw *et al.*, 2017). The root N pool at the beginning of the vegetative season is related to the yield of the previous year and to vine age (Löhnertz, 1988).

2. Nitrogen monitoring

Grapevine N status not only influences plant vigour and yield, but also grape composition and subsequent wine quality. By monitoring plant N status, agronomic practices and fertilisation can be adjusted to meet production objectives. This section reviews the indicators of plant N status and highlights their advantages and drawbacks.

2.1. Soil analysis

N fertiliser recommendations are usually based on the soil measurement of mineral N; *i.e.*, the form in which N is directly available to plants. Mineral N is mostly present in soils as nitrate (NO₃⁻), because NH₄⁺ is quickly nitrified, except when soil pH is very low. Mineral N, however, represents only a small fraction of total soil N, and its amount varies significantly depending on the rates of N mineralisation, plant N uptake and soil N losses (*i.e.*, leaching, denitrification, erosion and gaseous emission). The size of the mineral N pool can vary from a few tenths of kilograms to a few hundreds of kg/ha.



FIGURE 4. Factors influencing the mineralisation of soil organic matter.

Total N (mineral + organic) in the soil is not a good indicator of plant N status, because organic material needs to be broken down by soil microflora before being accessible to plants. Hence, the factors involved in the mineralisation of soil organic matter greatly influence the size of the mineral N pool available to the grapevine over time (Figure 4). Moreover, the sampling method used - particularly in terms of location and depth - can greatly affect results and interpretations. Consequently, a soil analysis can provide a baseline for N fertiliser management, but it is not sufficient on its own, as it does not reflect the dynamics of available soil N over the season. Moreover, it does not take into account grapevine N requirements, which also depend on yield and quality targets. Recommendations regarding N fertiliser supply can change on a yearly basis, especially with varying weather conditions (Van Cleemput et al., 2008).

2.2. Leaf and petiole analysis

Leaf petiole and leaf blade analyses can be used to monitor plant nutrition status during the season mainly for macro elements (N, phosphor, potassium, calcium, magnesium) (Gaudillère et al., 2003), for which results are expressed in percent of dry weight (% DW). Leaf N concentration is well-correlated with the chlorophyll index (Spring, 1999). N content in the leaf blade is very different to that in the petiole: petiole N content is more sensitive to variations in N nutrition than leaf blade N content, which is more constant (Delas, 2010). Consequently, the chosen analysis (*i.e.*, on either the leaf blade or petiole, or both together) will greatly affect the results and require adapted interpretation thresholds (Table 2). The interpretation may be refined with the ratios of N/P and N/K (Crespy, 2007) (Table 3).

TABLE 2. Threshold values for the interpretation of grapevine leaf and petiole N content with regard to vine N statu
--

N concentration (% DW)	Very low	Adequate	Very high	Reference
Leaf blade + petiole	< 1.8	2.0 - 2.3	> 2.5	Spring and Verdenal (2017)
Petiole	< 0.4	0.4 - 0.6	> 0.6	Champagnol (1984)

Measurements are implemented at veraison on leaves (either leaf blade + petiole or petiole only) from the main shoots of the bunch area. Results are expressed as % DW.

TABLE 3. Thresholds for the ratios N/P and N/K for the interpretation of grapevine leaf and petiole analysis with regard to vine N status.

		Very low	Adequate	Very high
Lasfhlada natiala	N / P	< 9.7	10.7 - 12.8	> 13.9
Leaf blade + petiole	N / K	< 1.0	1.1 - 1.3	> 1.4
D-4-1-	N / P	< 2.5	2.5 - 3.5	> 3.5
reuole	N / K	< 0.2	0.2 - 0.4	> 0.4

Measurements are implemented at veraison on adult leaves (either leaf blade + petiole or petiole only) from the bunch area. Results are expressed in % DW.

The limitations of plant N assessment through tissue analysis for fertilisation purposes have long been acknowledged, and the interpretation of results should be carried out with care (Perez and Kliewer, 1982; Delas, 2010). The results are mainly used as a complement to other observations. Before making any decision on fertiliser application, it is recommended to complete the diagnosis with visual observations of plant morphology. High vigour, dense canopies and high yields are generally indicators of high vine N status. Leaf and petiole analyses are essentially used for research purposes to observe the impact of a particular practice on plant composition or to confirm a nutrition problem in the plant.

2.3. Chlorophyll index

Various tools have been developed for plant-based N status assessment. These are usually based on indirect and non-destructive measurements, such as chlorophyll concentration. Examples of hand-held chlorophyll meters used for diagnosis purposes are the N-Tester (Yara, Oslo, Norway), SPAD 502 (Konica Minolta, Nieuwegein, Netherlands) and Dualex (Force A, Orsay, France). Chlorophyll meter readings reflect the intensity of the green colour of the foliage, and are thus wellcorrelated with leaf chlorophyll and N concentrations (Spring and Zufferey, 2000; Cerovic et al., 2015; Aranguren et al., 2018; Vrignon-Brenas et al., 2019). Therefore, chlorophyll content can be used to diagnose plant N status, making such readings effective tools for N monitoring. Knowledge of growth stage and sampling method is critical for a reliable estimation of grapevine N status in the vineyard. Interpretation thresholds have been proposed for measurements taken with the N-Tester for the cultivars Chasselas, Pinot noir and Gamay at the phenological stage of veraison (Table 4). Measurements taken earlier in the season are not recommended due to higher variability of the readings, since they are greatly influenced by cultivar, water status (e.g., severe drought), deficiency of other nutrients (e.g., magnesium, iron), disease symptoms on the leaves and canopy management (Cerovic et al., 2015; Friedel et al., 2020). Thresholds are currently lacking, but ideally, they should be available for every cultivar, and even for every cultivar-rootstock combination. Ongoing research is aiming to remotely characterise vine physiology and berry composition with the Normalised Difference Vegetation Index (NDVI) (Taskos et al., 2015; Kotsaki et al., 2020a; Kotsaki et al., 2020b). The NDVI is well-adapted to assessing the spatial variability of vine N status, and it can fine-tune agronomic practices in specific areas within a vineyard. However, NDVI has the drawback of combining information; for example, leaf density (related to vine vigour, which does not depend on vine N status alone) and leaf colour intensity (related to vine N status and, to a lesser extent, the variety).

2.4. N isotope composition

N dynamics in grapevine can be monitored by analysing isotopes for research and development purposes. Elemental N has two stable isotopes (¹⁴N and ¹⁵N); *i.e.*, atoms with the same number of protons (seven protons for N) and different numbers of neutrons. Both are present in nature at the natural abundance of 99.634 and 0.366 atom % respectively (Deléens *et al.*, 1997). The stable N isotope composition of a sample is determined by isotope ratio mass spectrometry (IRMS). It is reported as a δ^{15} N value, which is the relative deviation of the sample heavy-to-light isotope ratio ¹⁵N/¹⁴N (R_{sample}) from an international reference ($R_{standard}$ of atmospheric N₂) (Coplen, 2011):

$$\delta^{15} N_{\text{sample}} = \frac{R \left(\frac{^{15}N}{^{14}N}\right)_{\text{sample}}}{R \left(\frac{^{15}N}{^{14}N}\right)_{\text{standard}}} - 1 \quad (1)$$

The δ unit is milliurey (mUr) as defined by the International System of Units (Coplen, 2011). A review (Santesteban *et al.*, 2014) and two studies (Durante *et al.*, 2016; Paolini *et al.*, 2016) have described variations in ${}^{15}N/{}^{14}N$ isotope ratios in

TABLE 4. Thresholds for the interpretation of N-Tester index with regard to vine N status for Chasselas, Pinot noir and Gamay.

	N-Tester index		Corresponding
Chasselas	Pinot noir	Gamay	grapevine N status
< 420	< 460	< 380	Very low
420 - 460	460 - 500	380 - 430	Low
460 - 540	500 - 580	430 - 530	Normal
540 - 570	580 - 620	530 - 580	High
> 570	> 620	> 580	Very high

Measurements are implemented at veraison on adult leaves in the bunch area (Spring and Verdenal, 2017).

natural abundance from soil to wine. Several isotope fractionations occur during the soil N cycle, and then to a lesser extent through grapevine N metabolism (Santesteban et al., 2014). δ^{15} N values observed in plant tissues are mainly related to N source, with lower δ^{15} N values (*i.e.*, 0.2 mUr on average) for inorganic fertilisers than for organic matter (8.1 mUr on average) (Santesteban *et al.*, 2014). Grape $\delta^{15}N$ values are usually less than soil δ^{15} N values (Durante *et al.*, 2016). After grapevine N assimilation, a ¹⁵N enrichment can be observed from roots (6.6 mUr on average) to must (33.7 mUr on average) (Verdenal et al., 2020). N fractionation is related to several factors, such as water availability and fruit load. The water constraint that a grapevine can face during the vegetative season will negatively influence wine δ^{15} N values (Spangenberg and Zufferey, 2018). Conversely, fruit load will positively influence must $\delta^{15}N$ values; *i.e.*, from 19.5 mUr on average under low-yielding conditions to 33.7 mUr under high-yielding conditions (Verdenal et al., 2020). Winemaking processes do not change $\delta^{15}N$ values from must to wine (Durante et al., 2016). Despite multiple isotope fractionations from soil to grape, $\delta^{15}N$ values for leaves, grapes and wines conserve the variability of $\delta^{15}N$ found in the corresponding soil (Paolini et al., 2016; Spangenberg and Zufferey, 2018).

In contrast to natural abundance, N labelling consists of applying an N source to the grapevine with a known ¹⁵N abundance; *i.e.*, ¹⁵N is artificially substantially enriched or depleted (e.g., 10 atom %). Such a high concentration of ¹⁵N is easily detectable and quantified in the plant organs. When studying N metabolism, this method allows the labelled N, which has accumulated in specific organs to be traced and quantified, and it provides an insight into the fate of crop-applied N in terms of its uptake, assimilation, distribution and release (Van Cleemput et al., 2008). Variations in the natural abundance of $\delta^{15}N$ and possible isotope fractionation are considered negligible compared to the ¹⁵N content of the labelled source (Verdenal et al., 2016a). Once the plant has assimilated the labelled N, each fraction of the plant can be analysed separately as described hereafter.

The absolute abundance of ${}^{15}N$ (A %, atom percent) is the proportion of heavy isotopes per 100 N atoms (Cliquet *et al.*, 1990):

$$A\% = \frac{R}{R+1} \times 100 \quad (2)$$

Relative specific abundance (*RSA*, atom percent) is the proportion of newly incorporated N atoms originating from the labelling relative to total N in the sample (Cliquet *et al.*, 1990). The *RSA* also represents organ

sink strength, which is independent of organ size (Deléens *et al.*, 1997):

$$RSA = \frac{A\%_{sample} \text{ excess}}{A\%_{N} \text{ supplied} \text{ excess}} = \frac{A\%_{sample} - A\%_{non labelled control}}{A\%_{N} \text{ supplied} - A\%_{non labelled control}}$$
(3)

The new N pool, which has originated from the labelling, can be quantified in each plant fraction and the partitioning (% P) can subsequently be calculated (Cliquet *et al.*, 1990):

$$new N pool_{fraction} = RSA_{fraction} \times NQ_{fraction}$$
(4)

where NQ is the total N quantity

% P =
$$\frac{\text{new N pool}_{\text{fraction}}}{\text{new N pool}_{\text{whole plant}}} \times 100$$
 (5)

The overall net N uptake can then be calculated:

net N uptake =
$$\frac{\text{new N pool}_{\text{whole plant}}}{\text{labelled NQ}_{\text{supplied}}} \times 100$$
 (6)

Exclusively used for research purposes, the isotope labelling method has been applied on grapevine since the 1980s in order to study plant N metabolism (Conradie, 1983; Glad *et al.*, 1994; Morinaga *et al.*, 2003; Zapata *et al.*, 2004a; Zapata *et al.*, 2004b; Iandolino and Williams, 2014; Clarke *et al.*, 2015; Williams, 2015; Verdenal *et al.*, 2015; Verdenal *et al.*, 2016a; Verdenal *et al.*, 2020; Brunetto *et al.*, 2016; Hannam *et al.*, 2016).

2.5. Grape analysis

The analysis of grape N content at harvest gives an overall picture of plant N status over the entire season, including the ripening period, making it probably the most accurate indicator of grapevine N status. Conversely, the previously mentioned plant indicators (*i.e.*, leaf N content and chlorophyll index) are usually obtained at either the phenological stage of veraison (BBCH 85) or before. They consequently only give an integrative view of N metabolism until veraison. Van Leeuwen et al. (2000) have compared the performance of several indicators of grapevine N status. Both total N content and yeast assimilable N (YAN) in grape must were found to be correlated and highly responsive to fertilisation practices. YAN is the part of must N compounds that is assimilable by yeasts during alcoholic fermentation (AF), and it comprises ammonium (NH_4^+) and AAs (excluding proline and hydroxyproline); it also informs the winemaker about the must's fermentability. Low grape N concentration at harvest can be a sign of unbalanced vine nutrition. However, N fertilisation is not always the suitable solution. During grape development and ripening, berry N nutrition may be restrained by numerous biotic and

abiotic factors, which may alter both N assimilation and partitioning in the plant, despite soil N abundance (Delas, 2010; Spring et al., 2012). Consequently, low correlations are often observed between plant-based N indicators and grape N content at harvest, particularly if the grapevine N metabolism has been restrained during the ripening period. van Leeuwen et al. (2000) established a threshold at 180 mg/L of YAN, above which the grapevine N requirements are fulfilled and N fertilisation should be interrupted during the following year. This threshold may be lower in some situations for the production of red wine (van Leeuwen et al., 2018). To interprete YAN at harvest as an indicator of plant N status, thresholds need to be determined. Since the YAN concentration is also related to grape variety, specific thresholds are required for each variety.

Early determination of must YAN content can potentially be used for the purpose of N fertilisation at the beginning of grape ripening, with the aim of increasing must YAN content at harvest. At veraison, grapes are already rich in N, mainly in the form of NH₄⁺. YAN concentration generally decreases during grape ripening due to the decrease in NH⁺, while AA concentration remains relatively stable (Nisbet et al., 2014). A large database produced by the Agroscope Institute highlights the correlation between grape N content both at veraison and at harvest across 16 vintages (1997-2012), five cultivars and three experimental vineyards (240 data points, Lorenzini et al., unpublished data, 1997-2012). Each year, ripening was monitored in selected plots of the main cultivars every week until harvest, as an indication of N for the grape growers. Approximately, 80 % of the situations had equivalent N concentrations at veraison and harvest (Figure 5). This confirmed the results of Nisbet *et al.* (2014), who also found a strong correlation between YAN content at veraison and at harvest ($r^2 = 0.82$). When initial N content was higher than 140 mg N/L, N content at harvest was still above that deficiency threshold in 70 % of the cases, and when initial N was deficient, N deficiency was confirmed at harvest in 90 % of the cases. N analysis at veraison is too variable for a precise prediction of N content at harvest, but it still gives a good indication of N deficiency.

To conclude, there is no unique indicator to determine vine N status. In most cases, the absence of universal thresholds is limiting, as the desired N status in both plant and grape is relative to grape variety, yield and production objectives. Plant N status can be assessed by both applying routine dosage of YAN at harvest and observing plant physiology (vigour, leaf colour and bud fruitfulness). With this information, N fertilisation and agronomic practices can be fine-tuned to obtain optimum plant N status. A combination of several indicators will increase the reliability of a diagnosis of vine N status.

3. Nitrogen metabolism

Grapevine N restriction affects fruit N accumulation, altering the abundance of certain AAs more than others, and thus changing the fruit AA profile (Schreiner *et al.*, 2014). Organic N solutions available in industry to manipulate AA concentrations in musts are still expensive, and they have less impact on wine



FIGURE 5. Linear regression between the concentrations of YAN at veraison (onset of ripening) and at harvest. Data collected on Pinot noir from three different vineyards from 1997 to 2012. Risk of incomplete fermentation: green = none; orange = moderate; red = strong (Lorenzini *et al.*, unpublished data, 1997-2012, Agroscope, Switzerland).

aromas than vineyard management practices. A wine sensory profile will mainly depend on the initial grape composition at harvest, which has to be managed at vineyard level, despite the substantial influence of the winemaking process (Gutiérrez-Gamboa et al., 2019). In most vineyards, N availability is often limiting, which largely affects plant physiology, such as canopy expansion, root morphology, floral induction and seed dormancy (Hachiya and Sakakibara, 2016). A balanced grapevine N metabolism is thus required to achieve optimal N accumulation in the grapes and, ultimately, the desired wine flavour. Understanding N use efficiency (NUE) is critical for optimising the parameters involved in N metabolism to obtain both optimal production and composition of grapes at harvest, while reducing N fertilisation and environmental impacts (Masclaux-Daubresse et al., 2010).

3.1. Nitrogen use efficiency

It is commonly admitted that nearly 60-70 % of N applied to crops through fertilisation is actually lost, mainly by soil leaching and by gaseous emission (Masclaux-Daubresse et al., 2010: Reddy and Ulaganathan, 2015). Optimising grapevine N use with the aim of improving grape quality, while reducing the use of fertilisers and minimising N runoff into the environment, is critical for both the grower and the environment. The concept of NUE has been developed by several researchers (Lea and Azevedo, 2006; Masclaux-Daubresse et al., 2010; Xu et al., 2012). Crop NUE is usually represented by total yield produced per unit of fertiliser N applied (Xu et al., 2012). The definition of NUE differs, however, depending on whether crops are cultivated for biomass or grain (Masclaux-Daubresse et al., 2010). In the case of wine production, maximum grape yield is generally not the main target. Optimal grapevine NUE is not only a case of balancing N status between vegetative and reproductive growth, but also of favouring the accumulation in grapes of AAs and subsequent metabolites known to enhance wine quality (Schreiner et al., 2018). Optimal NUE can also contribute to a reduction in N input, and thus environmental impact. NUE is the combination of two parameters: 1) assimilation efficiency (i.e., uptake and assimilation), and 2) utilisation efficiency (i.e., allocation and remobilisation) (Kant et al., 2011). N uptake and N assimilation refer to two different processes: N uptake is the process of collecting inorganic N from the environment, from soil in particular; N assimilation is the formation of organic N compounds necessary for growth and development (e.g., the AAs). In order to provide favourable conditions by adapting agricultural practices, it is first necessary to understand the agronomic traits that influence the efficiency of assimilation and utilisation; this would help to either

enhance grape composition with the same N input, or maintain grape composition with lower N input (Kant *et al.*, 2011).

3.2. N uptake

Grapevines assimilate neither atmospheric dinitrogen (N_2) nor N bound to the organic matter present in the soil. Soil NO₃⁻ and NH₄⁺ are the primary N source for grapevines, but they can also take up organic N (urea, AAs and peptides) to a lesser extent (Keller, 2015; Hachiya and Sakakibara, 2016).

Root uptake is an active process (energy consuming) principally occurs in the fine roots which (Zapata et al., 2004b). NO₂⁻ uptake initially consists of a radial diffusion along both symplastic (interconnected cytoplasm) and apoplastic (intercellular spaces) routes: ions move through the root epidermis up to the endodermis. The endoderm plays a boundary role in the selection and regulation of ions. Energy from adenosine triphosphate consumption is used to 'pump' protons out of the root cells into the soil; protons diffuse back into the cells, carrying negatively-charged NO₃⁻ with them (Keller, 2015). The soil NO_3^- concentration is highly variable. The complex processes of active uptake by the roots allows the plant to adjust nutrient uptake according to its needs and to soil N availability. NO3⁻ assimilation depends on both soil and plant N status and involves hormonal controls and interactions with carbon (C) metabolism and status. Root elongation is stimulated by soil N deficiency (Xu et al., 2012). Numerous genes (> 20) are involved in regulating membrane transport (Morot-Gaudry et al., 2017). N uptake rate is affected by root architecture, morphology and transporter activity on one hand, and by N form and concentration in the soil on the other (Xu et al., 2012; Morot-Gaudry et al., 2017).

Leaves can take up nutrients through their cuticle and stomata. Over the past decade, scientific progress has improved knowledge of plant response to foliar fertilisation, resulting in an increase of this practice in agriculture (Fernández and Eichert, 2009; Fernández and Brown, 2013). Leaf uptake is nonselective, in contrast to root uptake (Eichert, 2013). Nutrients penetrate the leaf cuticle and the stomata depending on the concentration gradient at the leaf surface. Janzen and Bruinsma (1989) demonstrated that up to 30 % of N present in wheat shoot tissues derives from atmospheric ammonia (NH₃). Furthermore, the application of foliar urea at veraison efficiently increases grape N content without influencing plant vigour, when all other management measures to optimise N status have failed or been insufficient (Lasa et al., 2012; Hannam et al., 2016). Urea is hydrophilic, and resulting N metabolites are easily transported from the leaves

$$\underbrace{\text{CO(NH}_2)_2}_{\text{urea}} + \text{H}_2\text{O} \xrightarrow{\text{urease}} \underbrace{\text{NH}_3}_{\text{ammonia}} + \underbrace{\text{H}_2\text{NCOOH}}_{\text{carbamic acid}} \rightarrow 2\text{NH}_{3(\text{gas})} + \text{CO}_{2(\text{gas})}$$
(7)

to the sink organs. After application, urea is rapidly hydrolysed into NH_3 and carbon dioxide (CO₂) as follows (see equation 7 above) (Krogmeier *et al.*, 1989).

 NH_3 cannot be directly assimilated by grapevine and will volatise into the atmosphere unless it reacts with water to form NH_4^+ . The reaction depends on ambient temperature and humidity; wetter and cooler conditions are usually favourable for limiting NH_3 volatilisation and increasing foliar fertilisation efficiency.

$$NH_3 + H_2O \rightarrow NH_4^+ + HO^-$$
 (8)

The combined formation of hydroxide (HO⁻) raises the pH locally, which further increases NH₃ volatilisation. When foliar applications are necessary due to low vine N status, a supply of 10 to 20 kg N/ha is usually recommended at veraison, split into two to four weekly applications, to prevent symptoms of toxicity due to temporarily high concentrations of NH₃ and NH₄⁺ (Figure 6) (Krogmeier *et al.*, 1989).



FIGURE 6. Leaf symptoms of NH_4^+ toxicity due to an excess of foliar urea.

3.3. Assimilation, transport and storage

Nitrate assimilation takes place in both the roots and leaves depending on N availability and supply (Llorens *et al.*, 2002). Once inside the root cells, nitrates can either be temporarily stored in the cell vacuoles for later use (buffer role), assimilated into organic compounds (*i.e.*, AAs), or transported to the leaves by the sap flow via the xylem vessels (Loulakakis *et al.*, 2009). Before assimilation, nitrates must be reduced into NH₄⁺ in a two-step process: nitrate is first reduced to nitrite (NO₂⁻) by the enzyme nitrate reductase, and then to ammonium by the enzyme nitrite reductase.

$$NO_3^- \xrightarrow{\text{reductase}} NO_2^- \xrightarrow{\text{nitrite}} NH_4^+$$
(9)

Xylem and phloem are efficient transport vessels in vascular plants. Xylem transports water and nutrients from the roots through the entire plant, while the phloem mainly transports organic compounds from the shoots and leaves to the rest of the plant. Glutamine and glutamic acid are the predominant AAs in the xylem sap, while arginine and glycine are predominant in the phloem (Gourieroux *et al.*, 2016). Over short distances, nutrients can also be simply diffused through unspecialised cell membranes and cytoplasm due to their charge (lipid and hydrophobic membranes) (Morot-Gaudry *et al.*, 2017). Figure 7 summarises N uptake and assimilation in grapevine.

In contrast to NO_3^- , NH_4^+ is toxic for plant tissues and is rapidly assimilated into AAs. Ammonium assimilation is catalysed by two enzymes: glutamine synthetase (GS) and glutamate synthase (GOGAT). The sequential action of the coupled GS/GOGAT has been found to play a predominant role in the assimilation of ammonium in higher plants (Loulakakis *et al.*, 2009).

glutamate + NH₄⁺
$$\xrightarrow{\text{GS}}$$
 glutamine (10)
(11)

glutamine + 2 oxoglutarate
$$\longrightarrow$$
 2 glutamates

An alternative pathway for ammonium assimilation involves the enzyme glutamate dehydrogenase (GDH). The main role of GDH seems to be different, however, as the reaction can be reversed, thus oxidising glutamate (Keller, 2015). (12)

$$NH_4^+ + 2 \text{ oxoglutarate} \xrightarrow{\text{GDH}} \text{glutamate} + H_2O$$

The accumulation of glutamine is the main source of organic N in grape; the synthesis of the other AAs occurs with the transfer of the glutamate amino group by different aminotransferases (Xu *et al.*, 2012; Ribéreau-Gayon *et al.*, 2017). AAs are the major form of organic N for transport and storage in the plant. The AAs are distributed throughout the entire plant via the phloem and the xylem. After harvest and before leaf fall, the major part of organic N is transferred and stored in the roots in the form of AAs - mostly arginine - and proteins (Zapata *et al.*, 2004a; Zapata *et al.*, 2004b).

3.4. N efflux

Net N uptake refers to total N influx minus total N efflux (Hachiya and Sakakibara, 2016). Plant N losses must be included in the N budget to avoid an overestimation of N losses in soil and an underestimation of plant N uptake (Xu *et al.*, 2012). Knowledge about the amount and composition of organic compounds released into the soil by plant roots is incomplete and



FIGURE 7. N uptake and assimilation in grapevine. NO₄, nitrate; NO₄, nitrite; NH₄, ammonium; AA, amino acid.

not even available for grapevine, largely because of methodology limitations.

Nitrate, ammonium and AAs can be released by the roots into the soil, as a result of root activity and root life span. The rhizosphere is a site of intense interactions between roots and soil; organic components released from the roots influence the solubility and transport of nutrients and the decomposition of organic materials, as well as the activity and turnover of microorganisms (Reining et al., 1995). Zapata et al. (2004a) showed that about 60 % of grapevine root N is lost from the perennial tissues between bud break and the onset of flowering. However, this amount does not correspond to the increase in N content in the annual tissues. This increase is only around 40 %, suggesting that approximately 20 % of the N reserve is lost early in the season via grapevine root necrosis (fine roots in particular) and to a lesser extent sap bleeding. Reining et al. (1995) investigated this issue in wheat: using a split-root experimental design with labelled N supply on one side, they showed that approximately 7 % of assimilated N was released into the soil of the unlabelled compartment. Merbach et al. (1999) confirmed the

release of 5-6 % of ¹⁵N previously assimilated by wheat, which represents 15 kg N ha-1 of N released by roots into the soil. Of the N exudates, 60 % was found in the soluble organic N pool and 9 % in the inorganic N pool (Janzen and Bruinsma, 1989). Ammonium efflux from the roots inhibits root cell elongation (Li et al., 2010; Reddy and Ulaganathan, 2015). A nitrate efflux transporter has been identified in Arabidopsis roots, but its physiological role still needs to be determined (Xu et al., 2012). Reddy and Ulaganathan (2015) have explained that plants release ammonium into the soil to maintain N homeostasis, because a high internal NH_4^+ concentration is toxic to the plant and reduces N uptake efficiency. The decomposition rate and the release of N compounds by *Quercus* fine roots are not only functions of environmental temperature, rainfall and humidity, but also of initial soil composition and root diameter (Usman et al., 2000). In the case of Pinus, both the decomposition rate and the release of N compounds are negatively correlated to initial soil N content (Jing et al., 2019). Changes in chemical traits of fine roots affect fine root decomposition to a greater extent than do changes in soil N availability (Gang et al., 2019).

To a lesser extent, photorespiration also induces N losses through the emission of NH, by leaves (Kumagai et al., 2011). Differences in NH₃ losses between rice cultivars are a result of their different GS activities, which result in different capacities for the reassimilation of photorespiratory NH₂. Kumagai et al. (2011) also suggested that NH, emissions in rice leaves are not directly controlled by transpiration and stomatal conductance. The main factor for N losses (in the form of NH₂) from the aboveground parts is the excess of N accumulation in the tissues compared to N assimilation (Xu et al., 2012). Leaf senescence is also a cause of N loss, even if most of the soluble N components are translocated to other organs via the phloem before leaf fall. However, the leaves fall on the ground and are a potential source of nutrients. Similar soil/roots and atmosphere/leaves interactions are likely in the case of grapevine, but their proportions are still unknown. Research on this subject is of critical importance to obtain a complete picture of N dynamics in grapevine.

3.5. Synergy between C and N metabolisms

The assimilation of NO_3^- and NH_4^+ into AAs is a dynamic process that is regulated by both internal factors (C and N metabolism) and external factors (environmental conditions) (Keller and Koblet, 1995). Besides water availability, C-N interaction is a cornerstone of optimal biomass production. Vrignon-Brenas *et al.* (2019) demonstrated the preponderant role of plant N status in C balance related to both gain and storage. Indeed, both biomass production and photosynthesis activity require N supply, which, in turn, depends on photosynthetised-C compounds for nitrate assimilation (Gauthier *et al.*, 2010). Stitt and Krapp (1999) published a detailed review describing the interaction between elevated CO₂ and N nutrition. Nitrate reduction requires a parallel C oxidation via the respiration process (Xu et al., 2012). The C-skeletons and energy from starch and sucrose are essential for the biosynthesis of glutamine (Masclaux-Daubresse, 2010). In other words, C can be viewed as a substrate for N assimilation. Consequently, grapevine C status strongly influences N assimilation, which is fast when C status is high (Keller and Koblet, 1995). Conversely, under adverse environmental conditions, which restrict photosynthetic activity, N assimilation is reduced and AA synthesis is consequently limited. Higher N status stimulates both light-saturated photosynthesis activity and respiration rate. Under high N availability and proper light intensity, grapevine N demand is met, and assimilated N is accumulated in the root reserves, inducing lower N uptake (Keller, 2015). When subjected to low N supply and high irradiance, grapevine exhibited the highest root-to-shoot ratio (Grechi et al., 2007). The regulation of N uptake and assimilation by photosynthesis ensures that N and C uptakes are correlated (Masclaux-Daubresse et al., 2010).

4. Winemaking

4.1. Grape N composition and yeast assimilable N

Approximately 50 % of grape N is found in the seeds and skin, 8 % in the stem and 40 % in the must (Hernández-Orte *et al.*, 1999). Figure 8 illustrates the average must N composition at harvest. Free AAs are the main N form in the must, representing 60-80 % of total N (Aerny, 1996). There are two categories of free AAs depending on their molecule structure: AAs with a primary amine (-NH₂), representing 50-90 % of total AAs; AAs with a secondary amine (-NH-)



FIGURE 8. Average grape must N composition. Grey = inorganic N; white = organic N; tiled pattern = yeast assimilable N; AA, amino acid.

(Bell and Henschke, 2005). Other organic N forms are peptides (10-30 %), proteins (2-10 %) and trace amounts of vitamins, amines and nucleotides (< 5 %). Inorganic N forms are ammonium (5-20 %) and nitrate (< 5 %) (Henschke and Jiranek, 1993; Aerny, 1996; Bell and Henschke, 2005).

Yeasts play a major role in winemaking. For their growth and development, they assimilate soluble sugars, their major source of carbon, along with a mixture of nutrients, including lipids and N compounds (Ugliano and Henschke, 2009). Under the usual winemaking conditions, AAs with a secondary amine are not assimilable; *i.e.*, proline and hydroxyproline. Consequently, YAN is the sum of AAs with primary amine (organic) and ammonium (inorganic) (Figure 8).

$YAN = AA(primary amine) + NH_4^+$ (13)

For oenological purposes, YAN is usually measured on a centrifuged must sample collected at harvest and does not consider the grape solids. YAN content is an indicator of the must fermentability and quality potential (Martínez-Gil et al., 2012). Knowing YAN concentration in grapes before harvest can help winemakers to anticipate vinification conditions. Given the major role of YAN in winemaking, it is surprising that it is not always included in the must analyses to determine grape quality potential at harvest, along with the total soluble sugars, titratable acidity and pH. The assimilation order of the AAs during AF reflects both the initial must AA profile and the yeast strain preferences (Henschke and Jiranek, 1993). Yeasts select 'preferred' N sources that are rapidly assimilated into key components for their metabolism (Bell and Henschke, 2005; Crépin et al., 2017). However, Gobert et al. (2019) mentioned in their review that the 'preferred' and 'non-preferred' categories for YAN sources can widely vary depending on study conditions.

Oenological practices have major consequences for grape N extraction and, in turn, for must composition. N is present in the entire berry, but its distribution is uneven across berry fractions. Berry skin plays a central role in the synthesis of many compounds essential to wine quality, such as anthocyanins and aroma compounds (González-Barreiro *et al.*, 2015). During winemaking, the skin contact with must results in the extraction of the skin compounds and usually increases YAN content (Stines *et al.*, 2000). In the case of white wine making, cold racking is generally implemented before AF, and skins are not macerated in the must. Both actions are restrictive to YAN concentration in the must, which could explain why white wines are so sensitive to N restriction in the must.

4.2. Fermentation kinetics and must N correction

Must YAN concentration is often suboptimal, and this consequently restricts yeast growth and AF rate (Vilanova et al., 2007: Hannam et al., 2016). Below 200 mg YAN/L in the must, AF duration is negatively correlated to the concentration of YAN for a clarified must with average sugar concentration. Below 140 mg YAN/L, there is a major risk of stuck AF (Table 5)(Bell and Henschke, 2005; Torrea et al., 2011). This threshold is lower in the case of red must, because grape N extraction is higher due to longer skin contact; for example, the Swiss cultivar Humagne rouge often has very low must YAN concentration at harvest (< 100 mg/L), and AF is still properly completed in most cases. Similar observations have been reported for Pinot noir (Schreiner et al., 2018) and Merlot (Stockert et al., 2013). The Australian Wine Research Institute recommends a minimum of 100 mg/L YAN for red must (AWRI, 2020).

TABLE 5. YAN concentration thresholds to guarantee proper alcoholic fermentation kinetics in white grape must.

Must YAN content (mg/L)	Risk of incomplete fermentation of clarified must
> 200	None
140 < < 200	Moderate
< 140	Strong

To limit any risks related to must N deficiency, N supply to the must at the onset of AF - mainly in the form of diammonium phosphate (DAP) - has become a widespread practice. Bisson and Butzke (2000) recommended a YAN adjustment depending on °Brix degree: 200 mg/L at 21 °Brix, 250 mg/L at 23 °Brix, 300 mg/L at 25 °Brix and 350 mg/L at 27 °Brix. Martínez-Moreno *et al.* (2012) further demonstrated that the addition of a mixture of AAs increases AF kinetics and maximises sugar consumption more than DAP does. Rollero *et al.* (2016) highlighted the strong impact of yeast strain on the assimilation of N compounds and the formation of aromas during the AF.

Lactic acid bacteria require less N than yeasts during malo-lactic fermentation (MLF). N is assimilable to bacteria mainly in the form of AAs and, to a lesser extent, peptides (Ribéreau-Gayon *et al.*, 2017). A comparison of a wine's AA before and after MLF showed a very small decrease in their concentrations, probably due to the autolyse of

yeasts and bacteria (Alcaide-Hidalgo *et al.*, 2007). Despite the observation of temporary N deficiency during AF due to rapid yeast growth, N deficiency rarely occurs at the end of AF and is not responsible for the difficulties in MLF kinetics.

4.3. Flavour development related to N metabolism

Wine flavours are the result of a complex mixture of volatile and non-volatile compounds. Their interactions have physicochemical effects on the release of aroma (Robinson *et al.*, 2014). It is beyond the scope of this review to describe all the grape and wine flavour-active compounds and their metabolisms; abundant literature can be found on this topic (Rapp and Mandery, 1986; Henschke and Jiranek, 1993; Bell and Henschke, 2005; Swiegers *et al.*, 2005; Dunlevy *et al.*, 2009; Styger *et al.*, 2011; Robinson *et al.*, 2014; González-Barreiro *et al.*, 2015; Alem *et al.*, 2019). This review focuses on the role of N - particularly AAs - in the formation of the flavour compounds and their precursors.

The characterisation of AA composition in grape is of major interest because AAs are precursors of a large number of metabolites in grape and wine, particularly volatile compounds (Jackson, 2008; Garde-Cerdán *et al.*, 2018). In terms of flavour development, the initial N pool contributes either directly or indirectly to the following (Figure 9):

- ▷ Non-restricted C metabolism, involved in the synthesis of organic compounds responsible for varietal aromas, such as some aldehydes, terpenes and thiols;
- ➡ The accumulation of aroma precursors (*i.e.*, glyco-, glutathione- or cysteine-conjugates) which release their flavour-active compounds via yeast metabolism; and
- ⇒ The accumulation of nutrients essential for yeast metabolism (*i.e.*, YAN). This greatly influences the biosynthesis of flavour constituents (*e.g.*, organic acids, higher alcohols, aldehydes and phenols) during the AF (Hernández-Orte *et al.*, 2006; Jackson, 2008).

Grape development and composition define the potential of wine aroma, which later develops during winemaking. Grape N accumulation starts with berry set. During the ripening phase (from veraison to harvest), the synergy between C and N metabolisms enhances AA accumulation and the biosynthesis of aroma compounds and their precursors. Hernández-Orte *et al.* (2002) demonstrated that the characteristic aroma of some varieties are partially related to the AA composition of the must. Martínez-Gil *et al.* (2012) confirmed that it is possible to estimate the concentration of esters in wines from the must N concentration. Grape aroma compounds can be found in either volatile ('free')





or bound forms, such as glyco-, glutathione- and cysteine-conjugates (González-Barreiro *et al.*, 2015; Santamaría *et al.*, 2015). The bound form of these compounds is non-aromatic. As a result of the hydrolysis of glycoside, glutathione or cysteine, these compounds may then become volatile and thus aroma-active (Hjelmeland and Ebeler, 2015).

Terpenes, particularly monoterpenes and sesquiterpenes, are responsible for the characteristic aromas of varieties such as Gewürztraminer, Muscat and Riesling (Rapp and Versini, 1995; Robinson *et al.*, 2014). They are present in the grape in both free forms and non-aromatic glycoside precursors, and in variable proportions depending on the grape varieties. During winemaking, terpenes are released by the action of the glycosidase enzymes produced by grape, yeast and bacteria (Swiegers *et al.*, 2005). The presence of terpenes in wine is stimulated by higher YAN concentration in must (Hjelmeland and Ebeler, 2015).

Thiols (*e.g.*, mercaptohexanols) are another major group of wine aroma compounds, some of which give the characteristic aroma to varieties such as Sauvignon blanc and Petite Arvine. They are mainly present in the grape must as non-aromatic precursors. Helwi *et al.* (2016) demonstrated the positive impact of vine N status on the concentration of volatile thiols in wine through the increase in corresponding nonaromatic precursors in grape. Methoxypyrazines are N compounds naturally present in berries and associated with 'bell pepper' aroma, characteristic of several varieties, in particular Cabernet-Sauvignon (González-Barreiro *et al.*, 2015). Their concentration decreases during grape ripening. However, vine N status does not influence the concentration of this metabolite in grape at harvest, which is affected by the modification of the bunch-zone microclimate (Robinson *et al.*, 2014; Helwi *et al.*, 2015).

Phenolic compounds form another diverse family related to the composition and concentration of grape AAs. The total phenolic content of grape must has been reported to be negatively correlated with the N treatment given to vines (Bell and Henschke, 2005; Choné et al., 2006). However, Portu et al. (2015) reported increasing anthocyanin and flavonol concentration in wine after foliar treatment with phenylalanine. Phenylalanine is essential as a precursor in the flavonoid pathway for the synthesis of most phenolic compounds (Santamaría et al., 2015). In contrast, the application of different forms of N (i.e., urea, urea+sulphur and arginine) to Cabernet-Sauvignon decreased flavonoid concentration in wine (Gutiérrez-Gamboa et al., 2017a). Similar results have been reported for Chasselas wine, for which suboptimal must YAN was correlated with increasing flavonol concentration in wine, but no effect on total phenol



FIGURE 10. Synthesis of aroma-active compounds (in grey) through the yeast metabolism of AAs and ammonium.

AA	Chemical family	Compound	Concentration(mg/L)	Odour description	Odour threshold (mg/L)	References
Alanine	α-keto acid	a-ketopropionic acid				
	Aldehyde	Acetaldehyde	< 211	Fruity, rotting apple	100	1
Threonine	α-keto acid	a-ketobutyrate				
	Higher alcohol	1-propanol	13-125	Alcohol, fusel, sweet fruity, apple, pear	306	2, 3
		Methylglyoxal	0.1-1	Pungent	NR	1, 4
		Aminopropanol	NR	Fishy	NR	4
Valine	α-keto acid	α-ketoisovalerate				
	Aldehyde	Isobutyraldehyde	0.001-0.2	Apple, banana, melon, varnish, cheese	0.006	1, 4
	Higher alcohol	Isobutanol	9-174	Fruity, alcohol, solvent, green	40-75	4, 5, 6, 7
	Acetate ester	Isobutyl acetate	0.01-1.6	Banana, fruity	3.4 in 34 % spirit	4, 5, 3
	Fatty acid	Isobutyric acid	NR	Sweet, apple, rancid, butter, cheese	2.3	4,6
	Ethyl ester	Ethyl isobutyrate	NR	Sweet, rubber	0.015 in 10 % ethanol	8, 9
Isoleucine	α-keto acid	α-keto-β-methylvaleate				
	Aldehyde	2-methylbutyraldehyde	0.003-0.1	Green (herbaceous), malty, fruity	0.016	1, 4
	Higher alcohol	2-methyl-1-butanol	15-150	Marzipan (almond), malty	1.2 in water	4, 7, 10
	Fatty acid	2-methylbutanoic acid	NR	Fruity, waxy, sweaty fatty acid	2.2 in water	4, 10
	Ethyl ester	Ethyl 2-methylbutanoate	0-0.9	Strawberry, pineapple	0.001 in 10 % ethanol	4, 9
Leucine	α-keto acid	a-ketoisocaproate				
	Aldehyde	Isovaleraldehyde	0.04-0.25	Fruity, nut, banana, apple, cheese, amylic	0.004	1,4
	Higher alcohol	3-Methyl-1-butanol	45-490	Alcohol, nail polish, cheese	30	4, 5, 6, 8, 7
	Acetate ester	Isoamyl acetate	0.03-8.1	Banana, pear	0.03	4, 5, 6
	Fatty acid	Isovaleric acid	< 3	Rancid, cheese, rotten fruit, sweat	0.49 in water	4, 8, 10
	Ethyl ester	Ethyl isovalerate	0-0.7	Apple, fruity	0.0001 in water	4, 8, 10
Phenylalanine	α-keto acid	Phenylpyruvate				
	Aldehyde	Phenylacetaldehyde	0.0025-0.130	Floral	0.001	1
	Higher alcohol	2-Phenylethanol	4.0-197	Floral, rose, honey	10	5, 6, 8, 7
	Acetate ester	2-Phenylethyl acetate	0.01-4.5	Floral, rose, honey	0.25 in 10 % ethanol	5, 6, 8, 9
Methionine	α-keto acid	a-ketobutyrate				
	Aldehyde	3-methylthiopropanal	0.0005-0.080	Cooked potato, cabbage	0.250 in beer	1, 11
	Higher alcohol	3-methylthiopropanol	0.02-5	Cabbage, cooked vegetable	1	6, 7, 11
	Thiol	Methanethiol	NR	Rotten egg, cabbage	0.002-0.012 in beer	7, 11
	Ester	S-methyl thioacetate	0.002-0.016	Rotten vegetable	0.3 in beer	11

content was observed (Dienes-Nagy *et al.*, 2020). The effect of N nutrition on the phenolic compound content of grape is not yet fully understood and needs further investigation.

Winemaking strongly influences the development of wine aromas. Must N composition not only affects AF kinetics, but also the formation of aromatic compounds (Ugliano et al., 2007; Styger et al., 2011). The metabolism of yeasts releases a large number of aroma-active compounds; major volatile compounds derived from yeast metabolism include aldehydes, higher alcohols, esters and sulphur (S) compounds. all influencing wine flavour (Lambrechts and Pretorius, 2000; Santamaría et al., 2015; Garde-Cerdán et al., 2018) (Figure 10).

Ethanol, glycerol, fatty acids, acetic acid and carbon dioxide are only indirectly influenced by N metabolism. Crépin *et al.* (2017) studied aroma metabolism in *Saccharomyces cerevisiae*, and demonstrated that, contrary to what is generally acknowledged, only a limited fraction of the consumed AAs are directly incorporated by yeasts into proteins. Under the action of transaminases and deaminases, amine groups are collected from ammonium and AAs, and then are redistributed for *de novo* AA synthesis (Crépin *et al.*, 2017). The AAs can be further metabolised into higher alcohols through the Ehrlich pathway as follows (see equation 14 below) (Lilly *et al.*, 2006;Styger *et al.*, 2011):

The catabolism of AAs leads to the formation of α -keto acids and their corresponding aldehydes, which can be further reduced in 'higher alcohols' (Table 6).

The term higher alcohol refers to alcohols that possess more than two C atoms and have a higher molecular weight and boiling point than ethanol. Their concentration is usually positively correlated to must YAN concentration (Swiegers et al., 2005). However, Henschke and Jiranek (1993) reported a negative correlation between the YAN concentration in must and the content of 2- and 3-methyl-1butanol and 2-phenylethanol in wine. This may have resulted from the modified balance under N-deficient conditions between the reduced activity of the Ehrlich pathway and the increased activity of the biosynthetic pathway of branched-chain AAs from sugar metabolism (Swiegers et al., 2005). At moderate concentrations (i.e., below 300 mg/L), higher alcohols are desirable aroma compounds which contribute to the complexity of the wine fermentation bouquet. However, in high concentrations, 2- and 3-methyl-1butanol has been shown to have a negative impact on wine bouquet, masking the fruity notes in red wine (Cameleyre *et al.*, 2015; de-la-Fuente-Blanco *et al.*, 2016).

The formation of esters is related to the availability of both higher alcohols and fatty acid precursors. In fact, two major groups of esters are formed during fermentation: the acetate esters and the ethyl esters (Figure 10). Acetyl-CoA is condensed with higher alcohols to form acetate esters, and fatty acids are condensed with ethanol to form ethyl esters as a result of enzymatically catalysed reactions (Bell and Henschke, 2005). Despite their formation not being directly related to AAs, their concentration wine is often positively correlated to must in concentration (Bell and Henschke, Ν 2005; Ugliano et al., 2007; Barbosa et al., 2009). Most esters contribute significantly to the fermentation bouquet. Acetate esters have been found in wine in a concentration range of 0-18.5 mg/L, often above their detection threshold (Swiegers et al., 2005). Ethyl esters of branched chain fatty acids are only present in wine in concentrations below 1 mg/L. They are related to AAs, because they are formed from the oxidation of the aldehyde formed from α -keto acids during AA metabolism (Table 6). Swiegers et al. (2005) observed a synergy between grape and yeast metabolisms during the formation of characteristic ester profiles of grape varieties such as Chardonnay.

Suboptimal must YAN composition and concentration restrain yeast metabolism, including the sugar, N and S pathways. The production of both non-volatile and volatile metabolites is consequently affected and has sensory implications (Ugliano and Henschke, 2009). The increase in 2- and 3-methyl-1-butanol and 2-phenylethanol formation in these conditions demonstrates that modifications occur during yeast metabolism, and that there is also an increase in the formation of succinic acid and, consequently, in the succinic ester content of wine (Henschke and Jiranek, 1993; Garde-Cerdán and Ancín-Azpilicueta, 2008; Dienes-Nagy et al., 2020). The formation of free hydrogen sulphide (H₂S) ('rotten egg') and mercaptan ('onion') can increase in the event of YAN starvation during AF, which is deleterious to the wine bouquet. H₂S is a by-product of the biosynthesis of S-containing compounds, including AAs, methionine and cysteine. N supplementation during AF rapidly suppresses the accumulation of H₂S (Henschke and Jiranek, 1993), which is highly reactive and takes part in the formation



of other positive aroma-active S compounds, such as dimethyl sulphide (DMS) ('asparagus', 'truffle') (Swiegers et al., 2005). Although DMS does not give fruity aromas, it is indirectly involved in their development in wine (De Royer Dupré et al., 2014; Lytra et al., 2014; Lytra et al., 2016). The formation of 2-aminoacetophenone (2-APP) under low YAN conditions has been identified as being responsible for the atypical aging off-flavours in wines which are usually accompanied by an undesirable astringent and bitter flavour (Hoenicke et al., 2002; Linsenmeier et al., 2007). However, there is no clear correlation between the concentration of 2-AAP (or its precursor, indol-3-acetic acid) and the sensory perception of atypical aging. Schneider (2014) published a review about the atypical aging defect, discussing sensory discrimination, viticultural causes and oenological consequences, and thus illustrating the complexity of this problem. In contrast to N restriction, residual N in wine due to excessive supplementation lead can to precipitation (protein breakdown) and the formation of biogenic amines (allergen) and ethyl carbamate (carcinogenic) (Vincenzini et al., 2017). N excess may also lead to the development of undesirable microorganisms, such as Brettanomyces, responsible for wine spoilage (Bell and Henschke, 2005).

Suboptimal must YAN is usually corrected in the cellar with the addition of N to prevent sluggish AF. Aroma production in wine is affected by both the timing of N addition and the composition of the N source (Seguinot et al., 2018). The DAP supply to the must only increases the ammonium concentration, while a balanced must contains a complex mixture of ammonium and AAs. However, no clear correlation has been established between the impact of DAP supply and the wine sensory profile (Torrea et al., 2011). Conversely, many studies have demonstrated the positive influence of adding AA directly to the must on the formation of volatile compounds and, ultimately, on the development of wine aroma (Hernández-Orte et al., 2006; Garde-Cerdán and Ancín-Azpilicueta, 2008; Torrea et al., 2011). Fairbairn et al. (2017) investigated the effects of single AAs additions on the production of major volatile compounds in wine, which resulted in a predictable production of aromatic compounds with linear correlations. However, these correlations were lost as the complexity of the N sources increased. The choice of N source also affects the formation of glycerol and organic acids (Ugliano and Henschke, 2009). Several studies have demonstrated that the following AAs have a positive influence on flavour development during AF: threonine, phenylalanine, alanine and aspartic acid (Hernández-Orte *et al.*, 2006).

Understanding the fate of N sources during winemaking and their impact on the development of wine flavours could certainly help improve NUE. Controlling the development of wine flavours would then be possible by modifying the amount, type and timing of N sources. Moreover, the production of grapes rich and naturally balanced in AA compounds offers the winemaker high potential for making good quality wine.

THE IMPACT OF ENVIRONMENTAL CONDITIONS AND AGRONOMIC PRACTICES

Research on wine flavours has focused on AF conditions, since the majority of wine flavour compounds appear during winemaking as a result of yeast and bacteria metabolism (Robinson et al., 2014). However, since most of the substrates (particularly the N compounds) are grape-derived, the production of flavour compounds is strongly related to grape composition (Robinson et al., 2014). Plant physiology and grape composition depend on climate conditions and soil characteristics before and during berry development; they can be managed to some extent by optimising agronomic practices (Masclaux-Daubresse et al., 2010; Sweetman et al., 2009). The following section reviews the parameters, which influence grapevine N metabolism, distinguishing between the impact of the environment inherent to the vineyard and the agronomic management practices of the grape grower.

1. The environmental conditions of the vineyard

There are environmental conditions specific to the vineyard site which impact plant water and nutrient uptake, as well as leaf gas exchange and photosynthetic activity. Water, N and C are the three major components that significantly affect plant N metabolism, apparently following Liebig's law of the minimum. Any factor that either directly or indirectly influences water, C or N availability to the plant will potentially affects its N metabolism. The impacts of environmental conditions on grapevine N metabolism are summarised in Figure 11.

1.1.Climate and soil

The influence of climate on the plant metabolism can be considered at a regional scale (macroclimate), vineyard scale (mesoclimate) or plant scale (microclimate). In long-term experiments, the climate is also considered in terms of the 'year' effect.

Edaphic conditions (*i.e.*, soil depth, structure, temperature, water availability, pH, organic matter



FIGURE 11. Impacts of environmental conditions (i.e., climate and soil) on grapevine N metabolism.



FIGURE 12. Variability of yeast assimilable N in grape must at harvest.

Map obtained by ordinary kriging method based on a regular grid of eight samples per ha. Merlot, 2018, Saint-Julien, Bordeaux, France (van Leeuwen *et al.*, unpublished data).

content, limestone content and C/N ratio) highly influence the soil N cycle (turnover) and the subsequent N availability to the vine (van Leeuwen *et al.*, 2000; Hardarson *et al.*, 2008; Masclaux-Daubresse *et al.*, 2010; Marschner and Rengel, 2012). Consequently, grapevine N status (represented by must YAN at harvest) can vary considerably over short distances due to soil heterogeneity. To optimise vineyard management, it is important to visualise this spatial variability, which can be obtained by measuring YAN on a regular grid in a vineyard (Figure 12).

Soil temperature plays a major role in plant N uptake and metabolism: high temperatures (without water restriction) increase soil microbial activity and thus enhance organic matter mineralisation (Molina and Smith, 1997); furthermore, they increase root growth (higher fine root density) and thus favour N uptake (Clarke et al., 2015). Cold periods during springtime are a major cause of low N availability and uptake. However, excessively high air temperatures (e.g., above 40 °C) can also limit root N assimilation, partly due to lower photosynthesis and lower C availability: in response to heat stress the plant limits water consumption by closing stomata, which in turn reduces photosynthesis activity (Zufferey et al., 2017). Optimum temperature depends on grape variety, light intensity and phenological stage, and it is generally considered to be within the range of 10-35 °C (Hunter and Bonnardot, 2011; Keller, 2015). Temperatures out of this range can become a limiting factor for N metabolism. Global warming is a major concern in agriculture, as it also affects ambient CO₂ and solar radiation. It is generally projected that plant growth will increase under higher concentrations of ambient CO₂, due to improved photosynthetic activity (Tegeder, 2014). Because C metabolism and N metabolism are highly correlated, a higher concentration of C metabolites can improve N assimilation through the action of the enzymes GS/GOGAT; consequently, plant vigour will increase under unrestricted N availability. However, in many situations, restrictive N conditions can limit this increased capacity for using additional C (Stitt and Krapp, 1999).

Light is another factor that influences N metabolism. Poor weather conditions (*e.g.*, cloudy weather) can cause a decrease in N status, in response to reduced solar radiation (Keller, 2015). Light intensity influences photosynthesis rate and subsequent availability of C metabolites required for N assimilation (Masclaux-Daubresse *et al.*, 2010). Several studies have reported a correlation between grape exposure and the concentration of free aroma compounds or their bound glycosylated precursors (Bureau *et al.*, 2000; Marais *et al.*, 2001; Meyers *et al.*, 2013; Kwasniewski *et al.*, 2010). However, the relation between sunlight exposure and grape N content has not yet been clearly established.

Water and nutrients exist together in close association, because sufficient water availability (without waterlogging) will lead to nutrient solubilisation and facilitates plant N uptake and transport in the plant (Keller, 2005; Wang *et al.*, 2017). Vine water status depends on both climate-related factors (evapotranspiration and precipitation) and soil water holding capacity (van Leeuwen *et al.*, 2004). The best soils for viticulture induce both mild water restriction and non-limiting nutrient conditions (Fayolle *et al.*, 2019). Soil structure, texture and depth greatly affect water and nutrient availability for the plant, as they influence the soil water holding



FIGURE 13. Year-to-year variability of YAN in grape must at harvest.

Average data from six vineyard blocks, located on three soils and planted with two grapevine varieties (Merlot and Cabernet franc) in Saint-Émilion, France (adapted from van Leeuwen *et al.*, 2004-2011, unpublished data).



FIGURE 14. Impact of year, site and fertilisation on plant behaviour and must composition of the white cultivar, Doral (Chasselas × Chardonnay) in five vineyards (same plant material and agricultural practices) in a terroir study over three years in Switzerland.

White shapes = non-fertilised control treatment; black shapes = foliar urea supply at veraison (20 kg/ha of N) (adapted from Verdenal et al., 2016b).

capacity and the potential for root development (van Leeuwen and Seguin, 2006). In shallow soils, grapevine often has low grape N concentration, usually attributed to limited root colonisation (Reynard et al., 2011; Reynard et al., 2012). Under non-limiting water conditions, the plant can easily absorb the mineral N required for its development. High plant sap flow is a result of high transpiration and photosynthesis (Zufferey and Murisier, 2007). However, water excess due to high quantities of precipitation may induce low N uptake, either because of soil N leaching or because of waterlogging, which reduces the amount of oxygen in the soil needed for microbial activity. Conversely, under hot and dry conditions (*i.e.*, during the growing season in summer). N availability decreases at the soil surface due to low water content. In these conditions, water is a limiting factor for microbial activity, N solubility, N mobility and N uptake (Marschner and Rengel, 2012). Grapevine can counterbalance lower N availability with higher organic N mobilisation from the root reserves, as has been shown in maize by Wang et al. (2017). Moreover, root growth is limited in these conditions. Excessive water restriction may further induce a lower rate of photosynthesis and a subsequent lower plant C status. Climatic water deficit (precipitations minus evapotranspiration) during vegetative development is consequently negatively correlated to the accumulation of YAN in grapes (Spring et al., 2012). In an 8-year study combining six vineyards, three soil types and two cultivars, van Leeuwen et al. (unpublished data) observed a wide range of YAN values at harvest (from 80 to 150 mg/L) over the eight years (Figure 13).

This variability was explained by the soil type (45 % of total variance explained), cultivar (17 %) and climatic conditions of each year (14 %). The two vintages 2008 and 2011 showed significantly lower YAN values. This was probably due to the particular climatic conditions of those years: spring 2008 was cool and rainy, while spring 2011 was warm and particularly dry Hernández-Orte *et al.* (1999) confirmed that the highest grape YAN accumulation was obtained in the years with mild temperatures and moderate rainfall during ripening.

The impacts of pedoclimatic conditions on berry composition was assessed by Echeverría et al. (2017), who found that the synthesis of primary compounds is mostly dependent on both the climate and the climate-soil interaction, while the synthesis of secondary compounds (e.g., phenols) mostly depends on the source-sink relationship and the climate. These processes are regulated by both internal (C and N availability) and external factors (light, soil structure and composition, and soil microbiological activity) (Keller, 2015). A study by Verdenal et al. (2016) highlighted the strong overall impact of both climate and soil on grapevine N status. Five homogeneous plots of the white cultivar Doral (same plant material and agricultural practices) were chosen in different vineyards and were divided into control and N-fertilised treatments. Figure 14 shows the hierarchy of the three factors of discrimination; *i.e.*, year, site and fertilisation. First, the year (*i.e.*, climate) was the most variable and discriminating factor in terms of maturity and grape composition at harvest (i.e., sugar content and acidity). Second, the soil had a very steady impact on grapevine vigour (*i.e.*, bud fruitfulness, leaf area, pruning weight, bunch weight, yield and YAN) with the same site differentiation every year of the study. Third, fertilisation had a relatively small and variable impact on grapevine physiology and grape composition, despite a considerable impact on must YAN concentration, which significantly improved the wine organoleptic profile ($R^2 = 0.70$). This example shows the hierarchy in the climate-soil-plant ecosystem and demonstrates the possibility of improving grape composition via cultural practices, despite the major influence of both the year-to-year variability of climatic conditions and spacial variability of soil composition.

1.2. Phenotypic plasticity

Dal Santo et al. (2016) and Dal Santo et al. (2018) focused on the phenotypic plasticity of grapevine and dissected the berry transcriptome in response to the environment. Using an innovative data mining and statistical method, they investigated the separate impacts of climate, soil and grape variety, as well as their interactions. They found that grapevine is highly sensitive to environmental conditions and is characterised by a broad phenotypic plasticity (Dal Santo et al., 2016). In a study on Arabidopsis, Sakakibara et al. (2006) demonstrated that plants have the ability to sense their internal and external N status and to adapt to changing conditions by modifying their gene expression and morphology accordingly. Vines grown under low N and high irradiance conditions had the highest root-to-shoot ratios, and those grown under low irradiance and high N had the lowest (Grechi et al., 2007). N deprivation was found to enhance root growth at the expense of aboveground growth, whereas canopy size was significantly greater under high N conditions (Grechi et al., 2007). The plant can modify its root architecture, locally increasing root proliferation to reach nutrient-rich soil patches. The presence of nitrate stimulates the formation of lateral roots when it is applied to small sections of the primary roots (Lea and Azevedo, 2006). Leaves grown under low humidity (high vapour pressure deficit) have been found to be smaller than those grown under high humidity, even in the absence of soil water deficit (Keller, 2015). Canopy development and density ultimately affect the grape microclimate, particularly in terms of solar radiation interception. The grape AA profile of a given variety is generally similar from year to year, while AA concentration can vary widely (Hernández-Orte et al., 1999).

The plant affects, in turn, the soil composition through the process of N uptake. The rhizosphere is locally alkalinised and acidified following the uptake of nitrate and ammonium respectively (Hachiya and Sakakibara, 2016). Microbial activity is inhibited by a lower pH, which affects the fraction of the cation-exchange capacity occupied by cations and subsequent soil fertility. The optimum pH for N uptake ranges from 5.5 to 8.0 (Longbottom, 2009). Plant nutrition can also be enhanced by symbiosis with soil microorganisms, such as mycorrhiza, which are considered as 'new organs' unifying root tissues with the fungus mycelium in a symbiotic relationship. Mycorrhiza have a high capacity for assimilating N in the soil, thus benefiting the plant 'host' (Trouvelot et al., 2015). In return, the plant provides the fungus with photoassimilates. Such symbiosis concerns 95 % of plant species (Morot-Gaudry et al., 2017). Krishna et al. (2005) confirmed that the inoculation of mycorrhiza increases grapevine N content, as well as many other metabolites, such as nitrate reductase, chlorophyll, phenolics and proline contents. Grapevine rootstocks differ very little in their ability to form mycorrhiza, but other factors, such as crop load and soil moisture, have a great influence on root colonisation by mycorrhiza (Schreiner, 2003). The mycorrhiza colonisation of grapevines has been found to be unaffected by the presence of a cover crop (Klodd et al., 2016).

1.3. The concept of terroir

Understanding the impact of environmental conditions on plant N status helps make technical choices that will ensure and improve wine quality and sustainability. The International Organization of Vine and Wine defines the terroir as 'a concept that refers to an area in which collective knowledge of the interactions between the identifiable physical and biological environment and applied viti-vinicultural practices develops, providing distinctive characteristics for the products originating from this area. The terroir includes specific soil, topography, climate, landscape characteristics and biodiversity features' (Resolution OIV/VITI 333/2010). Vine growers must understand the intrinsic conditions of their vineyard in order to use the environmental conditions to their advantage (van Leeuwen et al., 2018). In order to reach a desired crop quality, it is necessary to integrate the optimisation of NUE into management practices, thereby modulating the influence of the environmental conditions (Figure 15).

1.4. Agronomic choices

No vineyard would exist without human intervention (van Leeuwen and Seguin, 2006). Reynolds (2010) summarised the common goals of human agronomic practices in cool climate conditions in four points: 1) keep the fruits warm, 2) keep the leaves exposed to light, 3) achieve vine balance between vegetative and reproductive organs, and 4) avoid water stress.



FIGURE 15. Illustration of the terroir concept, showing the influence of climate, soil and agronomic practices on grapevine N metabolism.



FIGURE 16. Agronomic practices influencing grape N metabolism.

However, there is no universal recipe, and vine growers must adapt their practices to their local environmental conditions in order to obtain optimal must composition. Habran *et al.* (2016) summarised the situation as follows: mild water deficit and moderate N availability can result in the metabolic synthesis of phenolic and aromatic compounds in berries, while surplus N can induce excessive vigour and exacerbate sensitivity to fungus. Consequently, N supply should be managed in such a way as to obtain a balance between vegetative and reproductive growth while preventing N deficiency. The objective is to optimise the grape N pool at veraison in order to enhance the biosynthesis of AAs and other aroma precursors in the must during grape maturation, while preserving vine balance and adequate ripening conditions. Several reviews have reported the influence of agricultural practices on the accumulation of aroma compounds and precursors in grapes (Poni *et al.*, 2018; Gutiérrez-Gamboa *et al.*, 2018), and on the development of aromas in wine (Robinson *et al.*, 2014; González-Barreiro *et al.*, 2015; Alem *et al.*, 2019). However, understanding how agronomic practices can specifically influence N metabolism would improve fruit quality control, as well as NUE and production sustainability (Boss *et al.*, 2014; González-Barreiro *et al.*, 2015). The following sections review the main agronomic choices that affect grape N metabolism. Four major factors are addressed: 1) plant material, 2) soil management, 3) vine balance, and 4) vineyard inputs (Figure 16).

2. Plant material

Ensuring that planting material is adapted to vineyard environmental conditions is a prerequisite for the production of quality grapes, and involves making choices regarding the rootstock, variety and clone.

2.1. Genetics and age

Grape varieties genetically differ from each other in terms of concentration and composition of N compounds in their fruits. Genetics has a great impact on grapevine NUE. Plants use several ways to sense environmental and internal N status. One is nitrate concentration, which regulates a wide variety of metabolic processes, including N and C metabolism (Sakakibara et al., 2006). The relative proportion of nitrate and ammonium in the soil influences N uptake. In rice, net nitrate uptake is inhibited by the presence of ammonium, compared to nitrate alone, while net ammonium uptake is enhanced by the presence of nitrate, compared to ammonium alone (Hachiya and Sakakibara, 2016). There is a general tendency across cultivars for increasing N uptake to induce lower leaf concentrations of K, P, Mg and boron (B) (Zamboni et al., 2016). Under non-limiting water and nutrient conditions, a significant correlation usually appears between plant vigour, plant N status and grape N concentration, with variations depending on the plant material; *i.e.*, rootstock, variety and/or clone.

The influence of genetics on N metabolism has been highlighted between the two varieties, Merlot and Pinot noir (Zapata et al., 2004b). In similar conditions, N uptake was higher in Pinot noir than in Merlot. Stines et al. (2000) suggested that the must AA profile is primarily genetically determined, whereas environmental conditions have a modifying effect. Several studies have shown a strong impact of grape varieties on the AA profile in grape must at harvest: the ratio of major AAs (proline, arginine, glutamine and histidine) to total AAs differed significantly across varieties (Hernández-Orte et al., 1999; Stines et al., 2000). Huang and Ough (1991) used the proline-to-arginine ratio to differentiate grape varieties. In Switzerland, a trial compared eight rootstocks over thirteen years, all grafted onto Pinot noir (clone RAC 12) and grown under homogeneous conditions (Spring et al., 2016a). The 13-year average leaf N content varied from 2.0 to 2.4 % DW, depending on the rootstock. It was correlated with vigour and must YAN content. The average YAN concentration greatly varied (from 132 to 224 mg/L) as a function of the rootstock. To a lesser extent, clones of the same variety also influence N metabolism, which has been shown in two studies. The first study compared 19 clones of Pinot gris (grafted onto 3309C) over seven years, while the second study compared 17 clones of Petite Arvine (grafted onto 5BB) over nine years, all grown under homogeneous conditions (Spring et al., 2016b; Spring et al., 2018). The average must YAN at harvest varied from 100 to 145 mg/L for Pinot gris, and 195 to 240 mg N/L for Petite Arvine, depending on the clone. Besides sensitivity to soil N content, the root mechanisms involved in N uptake are strongly affected by the variety-rootstock combination, which opens possibilities for adjusting grape composition via choice of planting material (Tomasi et al., 2015; Habran et al., 2016). Kant et al. (2011) reviewed the different genetic approaches for the improvement of NUE, starting with a description of the regulatory mechanisms involved in the plant response to N deficiency conditions. N uptake and remobilisation seem to be independently inherited traits; therefore, it is possible to combine favourable alleles when breeding for high NUE (Xu et al., 2012).

Plant material has long-term repercussions on wine style and quality and it must be determined with care, since not every vineyard can produce any possible wine style. First, the plant material must be chosen according to local climate to guarantee full ripeness of the grapes at the end of the season (van Leeuwen and Seguin, 2006). Climatic indices, such as the heliothermal index (Huglin, 1978), or the Grapevine Sugar Ripeness model (Parker et al., 2020) can be used for this purpose. Second, the plant material should be chosen according to soil N availability to guarantee balanced N nutrition. It should be kept in mind that grape N requirements are lower for red wine, compared to white wine, due to differences in the winemaking process; consequently, the producer might prefer to grow red varieties in vineyards, which have recurrent low N availability.

Moreover, grapevine age influences N metabolism. Using three white (Pinot blanc, Chasselas and Arvine) and three red cultivars (Gamay, Syrah and Humagne rouge), Zufferey and Maigre (2007) and Zufferey and Maigre (2008) compared the grapevine physiology and must composition of 4-8 years-old vines versus grapevines of 25 years of age and older. The young vines (< 8-years-old) were more susceptible to water stress and N deficiency due to their smaller and more superficial root system, and they had lower vigour, lower N status and lower grape YAN concentration. During the first years after planting, both root N reserves and N uptake restricted photosynthesis. Despite controlled and comparable yields, the red wines from older vines generally had higher quality aromas and a less astringent mouthfeel. Gamay wines showed no differences, which was probably due to the higher plasticity of the cultivar. No differences were found for white wines in terms of mouthfeel, and only a small preference for the aromas of wines from old vines was shown for Pinot blanc.

2.2. Maturity level

Grape maturity highly influences the berry AA profile. The accumulation of organic N and the formation of secondary metabolites within the berry, such as flavour-active compounds and their precursors, are affected by level of maturity (Hilbert et al., 2003; Robinson et al., 2014). Changes in AA profile during grape berry ripening have been demonstrated in several studies (Stines et al., 2000; Hilbert et al., 2003; Garde-Cerdán et al., 2009; Garde-Cerdán et al., 2018). Accumulation of grape YAN appears to differ significantly from metabolites other (González-Barreiro et al., 2015). Berry N accumulation starts as soon as berry set starts (BBCH 71). At the onset of ripening (veraison, BBCH 85), the berry YAN pool is mainly composed of glutamine and NH_4^+ , which both decline during grape ripening due to their conversion into other AAs (Stines et al., 2000). Overall, NH,⁺ concentration decreases while free AA concentration usually increases (Garde-Cerdán et al., 2018). Arginine accumulation in grape starts before veraison, while proline mostly accumulates during post-veraison (Stines et al., 2000). The accumulation of both arginine and proline seem to be developmentally regulated (Stines et al., 2000). Proline accumulation in vegetative tissues is often associated with osmotic stress during the post-veraison period (e.g., high concentration of sugars). However, Stines et al. (2000) argued that proline accumulation is part of normal fruit development, as in many other plant species, while the factors influencing the proline-to-arginine ratio remain unknown.

To monitor grape ripening, parameters such as sugars usually provide the most basic information about quality potential (González-Barreiro et al., 2015). A strong correlation was observed in the must between arginine accumulation and soluble sugar accumulation (Hernández-Orte et al., 1999; Garde-Cerdán et al., 2009). Garde-Cerdán et al. (2018) reported that technological maturity (i.e., optimal sugar content) coincides with the highest concentration of organic N compounds at 25 °Brix. Hence, they introduced the term, 'nitrogenous maturity'. González-Barreiro et al. (2015) confirmed that maximum flavour-active compound content is reached at maturity and remains constant over the following weeks. They described the aroma development in red grape as follows: esters characterise the beginning of ripening period, aldehydes the middle and alcohols the end. Consequently, they suggested using the alcohol-to-aldehyde ratio to optimise on the harvest date and to maximise grape aroma. However, the accuracy of this index seems to be low

for white varieties. The late formation of alcohols is desirable as they are precursors to the formation of esters in the presence of carboxylic acids during AF (González-Barreiro *et al.*, 2015). In view of the major role of must N (particularly YAN) in AF kinetics and in the development of wine flavour, must YAN concentration before and at harvest could be used as an indication of grape quality. In any case, must YAN should be routinely analysed for winemaking purposes, on the same basis as sugars and acids.

3. Soil management

Soil maintenance has a direct impact on grapevine root development and nutrition, with further consequences on must N composition and wine sensory profile (Bouzas-Cid *et al.*, 2018a). Proper soil maintenance guarantees sustainable soil fertility with proper N mineralisation and availability of mineral N for the plant. However, vineyard soil must be prepared before planting in order to relieve soil compaction and optimise soil structure. If necessary, an initial manuring can be applied. After planting, soils are usually managed through tillage, herbicides and/or cover crop.

Cover cropping is a common practice in vineyards which greatly affects soil N availability (Spring, 2001). The presence of a cover crop offers many advantages, such as reduced maintenance, reduced herbicide use, better soil stability, higher soil bearing capacity and permeability, and lower erosion. It also reduces plant N status and, consequently, overall grapevine vigour by limiting N availability (Tesic et al., 2007; Reeve et al., 2016). Depending on the cover-crop mixture, N competition between grapevine and cover crop can be exacerbated under low water availability (Celette et al., 2009). The implantation of legume (e.g., Trifolium subterraneum), which have the capacity of fixing N from the atmosphere, is an interesting alternative for limiting such competition (Spring, 2002). Both temporary and permanent cover crops decrease soil N mineralisation, due to a faster drying of the superficial soil layers (Celette et al., 2009). Grapevines may adjust their root development to access deeper water resources, although deeper lavers contain less mineral N (Celette et al., 2009). Vegetative development is limited, thus improving the grape microclimate (better sun exposure and higher temperature) (Maigre and Aerny, 2001a; Reeve et al., 2018). Lower N availability has been found to be related to a higher concentration of higher alcohols and phenolic compounds in wine (Choné et al., 2001; Maigre and Aerny, 2001b). However, over four years of experimenting on Gamay, researchers found that the wines produced from vines with bare soil treatment were usually preferred to those from vines with cover crop treatment, due to



July

November

FIGURE 17. Trial of *Hordeum murinum* as a cover crop. Sowing in 2007 and pictures taken in 2008. Epesses, Switzerland (Spring, 2008).

increased varietal aromas and reduced astringency (Maigre and Aerny, 2001b). It is difficult to control vine vigour exclusively via cover cropping. An excess of competition for N and water between the grapevines and the cover crop can damage the yield and the wine quality. In the 1980s, cover cropping was widely developed in Swiss vineyards and the winemakers started observing difficulties in AF kinetics, with the development of off-flavours, particularly in white wines. The lower N content in berries was explained by the reduced availability of soil N due to cover cropping (Gouthu et al., 2012). Cover crop affects grapevine N status in the long-term, as it also affects the perennial reserve of N build-up necessary for the next year (Celette et al., 2009; Gouthu et al., 2012). Celette and Gary (2013) further showed that the dynamics of water and N availability for the grapevine are partially uncoupled.

The cover crop must be adapted to soil conditions, as there is no universal cover crop suitable for all vineyards. In a situation of excessive grapevine vigour, the use of a competitive cover crop can be an effective strategy for limiting vine growth and yield, although water availability and grape YAN content should be monitored (Reeve *et al.*, 2016). To minimise competition with grapevine, a temporary cover crop can otherwise be recommended. The cover crop can also be limited to the row spacing (80 % of the surface, weeding under the row) and even

to every other row (only 40 % of the surface). The choice of the cover crop species is essential. The ideal cover crop species has the following characteristics (Delabays *et al.*, 2000): quick development, low vigour during summer, strong allelopathy towards other species, winter covering and frost resistance, and spontaneous seeding and regeneration. Ideally, the cover crop should grow during spring and autumn and dry during the summer, thus inducing lower competition for N and water and promoting grapevine development, as in the case of *Hordeum murinum* (Figure 17).

The use of the legume, Trifolium subterraneum, as a cover crop (every other row) increased the soil N content during the summer and increased the YAN content of Chasselas grapes at harvest in Switzerland (Spring, 2001). Consequently, AF was faster and the wines were significantly preferred (better aroma and mouthfeel, lower bitterness), in comparison to a mix of perennial and competitive grasses, such as Festuca rubra, Festuca ovina, Poa pratensis, Poa compressa, which reduced soil N availability (Spring, 2002). However, these results contradict those of Bouzas-Cid et al. (2018a) obtained from the cultivar, Mencia, under humid conditions in Spain. Depending on the environmental conditions, an adapted cover crop could be a sustainable solution for soil management and an option for modulating must composition and wine sensory profiles.

4. Vine balance

Vine balance is a common term used to express the balance between the vegetative growth and reproductive development of a plant. A balanced vine has the appropriate capacity for producing fully ripened grapes, while building nutrient reserves for the following year (Howell, 2001; Lakso and Sacks, 2009). To reach this balance, both canopy size and crop load have to be controlled. Clingeleffer (2009) highlighted a trend over the last century towards lower planting density, larger canopy size and higher crop load per vine. Larger trellis systems have been created to accommodate the larger number of shoots (*e.g.*, Geneva double curtain).

4.1. Canopy management

Grapevine trellising and canopy management (i.e., pruning, defoliation and hedge trimming) affect plant growth, fruit zone microclimate and consequently fruit composition (Azuma et al., 2012). It can also affect N nutrition. Rühl and Clingeleffer (1993) observed that N accumulation in roots and wood can vary from 88 to 139 kg/ha, depending on the pruning system, with spur-pruning resulting in higher N accumulation than minimal-pruning. An ideal canopy maximises light interception and guarantees a nonlimiting source of carbohydrates for the grapes through optimum photosynthesis activity. An abundance of carbohydrates contributes to non-limited N assimilation in leaves and roots. Light exposure enhances N reductase activity in leaves (Perez and Kliewer, 1982). A large canopy also guarantees adequate refilling of root N reserves, mainly in the form of AAs, in prevision for the following year (Zufferey et al., 2015; Verdenal et al., 2016a). Furthermore, an ideal canopy creates an optimal bunch microclimate, favouring the formation of secondary metabolites, such as phenolic compounds (Keller, 2015).

Plant N content and vigour are usually correlated (Verdenal et al., 2020). An oversized canopy can, however, induce fruit N deficiency uncoupled from plant vigour (i.e., due to improper canopy management), despite unlimited N resources for the plant (Spring et al., 2012). A strong negative correlation between grape N concentration and canopy trimming height has been shown for Chasselas and Pinot noir, despite unchanged fruit load, as if the N content were 'diluted' within the volume of the biomass (Spring et al., 2012). Verdenal et al. (2016a) observed that an oversized canopy (+31 % DW) induced a decrease in grape YAN concentration of up to 53 %. This situation can occur in vigorous grapevines in the absence of water restriction, and can strongly affect grape YAN concentration. Conversely, researchers found that a smaller canopy (due to either severe

pruning, shorter height or removal of lateral shoots) induced higher grape YAN concentration, but then full ripeness was difficult to attain in unfavourable years due to restricted carbon supply (Weyand and Schultz, 2006; Spring *et al.*, 2012).

Leaf removal in the bunch area induces better light penetration through the canopy, thus increasing bunch exposure and promoting grape ripening. defoliation Early reduces methoxypyrazine accumulation in the grape (Ryona et al., 2008; Serra-Stepke, 2010). Correlations between natural bunch exposure variability and the development of aromas is generally weaker than in situations in which differences are induced through imposed treatments, such as leaf removal (Meyers et al., 2013). Kwasniewski et al. (2010) showed that the timing of leaf removal also had an impact on C₁₃-norisoprenoids in resulting wines. However, no constant relationship with grape N content could be highlighted across years and cultivars (Verdenal et al., 2019).

4.2. Fruit load regulation

Bunch thinning (*i.e.*, crop load limitation by removing a proportion of fruits early in the season) is a worldwide practice for enhancing fruit maturation. Several studies have reported the influence of fruit load on C partitioning (Chaves, 1984; Morinaga et al., 2003; Dai et al., 2011; Dayer et al., 2017), but it is still unclear how fruit load influences grape N accumulation and composition. Under high yield conditions, grape AAs originate in the leaves (Rossouw et al., 2017). Root N reserves also play a major role in balancing grape N content. Root N accumulation in reserves is restricted by the presence of fruit before and after veraison (Rodriguez-Lovelle and Gaudillère, 2002; Rossouw et al., 2017). In response to a higher fruit load, vines extract more C and N from reserves mainly located in the storage organs, to match the demand of the maturing fruits (Howell, 2001). Overproduction can potentially induce a significant reduction in reserves in the long term, which may Ν affect vigour, bud fruitfulness and even plant sustainability. As compensation, N uptake is generally higher under high-yielding conditions (Treeby and Wheatley, 2006). The modulation of both reserve N mobilisation and N uptake contributes to a relatively constant grape N concentration, despite a large crop load variation (Verdenal et al., 2020). Grape AA profile has been found to change despite unchanged overall concentration, with vield conditions affecting certain AAs more than others (Figure 18) (Verdenal et al., 2020). Several authors have confirmed changes in volatile compounds in response to bunch thinning (Rutan et al., 2018; Wang et al., 2019). Lin et al. (2018)



FIGURE 18. Impact of crop load on must AA composition. Principal component analysis (PCA) of must AA profiles (AA proportions in %) at harvest.

Black = high-yielding conditions (HYC, n = 12); grey = low-yielding conditions (LYC, n = 9); circles = control vines (n = 11); squares = N-fertilized vines (n = 10). The PCA discriminates the vines under HYC from those under LYC, independently of the fertilisation treatment. Chasselas, 2017, Pully, Switzerland (from Verdenal *et al.*, 2020).

observed differential expressions of AA decarboxylase in relation to fruit load; *i.e.*, the enzyme regulating the concentration of aroma-active 2-phenylethanol. Based on this result, they further recommended a yield range at harvest for the cultivar Vidal for optimum aroma expression.

4.3. Leaf-to-fruit ratio

Production is at a maximum when the supply of resources equals or exceeds plant demand (Lawlor, 2002). In fact, several studies have shown an inconsistent impact of bunch thinning on fruit composition, highlighting the prevailing role of the leaf-to-fruit ratio (Jackson and Lombard, 1993; Keller et al., 2005; Parker et al., 2014; Parker et al., 2015; Verdenal et al., 2016b; Mawdsley et al., 2018; Wang et al., 2018). Indeed, bunch thinning may not alter the leaf area-to-fruit weight ratio enough to overcome carbon supply limitations (Reeve et al., 2018). Howell (2001) wrote a detailed review on the growth-to-yield relationship for sustainable viticulture. Vine balance is usually understood in terms of the principles of vine C balance (Howell, 2001). It has been found that maintaining a sufficient leaf area-to-fruit weight ratio (above 1 m² of exposed leaf area per kg of fruit) promotes grape development and maturation by providing a non-limiting source of photosynthetic carbohydrates (Kliewer and Dokoozlian, 2005; Zufferey et al., 2015; Gutiérrez-Gamboa et al., 2019). Vine balance may also be expressed using the Ravaz index (*i.e.*, the fruit-to-pruning wood ratio) as the wood quantity is closely related to the leaf area (Howell, 2001). To summarise, under cool-climate conditions, a leaf-to-fruit ratio of 1.0 to $1.2 \text{ m}^2/\text{kg}$ is recommended to promote both grape maturity and must YAN accumulation, while the root N reserve is replenished, which guarantees sustainability (Murisier and Zufferey, 1997; Verdenal *et al.*, 2016a).

4.4. Root restriction

Root restriction is an efficient method for controlling nutrient uptake and plant vigour, as it impacts both root development and activity. Root development can be limited by either root-zone limitation, partial rootzone drying or root pruning. Yang et al. (2007) studied the impact of root restriction on nitrate uptake kinetics using two pot sizes (2 and 12 L); they observed that root-zone limitation efficiently inhibited shoot and root development, while decreasing the amount of net N uptake. Root-zone limitation has further consequences on ascorbic acid and carotenoid pathways, among others, in plant metabolism (Leng et al., 2017). Partial root-zone drying due to localised irrigation (50 % evapotranspiration) was found to limit both root development and canopy development, in comparison to both full irrigation (100 % evapotranspiration) and deficit irrigation (50 % evapotranspiration) (Santos et al., 2005). Root pruning is a common practice in fruit production for limiting vigour; this practice affects the size of the root N reserve. Root pruning performed on grapevine after bud burst was shown to reduce both pruning weight (-8 %), petiole N content (-11 %) and must YAN content (-13 %) (Giese *et al.*, 2015). However, the long-term impact of these practices on grapevine physiology is still unknown.

5. Vineyard inputs

5.1. Irrigation

Under limited water conditions, vine growers may irrigate their vineyards. Depending on the water constraint, quantity of water applied and timing of application, irrigation may influence soil N availability and plant N uptake, with further consequences on plant vigour and grape ripening (Keller, 2005; White et al., 2007; Iandolino and Williams, 2014; Ortega-Heras et al., 2014). Bouzas-Cid et al. (2018b) observed only minor variations in must AA concentration following irrigation treatments. However, their trial involved only a null to mild water restriction (average stem water-potential -0.63 MPa). The method of irrigation also influences N uptake. Drip versus furrow irrigation methods were compared in a trial (Williams, 2015). Plant N uptake was increased by only 12 % for furrow irrigation conditions, in comparison to 40 % for drip irrigation.

The amount of water the vine receives (from both rainfall and irrigation) and its temporal distribution affect the quality of red and white wines differently. For instance, deficit irrigation can be applied along with limited N supply to control vegetative development, yield and fruit composition (Keller, 2005). Zufferey et al. (2017, 2018) observed that the absence of water deficit negatively affects the quality of red wines (cv. Pinot noir), while it slightly enhances the quality of white wines (cv. Chasselas). Moreover, moderate water restriction is desirable when growing red grape (White et al., 2007). Pinot noir wines produced from vines under moderate water restriction had a higher concentration of sugars, polyphenols and anthocyanins; they were thus found to be full-bodied, and to have better mouthfeel and higher-quality tannins (Zufferey et al., 2017; Kotsaki et al., 2020b). Conversely, irrigated Chasselas wines (no water restriction) were mostly preferred for their better mouthfeel and lower bitterness (Zufferey et al., 2018). Moderate water restriction enhances grape maturation (Zufferey et al., 2017), while it can also simultaneously induce lower N content in the plant and in must. Accumulations of C and N in grapes follow different pathways: under water restriction, non-structural reserve carbohydrate are remobilised, contributing to berry sugar accumulation, while fruit N accumulation can be affected due to lower N availability

(Rossouw *et al.*, 2017; Zufferey *et al.*, 2018). However, it is not easy to separate the effect of water and N restriction in these trials.

5.2. Fertilisation

N fertilisation is an efficient practice for manipulating grape must composition, particularly in terms of pH, malic acid and potassium (Rühl et al., 1992). N fertilisation purposely enhances N availability for the plant and increases N uptake. However, net N uptake from an applied fertiliser is usually as low as 30-40 %, mainly due to surface run-off, leaching or gaseous emissions (Van Cleemput et al., 2008; Williams, 2015). Fertilisation efficiency largely depends on NUE (Porro et al., 2010). The limiting factors for maximising NUE are different at high and low N supply, and NUE is generally higher under low N conditions (Xu et al., 2012). The only consistent effect of vineyard N application on grape metabolites is an increase in total N compounds (Bell and Henschke, 2005). N fertilisation is usually applied to the soil surface between bud burst and flowering, which corresponds to the first period of high root N uptake. As a result, grapevine vegetative development and berry set are generally improved.

Excessive fertilisation is highly detrimental to both grape composition and grape sanitary status and to the environment. The negative impact on grape composition often manifests itself through an excessive increase in vigour. Many studies comparing different levels of N supply have demonstrated the negative consequences of excessive N supply on berry composition (Delas et al., 1991; Hilbert et al., 2003; Schreiner et al., 2014; Soubeyrand et al., 2014). In some cases, N supply was extremely high (i.e., above 100 kg/ha), in which cases, vine vigour was exacerbated, while bud fruitfulness and leaf area increased. Berry set was lower and bunch rot sensitivity increased (both negatively affecting yield in extreme cases). Fruit maturity was delayed; the must at harvest contained less sugar, had higher concentrations of organic acids and a higher pH. Furthermore, it was found that, while progressively reducing the quantity of N supply, vegetative growth will decrease prior to a reduction in fruit load, thus further impacting must YAN (Schreiner et al., 2014). It has been established that excessive N supply also induces lower anthocyanin and tannin content in red grapes, independently from phenylalanine content (Choné et al., 2001; Hilbert et al., 2003; Schreiner et al., 2018). Further investigation is necessary to understand all the mechanisms related to N content and involved in the synthesis of polyphenols. One limiting factor is the higher C quantity required for N assimilation, to the detriment of the flavonoid pathway (Dai et al., 2011; Soubeyrand *et al.*, 2018). Another negative factor related to flavonoid metabolism is the resulting excessive vigour of the canopy, which reduces fruit exposure to sunlight due to bunch shading (Stamatiadis *et al.*, 2007; Jackson, 2008). At a molecular level, genes involved in the flavonoid pathway (encoding phenylalanine ammonia-lyase, chalcone synthase, flavonoid30, 50hydroxylase, dihydroflavonol4reductase and leucoanthocyanidin dioxygenase) revealed a lower transcript level in berries under excessive N fertilisation (*i.e.*, 120 kg/ha of N), in comparison to a non-fertilised control treatment (Soubeyrand *et al.*, 2014).

Foliar fertilisation in viticulture has been implemented worldwide. A complete review has summarised the influence of foliar-fertiliser formulations and biostimulants (i.e., elicitors and resistance inducers) on grape composition (Gutiérrez-Gamboa et al., 2019). Amongst them, the application of urea at veraison is the most common, due to its low price and fast uptake by plants. Whether applied alone or with S (which facilitates urea uptake by the leaves), it efficiently increases the concentrations of NH4+, AAs, glycosides and glutathione in grapes (Lacroux et al., 2008; Hannam et al., 2016; Gutiérrez-Gamboa et al., 2017a). Portu et al. (2015) even found a positive impact on anthocyanin and flavanol content, in opposition to the usual impact of soil N fertilisation. The direct addition of AAs on the canopy (i.e., phenylalanine, proline and arginine) showed a lower efficiency (Garde-Cerdán et al., 2014; Gutiérrez-Gamboa et al., 2017a).

The localisation of fertilisation is also very important. N is usually applied to the soil before flowering. Soil fertilisation inevitably stimulates cover crop development, which consequently competes with the grapevine for access to water and nutrients (Maigre and Aerny, 2001a). The exclusive application of N under the row, instead of to the entire soil surface, significantly increases fertilisation efficiency. inducing lower competition and improved N uptake by the grapevine (Spring, 2003). In terms of foliar application, no differences have been found between applying urea exclusively to the top, bottom or entire canopy (Verdenal et al., 2017). However, the authors recommended spreading urea over the entire canopy to limit the amount of urea per leaf surface unit, and to avoid necrosis symptoms due to a temporary excess of NH_4^+ in the leaves.

The timing of fertilisation can significantly influence the quantity of N uptake and N partitioning in the plant. Conradie (2005) summarises the different periods for optimum fertilisation efficiency, highlighting the impacts of climate, soil and plant genetics. For instance, in warmer countries such as South Africa, the long post-harvest period (several months) is effective for N application, while in cooler countries, little N is absorbed during that period (few weeks only) (Conradie, 1992). The application to soil of 60 kg/ha of N at berry set in N deficient vines was found to increase vigour and grape YAN content, as well as cysteine-conjugated compounds and glutathione, but it decreased phenolic compounds (Choné et al., 2006). N supply was also found to increase grape aroma precursors; volatile thiols in wine were better preserved due to lower phenolic and higher glutathione levels (Choné et al., 2006). Grapes benefit more from a late foliar N application than an application at the flowering stage (Porro et al., 2010; Verdenal et al., 2015). Foliar fertilisation during the period of veraison (in the form of urea) has often been shown to be a reliable and efficient way of increasing YAN concentration must without affecting grapevine vigour in (Nisbet et al., 2014; Hannam et al., 2016; Alem et al., 2019; Gutiérrez-Gamboa et al., 2019). It is particularly recommended for promoting the development of aromas in white and rosé wines. The impact of late foliar urea supply also improves the sensory profile of red wine, inducing a lower astringency (Reynard et al., 2012; Verdenal et al., 2016c). Conversely, post-harvest N application has a negligible impact on grape YAN concentration in the following season (Holzapfel and Treeby, 2007).

Varying N applications according to vine N status across a vineyard block is an appropriate method of homogenising vine vigour, yield and grape composition. Vigour variations are generally related to vine N status and can be remotely determined using the NDVI. Using the NDVI, Gatti *et al.* (2018) applied three levels of fertilisation in their field trial depending on grapevine vigour and N status. Despite the fact that the NDVI is also related to other factors (*i.e.*, water availability and rootstock vigour), the homogeneity in terms of vigour was significantly increased within four years. This result should encourage further research on this important issue in vineyard management.

5.3. Other inputs

Copper (Cu) is widely used in viticulture, especially in organic production. It is the base component of the Bordeaux mixture used to control downy mildew. Copper formulations have been shown to affect grape AA concentration. Both the Bordeaux mixture and copper hydroxide decreased the content of AAs in grapes, compared to control samples (Garde-Cerdán*etal.*,2017).Oliva*etal.*(2011)studiedthe impact of several fungicides (famoxadone, fenhexamid, fluquinconazole, kresoxim-methyl, quinoxyfen and trifloxystrobin) on grape N composition. These

fungicides induced an overall lower N concentration with different quantitative and qualitative effects on grape AA composition, depending on the fungicide. It is not clear whether the impact of fungicide is due to a lower biosynthesis of AAs, or to a decrease in their precursors (Oliva et al., 2011). Gutiérrez-Gamboa et al. (2019) have reviewed several studies, which have experimented on the use of biostimulants on grapevine. While chitosan, laminarin and yeast extracts decreased must AA content, methyl jasmonate, abscisic acid, riboflavin and seaweed extracts had a positive impact on AA accumulation in grape (Ju et al., 2016; Garde-Cerdán et al., 2017; Gutiérrez-Gamboa et al., 2017b; González-Santamaría et al., 2018; Gutiérrez-Gamboa et al., 2020b). This list is not exhaustive, however.

To conclude, vineyard inputs greatly influence N availability for the plant, despite the risks of excessive supply and pollution of the environment. The variability of environmental conditions also play a major role in the efficiency of the input. An integrative view of the vineyard would be conducive to the sustainable optimisation of agronomic practices, in order to minimise the need for external inputs.

CONCLUSION AND PERSPECTIVES

This review emphasises the importance of N in viticulture and winemaking. Mineral N is assimilated into AAs, which are further involved in many metabolic pathways, from protein synthesis to the formation of grape aroma-active compounds. Grape AA content also influences the winemaking process, including both the fermentation kinetics and the development of wine flavours. Vineyard N status management should be based on the knowledge that N excess is as detrimental to wine quality as N depletion. Plant N demand is driven by vegetative development and N removal is related to crop load. While the amount of N exported from the vineyard is quite easy to establish, determining the soil mineral N availability is more complex, as it is influenced by environmental conditions. The influence of both the environment (i.e., climate and soil) and plant genetics creates a myriad of unique situations to which growers must adapt their practices, in order to produce grapes of suitable quantity and quality.

The complexity of the processes involved requires an integrative approach to managing grapevine N nutrition. When necessary, N fertilisation can be carried out on the ground between bud burst and flowering to improve vegetative development, while a foliar application can be realised at veraison stage to enhance grape YAN concentration for winemaking purposes. Taking environmental conditions into account, the grape grower can also adapt plant material, soil management

and vine balance to improve NUE and minimise N inputs in the vineyard. Grapevine N balance depends on canopy size, fruit load and annual replenishment of root N reserves. The major role of the roots in vine balance has been highlighted over the past decades, thanks to methods such as isotope labelling. The strong correlation between must YAN concentration and wine quality clearly shows a need for further research. Early assessment of grape N content during the season would help to justify late foliar N application in order to prevent grape YAN deficiency for winemaking. Recent research has shown that grape YAN content is a potential criterion for grape maturity and quality potential. It could also be a selective criterion for grapevine breeding. Further sustainable strategies for high-quality viticulture and wine production include improving plant material and fine-tuning agronomic practices to balance vine N status.

Acknowledgments: The authors would also like to thank Virginie Duquette for her artwork (www.gravir. ch). This research did not receive any specific funding.

REFERENCES

Aerny, J. (1996). Composés azotés des moûts et des vins. *Revue suisse de Viticulture, Arboriculture et Horticulture, 28*(3), 161-165.

Alcaide-Hidalgo, J. M., Moreno-Arribas, M. V., Martín-Álvarez, P. J., & Polo, M. C. (2007). Influence of malolactic fermentation, postfermentative treatments and ageing with lees on nitrogen compounds of red wines. *Food Chemistry*, *103*(2), 572-581. doi:10.1016/j.foodchem.2006.09.002

Alem, H., Rigou, P., Schneider, R., Ojeda, H., & Torregrosa, L. (2019). Impact of agronomic practices on grape aroma composition: a review. *Journal of the Science of Food and Agriculture, 99*(3), 975-985. doi:10.1002/jsfa.9327

Aranguren, M., Castellón, A., & Aizpurua, A. (2018). Topdressing nitrogen recommendation in wheat after applyingorganic manures: the use of field diagnostic tools. *Nutrient Cycling in Agroecosystems, 110*(1), 89-103. doi:10.1007/s10705-017-9865-7

Autralian Wine Research Institute, 2020. Yeast Assimilable Nitrogen. https://www.awri.com.au/ industry_support/winemaking_resources/wine_ fermentation/yan/ visited on the 24/08/2020.

Azuma, A., Yakushiji, H., Koshita, Y., & Kobayashi, S. (2012). Flavonoid biosynthesis-related genes in grape skin are differentially regulated by temperature and light conditions. *Planta*, 236(4), 1067-1080. doi:10.1007/s00425-012-1650-x

Barbosa, C., Falco, V., Mendes-Faia, A., & Mendes-Ferreira, A. (2009). Nitrogen addition influences formation of aroma compounds, volatile acidity and ethanol in nitrogen deficient media fermented by *Saccharomyces cerevisiae* wine strains. *Journal of Bioscience and Bioengineering*, *108*(2), 99-104. https://doi.org/10.1016/j.jbiosc.2009.02.017

Bates, T. R., Dunst, R. M., & Joy, P. (2002). Seasonal dry matter, starch, and nutrient distribution in 'Concord' grapevine roots. *HortSicence*, *37*(2), 313-316. doi:10.21273/hortsci.37.2.313

S.-J., & Henschke, (2005). Bell. Р. Α. Implications of nitrogen nutrition for grapes, fermentation and wine. Australian Journal of Grape and Wine Research, 11, 242-295. doi:10.1111/j.1755-0238.2005.tb00028.x

Bisson, L. F., & Butzke, C. E. (2000). Diagnosis and rectification of stuck and sluggish fermentations. *American Journal of Enology and Viticulture*, *51*(2), 168-177.

Boss, P. K., Bottcher, C., & Davies, C. (2014). Various Influences of Harvest Date and Fruit Sugar Content on Different Wine Flavor and Aroma Compounds. *American Journal of Enology and Viticulture, 65*(3), 341-353. doi:10.5344/ajev.2014.13137

Bouzas-Cid, Y., Trigo-Córdoba, E., Orriols, I., Falqué, E., & Mirás-Avalos, M. J. (2018a). Influence of soil management on the red grapevine (*Vitis vinifera* L.) Mencía must amino acid composition and wine volatile and sensory profiles in a humid region. *Beverages*, *4*, 76. doi:10.3390/beverages4040076

Bouzas-Cid, Y., Díaz-Losada, E., Trigo-Córdoba, E., Falqué, E., Orriols, I., Garde-Cerdán, T., & Mirás-Avalos, J. M. (2018b). Effects of irrigation over three years on the amino acid composition of Albariño (*Vitis vinifera* L) musts and wines in two different terroirs. *Scientia Horticulturae*, 227, 313-325. https://doi.org/10.1016/j.scienta.2017.05.005

Brunetto, G., Ceretta, C.A., de Melo, G. W. B., Girotto, E., Ferreira, P. A. A., Lourenzi, C. R., da Rosa Couto R., Tassinaria, A., Knevitz Hammerschmitt R., da Silva, L. O. S., Lazzaretti B. P., de Souza Kulmann M. S. & Carranca, C., (2016). Contribution of nitrogen from urea applied at different rates and times on grapevine nutrition. *Scientia Horticulturae*, 207, 1-6. doi:10.1016/j.scienta.2016.05.002

Bureau, S. M., Baumes, R. L., & Razungles, A. J. (2000). Effects of vine or bunch shading on the glycosylated flavor precursors in grapes of *Vitis vinifera* L. cv. Syrah. *Journal of Agricultural and Food Chemistry*, 48(4), 1290-1297. doi:10.1021/jf990507x

Cameleyre, M., Lytra, G., Tempere, S., & Barbe, J.-C. (2015). Olfactory impact of higher alcohols on red wine fruity ester aroma expression in model solution. *Journal of Agricultural and Food Chemistry*, 63(44), 9777-9788. doi:10.1021/acs.jafc.5b03489

Celette, F., Findeling, A., & Gary, C. (2009). Competition for nitrogen in an unfertilized intercropping system: The case of an association of grapevine and grass cover in a Mediterranean climate. *European Journal of Agronomy*, *30*(1), 41-51. doi:10.1016/j.eja.2008.07.003

Celette, F., & Gary, C. (2013). Dynamics of water and nitrogen stress along the grapevine cycle as affected by cover cropping. *European Journal of Agronomy*, *45*, 142-152. https://doi.org/10.1016/j.eja.2012.10.001

Cerovic, Z. G., Ghozlen, N. B., Milhade, C., Obert, M., Debuisson, S., & Moigne, M. L. (2015). Nondestructive diagnostic test for nitrogen nutrition of grapevine (*Vitis vinifera* L.) based on dualex leaf-clip measurements in the field. *Journal of Agricultural and Food Chemistry*, *63*(14), 3669-3680. doi:10.1021/acs.jafc.5b00304

Chaves, M. M. (1984). Photosynthesis and assimilate partition in fruiting and non-fruiting grapevine shoots. In C. Sybesma (Ed.), Advances in photosynthesis research: proceedings of the VIth international congress on photosynthesis, Brussels, Belgium, August 1–6, 1983. Volume IV (pp. 145-148). Dordrecht: Springer Netherlands. https://doi.org/10.1007/978-94-017-4971-8_32

Choné, X., van Leeuwen, C., Chéry, P., & Ribéreau-Gayon, P. (2001). Terroir influence on water status and nitrogen status of non-irrigated Cabernet sauvignon (*Vitis vinifera* L.). Vegetative development, must and wine compositions (example of a Medoc top estate vineyard, Saint-Julien area, Bordeaux, 1997). *South African Journal of Enology and Viticulture*, 22(1), 8-15. doi:10.21548/22-1-2159

Choné, X., Lavigne-Cruège, V., Tominaga, T., van Leeuwen, C., Castagnède, C., Saucier, C., & Dubourdieu, D. (2006). Effect of vine nitrogen status on grape aromatic potential: falvor precursors (S-cysteine conjugates), glutathione and phenolic content in *Vitis vinifera* L. cv. Sauvignon blanc grape juice. *Journal International des Sciences de la Vigne et du Vin,* 40(1), 1-6. doi:10.20870/oeno-one.2006.40.1.880

Clarke, S. J., Lamont, K. J., Pan, H. Y., Barry, L. A., Hall, A., & Rogiers, S. Y. (2015). Spring root-zone temperature regulates root growth, nutrient uptake and shoot growth dynamics in grapevines. *Australian Journal of Grape and Wine Research*, *21*(3), 479-489. doi:10.1111/ajgw.12160

Cliquet, J.-B., Deléens, E., Bousser, A., Martin, M., Lescure, J.-C., Prioul, J.-L., Mariotti, A., & Morot-Gaudry, J.-F., (1990). Estimation of carbon and nitrogen allocation during stalk elongation by ¹³C and ¹⁵N tracing in Zea mays L. *Plant Physiology*, *92*(1), 79-87. doi:10.1104/pp.92.1.79

Clingeleffer, P. R. (2009). Influence of canopy management systems on vine productivity and fruit composition. In: *International symposium on grapevine canopy management*, California (Ed. University of California Davis), 13-20.

Conradie, W. J. (1980). Seasonal uptake of nutrients by Chenin blanc in sand culture. *South African Journal of Enology and Viticulture, 1*(1), 59-65. doi:10.21548/1-1-2414

Conradie, W. J. (1983). The uptake and distribution of ¹⁵N enriched nitrate by three rootstock cultivars grafted to Chenin blanc. *South African Journal of Enology and Viticulture*, 4(2), 45-47. https://doi.org/10.21548/4-2-2369 Conradie, W. J. (1991). Distribution and translocation of nitrogen absorbed during early summer by two-year-old grapevines grown in sand culture. *American Journal of Enology and Viticulture*, 42(3), 180-190.

Conradie, W. J. (1992). Partitioning of nitrogen in grapevines during autumn and the utilisation of nitrogen reserves during the following growing season. *South African Journal of Enology and Viticulture, 13*(1), 45-51. https://doi.org/10.21548/13-1-2198

Conradie, W. J. (2005). *Partitioning of Mineral Nutrients and Timing of Fertilizer Applications for Optimum Efficiency.* Paper presented at the Soil Environment and Vine Mineral Nutrition, Davis.

Cookson, S. J., Clemente Moreno, M. J., Hevin, C., Nyamba Mendome, L. Z., Delrot, S., Trossat-Magnin, C., & Ollat, N. (2013). Graft union formation in grapevine induces transcriptional changes related to cell wall modification, wounding, hormone signalling, and secondary metabolism. *Journal of Experimental Botany*, 64(10), 2997-3008. doi:10.1093/jxb/ert144

Coplen, T. B. (2011). Guidelines and recommended terms for expression of stable-isotope-ratio and gas-ratio measurement results. *Rapid Communications in Mass Spectrometry*, 25(17), 2538-2560. doi:10.1002/rcm.5129

Crépin, L., Truong, N. M., Bloem, A., Sanchez, I., Dequin, S., & Camarasa, C. (2017). Management of Multiple Nitrogen Sources during Wine Fermentation by *Saccharomyces cerevisiae*. *Applied and Environmental Microbiology*, *83*(5), e02617-02616. doi:10.1128/aem.02617-16

Crespy, A. (2007). *Manuel pratique de fertilisation, qualité des moûts et des vins* (Oenoplurimedia Ed.).

Czerny, M., Christlbauer, M., Christlbauer, M., Fischer, A., Granvogl, M., Hammer, M., Hartl, C., Moran Hernandez, N., & Schieberle, P. (2008). Re-investigation on odour thresholds of key food aroma compounds and development of an aroma language based on odour qualities of defined aqueous odorant solutions. *European Food Research and Technology*, 228(2), 265-273. doi:10.1007/s00217-008-0931-x

Dai, Z. W., Ollat, N., Gomès, E., Decroocq, S., Tandonnet, J.-P., Bordenave, L., Pieri, P., Hilbert, G., Kappel, C., van Leeuwen C., Vivin, P. & Delrot, S., (2011). Ecophysiological, genetic, and molecular causes of variation in grape berry weight and composition: a review. *American Journal of Enology and Viticulture*, *62*(4), 413-425. doi:10.5344/ajev.2011.10116

Dal Santo, S., Fasoli, M., Negri, S., D'Incà, E., Vicenzi, N., Guzzo, F., Tornielli, G. B., Pezzotti, M., & Zenoni, S., (2016). Plasticity of the berry ripening program in a white grape variety. *Frontiers in Plant Science*, *7*(970). doi:10.3389/fpls.2016.00970

Dal Santo, S., Zenoni, S., Sandri, M., De Lorenzis, G., Magris, G., De Paoli, E., Di Gaspero, G., Del Fabbro, C., Morgante, M., Brancadoro, L., Grossi D., Fasoli, M., Zuccolotto P., Tornielli G. B., & Pezzotti, M. (2018). Grapevine field experiments reveal the contribution of genotype, the influence of environment and the effect of their interaction (G×E) on the berry transcriptome. *The Plant Journal*, 93(6), 1143-1159. doi:10.1111/tpj.13834

Dayer, S., Peña, J. P., Gindro, K., Torregrosa, L., Voinesco, F., Martínez, L., Pietro, J. A. & Zufferey, V. (2017). Changes in leaf stomatal and conductance, petiole hydraulics vessel morphology in grapevine (Vitis vinifera cv. Chasselas) under different light and irrigation regimes. Functional Plant Biology, 44. 679-693. doi:10.1071/FP16041

Delabays, N., Spring, J. L., Ançay, A., Mosimann, E., & Schmid, A. (2000). Sélection d'espèces pour l'enherbement des cultures spéciales. *Revue suisse de Viticulture, Arboriculture et Horticulture, 32*(2), 95-104

de-la-Fuente-Blanco, A., Sáenz-Navajas, M.-P., & Ferreira, V. (2016). On the effects of higher alcohols on red wine aroma. *Food Chemistry*, *210*, 107-114. https://doi.org/10.1016/j.foodchem.2016.04.021

Delas, J., Molot, C. & Soyer J.P. (1991). Effects of nitrogen fertilization and grafting on the yield and quality of the crop of *Vitis vinifera* cv. Merlot. International Symposium, Jun 1991, Seattle, United States.

Delas, J. (2010). *Fertilisation de la vigne, 2e edition*. Ed Féret, Bordeaux, 165 p.

Deléens, E., Morot-Gaudry, J.-F., Martin, F., Thoereux, A., & Gojon, A. (1997). Méthodologie ¹⁵N. In *Assimilation de l'azote chez les plantes* (pp. 265-280). Paris: Inra.

De Royer Dupré, N., Schneider, R., Payan, J. C., Salançon, E., & Razungles, A. (2014). Effects of vine water status on dimethyl sulfur potential, ammonium, and amino acid contents in Grenache noir grapes (*Vitis vinifera*). J Agric Food Chem, 62(13), 2760-2766. doi:10.1021/jf404758g

Dienes-Nagy, Á., Marti, G., Breant, L., Lorenzini, F., Fuchsmann, P., Baumgartner, D., Zufferey, V., Spring, J.-L., Gindro, K., Viret, O., Wolfender, J.-L. & Rösti, J. (2020). Identification of putative chemical markers in white wine (Chasselas) related to nitrogen deficiencies in vineyards. *Oeno One*, *54*(3), 583-599. doi:10.20870/oeno-one.2020.54.3.3285

Dunlevy, J. D., Kalua, C. M., Keyzers, R. A., & Boss, P. K. (2009). The Production of Flavour & Aroma Compounds in Grape Berries. *In* K. A. Roubelakis-Angelakis (Ed.), *Grapevine Molecular Physiology & Biotechnology* (pp. 293-340). Dordrecht: Springer Netherlands. https://doi.org/10.1007/978-90-481-2305-6_11

Durante, C., & Bertacchini, L., Bontempo, L., Camin, F., Manzini, D., Lambertini, P., Marchetti, A., & Paolini, M. (2016). From soil to grape and wine: Variation of light and heavy elements isotope ratios. *Food Chemistry*, *210*, 648-659. doi:10.1016/j.foodchem.2016.04.108 Echeverría, G., Ferrer, M., & Mirás-Avalos, J. (2017). Quantifying the relative impact of physical and human factors on the viticultural expression of terroir. *International Journal of Environmental & Agriculture Research*, 3(4), 12-25.

Eichert, T. (2013). *Foliar Nutrient Uptake - of Myths and Legends*. Paper presented at the VIIth ISon Mineral Nutrition of Fruit Crops, Bonn, Germany. https://doi.org/10.17660/ActaHortic.2013.984.5

Fairbairn, S., McKinnon, A., Musarurwa, H. T., Ferreira, A. C., & Bauer, F. F. (2017). The impact of single amino acids on growth and volatile aroma production by *Saccharomyces cerevisiae* Strains. *Frontiers in Microbiology, 8*(2554). doi:10.3389/fmicb.2017.02554

Fayolle, E., Follain, S., Marchal, P., Chéry, P., & Colin, F. (2019). Identification of environmental factors controlling wine quality: A case study in Saint-Emilion Grand Cru appellation, France. *Science of The Total Environment, 694*, 133718. https://doi.org/10.1016/j.scitotenv.2019.133718

Fernández, V., & Eichert, T. (2009). Uptake of hydrophilic solutes through plant leaves: current state of knowledge and perspectives of foliar fertilization. *Critical Reviews in Plant Sciences, 28*(1-2), 36-68. doi:10.1080/07352680902743069

Fernández, V., & Brown, P. H. (2013). From plant surface to plant metabolism: the uncertain fate of foliarapplied nutrients. *Frontiers in Plant Science*, *4*(289). doi:10.3389/fpls.2013.00289

Fleet, G. H. (1993). *Wine microbiology and biotechnology*. London: Taylor and Francis.

Friedel, M., Hendgen, M., Stoll, M., & Löhnertz, O. (2020). Performance of reflectance indices and of a handheld device for estimating in-field the nitrogen status of grapevine leaves. *Australian Journal of Grape and Wine Research*, *26*(2), 110-120. doi:10.1111/ajgw.12424

Gang, Q., Chang, S. X., Lin, G., Zhao, Q., Mao, B., & Zeng, D.-H. (2019). Exogenous and endogenous nitrogen differentially affect the decomposition of fine roots of different diameter classes of Mongolian pine in semiarid northeast China. *Plant and Soil, 436*(1), 109-122. doi:10.1007/s11104-018-03910-0

Gaudillère, J.-P., Chantelot, E., Soyer, J.-P., Molot, C., & Milin, S. (2003). Leaf and must nitrogen content two complementary indicators of grapevine nitrogen status. *Journal International des Sciences de la Vigne et du Vin*, *37*(2), 91-101. doi:10.20870/oeno-one.2003.37.2.943

Garde-Cerdán, T., & Ancín-Azpilicueta, C. (2008). Effect of the addition of different quantities of amino acids to nitrogen-deficient must on the formation of esters, alcohols, and acids during wine alcoholic fermentation. *LWT - Food Science and Technology*, 41(3), 501-510. doi:10.1016/j.lwt.2007.03.018

Garde-Cerdán, T., Lorenzo, C., Lara, J. F., Pardo, F., Ancin-Azpilicueta, C., & Salinas, M. R. (2009). Study of the evolution of nitrogen compounds during grape ripening. application to differentiate grape varieties and cultivated systems. *Journal of Agricultural and Food Chemistry*, *57*, 2410-2419. doi:10.1021/jf8037049

Garde-Cerdán, Т., Lopez, R., Portu, J., L., Lopez-Alfaro, I., Gonzalez-Arenzana, & Santamaria, P. (2014). Study of the effects of proline, phenylalanine, and urea foliar application to Tempranillo vineyards on grape amino acid content. Comparison with commercial nitrogen fertilisers. Food Chemistry. 163. 136-141. doi:10.1016/j.foodchem.2014.04.101

Garde-Cerdán, T., Mancini, V., Carrasco-Quiroz, M., Servili, A., Gutiérrez-Gamboa, G., Foglia, R., Pérez-Álvarez, E. P., & Romanazzi, G. (2017). Chitosan and laminarin as alternatives to copper for plasmopara viticola control: effect on grape amino acid. *Journal of Agricultural and Food Chemistry*, *65*(34), 7379-7386. doi:10.1021/acs.jafc.7b02352

Garde-Cerdán, T., Gutiérrez-Gamboa, G., Fernández-Novales, J., Pérez-Álvarez, E. P., & Diago, M. P. (2018). Towards the definition of optimal grape harvest time in Grenache grapevines: Nitrogenous maturity. *Scientia Horticulturae*, 239, 9-16. doi:10.1016/j.scienta.2018.05.014

Gatti, M., Squeri, C., Garavani, A., Vercesi, A., Dosso, P., Diti, I., & Poni, S. (2018). Effects of variable rate nitrogen application on cv. Barbera performance: vegetative growth and leaf nutritional status. *American Journal of Enology and Viticulture, 69*(3), 196-209. doi:10.5344/ajev.2018.17084

Gauthier, P. P. G., Bligny, R., Gout, E., Mahé, A., Nogués, S., Hodges, M., & Tcherkez, G. G. B. (2010). In folio isotopic tracing demonstrates that nitrogen assimilation into glutamate is mostly independent from current CO_2 assimilation in illuminated leaves of Brassica napus. *New Phytologist, 185*(4), 988-999. doi:10.1111/j.1469-8137.2009.03130.x

Giese, W. G., Wolf, T. K., Velasco-Cruz, C., & Roberts, L. (2015). Cover crop and root pruning effects on the rooting pattern of SO4 rootstock grafted to Cabernet-Sauvignon. *American Journal of Enology and Viticulture*. doi:10.5344/ajev.2015.15066

Glad, C., Farineau, J., Regnard, J.-L., & Morot-Gaudry, J.-F. (1994). The relative contribution of nitrogen originating from two seasonal 15N supplies to the total nitrogen pool present in the bleeding sap and in whole *Vinifera* cv. Pinot noir grapevines at bloom time. *American Journal of Enology and Viticulture*, 45(3), 327-332.

Gobert, A., Tourdot-Maréchal, R., Sparrow, C., Morge, C., & Alexandre, H. (2019). Influence of nitrogen status in wine alcoholic fermentation. *Food Microbiology*, *83*, 71-85. https://doi.org/10.1016/j.fm.2019.04.008

González-Barreiro, C., Rial-Otero, R., Cancho-Grande, B., & Simal-Gándara, J. (2015). Wine aroma compounds in grapes: a critical review. *Critical Reviews in Food Science and Nutrition*, 55(2), 202-218. doi:10.1080/10408398.2011.650336

González-Santamaría, R., Ruiz-González, R., Nonell, S., Garde-Cerdán, T., & Pérez-Álvarez, E. P. (2018). Influence of foliar riboflavin applications to vineyard on grape amino acid content. *Food Chemistry, 240*, 601-606. doi:10.1016/j.foodchem.2017.07.115

Gourieroux, A. M., Holzapfel, B. P., Scollary, G. R., McCully, M. E., Canny, M. J., & Rogiers, S. Y. (2016). The amino acid distribution in rachis xylem sap and phloem exudate of *Vitis vinifera* 'Cabernet-Sauvignon' bunches. *Plant Physiology and Biochemistry*, *105*, 45-54. doi:10.1016/j.plaphy.2016.04.010

Gouthu, S., Skinkis, P. A., Morre, J., Maier, C. S., & Deluc, L. G. (2012). Berry nitrogen status altered by cover cropping: Effects on berry hormone dynamics, growth and amino acid composition of Pinot noir. *Food Chemistry*, *135*(1), 1-8. doi:10.1016/j.foodchem.2012.04.019

Grechi, I., Vivin, P., Hilbert, G., Milin, S., Robert, T., & Gaudillère, J. P. (2007). Effect of light and nitrogen supply on internal C:N balance and control of root-to-shoot biomass allocation in grapevine. *Environmental and experimental Botany, 59*, 139-149. doi:10.1016/j. envexpbot.2005.11.002

Guilpart, N., Metay, A., & Gary, C. (2014). Grapevine bud fertility and number of berries per bunch are determined by water and nitrogen stress around flowering in the previous year. *European Journal of Agronomy*, 54, 9-20. doi:10.1016/j.eja.2013.11.002

Gutiérrez-Gamboa, G., Garde-Cerdán, T., Portu, J., Moreno-Simunovic, Y., & Martínez-Gil, A. M. (2017a). Foliar nitrogen application in Cabernet-Sauvignon vines: Effects on wine flavonoid and amino acid content. *Food Research International*, *96*, 46-53. doi:10.1016/j.foodres.2017.03.025

Gutiérrez-Gamboa, G., Portu, J., Santamaría, P., López, R., & Garde-Cerdán, T. (2017b). Effects on grape amino acid concentration through foliar application of three different elicitors. *Food Research International*, 99, 688-692. doi:10.1016/j.foodres.2017.06.022

Gutiérrez-Gamboa, G., Carrasco-Quiroz, M., Martínez-Gil, A. M., Pérez-Álvarez, E. P., Garde-Cerdán, T., & Moreno-Simunovic, Y. (2018). Grape and wine amino acid composition from Carignan noir grapevines growing under rainfed conditions in the Maule Valley, Chile: Effects of location and rootstock. *Food Research International*, *105*, 344-352. doi:10.1016/j.foodres.2017.11.021

Gutiérrez-Gamboa, G., Romanazzi, G., Garde-Cerdán, T., & Pérez-Álvarez, E. P. (2019). A review of the use of biostimulants in the vineyard for improved grape and wine quality: effects on prevention of grapevine diseases. *J Sci Food Agric, 99*(3), 1001-1009. doi:10.1002/jsfa.9353

Gutiérrez-Gamboa, G., Garde-Cerdán, T., Rubio-Bretón, P., & Pérez-Álvarez, E. P. (2020a). Seaweed foliar applications at two dosages to Tempranillo blanco (*Vitis vinifera* L.) grapevines in two seasons: Effects on grape and wine volatile composition. *Food Research International, 130*, 108918. doi:10.1016/j.foodres.2019.108918

Gutiérrez-Gamboa, G., Garde-Cerdán, T., Rubio-Bretón, P., & Pérez-Álvarez, E. P. (2020b). Biostimulation to Tempranillo grapevines (*Vitis vinifera* L.) through a brown seaweed during two seasons: Effects on grape juice and wine nitrogen compounds. *Scientia Horticulturae*, 264, 109177. doi:10.1016/j.scienta.2020.109177

Habran, A., Commisso, M., Helwi, P., Hilbert, G., Negri, S., Ollat, N., Gomès, E., van Leeuwen, C., Guzzo, F., & Delrot, S. (2016). Roostocks/scion/ nitrogen interactions affect secondary metabolism in the grape berry. *Frontiers in Plant Science*, *7*(1134). doi:10.3389/fpls.2016.01134

Hachiya, T., & Sakakibara, H. (2016). Interactions between nitrate and ammonium in their uptake, allocation, assimilation, and signaling in plants. *Journal of Experimental Botany*, *68*(10), 2501-2512. doi:10.1093/jxb/erw449

Hannam, K. D., Neilsen, G. H., Neilsen, D., Midwood, A. J., Millard, P., Zhang, Z., Thornton, B., & Steinke, D. (2016). Amino acid composition of grape (*Vitis vinifera* L.) juice in response to applications of urea to the soil or foliage. *American Journal of Enology and Viticulture*, *67*(1), 47-55. doi:10.5344/ajev.2015.15015

Hardarson, G., Hood-Nowotny, R., Jensen, E. S., Peoples, M. B., Van Cleemput, O., van Kessel, C., ... Zapata, F. (2008). Guidelines on nitrogen management in agricultural systems. *International Atomic Energy Agency, Vienna, IAEA-TCS-29*.

Haynes, R. J. (1986). Uptake and assimilation of mineral nitrogen by plants. In R. J. Haynes (Ed.), *Mineral nitrogen in the plant-soil system* (pp. 303-378). Orlando, Florida, USA: Academic Press. https://doi.org/10.1016/B978-0-12-334910-1.50010-8

Helwi, P., Habran, A., Guillaumie, S., Thibon, C., Hilbert, G., Gomès, E., Delrot, S., Darriet, P. & van Leeuwen, C. (2015). Vine nitrogen status does not have a direct impact on 2-methoxy-3-isobutylpyrazine in grape berries and wines. *J Agric Food Chem*, 63(44), 9789-9802. doi:10.1021/acs.jafc.5b03838

Helwi, P., Guillaumie, S., Thibon, C., Keime, C., Habran, A., Hilbert, G., Gomès, E., Darriet, P., Delrot, S., & van Leeuwen, C. (2016). Vine nitrogen status and volatile thiols and their precursors from plot to transcriptome level. *BMC Plant Biology*, *16*(1), 173. doi:10.1186/s12870-016-0836-y Henschke, P. A., & Jiranek, V. (1993). Yeasts – metabolism of nitrogen compounds. *In* G. H. Fleet (Ed.), *Wine microbiology and biotechnology* (pp. 77–164). Chur, Switzerland: Harwood Academic Publishers.

Hernández-Orte, P., Guitart, A., & Cacho, J. (1999). Changes in the concentration of amino acids during the ripening of *Vitis vinifera* Tempranillo variety from the *Denomination d'Origine* Somontano (Spain). *American Journal of Enology and Viticulture, 50*(2), 144-154.

Hernández-Orte, P., Cacho, J. F., & Ferreira, V. (2002). Relationship between varietal amino acid profile of grapes and wine aromatic composition. Experiments with model solutions and chemometric Study. *Journal of Agricultural and Food Chemistry*, *50*(10), 2891-2899. doi:10.1021/jf0113950

Hernández-Orte, P., Ibarz, M. J., Cacho, J., & Ferreira, V. (2006). Addition of amino acids to grape juice of the Merlot variety: Effect on amino acid uptake and aroma generation during alcoholic fermentation. *Food Chemistry*, *98*(2), 300-310. https://doi.org/10.1016/j.foodchem.2005.05.073

Hjelmeland, A. K., & Ebeler, S. E. (2015). Glycosidically bound volatile aroma compounds in grapes and wine: A Review. *American Journal of Enology and Viticulture, 66*(1), 1-11. doi:10.5344/ajev.2014.14104

Hilbert, G., Soyer, J.-P., Giraudon, J., Milin, S., & Gaudillère, J. P. (2003). Effects of nitrogen supply on must quality and anthocyanin accumulation in berries of cv. Merlot. *Vitis*, *42*(2), 69-76.

Hoenicke, K., Simat, T. J., Steinhart, H., Christoph, N., Geßner, M., & Köhler, H.-J. (2002). 'Untypical aging off-flavor' in wine: formation of 2-aminoacetophenone and evaluation of its influencing factors. *Analytica chimica Acta*, 458(1), 29-37. http://dx.doi.org/10.1016/S0003-2670(01)01523-9

Holzapfel, B. P., & Treeby, M. T. (2007). Effects of timing and rate of N supply on leaf nitrogen status, grape yield and juice composition from Shiraz grapevines grafted to one of three different rootstocks. *Australian Journal of Grape and Wine Research*, *13*(1), 14-22. doi:10.1111/j.1755-0238.2007.tb00067.x

Holzapfel, B. P., Smith, J. P., & Field, S. K. (2019). Seasonal vine nutrient dynamics and distribution of Shiraz grapevines. *Oeno One*, *53*(2). doi:10.20870/oeno-one.2019.53.2.2425

Howell, G. S. (2001). Sustainable grape productivity and the growth-yield relationship: a review. *American Journal of Enology and Viticulture*, *52*(3), 165-174.

Huang, Z., & Ough, C. S. (1991). Amino acid profiles of commercial grape juices and wines. *American Journal of Enology and Viticulture*, *42*(3), 261-267.

Huglin, P., and Schneider, C., 1998. *Biologie et écologie de la vigne*. Lavoisier Tec. et Doc., Paris.

Hunter, J. J., & Bonnardot, V. (2011). Suitability of some climatic parameters for grapevine cultivation in South Africa, with focus on key physiological processes. *South African Journal of Enology and Viticulture*, *32*(1), 18. doi:10.21548/32-1-1374

Iandolino, A. B., & Williams, L. E. (2014). Recovery of 15N labeled fertilizer by *Vitis vinifera* L. cv. Cabernet Sauvignon: effects of N fertilizer rates and applied water amounts. *American Journal of Enology and Viticulture*, 65(2), 189-196. doi:10.5344/ajev.2014.11019

Jackson, D. I., & Lombard, P. B. (1993). Environmental and management practices affecting grape composition and wine quality - a review. *American Journal of Enology and Viticulture, 44*(4), 409-430.

Jackson, R. S. (2008). *Wine Science* (3rd édition ed.). San Diego: Academic Press.

Janzen, H. H., & Bruinsma, Y. (1989). Methodology for the quantification of root and rhizosphere nitrogen dynamics by exposure of shoots to 15N-labelled ammonia.*SoilBiologyandBiochemistry*, *21*(2), 189-196. https://doi.org/10.1016/0038-0717(89)90094-1

Jing, H., Zhang, P., Li, J., Yao, X., Liu, G., & Wang, G. (2019). Effect of nitrogen addition on the decomposition and release of compounds from fine roots with different diameters: the importance of initial substrate chemistry. *Plant and Soil, 438*(1), 281-296. doi:10.1007/s11104-019-04017-w

Ju, Y.-L., Liu, M., Zhao, H., Meng, J.-F., & Fang, Y.-L. (2016). Effect of exogenous abscisic acid and methyl jasmonate on anthocyanin composition, fatty acids, and volatile compounds of Cabernet-Sauvignon (*Vitis vinifera* L.) grape berries. *Molecules*, *21*, 1354. doi:10.3390/molecules21101354

Kant, S., Bi, Y.-M., & Rothstein, S. J. (2011). Understanding plant response to nitrogen limitation for the improvement of crop nitrogen use efficiency. *Journal of Experimental Botany*, 62(4), 1499-1509. doi:10.1093/jxb/erq297

Keller, M., & Koblet, W. (1995). Dry matter and leaf area partitioning, bud fertility and second season growth of *Vitis vinifera* L.: responses to nitrogen supply and limiting irradiance. *Vitis*, *34*(2), 77-83.

Keller, M. (2005). Deficit irrigation and vine mineral nutrition. *American Journal of Enology and Viticulture*, *56*(3), 267-283.

Keller, M., Mills, L. J., Wample, R. L., & Spayd, S. E. (2005). Cluster thinning effects on three deficit-irrigated *Vitis vinifera* cultivars. *American Journal of Enology and Viticulture*, *56*(2), 91-103.

Keller, M. (2015). *The science of grapevines* (M. Keller Ed. 2nd edition): Elsevier Inc.

Kliewer, W. M., & Dokoozlian, N. (2005). Leaf area/crop weight ratios of grapevines: influence on fruit composition and wine quality. *American Journal of Enology and Viticulture, 56*(2), 170-181. Klodd, A. E., Eissenstat, D. M., Wolf, T. K., & Centinari, M. (2016). Coping with cover crop competition in mature grapevines. *Plant and Soil*, *400*(1), 391-402. doi:10.1007/s11104-015-2748-2

Kotsaki, E., Reynolds, A. G., Brown, R., Lee, H.-S., & Aubie, E. (2020a). Proximal sensing and relationships to soil and vine water status, yield, and berry composition in Ontario vineyards. *American Journal of Enology and Viticulture*, 71(2), 114-131. doi:10.5344/ajev.2019.19018

Kotsaki, E., Reynolds, A. G., Brown, R., Lee, H.-S., & Jollineau, M. (2020b). Spatial variability in soil and vine water status in Ontario vineyards: relationships to yield and berry composition. *American Journal of Enology and Viticulture*, *71*(2), 132-148. doi:10.5344/ajev.2019.19019

Krishna, H., Singh, S. K., Sharma, R. R., Khawale, R. N., Grover, M., & Patel, V. B. (2005). Biochemical changes in micropropagated grape (*Vitis vinifera* L.) plantlets due to arbuscular-mycorrhizal fungi (AMF) inoculation during ex vitro acclimatization. *Scientia Horticulturae*, *106*(4), 554-567. doi:10.1016/j.scienta.2005.05.009

Krogmeier, M. J., McCarty, G. W., & Bremner, J. M. (1989). Phytotoxicity of foliar-applied urea. *Proceedings of the National Academy of Sciences*, *86*(21), 8189-8191. doi:10.1073/pnas.86.21.8189

Kumagai, E., Araki, T., Hamaoka, N., & Ueno, O. (2011). Ammonia emission from rice leaves in relation to photorespiration and genotypic differences in glutamine synthetase activity. *Annals of Botany*, *108*(7), 1381-1386. doi:10.1093/aob/mcr245

Kwasniewski, M. T., Vanden Heuvel, J. E., Pan, B. S., & Sacks, G. L. (2010). Timing of cluster light environment manipulation during grape development affects C13 norisoprenoid and carotenoid concentrations in Riesling. *Journal of Agricultural and Food Chemistry*, 58(11), 6841-6849. doi:10.1021/jf904555p

Lacroux, F., Tregoat, O., van Leeuwen, C., Pons, A., Tominaga, T., Lavigne-Cruège, V., & Dubourdieu, D. (2008). Effect of foliar nitrogen and sulfur application on aromatic expression of *Vitis vinifera* L. cv. Sauvignon blanc. *Journal International des Sciences de la Vigne et du Vin, 42*(3), 125-132. doi:10.20870/oeno-one.2008.42.3.816

Lakso, A. and Sacks, G. (2009). Vine balance: what is it and how does it change over the season? In: *International symposium on grapevine canopy management, California* (Ed. University of California Davis), 21-25.

Lambrechts, M., & Pretorius, I. (2000). Yeast and its importance to wine aroma-a review. *South African Journal of Enology and Viticulture, 21*(1), 97-129. doi:10.21548/21-1-3560

Lasa, B., Menendez, S., Sagastizabal, K., Calleja-Cervantes, M., Irigoyen, I., Muro, J., Aparicio-Tejo, P., & Ariz, I. (2012). Foliar application of urea to "Sauvignon blanc" and "Merlot" vines: doses and time of application. *Plant Growth Regulation*, 67(1), 73-81. doi:10.1007/s10725-012-9667-5

Lawlor, D. W. (2002). Carbon and nitrogen assimilation in relation to yield: mechanisms are the key to understanding production systems. *Journal of Experimental Botany*, *53*(370), 773-787. doi:10.1093/jexbot/53.370.773

Lea, P. J., & Azevedo, R. A. (2006). Nitrogen use efficiency. 1. Uptake of nitrogen from the soil. *Annals of Applied Biology*, *149*(3), 243-247. doi:10.1111/j.1744-7348.2006.00101.x

Leng, F., Tang, D., Lin, Q., Cao, J., Wu, D., Wang, S., & Sun, C. (2017). Transcriptomic analyses of ascorbic acid and carotenoid metabolites influenced by root restriction during grape berry development and ripening. *Journal of Agricultural and Food Chemistry*, *65*(9), 2008-2016. doi:10.1021/acs.jafc.6b05322

Li, Q., Li, B.-H., Kronzucker, H. J., & Shi, W.-M. (2010). Root growth inhibition by NH_4^+ in Arabidopsis is mediated by the root tip and is linked to NH_4^+ efflux and GMPase activity. *Plant, Cell & Environment, 33*(9), 1529-1542. doi:10.1111/j.1365-3040.2010.02162.x

Lilly, M., Bauer, F. F., Styger, G., Lambrechts, M. G., & Pretorius, I. S. (2006). The effect of increased branchedchain amino acid transaminase activity in yeast on the production of higher alcohols and on the flavour profiles of wine and distillates. *FEMS Yeast Research*, *6*(5), 726-743. doi:10.1111/j.1567-1364.2006.00057.x

Lin, Y., Jiang, C., Guo, Y., Li, K., Liu, Z., Lin, Z., Li, X., Yue, G., Fu, Q., Li, W., Zheng, J., Dou, Y., & Guo, X. (2018). Relationship between 2-phenylethanol content and differential expression of L-amino acid decarboxylases (AADC) in (*Vitis vinifera*) vidal wine grape at different loads. *Pakistan Journal of Botany*, *50*(2), 661-666.

Linsenmeier, A., Rauhut, D., Kürbel, H., & Schubert, S. (2007). Untypical ageing off-flavour and masking effects due to long-term nitrogen fertilization. *Vitis-Geilweilerhof, 46*(1), 33-38.

Linsenmeier, A. W., Loos, U., & Löhnertz, O. (2008). Must composition and nitrogen uptake in a long-term trial as affected by timing of nitrogen fertilization in a cool-climate Riesling vineyard. *American Journal of Enology and Viticulture, 59*(3), 255-264.

Llorens, N., Arola, L., Bladé, C., & Mas, A. (2002). Nitrogen metabolism in a grapevine in vitro system. *Oeno One*, *36*(3), 157-159. doi:10.20870/oenoone.2002.36.3.968

Löhnertz, O. (1988). Untersuchungen zum zeitlichen Verlauf der Nährstoffaufnahme bei Vitis vinifera (cv. Riesling). Geisenheim, Universität Giessen, 228 p.

Longbottom, M. (2009). *Managing grapevine nutrition in a changing environment*: The Australian Wine Research Institute Ltd., p.144 Loulakakis, K. A., Morot-Gaudry, J. F., Velanis, C. N., Skopelitis, D. S., Moschou, P. N., Hirel, B., & Roubelakis-Angelakis, K. A. (2009). Advancements in nitrogen metabolism in grapevine. *In* K. A. Roubelakis-Angelakis (Ed.), *Grapevine Molecular Physiology & Biotechnology* (pp. 161-205). Dordrecht: Springer Netherlands. https://doi.org/10.1007/978-90-481-2305-6 7

Lytra, G., Tempère, S., Zhang, S., Marchand, S., de Revel, G., & Barbe, J.-C. (2014). Olfactory impact of dimethyl sulfide on red wine fruity esters aroma expression in model solution. *Oeno One*, *48*(1), 75-85. doi:10.20870/oeno-one.2014.48.1.1660

Lytra, G., Tempère, S., Marchand, S., de Revel, G., & Barbe, J.-C. (2016). How do esters and dimethyl sulphide concentrations affect fruity aroma perception of red wine? Demonstration by dynamic sensory profile evaluation. *Food Chemistry*, *194*, 196-200. doi:10.1016/j.foodchem.2015.07.143

Maigre, D., & Aerny, J. (2001a). Enherbement permanent et fumure azotée sur cv. Gamay dans le Valais central. 1. Résultats agronomiques. *Revue suisse de Viticulture, Arboriculture et Horticulture, 33*(3), 145-150.

Maigre, D., & Aerny, J. (2001b). Enherbement et fumure azotée sur cv. Gamay dans le Valais central. 2. Résultats analytiques et organoleptiques. *Revue suisse de Viticulture, Arboriculture et Horticulture, 33*(4), 199-204.

Marais, J., Calitz, F., & Haasbroek, P. D. (2001). Relationship between microclimatic data, aroma component concentrations and wine quality parameters in the prediction of Sauvignon blanc wine quality. *South African Journal of Enology and Viticulture, 22*(1), 22. doi:10.21548/22-1-2161

Martínez-Gil, A. M., Garde-Cerdán, T., Lorenzo, C., Félix Lara, J., Pardo, F., & Rosario Salinas, M. (2012). Volatile compounds formation in alcoholic fermentation from grapes collected at 2 maturation stages: influence of nitrogen compounds and grape variety. *Journal of food science*, 77(1), C71-C79. doi:10.1111/j.1750-3841.2011.02441.x

Martínez-Moreno, R., Morales, P., Gonzalez, R., Mas, A., & Beltran, G. (2012). Biomass production and alcoholic fermentation performance of *Saccharomyces cerevisiae* as a function of nitrogen source. *FEMS Yeast Research*, *12*(4), 477-485. doi:10.1111/j.1567-1364.2012.00802.x

Marschner, P., & Rengel, Z. (2012). Chapter 12 -Nutrient availability in soils. *In* P. Marschner (Ed.), *Marschner's Mineral Nutrition of Higher Plants (Third Edition)* (pp. 315-330). San Diego: Academic Press. doi:10.1016/B978-0-12-384905-2.00012-1 Masclaux-Daubresse, C., Daniel-Vedele, F., Dechorgnat, J., Chardon, F., Gaufichon, L., & Suzuki, A. (2010). Nitrogen uptake, assimilation and remobilization in plants: challenges for sustainable and productive agriculture. *Annals of Botany*, *105*(7), 1141-1157. doi:10.1093/aob/mcq028

Mawdsley, P. F. W., Dodson Peterson, J. C., & Casassa, L. F. (2018). Agronomical and Chemical Effects of the Timing of Cluster Thinning on Pinot Noir (Clone 115) Grapes and Wines. *Fermentation*, *4*(3), 60. doi:10.3390/fermentation4030060

Mengel, K., & Pilbeam, D. J. (1992). Nitrogen metabolism of plants. *In: Proceedings of the phytochemical society of Europe* 33 New York: Oxford University Press.

Merbach, W., Mirus, E., Knof, G., Remus, R., Ruppel, S., Russow, R., Gransee, A., & Schulze, J. (1999). Release of carbon and nitrogen compounds by plant roots and their possible ecological importance⁺. *Journal of Plant Nutrition and Soil Science*, *162*(4), 373-383.

Metay, A. l., Magnier, J., Guilpart, N., & Christophe, A. l. (2014). Nitrogen supply controls vegetative growth, biomass and nitrogen allocation for grapevine (cv. Shiraz) grown in pots. *Functional Plant Biology*, *42*(1), 105. doi:10.1071/fp14062

Meyers, J. M., Sacks, G. L., & Vanden Heuvel J. E. (2013). Glycosylated aroma compound responses in 'Riesling' wine grapes to cluster exposure and vine yield. 23(5), 581. doi:10.21273/horttech.23.5.581

Miller, G. H. (2019). *Whiskey science*. Cham, Switzerland: Springer, 529 p. doi:10.1007/978-3-030-13732-8

Molina, J.-A. E., & Smith, P. (1997). Modeling carbon and nitrogen processes in soils. *In* D. L. Sparks (Ed.), *Advances in Agronomy* (Vol. 62, pp. 253-298): Academic Press. doi:10.1016/S0065-2113(08)60570-2

Morinaga, K., Imai, S., Yakushiji, H., & Koshita, Y. (2003). Effects of fruit load on partitioning of ¹⁵N and ¹³C, respiration, and growth of grapevine roots at different fruit stages. *Scientia Horticulturae*, *97*, 239-253. doi:10.1016/S0304-4238(02)00199-1

Morot-Gaudry, J.-F., Moreau, F., Prat, R., Maurel, C., & Sentenac, H. (2017). *Biologie végétale: Nutrition et métabolisme-3e éd*: Dunod.

Murisier, F., & Zufferey, V. (1997). Rapport feuille-fruit de la vigne et qualité du raisin. *Revue suisse de Viticulture, Arboriculture et Horticulture, 29*(6), 355-362.

Nisbet, M. A., Martinson, T. E., & Mansfield, A. K. (2014). Accumulation and prediction of yeast assimilable nitrogen in New York winegrape cultivars. *American Journal of Enology and Viticulture, 65*(3), 325-332. doi:10.5344/ajev.2014.13130

Nykänen, L. (1986). Formation and occurrence of flavor compounds in wine and distilled alcoholic beverages. *American Journal of Enology and Viticulture, 37*(1), 84-96.

Oliva, J., Garde-Cerdán, T., Martínez-Gil, A. M., Rosario Salinas, M., & Barba, A. (2011). Fungicide effects on ammonium and amino acids of Monastrell grapes. *Food Chemistry*, *129*(4), 1676-1680. doi:10.1016/j.foodchem.2011.06.030

Ong, P. K. C., & Acree, T. E. (1999). Similarities in the aroma chemistry of Gewürztraminer variety wines and lychee (Litchi chinesis Sonn.) fruit. *Journal of Agricultural and Food Chemistry*, 47(2), 665-670. doi:10.1021/jf980452j

Ortega-Heras, M., Pérez-Magariño, S., Del-Villar-Garrachón, V., González-Huerta, C., Moro Gonzalez, L. C., Guadarrama Rodríguez, A., . . . Martín de la Helguera, S. (2014). Study of the effect of vintage, maturity degree, and irrigation on the amino acid and biogenic amine content of a white wine from the Verdejo variety. *Journal of the Science of Food and Agriculture*, *94*(10), 2073-2082. doi:10.1002/jsfa.6526

Paolini, M., Ziller, L., Bertoldi, D., Bontempo, L., Larcher, R., Nicolini, G., & Camin, F. (2016). δ^{15} N from soil to wine in bulk samples and proline. *Journal* of Mass Spectrometry, 51(9), 668-674. doi:10.1002/ jms.3824

Parker, A. K., Hofmann, R. W., van Leeuwen, C., McLachlan, A. R. G., & Trought, M. C. T. (2014). Leaf area to fruit mass ratio determines the time of veraison in Sauvignon blanc and Pinot noir grapevines. *Australian Journal of Grape and Wine Research*, 20(3), 422-431. doi:10.1111/ajgw.12092

Parker, A. K., Hofmann, R. W., van Leeuwen, C., McLachlan, A. R. G., & Trought, M. C. T. (2015). Manipulating the leaf area to fruit mass ratio alters the synchrony of total soluble solids accumulation and titratable acidity of grape berries. *Australian Journal of Grape and Wine Research*, 21(2), 266-276. doi:10.1111/ ajgw.12132

Parker, A. K., García de Cortázar-Atauri, I., Gény, L., Spring, J.-L., Destrac, A., Schultz, H., Molitor, D., Lacombe, T., Graça, A., Monamy, C., Stoll M., Storchi, P., Trought, M. C. T., Hofmann, R. W. & van Leeuwen, C. (2020). Temperature-based grapevine sugar ripeness modelling for a wide range of *Vitis vinifera* L. cultivars. *Agricultural and Forest Meteorology, 285-286*, 107902. https://doi.org/10.1016/j.agrformet.2020.107902

Petrie, P. R., & Clingeleffer, P. R. (2006). Crop thinning (hand versus mechanical), grape maturity and anthocyanin concentration: outcomes from irrigated Cabernet-Sauvignon (*Vitis vinifera* L.) in a warm climate. *Australian Journal of Grape and Wine Research*, *12*(1), 21-29. doi:10.1111/j.1755-0238.2006. tb00040.x

Poni, S., Gatti, M., Palliotti, A., Dai, Z., Duchêne, E., Truong, T.-T., Ferrara, G., Stella Matarrese A. M., Gallotta A., Bellincontro A., Mencarelli, F., & Tombesi, S. (2018). Grapevine quality: A multiple choice issue. *Scientia Horticulturae, 234*, 445-462. doi:10.1016/j. scienta.2017.12.035

Portu, J., Gonzalez-Arenzana, L., Hermosin-Gutierrez, I., Santamaria, P., & Garde-Cerdan, T. (2015). Phenylalanine and urea foliar applications to grapevine: effect on wine phenolic content. *Food Chemistry, 180*, 55-63. doi:10.1016/j.foodchem.2015.02.008

Porro, D., Dorigatti, C., Zatelli, A., Ramponi, M., Stefanini, M., & Policarpo, M. (2007). Partitioning of dry matter in grapevines during a season: estimation of nutrient requirement. *Proceedings of the XV International Symposium "GESCO 2007"*, Poreč – Croatia, Vol. 1: 403-411.

Porro, D., Stefanini, M., & Dorigatti, C. (2010). Nitrogen Foliar Uptake and Partitioning in "Cabernet Sauvignon" Gravevines. *Proceedings of the Proc. VIth IS on Mineral Nutrition*, Faro, Portugal. https://doi.org/10.17660/ActaHortic.2010.868.21

Rapp, A., & Mandery, H. (1986). Wine aroma. *Experientia*, 42(8), 873-884. doi:10.1007/bf01941764

Rapp, A., & Versini, G. (1995). Influence of nitrogen compounds in grapes on aroma compounds of wines. In C. George (Ed.), *Developments in Food Science* (Vol. 37, pp. 1659-1694): Elsevier. https://doi.org/10.1016/S0167-4501(06)80257-8

Reddy, M. M., & Ulaganathan, K. (2015). Nitrogen nutrition, its regulation and biotechnological approaches to improve crop productivity. *American Journal of Plant Sciences*, *6*, 2745-2798. doi:10.4236/ajps.2015.618275

Reeve, A. L., Skinkis, P. A., Vance, A. J., Lee, J., & Tarara, J. M. (2016). Vineyard floor management influences 'Pinotnoir' vine growth and productivity more than cluster thinning. *HortScience*, *51*(10), 1233-1244. doi:10.21273/hortsci10998-16

Reeve, A. L., Skinkis, P. A., Vance, A. J., McLaughlin, K. R., Tomasino, E., Lee, J., & Tarara, J. M. (2018). Vineyard floor management and cluster thinning inconsistently affect 'Pinot noir' crop load, berry composition and wine quality. *HortScience*, *53*(3), 318-328. doi:10.21273/hortsci12682-17

Reining, E., Merbach, W., & Knof, G. (1995). ¹⁵N Distribution in wheat and chemical fractionation of root-borne ¹⁵N in the soil. *Isotopes in Environmental and Health Studies*, *31*(3-4), 345-349. doi:10.1080/10256019508036280

Reynard, J. S., Zufferey, V., Nicol, G. C., & Murisier, F. (2011). Soil parameters impact the vine-fruit-wine continuum by altering vine nitrogen status. *Journal International des Sciences de la Vigne et du Vin, 45*(4), 211-221. https://doi.org/10.20870/oeno-one.2011.45.4.1502

Reynard, J. S., Zufferey, V., Nicol, G. C., Pythoud, K., Aleid-Germanier, L., & Murisier, F. (2012). Adaptation du Gamaret aux terroirs viticoles vaudois. *Revue suisse de Viticulture, Arboriculture et Horticulture, 44*(5), 290-296.

Reynolds, A. G. (2010). Viticultural and vineyard management practices and their effects on grape and wine quality. In A. G. Reynolds (Ed.), *Managing Wine Quality* (pp. 365-444): Woodhead Publishing. https://doi.org/10.1533/9781845699284.3.365

Ribéreau-Gayon, P., Dubourdieu, D., Donèche, B., & Lonvaud, A. (2017). *Traité d'œnologie* (7^e edition ed. Vol. tome 1). Paris: Dunod.

Robinson, A. L., Boss, P. K., Solomon, P. S., Trengove, R. D., Heymann, H., & Ebeler, S. E. (2014). Origins of grape and wine aroma. Part 1. Chemical components and viticultural impacts. *American Journal of Enology and Viticulture*, *65*(1), 1-24. doi:10.5344/ajev.2013.12070

Rodriguez-Lovelle, B., & Gaudillère, J. P. (2002). Carbon and nitrogen partitioning in either fruiting or non-fruiting grapevines: effects of nitrogen limitation before and after veraison. *Australian Journal of Grape and Wine Research, 8*(86), 86-94. doi:10.1111/j.1755-0238.2002.tb00216.x

Rollero, S., Mouret, J.-R., Sanchez, I., Camarasa, C., Ortiz-Julien, A., Sablayrolles, J.-M., & Dequin, S. (2016). Key role of lipid management in nitrogen and aroma metabolism in an evolved wine yeast strain. *Microbial Cell Factories*, *15*(1), 32. doi:10.1186/ s12934-016-0434-6

Rossouw, G. C., Smith, J. P., Barril, C., Deloire, A., & Holzapfel, B. P. (2017). Implications of the presence of maturing fruit on carbohydrate and nitrogen distribution in grapevines under postveraison water constraints. *Journal of the American Society for Horticultural Science*, *142*(2), 71-84. doi:10.21273/JASHS03982-16

Roubelakis-Angelakis, K. A., & Kliewer, W. M. (1992). Nitrogen metabolims in grapevine. *In Horticultural reviews* (Vol. 14, pp. 407-452): John Wiley & Sons, Inc. https://doi.org/10.1002/9780470650523.ch9

Rühl, E. H., & Clingeleffer, P. R. (1993). Effect of minimal pruning and virus inoculation on the carbohydrate and nitrogen accumulation in Cabernet franc vines. *American Journal of Enology and Viticulture*, 44(1), 81-85.

Rühl, E., Fuda, A., & Treeby, M. (1992). Effect of potassium, magnesium and nitrogen supply on grape juice composition of Riesling, Chardonnay and Cabernet-Sauvignon [Victoria] vines. *Australian Journal of Experimental Agriculture, 32*(645-649). https://doi.org/10.1071/EA9920645

Rutan, T. E., Herbst-Johnstone, M., & Kilmartin, P. A. (2018). Effect of cluster thinning *Vitis vinifera* cv. Pinot noir on wine volatile and phenolic composition. *Journal of Agricultural and Food Chemistry*, *66*(38), 10053-10066. doi:10.1021/acs.jafc.8b04062

Ryona, I., Pan, B. S., Intrigliolo, D. S., Lakso, A. N., & Sacks, G. L. (2008). Effects of cluster light exposure on 3-isobutyl-2-methoxypyrazine accumulation and degradation patterns in red wine grapes (*Vitis vinifera* L. Cv. Cabernet Franc). *Journal of Agricultural and Food Chemistry*, *56*(22), 10838-10846. doi:10.1021/jf801877y

Sakakibara, H., Takei, K., & Hirose, N. (2006). Interactions between nitrogen and cytokinin in the regulation of metabolism and development. *Trends in Plant Science*, *11*(9), 440-448. doi:10.1016/j. tplants.2006.07.004

Santamaría, P., Lopez, R., Portu, J., Gonzalez-Arenzana, L., Lopez-Alfaro, I., & Garde-Cerdan, T. (2015). Role of phenylalanine in viticulture and enology. *In:* Elliamson D (ed) *Phenylalanine. Dietary, sources, functions and health effects.* Nova Publisher, New York, pp 49–70

Santesteban, L., Miranda, C., Barbarin, I., & Royo, J. (2014). Application of the measurement of the natural abundance of stable isotopes in viticulture: a review. *Australian Journal of Grape and Wine Research*, *21*(2), 157-167. doi:10.1111/ajgw.12124

Santos, T. P., Lopes, C. M., Rodrigues, M. L., De Souza, C. R., Ricardo-da-Silva, J. M., Maroco, J. P., Pereira, J.S., & Chaves, M. M. (2005). Effects of partial root-zone drying irrigation on cluster microclimate and fruit composition of field-grown Castelão grapevines. *Vitis*, 44(3), 117-125.

Schneider, V. (2014). Atypical aging defect: sensory discrimination, viticultural causes, and enological consequences. A review. *American Journal of Enology and Viticulture, 65*(3), 277-284. doi:10.5344/ajev.2014.14014

Schreiner, R. P. (2003). Mycorrhizal colonization of grapevine rootstocks under field conditions. *American Journal of Enology and Viticulture*, *54*(3), 143-149.

Schreiner, R. P., Scagel, C. F., & Lee, J. (2014). N, P, and K supply to Pinot noir grapevines: impact on berry phenolics and free amino acids. *American Journal of Enology and Viticulture*, *65*(1), 43-49. doi:10.5344/ ajev.2013.13037

Schreiner, R. P. (2016). Nutrient uptake and distribution in young Pinot noir grapevines over two seasons. *American Journal of Enology and Viticulture, 67*(4), 436-448. doi:10.5344/ajev.2016.16019

Schreiner, R. P., Osborne, J., & Skinkis, P. A. (2018). Nitrogen requirements of Pinot noir based on growth parameters, must composition, and fermentation behavior. *American Journal of Enology and Viticulture*, *69*(1), 45-58. doi:10.5344/ajev.2017.17043

Seguinot, P., Rollero, S., Sanchez, I., Sablayrolles, J.-M., Ortiz-Julien, A., Camarasa, C., & Mouret, J.-R. (2018). Impact of the timing and the nature of nitrogen additions on the production kinetics of fermentative aromas by *Saccharomyces cerevisiae* during winemaking fermentation in synthetic media. *Food Microbiology*, *76*, 29-39. doi:10.1016/j.fm.2018.04.005 Serra-Stepke, I. M. (2010). *Effect of soil parameters and canopy structure on root growth and distribution*. (Master degree), Stellenbosch University, Stellenbosch, 71pp.

Soubeyrand, E., Basteau, C., Hilbert, G., van Leeuwen, C., Delrot, S., & Gomes, E. (2014). Nitrogen supply affects anthocyanin biosynthetic and regulatory genes in grapevine cv. Cabernet-Sauvignon berries. *Phytochemistry*, *103*, 38-49. doi:10.1016/j.phytochem.2014.03.024

Soubeyrand, E., Colombié, S., Beauvoit, B., Dai, Z., Cluzet, S., Hilbert, G., Christel R., Christel R., Maneta-Peyret Lilly, Dieuaide-Noubhani M., Mérillon J.-M., Gibon, Y., Delrot S. & Gomès, E. (2018). Constraint-based modeling highlights cell energy, redox status and α -ketoglutarate availability as metabolic drivers for anthocyanin accumulation in grape cells under nitrogen limitation. *Frontiers in Plant Science*, 9(421). doi:10.3389/fpls.2018.00421

Spangenberg, J. E., & Zufferey, V. (2018). Changes in soil water availability in vineyards can be traced by the carbon and nitrogen isotope composition of dried wines. *Science of The Total Environment, 635*, 178-187. doi:10.1016/j.scitotenv.2018.04.078

Spring, J.-L. (1999). Indice *chlorophylliendu* feuillage et nutrition azotée du cépage Chasselas. *Revue suisse de Viticulture, Arboriculture et Horticulture, 31*(3), 141-145.

Spring, J.-L., & Zufferey, V. (2000). Intérêt de la détermination de l'indice chlorophyllien du feuillage en viticulture. *Revue suisse de Viticulture, Arboriculture et Horticulture, 32*(6), 323-328.

Spring, J. L. (2001). Influence du type d'enherbement sur le comportement de la vigne et la qualité des vins. Résultats d'un essai sur Chasselas dans le bassin lémanique. *Revue suisse de Viticulture, Arboriculture et Horticulture, 33*(5), 253-260.

Spring, J. L. (2002). Influence du type d'enherbement sur le comportement de la vigne et la qualité des vins. Résultats d'un essai sur Chasselas dans le bassin lémanique. 2. Résultats oenologiques. *Revue suisse de Viticulture, Arboriculture et Horticulture, 34*(2), 111-116.

Spring, J. L. (2003). Localisation de la fumure azotée sur l'intercep dans les vignes enherbées. *Revue suisse de Viticulture, Arboriculture et Horticulture, 35*(2), 113-119.

Spring, J. L., Verdenal, T., Zufferey, V., & Viret, O. (2012). Nitrogen dilution in excessive canopies of Chasselas and Pinot noir cvs. *Journal International des Sciences de la Vigne et du Vin, 46*(3), 233-240. doi:10.20870/oeno-one.2012.46.3.1520

Spring, J. L., Zufferey, V., Verdenal, T., & Viret, O. (2016a). Influence du porte-greffe sur le comportement du Pinot noir dans les conditions du Valais central. *Revue suisse de Viticulture, Arboriculture et Horticulture,* 48(2), 112-122.

Spring, J. L., Reynard, J. S., Zufferey, V., Verdenal, T., Duruz, P., & Viret, O. (2016b). Diversité intra-variétale et sélection clonale de la Petite Arvine. *Revue suisse de Viticulture, Arboriculture et Horticulture, 48*(3), 156-163.

Spring, J.-L., & Verdenal, T. (2017). Fertilisation en viticulture. *In:* S. Sinaj & W. Richner (Eds.), *Principes de fertilisation des cultures agricoles en Suisse (PRIF 2017)* (pp. 276).

Spring, J. L., Reynard, J. S., Zufferey, V., Verdenal, T., Duruz, P., Rösti, J., Lorenzini, F., & Favre, G. (2018). Nouveaux clones de Pinot gris sélectionnés à Agroscope. *Revue suisse de Viticulture, Arboriculture et Horticulture, 50*(3), 158-165.

Stamatiadis, S., Christofides, C., Tsdila, E., Taskos, D., & Schepers, J. S. (2007). Relationship of leaf stable isotopes (δ^{13} C and δ^{15} N) to biomass production in two fertilized Merlot vineyards. *American Journal of Enology and Viticulture*, *58*(1), 67-74.

Stines, A. P., Grubb, J., Gockowiak, H., Henschke, P. A., Hoj, P. B., & van Heeswijck, R. (2000). Proline and arginine accumulation in developing berries of *Vitis vinifera* L. in Australian vineyards: Influence of vine cultivar, berry maturity and tissue type. *Australian Journal of Grape and Wine Research*, *6*, 150-158. doi:10.1111/j.1755-0238.2000.tb00174.x

Stitt, M., & Krapp, A. (1999). The interaction between elevated carbon dioxide and nitrogen nutrition: the physiological and molecular background. *Plant, Cell & Environment, 22*(6), 583-621. doi:10.1046/j.1365-3040.1999.00386.x

Stockert, C. M., Bisson, L. F., Adams, D. O., & Smart, D. R. (2013). Nitrogen status and fermentation dynamics for Merlot on two rootstocks. *American Journal of Enology and Viticulture, 64*(2), 195-202. doi:10.5344/ajev.2013.12065

Styger, G., Prior, B., & Bauer, F. F. (2011). Wine flavor and aroma. *Journal of Industrial Microbiology & Biotechnology*, *38*(9), 1145. doi:10.1007/s10295-011-1018-4

Sweetman, C., Deluc, L. G., Cramer, G. R., Ford, C. M., & Soole, K. L. (2009). Regulation of malate metabolism in grape berry and other developing fruits. *Phytochemistry*, 70. doi:10.1016/j. phytochem.2009.08.006

Swiegers, J. H., Bartowsky, E. J., Henschke, P. A., & Pretorius, I. S. (2005). Yeast and bacterial modulation of wine aroma and flavour. *Australian Journal of Grape and Wine Research*, *11*(2), 139-173. doi:10.1111/j.1755-0238.2005.tb00285.x

Taskos, D. G., Koundouras, S., Stamatiadis, S., Zioziou, E., Nikolaou, N., Karakioulakis, K., & Theodorou, N. (2015). Using active canopy sensors and chlorophyll meters to estimate grapevine nitrogen status and productivity. *Precision Agriculture, 16*(1), 77-98. doi:10.1007/s11119-014-9363-8

Tegeder, M. (2014). Transporters involved in source to sink partitioning of amino acids and ureides: opportunities for crop improvement. *Journal of Experimental Botany*, *65*(7), 1865-1878. doi:10.1093/jxb/eru012

Tesic, D., Keller, M., & Hutton, R. J. (2007). Influence of vineyard floor management practices on grapevine vegetative growth, yield, and fruit composition. *American Journal of Enology and Viticulture, 58*(1), 1-11.

Thomidis, T., Zioziou, E., Koundouras, S., Karagiannidis, C., Navrozidis, I., & Nikolaou, N. (2016). Effects of nitrogen and irrigation on the quality of grapes and the susceptibility to *Botrytis* bunch rot. *Scientia Horticulturae, 212*, 60-68. doi:10.1016/j. scienta.2016.09.036

Tomasi, N., Monte, R., Varanini, Z., Cesco, S., & Pinton, R. (2015). Induction of nitrate uptake in Sauvignon Blanc and Chardonnay grapevines depends on the scion and is affected by the rootstock. *Australian Journal of Grape and Wine Research, 21*(2), 331-338. doi:10.1111/ajgw.12137

Torrea, D., Varela, C., Ugliano, M., Ancin-Azpilicueta, C., Leigh Francis, I., & Henschke, P. A. (2011). Comparison of inorganic and organic nitrogen supplementation of grape juice - Effect on volatile composition and aroma profile of a Chardonnay wine fermented with *Saccharomyces cerevisiae* yeast. *Food Chemistry*, *127*(3), 1072-1083. doi:10.1016/j.foodchem.2011.01.092

Treeby, M. T., & Wheatley, D. M. (2006). Effect of nitrogen fertiliser on nitrogen partitioning and pool sizes in irrigated Sultana grapevines. *Australian Journal of Experimental Agriculture, 46*(9), 1207-1215. doi:10.1071/EA05238

Trouvelot, S., Bonneau, L., Redecker, D., van Tuinen, D., Adrian, M., & Wipf, D. (2015). Arbuscular mycorrhiza symbiosis in viticulture: a review. *Agronomy for Sustainable Development*, *35*(4), 1449-1467. doi:10.1007/s13593-015-0329-7

Ugliano, M., Henschke, P. A., Herdrich, M. J., & Pretorius, I. S. (2007). Nitrogen management is critical for wine flavour and style. *Aus. N.Z. Wine Industry J.*, 22(6), 24-30.

Ugliano, M., & Henschke, P. A. (2009). Yeasts and Wine Flavour. *In* M. V. Moreno-Arribas & M. C. Polo (Eds.), *Wine Chemistry and Biochemistry* (pp. 313-392). New York, : Springer. https://doi.org/10.1007/978-0-387-74118-5_17

Usman, S., Singh, S. P., Rawat, Y. S., & Bargali, S. S. (2000). Fine root decomposition and nitrogen mineralisation patterns in *Quercus leucotrichophora* and *Pinus roxburghii* forests in central Himalaya. *Forest Ecology and Management*, *131*(1), 191-199. doi:https://doi.org/10.1016/S0378-1127(99)00213-3

Van Cleemput, O., Zapata, F., & Vanlauwe, B. (2008). Use of tracer technology in mineral fertilizer

management. In *Guidelines on nitrogen management in agricultural systems* (Vol. 29, pp. 19-125). Vienna: International Atomic Energy Agency.

van Leeuwen, C., Friant, P., Soyer, J.-P., Molot, C., Chone, X., & Dubourdieu, D. (2000). L'intérêt du dosage de l'azote total et de l'azote assimilable dans le moût comme indicateur de la nutrition azotée de la vigne. *Journal International des Sciences de la Vigne et du Vin, 34*(2), 75-82.

van Leeuwen, C., Friant, P., Choné, X., Tregoat, O., Koundouras, S., & Dubourdieu, D. (2004). Influence of Climate, Soil and Cultivar on Terroir. *American Journal of Enology and Viticulture*, *55*(3), 207-217.

van Leeuwen, C., & Seguin, G. (2006). The concept of terroir in viticulture. *Journal of Wine Research*, *17*(1), 1-10. doi:10.1080/09571260600633135

van Leeuwen, C., Roby, J.-P., & de Rességuier, L. (2018). Soil-related terroir factors: a review. *Oeno One, 52*(2), 173-188. doi:10.20870/oeno-one.2018.52.2.2208

Verdenal, T., Spangenberg, J. E., Zufferey, V., Lorenzini, F., Spring, J. L., & Viret, O. (2015). Effect of fertilisation timing on the partitioning of foliar-applied nitrogen in *Vitis vinifera* cv. Chasselas: a ¹⁵N labelling approach. *Australian Journal of Grape and Wine Research*, *21*(1), 110-117. doi:10.1111/ajgw.12116

Verdenal, T., Spangenberg, J. E., Zufferey, V., Lorenzini, F., Dienes-Nagy, A., Gindro, K., Spring J.-L., Viret, O. (2016a). Leaf-to-fruit ratio affects the impact of foliarapplied nitrogen on N accumulation in the grape must. *Journal International des Sciences de la Vigne et du Vin, 50*(1), 23-33. doi:10.20870/oeno-one.2016.50.1.55

Verdenal, T., Zufferey, V., Spring, J. L., Rösti, J., Dienes-Nagy, A., Lorenzini, F., Wolfender, J.-L., Spangenberg, J., Burgo, S., Gindro, K, & Viret, O. (2016b). Complémentation foliaire azotée du cépage *Vitis vinifera* Doral dans le canton de Vaud. *Revue suisse de Viticulture, Arboriculture et Horticulture,* 48(4), 238-245.

Verdenal, T., Zufferey, V., Spring, J. L., Rösti, J., Dienes-Nagy, A., Lorenzini, F., Wolfender, J.-L., Spangenberg, J., Burgo, S., Gindro, K, Viret, O. (2016c). Complémentation foliaire azotée du cépage *Vitis vinifera* Doral dans le canton de Vaud. *Revue suisse de Viticulture, Arboriculture et Horticulture,* 48(4), 238-245.

Verdenal, T., Zufferey, V., Spring, J. L., Dienes-Nagy, A., Belcher, S., & Lorenzini, F. (2017). Impact de la zone d'application de l'urée foliaire sur la teneur en azote des raisins. *Revue suisse de Viticulture, Arboriculture et Horticulture, 49*(6), 354-359.

Verdenal, T., Zufferey, V., Dienes-Nagy, A., Bourdin, G., Gindro, K., Viret, O., & Spring, J.-L. (2019). Timing and intensity of grapevine defoliation: an extensive overview on five cultivars in Switzerland. *American Journal of Enology and Viticulture, 70*(4), 427-434. doi:10.5344/ajev.2019.19002 Verdenal, T., Spangenberg, J. E., Zufferey, V., Dienes-Nagy, Á., Viret, O., van Leeuwen, C., & Spring, J.-L. (2020). Impact of crop load on nitrogen uptake and reserve mobilisation in *Vitis vinifera*. *Functional Plant Biology*, *47*, 744-756. doi:10.1071/FP20010

Vilanova, M., Ugliano, M., Varela, C., Siebert, T., Pretorius, I. S., & Henschke, P. A. (2007). Assimilable nitrogen utilisation and production of volatile and non-volatile compounds in chemically defined medium by *Saccharomyces cerevisiae* wine yeasts. *Applied Microbiology and Biotechnology*, 77(1), 145-157. doi:10.1007/s00253-007-1145-z

Vincenzini, M., Guerrini, S., Mangani, S., & Granchi, L. (2017). Amino acid metabolisms and production of biogenic amines and ethyl carbamate. *In* H. König, G. Unden, & J. Fröhlich (Eds.), *Biology of Microorganisms on Grapes, in Must and in Wine* (pp. 231-253). Cham: Springer International Publishing. https://doi.org/10.1007/978-3-319-60021-5 10

Vrignon-Brenas, S., Metay, A., Leporatti, R., Gharibi, S., Fraga, A., Dauzat, M., Roland G., & Pellegrino, A. (2019). Gradual responses of grapevine yield components and carbon status to nitrogen supply. *Oeno One, 53*(2). doi:10.20870/oeno-one.2019.53.2.2431

Wang, Y., Janz, B., Engedal, T., & Neergaard, A. (2017). Effect of irrigation regimes and nitrogen rates on water use efficiency and nitrogen uptake in maize. *Agricultural Water Management*, *179*, 271-276. doi:10.1016/j.agwat.2016.06.007

Wang, Y., He, Y.-N., Chen, W.-K., He, F., Chen, W., Cai, X.-D., Duan, C.-Q., & Wang, J. (2018). Effects of cluster thinning on vine photosynthesis, berry ripeness and flavonoid composition of Cabernet-Sauvignon. *Food Chemistry*, 248, 101-110. doi:10.1016/j.foodchem.2017.12.021

Wang, Y., He, Y.-N., He, L., He, F., Chen, W., Duan, C.-Q., & Wang, J. (2019). Changes in global aroma profiles of Cabernet-Sauvignon in response to cluster thinning. *Food Research International*, *122*, 56-65. doi:10.1016/j.foodres.2019.03.061

Waterhouse, A. L., Sacks, G. L., & Jeffery, D. W. (2016). *Understanding wine chemistry*: Wiley-Blackwell. https://doi.org/10.1002/9781118730720

Wermelinger, B. (1991). *Nitrogen Dynamics in Grapevine: Physiology and Modeling*. Paper presented at the International Symposium on Nitrogen in Grapes and Wine, Seattle WA, USA.

Weyand, K. M., & Schultz, H. R. (2006). Long-term dynamics of nitrogen and carbohydrate reserves in woody parts of minimally and severely pruned Riesling vines in a cool climate. *American Journal of Enology and Viticulture*, *57*(2), 172-182.

White, R., Balachandra, L., Edis, R., & Chen, D. (2007). The soil component of terroir. *Journal International des Sciences de la Vigne et du Vin*, 41(1), 9-18. https://doi.org/10.20870/oeno-one.2007.41.1.860

Williams, L. E. (2015). Recovery of ¹⁵N Labeled Fertilizer by Thompson Seedless Grapevines: Effects of N Fertilizer Type and Irrigation Method. *American Journal of Enology and Viticulture, 66*, 509-517. doi:10.5344/ajev.2015.14115

Xu, G., Fan, X., & Miller, A. J. (2012). Plant nitrogen assimilation and use efficiency. *Annual Review of Plant Biology*, *63*(1), 153-182. doi:10.1146/annurev-arplant-042811-105532

Zamboni, M., Garavani, A., Gatti, M., Vercesi, A., Parisi, M. G., Bavaresco, L., & Poni, S. (2016). Vegetative, physiological and nutritional behavior of new grapevine rootstocks in response to different nitrogen supply. *Scientia Horticulturae*, 202, 99-106. doi:10.1016/j.scienta.2016.02.032

Zapata, C., Deleens, E., Chaillou, S., & Magne, C. (2004a). Partitioning and mobilization of starch and N reserves in grapevine (*Vitis vinifera* L.). *J Plant Physiol*, *161*(9), 1031-1040. doi:10.1016/j.jplph.2003.11.009

Zapata, C., Deléens, E., Chaillou, S., & Magné, C. (2004b). Mobilisation and distribution of starch and total N in two grapevine cultivars differing in their susceptibility to shedding. *Functional Plant Biology*, *31*, 1127-1135. doi:10.1071/FP04028

Zufferey, V., & Maigre, D. (2007). Âge de la vigne. I. Influence sur le comportement physiologique des souches. *Revue suisse de Viticulture, Arboriculture et Horticulture, 39*(4), 257-261.

Zufferey, V., & Murisier, F. (2007). Assessment of plant hydraulics in grapevine on various "terroirs" in the canton de Vaud (Switzerland). *Journal International des Sciences de la Vigne et du Vin, 41*(2), 95-102. https://doi.org/10.20870/oeno-one.2007.41.2.854

Zufferey, V., & Maigre, D. (2008). Âge de la vigne II. Influence sur la qualité des raisins et des vins. *Revue suisse de Viticulture, Arboriculture et Horticulture,* 40(4), 241-245.

Zufferey, V., Murisier, F., Belcher, S., Lorenzini, F., Vivin, P., Spring, J. L., & Viret, O. (2015). Nitrogen and carbohydrate reserves in the grapevine (*Vitis vinifera* L. 'Chasselas'): the influence of the leaf to fruit ratio. *Vitis*, *54*, 183-188. doi:10.5073/vitis.2015.54.183-188

Zufferey, V., Spring, J.-L., Verdenal, T., Dienes, A., Belcher, S., Lorenzini, F., Koestel F., Rösti, H., Gindro, K., Spangenberg, J., & Viret, O. (2017). The influence of water stress on plant hydraulics, gas exchange, berry composition and quality of Pinot noir wines in Switzerland. *Oeno One*, *51*(1), 17-27. doi:10.20870/oeno-one.2017.51.1.1314

Zufferey, V., Verdenal, T., Dienes, A., Belcher, S., Lorenzini, F., Koestel, C., Gindro, K., Spangenberg, J.E., Viret, O. & Spring, J. L. (2018). The impact of plant water status on the gas exchange, berry composition and wine quality of Chasselas grapes in Switzerland. *Oeno One, 52*(4), 333-347. doi:10.20870/oeno-one.2018.52.4.2181