Iron Assimilation and Carbon Metabolism in ‘Concord’ Grapevines Grown at Different pHs

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ABSTRACT. ‘Concord’ grapevines (Vitis labruscana Bailey) are susceptible to lime-induced chlorosis, which decreases growth and productivity. In two separate experiments, we grew own-rooted vines in a peat–perlite medium adjusted to different pHs with CaCO3 to characterize how lime-induced Fe deficiency affects root and leaf ferric chelate reductase (FCR) and key enzymes and metabolites involved with glycolysis and the tricarboxylic acid (TCA) cycle in leaves. In addition, we measured the pH of the xylem sap as well as Fe, citrate, and malate concentrations. For both experiments, foliar levels of total Fe, active Fe (extracted in 0.1N HCl), and chlorophyll decreased as lime rate increased. An increase in root-medium pH from 5.8 to 7.5 resulted in a 10-fold increase in root FCR activity, whereas leaf FCR activity decreased 10-fold. An increase in root-medium pH did not raise xylem sap pH but decreased Fe and citrate to some extent. Xylem malate was highest at pH 6.6 and decreased both above and below this pH. Foliar data were evaluated in relation to active Fe content, because it is a better indicator of Fe nutritional status. Lower active Fe concentrations were monitored in ‘Concord’ grapevines, and Fe deficiency was more severe in plants grown at pH 6.6 and 7.0. Our results suggest that leaf FCR activity may limit Fe assimilation to a greater extent than root FCR activity. The decreased leaf aconitase activity under Fe deficiency is the most likely cause of the increase in citrate levels. Greater activity of the other glycolytic and TCA enzymes under Fe deficiency may help to funnel carbon into the mitochondria and enhance NAD(P) reduction. Citrate levels (and the citrate/malate ratios) in the xylem exudate and leaf were much lower when compared with other species and may be linked to Fe inefficiency of ‘Concord’.

Chlorosis from lime-induced Fe deficiency limits grapevine (Vitis L.) growth and productivity (Bavaresco et al., 2003; Gruber and Kosegarten, 2002; Mengel et al., 1984a). As soil pH increases, Fe solubility decreases, and an increase in soil pH by one unit decreases Fe3+ activity 1000-fold (Lindsay and Schwab, 1982). ‘Concord’ grapevines (V. labruscana) are particularly susceptible to lime-induced chlorosis, and this situation can be made worse under conditions of high soil moisture and low soil temperature (Davenport and Stevens, 2006).

When Fe is limiting, dicotyledons can facilitate Fe uptake by acidifying the rhizosphere to increase Fe solubility and exuding organic acids and phenolics to chelate Fe (de Vos et al., 1986; Marschner et al., 1986; Römheld and Marschner, 1983). In addition, roots can increase the activity of plasma membrane-bound ferric chelate reductase (FCR), which uses cytosolic NAD(P)H to cleave Fe(III)-chelates and reduce Fe3+ to Fe2+ before transport across the membrane by inducible Fe2+-transporters (Chaney et al., 1972; Vert et al., 2002). Reduction of Fe(III)-chelates is proposed to be the rate-limiting step for Fe assimilation in dicotyledons (Grotz and Guerinot, 2002), and root FCR activity is often higher in Fe-efficient vs. Fe-inefficient species (Brancadoro et al., 1995; Römheld and Marschner, 1981).

Iron is transported to the leaves primarily as Fe(III)-citrate, although there is some evidence that nicotianamine and carboxylic acids such as malate may play a role in Fe transport (Rombolá et al., 2000; Tiffon, 1970; von Wirén et al., 1999). Foliar Fe assimilation is dependent on FCR activity to reduce Fe(III)-chelates and provide Fe2+ for transport across the mesophyll plasma membrane (Brüggemann et al., 1993). Unlike root FCR, foliar FCR does not appear to be induced by Fe deficiency; however, foliar FCR activity is regulated by light (Brüggemann et al., 1993; de la Guardia and Alcantara, 1996; Gonzalez-Vallejo et al., 2000; Larbi et al., 2001).

It has been proposed that bicarbonate uptake increases the pH of xylem sap and leaf apoplast and interferes with foliar Fe utilization (Mengel et al., 1984a, 1984b), but this is not well-agreed on. For instance, the xylem sap pH of corn (Zea mays L.), seedlings increased 0.6 and 1.1 units when supplied with 5 to 20 mM HCO3 in the nutrient solution (Wegner and Zimmermann, 2004), and López-Millán et al. (2001b) found that the pH of the
apoplast in Fe-deficient pear (*Pyrus communis* L.) leaves was 6.5 to 6.6, whereas the apoplast pH of green leaves was lower, from 5.5 to 5.9. Yet on the other hand, Nikolic and Romheld (2002) found that the addition of 10 mM HCO$_3^-$ to the nutrient solution did not change the pH of the leaf apoplastic fluid in sunflower and that there was no difference in the leaf apoplastic pH between chlorotic and field-grown *Vitis vinifera* L. In addition to bicarbonate, nitrate may also influence the pH of the xylem sap and leaf apoplast and impact Fe assimilation, but this is not fully supported either (Kosegarten et al., 2001; Lucena, 2000; Nikolic and Romheld, 2003).

The accumulation of organic acids, primarily malate and citrate, commonly occurs in iron-deficient roots and leaves (Abadia et al., 2002). For example, citrate and malate increased in the root tips of four different grapevine genotypes in response to Fe limitation, and chlorosis-resistant genotypes contained higher levels of organic acids (Ollat et al., 2003). An increase in malate and citrate in response to Fe deficiency has a number of potential benefits. Increased root exudation of citric and malic acid will improve soil Fe availability both through chelation and rhizosphere acidification (Jones, 1998; Landsberg, 1981). In addition, greater concentrations of citrate could also aid in transport of Fe in the xylem (Tiffin, 1970).

It is not clear, however, why organic acids accumulate in Fe-deficient tissues. Citrate accumulation in Fe-deficient plants was originally thought to result from a decreased conversion of citrate to isocitrate by aconitase, which requires Fe as a cofactor (Bacon et al., 1961). However, aconitase activity is not consistently decreased under Fe deficiency (de Vos et al., 1986; López-Millán et al., 2001a). Alternatively, organic acid accumulation could result from increased carbon fixation by phosphoenolpyruvate (PEP) carboxylase when soil HCO$_3^-$ is high or Fe is limiting. According to this theory, high rates of H$^+$ efflux under Fe deficiency would increase the pH of the cytoplasm and activate PEP carboxylase (Felle, 1988; Rabotti et al., 1995). Carboxylation of PEP through PEP carboxylase would result in oxaloacetate (OAA) and malate synthesis and a lowering of cytosolic pH (Davies, 1986).

Considering the importance of FCR and organic acids in Fe assimilation, there were two main objectives for our work with low pH-tolerant (Fe-inefficient) ‘Concord’ grapevines. The first objective was to characterize root and leaf ferric chelate reductase activity in response to lime-induced Fe limitation. The second objective was to quantify how lime-induced Fe limitation affects key enzymes and metabolites involved with glycolysis and the tricarboxylic acid cycle in leaves. We hypothesized that Fe-deficient leaves would have lower aconitase activity and that this would result in an accumulation of citrate. We also hypothesized that an increase in CaCO$_3$ in the rooting medium would raise the xylem sap pH and negatively impact Fe utilization.

**Materials and Methods**

Two separate experiments were conducted using a peat/perlite medium (starting pH 4.5) that was adjusted to different pHs using CaCO$_3$. We provided the vines with all essential nutrients except Fe as a way of ensuring that the lime-induced chlorosis was primarily the result of Fe limitation. By omitting Fe from the nutrient solution, and providing other micro-nutrients as sulfate salts, we could avoid any complications from the use of synthetic chelates and allow the vines to mine for Fe from the peat and perlite.

**Expt. 1: Root and leaf ferric chelate reductase activity in ‘Concord’ grapevines grown at three different root-medium pHs**

**Plant culture and treatments.** Own-rooted 1-year-old ‘Concord’ grapevines were transplanted into 19.8-L plastic pots containing 80% perlite:20% perlite (v:v) (Conrad Fafard, Agawam, MA) and grown for 9 weeks in a greenhouse. Substrate pH was adjusted with 2.1, 2.8, or 16.3 kg m$^{-3}$ CaCO$_3$ (Grade R2; Mississippi Lime, St. Genevieve, MO), and the pH of a 1:2 (v:v) substrate:deionized H$_2$O mixture after 4 weeks was 5.8 ± 0.1, 6.8 ± 0.3, or 7.5 ± 0.1, respectively. There were five-three-plant replicates per pH treatment in a randomized complete block design.

Once a week, each plant received 1 L of a water-soluble fertilizer with all essential nutrients except Fe. Nutrients were blended based on Hoagland’s No. 2 solution by GreenCare Fertilizers (Chicago) and applied at a rate of (in mg L$^{-1}$) 210 N, 31 P, 235 K, 160 Ca, 49 Mg, 0.50 Mn, 0.05 Zn, 0.02 Cu, 0.01 Mo, and 0.49 B. Micronutrient cations were applied as sulfate salts, and solution was supplemented with 80 mg L$^{-1}$ K from KCl.

At budbreak, extra shoots were removed and two shoots were allowed to grow on each plant. Nine weeks after transplanting, the youngest fully expanded leaf from one shoot per replication was selected for foliar FCR, chlorophyll, and total and active Fe analysis. After collecting leaves, root tips from one plant per replication were collected for root FCR measurements.

**Leaf and root ferric chelate reductase assay and Fe analysis.** Foliar FCR was assayed according to Rombolá et al. (2000) and de la Guardia and Alcantara (1996) with some modifications. During midday, 12 leaf discs (0.5 cm$^2$ each) were punched from each leaf and immediately placed in 5 mL of ice-cold 50 mM MES-KOH pH 5.5, 500 mM sorbitol, 1 mM CaCl$_2$, and 1% bovine serum albumin (BSA) and kept on ice until all discs were punched. Discs were rinsed twice with 5 mL of ice-cold rinse buffer (50 mM MES-KOH pH 6.5, 500 mM sorbitol, 1 mM CaCl$_2$) and then were vacuum-infiltrated (5 kPa) for 20 min in 5 mL of rinse buffer. FCR activity was determined in 3 mL of 50 mM MES-KOH pH 6.5, 0.5 mM CaSO$_4$, 500 mM sorbitol, 1 mM KCl, and 750 µM BPDS. The reaction was initiated with 250 µM Fe(III)-ethylenediaminetetraacetic acid (EDTA), and the discs were shaken under 5 kPa vacuum and ~200 µmol m$^{-2}$ s$^{-1}$ light at sample level ($λ > 600$ nm filtered with red plastic) for 30 min at 25 °C. The absorbance of Fe(II)-BPDS was determined at 535 nm against a no-sample blank, and Fe$^{2+}$ concentration was calculated using a molar extinction coefficient of 22.14 mm$^{-1}$ cm$^{-1}$ (Chaney et al., 1972). To correct for nonenzymatic reduction, samples were also shaken in assay buffer without Fe(III)-EDTA. After 30 min, an aliquot was removed, mixed with 250 µM Fe(III)-EDTA, and placed in darkness for 30 min.

After leaf FCR measurements, chlorophyll was extracted from two leaf discs (1.2 cm$^2$ each) according to Arnon (1949). Leaves were then rinsed in deionized H$_2$O and dried at 70 °C for 7 d. Total and active Fe was determined according to Smith and Cheng (2006).

Root FCR was determined as described by Poonnachit and Darnell (2004). Root tips (~1 cm long) were cut and placed in 20 mL of ice-cold 0.5 mM CaCl$_2$ and 1% BSA and kept on ice until all roots were harvested. Root tips were rinsed in 10 mL of ice-cold 0.5 mM CaCl$_2$, and then ~100 mg were transferred to 5 mL of 10 mM MES-KOH pH 5.5, 0.2 mM CaSO$_4$, and 750 µM
BPDS. The reaction was initiated with 250 μM Fe(III)-EDTA, and samples were shaken in the dark for 60 min at 25 °C. Fe²⁺ concentration was determined as described for foliar FCR and corrected for nonenzymatic reduction by shaking root tips in assay buffer without Fe(III)-EDTA for 60 min in the dark and then mixing an aliquot with 250 μM Fe(III)-EDTA and shaking for an additional 60 min in the dark.

Expt. 2: Effect of root-medium pH on carbon metabolism in ‘Concord’ grapevines

**PLANT CULTURE AND TREATMENTS.** Own-rooted 1-year-old ‘Concord’ grapevines were transplanted on 5 May 2005 into 19.8-L plastic pots containing 80% peat:20% perlite (v:v) and grown outdoors from 17 May 2005 to 29 Sept. 2005. Substrate pH was adjusted with 1.4, 2.1, 2.8, 4.1, or 16.3 kg·m⁻³ CaCO₃, and the mean pH over 9 weeks (measured on weeks 1, 5, and 9) was 5.1 ± 0.1, 5.7 ± 0.1, 6.6 ± 0.0, 7.1 ± 0.0, or 7.5 ± 0.1, respectively. There were five three-plant replicates per pH treatment in a completely randomized design.

At budbreak, extra shoots were removed and two shoots were allowed to grow on each plant. Twice per week, each plant received 1 L of the same nutrient solution used in Expt. 1 with no applied Fe. Nine weeks after transplanting, the youngest fully expanded leaf from one shoot per replicate was chosen for analysis during midday, and the corresponding leaf on the second shoot was selected for analysis during the night. In addition, 30-cm shoot tips were harvested from another shoot per replicate, and xylem sap was expressed using a Scholander pressure bomb.

**XYLEM SAP COLLECTION AND ANALYSIS.** Approximately 500 μL of xylem sap was expressed at 2 MPa and collected in a 1.5-mL microcentrifuge tube by resting the pressure bomb on its side and allowing the sap to drip into the tube. The phloem was stripped from 1.5 cm of the shoot tip, and the tip was blotted several times with a paper towel before collection. The pH of the sap was measured immediately with a micro pH electrode (9802BN; Thermo Orion, Waltham, MA). Sap was stored at −80 °C until it was analyzed for total Fe, citrate, and malate.

Total Fe in the xylem sap was determined essentially as described for the analysis of active Fe in leaves (Smith and Cheng, 2006) with the exception that 50 μL of sap was mixed with 50 μL of 4% (v/v) HClO₄ and allowed to sit for 1 h before analysis. Citrate and malate in the xylem sap were assayed according to Passonneau and Lowry (1993) with some modifications. Citrate was quantified in a 1-mL mixture containing 200 mM HEPES [4-(2-Hydroxyethyl)piperazine-1-ethanesulfonic acid]-KOH pH 7.5, 200 μM ZnSO₄, 200 μM NADH, 5 units of malate dehydrogenase, 4 units of lactate dehydrogenase, 20 μL xylem sap, and 0.2 units of citrate lyase.

Malate concentration was determined in a 1-mL mixture containing 250 mM glycine-glycine-KOH pH 10.0, 40 mM glutamate-KOH pH 10.0, 2.7 mM NAD, 1 unit of glutamate-oxaloacetate transaminase, 5 units of malate dehydrogenase, and 10 μL of xylem sap. For all NAD(P)-linked assays used in these experiments, absorbance was determined at 340 nm and enzyme activity or metabolite concentration calculated using a molar extinction coefficient of 6.22 mm⁻¹·cm⁻¹.

**GAS EXCHANGE AND CHLOROPHYLL FLUORESCENCE MEASUREMENTS.** Gas exchange was measured with a CIRAS-1 system (PP Systems, Herts, UK) at ambient CO₂ (360 μmol·mol⁻¹) either at noon under a photon flux density of 1606 ± 57 μmol·m⁻²·s⁻¹ and a leaf temperature of 27.4 ± 0.8 °C or at midnight with a leaf temperature of 17.2 ± 0.6 °C. Chlorophyll fluorescence was measured with a pulse-modulated fluorometer (FMS2; Hansatech Instruments, Norfolk, UK) as described by Chen and Cheng (2003). Photosystem II quantum efficiency (proportion of light absorbed by PSII used in photochemistry) was measured at noon directly after gas exchange measurements and calculated as (Fₘ’ − F’)/Fₘ’ (Genty et al., 1989). Maximum photochemical efficiency of dark-adapted leaves was determined at midnight after gas exchange measurements and was calculated as Fₗ'/Fₘ’ (van Kooten and Snel, 1990).

**SAMPLE COLLECTION, CHLOROPHYLL AND Fe ANALYSIS.** After gas exchange and fluorescence measurements, leaf discs (1.2 cm²) were punched from each leaf and frozen in liquid N₂. Frozen leaf discs were stored at −80 °C until analysis. Chlorophyll was extracted from two leaf discs according to Arnon (1949), and protein was quantified from two discs according to Bradford (1976). The area of the remaining leaf was measured (corrected for leaf discs removed), and then leaves were rinsed in deionized H₂O and dried at 70 °C for 7 d. Total and active Fe was determined according to Smith and Cheng (2006). At the end of the season, all leaves were harvested from one plant per replication. The total leaf area per vine was measured, leaves were dried at 70 °C for 7 d, and then the total foliar dry weight was recorded.

**ASSAY OF ENZYMES IN CARBON METABOLISM.** For foliar enzyme assays, two discs were ground with a precooled mortar and pestle in 2.0 mL of 200 mM HEPES [adjusted to pH 7.5 with dry TRIS (tris (hydroxymethyl) aminomethane)], 1 mM EDTA, 5 mM MgSO₄, 5 mM dithiothreitol (DTT), 1% (w/v) Triton X-100 (Hach Co., Ames, IA), 30% (w/v) glycerol, 8% (w/v) polyvinylpyrrolidone (PVPP), and 1 mM phenylmethylsulfonyl fluoride (PMSF). Extract was centrifuged at 15,000 g, for 2 min to remove PVPP, the supernatant was centrifuged again at 15,000 g for 10 min, and the supernatant was used immediately for the measurement of enzyme activity. Aconitase, NAD-malate dehydrogenase (MDH), and NAD-malic enzyme were measured as described by Jenner et al. (2001) with some modifications. Aconitase was assayed in a 1-mL mixture containing 85 mM HEPES-KOH pH 7.5, 10 mM MgSO₄, 5 mM MnCl₂, 1 mM DTT, 1 mM EDTA, 250 mM NaF, and 0.5 mM ATP for 15 min at 25 °C and then stopped by boiling for 2 min. Pyruvate was determined in a 1-mL mixture containing 100 mM imidazole-HCl pH 7.0, 10 mM MgSO₄, 50 μM coenzyme A, 4 mM NAD, and 50 μL enzyme extract. The reaction was initiated with 20 mM malate and allowed to run for 15 min at 25 °C and then stopped by boiling for 2 min. Pyruvate was determined as described for the analysis of active Fe in leaves (Smith and Cheng, 2006) with the exception that 50 μL of mixture was used containing 200 mM HEPES [4-(2-Hydroxyethyl)piperazine-1-ethanesulfonic acid]-KOH pH 7.5, 200 μM ZnSO₄, 200 μM NADH, 5 units of malate dehydrogenase, 4 units of lactate dehydrogenase, 20 μL xylem sap, and 0.2 units of citrate lyase. Malate concentration was determined in a 1-mL mixture containing 250 mM glycine-glycine-KOH pH 10.0, 40 mM glutamate-KOH pH 10.0, 2.7 mM NAD, 1 unit of glutamate-oxaloacetate transaminase, 5 units of malate dehydrogenase, and 10 μL of xylem sap. For all NAD(P)-linked assays used in these experiments, absorbance was determined at 340 nm and enzyme activity or metabolite concentration calculated using a molar extinction coefficient of 6.22 mm⁻¹·cm⁻¹.

**SAMPLE COLLECTION, CHLOROPHYLL AND Fe ANALYSIS.** After gas exchange and fluorescence measurements, leaf discs (1.2 cm²) were punched from each leaf and frozen in liquid N₂. Frozen leaf discs were stored at −80 °C until analysis. Chlorophyll was extracted from two leaf discs according to Arnon (1949), and protein was quantified from two discs according to Bradford (1976). The area of the remaining leaf was measured (corrected for leaf discs removed), and then leaves were rinsed in deionized H₂O and dried at 70 °C for 7 d. Total and active Fe was determined according to Smith and Cheng (2006). At the end of the season, all leaves were harvested from one plant per replication. The total leaf area per vine was measured, leaves were dried at 70 °C for 7 d, and then the total foliar dry weight was recorded.
containing 100 mM Bicine-KOH pH 8.0, 5 mM MgSO₄, 5 mM MnCl₂, 0.5 mM NADP, and 75 µL of enzyme extract. The reaction was initiated with 5 mM malate. G6PDH was assayed in a 1-mL mixture containing 100 mM TRIS-HCl pH 8.0, 1 mM EDTA, 5 mM MgSO₄, 5 mM KCl, 0.5 mM NADP, and 40 µL enzyme extract and the reaction was initiated with 3 mM glucose-6-phosphate (G6P). NAD-IDH activity was measured in a 1-mL mixture containing 100 mM HEPES-KOH pH 7.5, 5 mM MnCl₂, 2 mM NAD, and 150 µL of sample. The reaction was initiated with 5 mM isocitrate. NADP-IDH was assayed in a 1-mL mixture containing 100 mM Bicine-KOH pH 8.0, 5 mM MgSO₄, 5 mM KCl, 0.5 mM NADP, and 50 µL of enzyme extract. The reaction was initiated with 5 mM isocitrate.

Pyruvate kinase (PK) and PEP phosphatase were assayed according to Plaxton (1990) with some modifications. Pyruvate kinase was determined in a 1-mL reaction containing 100 mM imidazole-HCl pH 7.0, 50 mM KCl, 10 mM MgCl₂, 0.05% BSA, 2 mM DTT, 150 µM NADH, 1 unit lactate dehydrogenase, 2 mM ADP, and 100 µL enzyme extract. The reaction was initiated with 2 mM PEP. PEP phosphatase was determined under the same conditions with the omission of ADP, and PK activity was corrected for PEP phosphatase activity. PEP carboxylase was determined according to Stitt et al. (1989) in a 250 µL mixture containing 100 mM Bicine-KOH pH 8.0, 2 mM [¹⁴C]K₂CO₃, 10 mM MgSO₄, 350 µM NADH, 10 units of malate dehydrogenase, and 2 mM PEP. The reaction was initiated with 25 µL enzyme extract and stopped after 15 min with the addition of 500 µL of 0.5 N HCl. The solution was dried overnight at 50 to 70 °C and redissolved in 200 µL H₂O and 3 mL of scintillation fluid.

**EXTRACTION AND ASSAY OF METABOLITES.** Metabolites were extracted from the leaves according to Leegood (1993). Eight discs were ground with a precooled mortar and pestle in 1.5 mL of ice-cold 4% (v/v) HClO₄. Extract was added to a microcentrifuge tube containing 50 mg HCl-washed activated charcoal. The mortar and pestle was rinsed with 0.5 mL of 4% (v/v) HClO₄ and added to the same tube. The extract sat on ice for 30 min and then was centrifuged at 15,000 g for 10 min. A 1.5-mL aliquot of the supernatant was removed and neutralized to pH 6 with 200 µL of 5 M KOH in 1 M triethanolamine. Sample was mixed with 50 µg PVPP and then allowed to sit on ice for 15 min to precipitate KCIO₄ and remove any remaining phenolics. Extract was centrifuged at 15,000 g for 10 min, and the supernatant was stored at −80 °C until analysis.

Before metabolite analysis, samples were thawed and centrifuged at 15,000 g for 10 min to remove any remaining KCIO₄. Fructose-6-phosphate (F6P), G6P, and 3-phosphoglycerate (PGA) concentration was determined according to Leegood (1993). G6P was measured in a 1-mL mixture containing 100 mM TRIS-HCl pH 8.0, 1 mM EDTA, 5 mM MgSO₄, 5 mM KCl, 500 µM NAD, 1 unit G6PDH, and 200 µL of sample. F6P was determined in the same cuvette with the addition of 1 unit of phosphoglucose isomerase. PGA was quantified in a 1-mL mixture containing 50 mM imidazole-HCl pH 7.0, 2 mM MgCl₂, 5 mM ATP, 20 mM NaCl, 2 mM β-mercaptoethanol, 200 µM NADH, 5 units gyceraldehyde-3-phosphate dehydrogenase, 5 units of PGA kinase, and 100 µL of extract. Citrate and malate were assayed as previously described. Unless otherwise indicated, all enzyme and metabolites were measured against a no-sample blank, and reactions containing sample were also run without substrate when necessary.

**STATISTICAL ANALYSIS.** Analysis of variance was performed using JMP (version 5.1; SAS Institute, Cary, NC), and regressions were performed using SigmaPlot (version 9.0; Systat Software, Point Richmond, CA). Open circles in the figures reflect data collected at noon, and black circles represent data collected at midnight. In situations in which data collection time was not significant (P > 0.05), treatments were grouped together and are indicated by gray circles. Error bars represent the standard error of the mean.

**Results.**

For the greenhouse experiment, vines grown at the lowest pH treatment were healthy and green, and an increase in root-medium pH decreased the concentration of Chl, total Fe, and active Fe in the leaves (Table 1). Leaf total Fe at the highest pH was approximately 50% lower than the amount found in vines grown at the lowest pH, whereas Chl and active Fe concentrations were 84% lower. As pH increased from 5.8 to 7.5, root FCR activity increased 10-fold, whereas leaf FCR activity decreased 10-fold (Table 1).

In the second experiment, as root-medium pH increased from 5.1 to 7.5, both the total leaf area and the total foliar dry weight per vine decreased curvilinearly (Fig. 1A). The total and active Fe content of recently matured leaves also decreased curvilinearly as pH rose with a greater decrease in active Fe compared with total Fe (Fig. 2A–B). Active Fe content is often a better indicator of foliar Fe nutritional status (Gezgin and Er, 2001; Oserkowsky, 1933; Smith and Cheng, 2005), and the remaining foliar data are presented in relation to leaf active Fe content.

As leaf active Fe content decreased, the area of the recently matured leaf selected for biochemical analysis also decreased (Fig. 3A). The Chl content of fresh leaves was highly correlated to active Fe content (R² = 0.88; Fig. 3B) and decreased as active Fe decreased. Protein decreased linearly as active Fe decreased (Fig. 3C).

At midday, CO₂ assimilation, gs, and PSII quantum efficiency decreased from maximum levels in leaves with a high active Fe content to very low levels in leaves with the lowest active Fe content (Fig. 4A, B, and D). Internal CO₂ increased curvilinearly as active Fe decreased (Fig. 4C). The maximum PSII efficiency of dark-adapted leaves was decreased only at

| pH   | Chl (mg g⁻¹) | Total Fe (µg g⁻¹) | Active Fe (µg g⁻¹) | Root FCR (nmol g⁻¹ min⁻¹) | Leaf FCR (nmol g⁻¹ min⁻¹) |
|------|--------------|-------------------|-------------------|--------------------------|---------------------------|
| 5.8  | 1.9 ± 0.1    | 93.7 ± 7.3        | 22.5 ± 1.7        | 0.8 ± 0.3                | 9.0 ± 1.9                  |
| 6.8  | 1.3 ± 0.1    | 59.4 ± 6.3        | 10.5 ± 0.6        | 2.8 ± 1.1                | 2.9 ± 0.6                  |
| 7.5  | 0.3 ± 0.1    | 46.0 ± 4.3        | 3.4 ± 0.9         | 8.1 ± 1.0                | 0.9 ± 0.5                  |

P** Analysis of variance results: P < 0.01 and 0.001, respectively.

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the lowest active Fe content, and night respiration rate decreased curvilinearly as active Fe decreased (Fig. 4E–F).

Activity of G6PDH, aconitase, and both NAD(P) isoforms of IDH and malic enzyme was greater at midnight vs. midday (Fig. 5A–D, F, and G), whereas sampling time did not affect the activity of NAD-MDH, PEP carboxylase, PEP phosphatase, or PK (Fig. 5E and H–J). With the exception of aconitase, all other enzymes responded similarly to active Fe. Enzyme activity increased as active Fe decreased below 1.0 mg/C1m–2, whereas above 1.0 mg/C1m–2, activity did not change or increased slightly (Fig. 5A and C–J). On the other hand, aconitase activity, and the difference between day and night measurements, decreased as active Fe decreased (Fig. 5B).

G6P, F6P, and PGA decreased curvilinearly as active Fe decreased; G6P content was greater at night, whereas F6P and PGA content was higher during the day (Fig. 6A–D, F, and G), whereas sampling time did not affect the activity of NAD-MDH, PEP carboxylase, PEP phosphatase, or PK (Fig. 5E and H–J). With the exception of aconitase, all other enzymes responded similarly to active Fe. Enzyme activity increased as active Fe decreased below 1.0 mg/C1m–2, whereas above 1.0 mg/C1m–2, activity did not change or increased slightly (Fig. 5A and C–J). On the other hand, aconitase activity, and the difference between day and night measurements, decreased as active Fe decreased (Fig. 5B).

In both experiments, total Fe and active Fe was lower in leaves with lime-induced chlorosis (Table 1; Fig. 2A–B). Previously, we reported that chlorotic leaves from ‘Concord’ vines can contain as much or more total Fe than green leaves and that active Fe provides a better relationship with Chl (Smith and Cheng, 2005, 2006). It is possible that the reason we observed a decrease in foliar total Fe in this study, but not in the other experiments, is related to a treatment effect. The treatment in this study was CaCO3, or root-medium pH, and we did not supply any chelated Fe, whereas in previous studies, we applied different rates of Fe at a given pH using the synthetic chelate Fe(III)-EDDHA [ferric ethylenediamine di-(α-hydroxyphenylacetic) acid]. In the current experiments, although leaf total Fe content did decrease in the chlorotic leaves, active Fe decreased to a greater
rates obtained for Fe-efficient grapevine and kiwifruit [Actinidia deliciosa (A.Chev.) C.F. Liang & A.R. Ferguson] cultivars (BavareSCO et al., 1991; Ksouri et al., 2006; Rombolà et al., 2002). Ksouri et al. (2006) found that root FCR activity did not increase in chlorosis-susceptible ‘Balta4’ grapevines in response to bicarbonate-induced chlorosis, but did increase three- to fourfold in the chlorosis-resistant ‘Khamri’. From comparison with these data, it appears that V. labruscana roots are Fe-efficient. Yet leaves from vines grown at high pH were severely chlorotic (Table 1) and this alone confirms that V. labruscana is not Fe-efficient. Our results for root and shoot FCR activity in ‘Concord’ grapevines suggest that lime-induced chlorosis may be the result of impaired Fe assimilation from decreased foliar FCR activity rather than from decreased Fe uptake at the root level. Further use of the leaf disc assay is needed to compare FCR activity between chlorosis-susceptible and chlorosis-resistant grapevine genotypes.

In addition to reducing total dry weight and leaf area (Fig. 1A–B), a decrease in active Fe corresponded to smaller leaves that contained less Chl and protein (Fig. 3A–C). The decrease in active Fe also resulted in a curvilinear decrease in CO₂ assimilation, gs, and PSII quantum efficiency and a curvilinear increase in the calculated intercellular CO₂ concentration (Fig. 4A–D). These trends are similar to those observed in previous experiments in which vines were treated with different rates of Fe(III)-EDDHA (Chen et al., 2004; Smith and Cheng, 2005, 2006), indicating that physiological responses to Fe deficiency from a lack of applied Fe (so-called “direct” or “true” Fe deficiency) are similar to those resulting from lime-induced (or indirect) Fe deficiency.

The decrease in night respiration, from –1.05 μmol·m⁻²·s⁻¹ at the highest active Fe content to –0.72 μmol·m⁻²·s⁻¹ at the lowest active Fe content (Fig. 4F), contrasts with most of the previous reports, which indicate that dark respiration is unaffected by Fe deficiency (Morales et al., 1998; Pushnik and Miller, 1989; Terry, 1983). Considering that Fe is a cofactor for aconitase (conversion of citrate to isocitrate) and is a key component of the mitochondrial electron transfer chain (Fe-S centers in complexes I, II, and III; heme prosthetic groups in cytochromes), it seems remarkable that Fe deficiency would not significantly impact respiration rates.

Pascal and Douce (1993) did find that Fe limitation decreased the maximum O₂ consumption in sycamore cells (Acer pseudoplatanus L.), and they concluded that this was most likely from a decrease in the formation of Fe-S centers in complex II (succinate:ubiquinone oxidoreductase). It is possible that nonphosphorylating bypasses such as rotenone-insensitive NAD(P)H dehydrogenase and alternative oxidase may play a greater role in mitochondrial electron transport under Fe deficiency. Although the function of the alternative oxidase (AOX) is not well understood, it has been shown to increase under nutrient stress, and when the rate of the tricarboxylic acid (TCA) cycle decreases (citrate and pyruvate activate AOX) (Moore et al., 2002; Siedow and Day, 2000). Electron flow through these alternative routes may help maintain (or increase) respiratory rates when the cytochrome pathway is inhibited or when respiratory substrates accumulate (Moore et al., 2002; Siedow and Day, 2000).

In addition, Pascal and Douce (1993) measured a 50% decrease in mitochondrial aconitase activity under Fe deficiency and found that this corresponded to impaired citrate oxidation. Early on, Bacon et al. (1961) proposed that Fe
deficiency would decrease aconitase activity and result in citrate accumulation, yet recently there has been mixed support for this hypothesis (de Vos et al., 1986; López-Millán et al., 2001a; McCluskey et al., 2004; Pich and Scholz, 1993). In fact, López-Millán et al. (2001a) measured a 1.7-fold increase in aconitase activity in Fe-deficient sugar beet (*Beta vulgaris* L.) leaves. We found a strong, linear relationship between active Fe and aconitase activity (Fig. 3B), and leaves with low active Fe had three- to fourfold lower rates of aconitase activity compared with leaves with the highest active Fe. We were not able to obtain linear slopes for aconitase using citrate as a substrate and measuring either cis-aconitate formation at 240 nm or NADP⁺ reduction at 340 nm (coupled with IDH) as described by de Vos et al. (1986). Nor were we able to successfully measure activity in the reverse direction using isocitrate as a substrate in aconitase (Bacon et al., 1961). We were able to obtain linear slopes using cis-aconitate as substrate and measuring NADP⁺ reduction at 340 nm in a coupled assay with IDH as described by Jenner et al. (2001). We are unsure why stable measurements could only be made using cis-aconitate, but suggest that aconitase determination in relation to Fe deficiency warrants further investigation.

A decrease in aconitase activity in response to lower active Fe corresponded to an increase in citrate content; however, there was no difference in foliar citrate content between day and night, although aconitase activity was higher in leaves sampled at night (Figs. 5B and 6E). The lack of a relationship between the diurnal changes in enzyme and substrate is unclear, but it is reasonable to assume that the accumulation of citrate in Fe-deficient leaves was the result of lower leaf aconitase activity. Citrate concentration in the xylem sap increased as soil pH increased (and leaf active Fe decreased); thus, loading of citrate in the xylem sap from the roots most likely did not contribute to high levels of citrate in the leaves in contrast to the proposal by López-Millán et al. (2001a).

In recent reviews, both Schmidt (1999) and Abadí et al. (2002) concluded that organic acid accumulation in Fe-deficient tissues was primarily from the increased activity of PEP carboxylase (particularly at the root level) and not related to lower aconitase activity. We found that foliar PEP carboxylase activity did increase as active Fe decreased (Fig. 5I); however, malate content did not follow the same trend (Fig. 6D). Both citrate content and PEP carboxylase activity did increase as active Fe decreased, but with the exception of aconitase, the activity of all of the other enzymes we measured also increased under Fe deficiency (Fig. 5A–J). Therefore, we agree with Bacon et al. (1961) and Pascal and Douce (1993) that Fe deficiency decreases aconitase activity in the leaves and that the accumulation of foliar citrate is the result of impaired citrate oxidation in the TCA cycle.

We suggest that the increased activity of the other enzymes we measured is a response to lower aconitase activity, helps funnel carbon into the mitochondria, and increases the capacity to reduce pyridine nucleotides as the TCA cycle slows. In C3 plants, PEP carboxylase activity, and the import of malate and OAA into the mitochondria, functions primarily in an anaplerotic role to replenish TCA cycle intermediates exported for ammonia assimilation and amino acid synthesis (Plaxton, 1996; Siedow and Day, 2000). In addition, the observed increase of NADP-malic enzyme, NADP-IDH, and G6PDH activity under Fe deficiency would increase the NADPH needed for biosynthetic reaction. It is also possible that the increase in G6PDH activity (and flux through the pentose-phosphate pathway) may not only provide NADPH, but may also bypass phosphofructokinase (PFK) and supply triose phosphates for conversion to PEP when PFK is inhibited by citrate (Givan, 1999).

In this study, we found that PGA, G6P, and F6P decreased as active Fe decreased, and the difference between day and night content of PGA and G6P also decreased, indicating that Fe-deficient leaves are carbon-limited. Interestingly, in previous

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**Fig. 4.** (A) CO₂ assimilation rate, (B) gs, (C) calculated intercellular CO₂ concentration, and (D) PSII quantum efficiency of leaves sampled at noon, and (E) maximum photochemical efficiency (Fv/Fm) and (F) respiration rate of leaves sampled at midnight (n = 5) in relation to foliar active Fe content of 'Concord' grapevines grown at five different pHs. Regression equations for (A) \( y = -5.20 + 33.81x - 10.58x^2 \) \( R^2 = 0.86, P < 0.001 \), (B) \( y = -87.23 + 948.67x - 316.48x^2 \) \( R^2 = 0.74, P < 0.001 \), (C) \( y = 348.25 - 137.15x + 46.02x^2 \) \( R^2 = 0.88, P < 0.001 \), (D) \( y = 0.07 + 0.29x - 0.08x^2 \) \( R^2 = 0.82, P < 0.001 \), and (F) \( y = -0.62 - 0.44x + 0.10x^2 \) \( R^2 = 0.56, P < 0.001 \). Statistical significance for (E) was determined by analysis of variance \( P < 0.001 \); data are means ± se.
Fig. 5. Activity of (A) glucose-6-phosphate dehydrogenase (G6PDH), (B) aconitase, (C) NAD-isocitrate dehydrogenase (NAD-IDH), (D) NAD-isocitrate dehydrogenase (NAD-IDH), (E) NAD-malate dehydrogenase (NAD-MDH), (F) NAD-malic enzyme, (G) NAD-malic enzyme, (H) phosphoenolpyruvate carboxylase (PEPC), (I) phosphoenolpyruvate phosphatase (PEPP), and (J) pyruvate kinase (PK) in relation to foliar active Fe content of 'Concord' grapevines grown at five different pHs (n = 5). Open circles in the figures reflect data collected at noon, and black circles represent data collected at midnight. In situations in which time of day was not significant (P > 0.05) using analysis of variance, treatments were grouped together and are indicated by gray circles (n = 10). Regression equations for (A) noon: y = 66.55–73.94x + 27.45x² (R² = 0.71, P < 0.001), midnight: y = 90.15–104.75x + 39.57x² (R² = 0.62, P < 0.001); (B) noon: y = 1.49 + 3.62x (R² = 0.57, P < 0.001), midnight: y = 1.72 + 6.67x (R² = 0.62, P < 0.001); (C) noon: y = 17.10–18.89x + 6.76x² (R² = 0.73, P < 0.001), midnight: y = 90.15–104.75x + 39.57x² (R² = 0.66, P < 0.001); (D) noon: y = 2.70–3.00x (R² = 0.56, P < 0.001), midnight: y = 29.32–18.14x (R² = 0.58, P < 0.001); (E) noon: y = 17.10–18.89x + 6.76x² (R² = 0.73, P < 0.001), midnight: y = 90.15–104.75x + 39.57x² (R² = 0.66, P < 0.001); (G) noon: y = 3.39–4.41x + 1.79x² (R² = 0.72, P < 0.001). Statistical significance for (D) NADP-MDH (P < 0.001), (E) NAD-MDH (P < 0.001), and (F) NADP-malic enzyme (P < 0.001) was determined by analysis of variance; data are means ± se.

In conclusion, root FCR activity is enhanced, whereas leaf FCR activity

work with 'Concord' grapevines grown with different rates of Fe-EDDHA, only PGA content decreased as active Fe decreased, whereas G6P and F6P content did not change (Chen et al., 2004). Espen et al. (2000) found elevated G6P and F6P levels in Fe-deficient cucumber (Cucumis sativus L.) roots, but there are little data for foliar hexose phosphate content in relation to Fe nutrition. Malate content in the leaves followed a similar pattern to the concentration in the xylem sap, and it is possible that root export of malate contributed to foliar malate levels more than PEP carboxylase activity in the leaves (Figs. 6D and 7D).

Contrary to what we hypothesized, xylem sap pH did not change in response to CaCO₃ application (Fig. 7A). It is possible that root PEP carboxylase may have fixed any HCO₃⁻ taken up by the roots (and regulated the pH), but citrate and malate content in the xylem sap does not reflect this assumption. It is interesting that citrate concentration in the xylem exudate was much lower than malate, a trend that was also similar in the leaves (citrate was an order of magnitude lower than malate), because Fe efficiency is often associated with high citrate levels (Brown and Ambler, 1970; Fournier et al., 1992). We could not find data to compare foliar citrate levels in Vitis spp.; however, Ollat et al. (2003) recently demonstrated the link between Fe efficiency and citrate in grapevine roots. When grown in Fe-deficient (0.5 μM Fe-EDTA) vs. Fe-sufficient (10 μM Fe-EDTA) conditions, citrate levels in root tips of high pH-tolerant Vitis berlandieri Planch. × V. vinifera ‘Fercal’ were 9.8 and 1.0 mg g⁻¹, respectively, whereas in the high pH-susceptible Vitis rupestris Scheele × Vitis riparia Michx. ‘101–14’, citrate was 0.7 and 0.4 mg g⁻¹, respectively. In the Fe-sufficient sugar beet leaves, malate and citrate were 0.9 and 0.8 mmol-m⁻² and increased to 6.1 and 9.9 mmol-m⁻² under Fe deficiency, respectively (López-Millán et al., 2001a). Thus, it is possible that lower citrate levels in 'Concord' vines are related to its susceptibility to lime-induced chlorosis.

In conclusion, root FCR activity is enhanced, whereas leaf FCR activity
is decreased under high pH-induced Fe deficiency in ‘Concord’ vines. Xylem sap pH did not change and does not appear to be related to the observed soil pH effects. The decreased leaf aconitase activity under Fe deficiency is the most likely cause of the increase in citrate levels. Greater activity of the other glycolytic and TCA enzymes under Fe deficiency may help to funnel carbon into the mitochondria and enhance NAD(P) reduction. Compared with other species, Concord vines have much lower levels of citrate (and the citrate:malate ratios) in the xylem exudate and leaves, which may be linked to Fe inefficiency. Reciprocal grafting experiments between *V. labruscana* and an Fe-efficient genotype will help determine whether the limiting factor in Fe assimilation at high pH is primarily at the root or leaf level.
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