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# Photorespiration and nitrate assimilation: a major intersection between plant carbon and nitrogen

#### **Abstract**

 $C_3$  carbon fixation has a bad reputation, primarily because it is associated with photorespiration, a biochemical pathway thought to waste a substantial amount of the carbohydrate produced in a plant. This review presents evidence collected over nearly a century that (1) Rubisco when associated with  $Mn^{2+}$  generates additional reductant during photorespiration, (2) this reductant participates in the assimilation of nitrate into protein, and (3) this nitrate assimilation facilitates the use of a nitrogen source that other organisms tend to avoid. This phenomenon explains the continued dominance of  $C_3$  plants during the past 23 million years of low  $CO_2$  atmospheres as well as the decline in plant protein concentrations as atmospheric  $CO_2$  rises.

# **Keywords**

photorespiration, C<sub>3</sub> carbon fixation, nitrate assimilation, photosynthesis, plant evolution, nitrogen sources

#### **Premise**

Plants, by most accounts, convert less than 6% of the incoming solar energy into useable chemical energy (Hall et al. 1999; Zhu et al. 2008). Efforts to improve this conversion rate have focused on the light-independent reactions of photosynthesis (e.g., Parry et al. 2013; Studer et al. 2014; Whitney et al. 2011; Zhu et al. 2010). "The light reactions are highly efficient, converting as much as 40% - 50% of the captured solar energy into energy carriers. The dark reactions are not developed for energy efficiency and it is here the energy is...lost" (Swedish Energy Agency 2003). In particular, Rubisco (ribulose-1,5-bisphosphate carboxylase/oxygenase; EC 4.1.1.39), the enzyme which catalyzes the first reaction of the  $C_3$  pathway and constitutes about half of the protein in leaves (Parry et al. 2003), has been identified as a target of opportunity.

# **Competing Reactions**

Rubisco exhibits opposing tendencies in that it catalyzes two different chemical reactions: one reaction combines a five-carbon sugar RuBP (ribulose-1,5-bisphosphate) with CO<sub>2</sub> (carboxylation), and the other reaction combines this same sugar with O<sub>2</sub> (oxygenation).

- The carboxylation reaction of RuBP produces a six-carbon compound that quickly divides into two molecules of a three-carbon compound, PGA (3-phosphoglycerate), hence the name *C₃ carbon fixation*. Six of these PGA molecules pass through an elaborate pathway that expends the energy of 18 ATP and 12 NADPH molecules, forms one molecule of fructose-6-phosphate, a six-carbon sugar, and regenerates six molecules of RuBP.
- The oxygenation reaction splits the RuBP into one molecule of a three-carbon PGA and one molecule of a two-carbon PG (2-phosphoglycolate), hence the name  $C_2$  pathway or, more commonly, photorespiration (Foyer et al. 2009). In total, photorespiration consumes 3.5 ATP and 2 NADPH per RuBP oxygenated and regenerated, but does not result in any net production of sugar (Bauwe et al. 2010; Tolbert 1994). Thus photorespiration seems to be largely a superfluous process, one thought to dissipate 76.3 kcal mol<sup>-1</sup> as waste heat (Frank et al. 2000).

The balance between  $C_3$  carbon fixation and photorespiration depends on the relative amounts of  $CO_2$  and  $O_2$  entering the active site of Rubisco and the specificity of the enzyme for each gas. Atmospheric concentrations of  $CO_2$  and  $O_2$  are currently 0.04% and 20.94%, respectively, yielding a  $CO_2$ : $O_2$  ratio of 0.0019. Gaseous  $CO_2$ , however, is much more soluble in water than  $O_2$ , and so the  $CO_2$ : $O_2$  ratio near the chloroplast, the part of a cell where these reactions occur, is about 0.026 at 25°C. Rubisco has about a 50-fold (cyanobacteria) to 100-fold (higher plants) greater specificity for  $CO_2$  than  $O_2$  (Galmes et al. 2005). Together, because of the relative concentrations of and specificity for  $CO_2$  over  $O_2$ , Rubisco catalyzes about two to three cycles of  $C_3$  carbon fixation for every cycle of photorespiration under current atmospheres (Sharkey 1988). Conditions that inhibit photorespiration—namely, high  $CO_2$  or low  $O_2$  atmospheric concentrations—stimulate carbon fixation in the short term by about 35%.

Temperature influences the balance between  $C_3$  carbon fixation and photorespiration in two ways. First, as temperature rises, the solubility of  $CO_2$  in water decreases more than the solubility of  $O_2$ , resulting in a lower  $CO_2$ : $O_2$  ratio. Second, the enzymatic properties of Rubisco shift with increasing temperature, stimulating the reaction with  $O_2$  to a greater degree than the one with  $CO_2$ . Warmer temperatures, therefore, favor photorespiration over  $C_3$  carbon fixation, and photosynthetic conversion of absorbed light into sugars becomes less efficient (Ehleringer et al. 1997). Based on the temperature response of Rubisco carboxylation

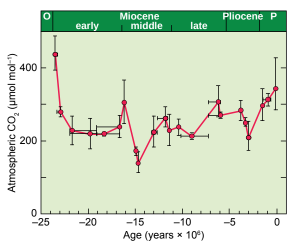
and oxygenation,  $C_4$  plants should be more competitive in regions where the mean monthly air temperature exceeds 22°C (Collatz et al. 1998).

Overall, Rubisco seems a vestige of the high  $CO_2$  and low  $O_2$  atmospheres under which plants first evolved (Wingler et al. 2000). To compensate for the shortcomings of Rubisco, some plants employ  $CO_2$  pumping mechanisms such as  $C_4$  carbon fixation that elevate  $CO_2$  concentrations at the active site of the enzyme. The  $C_4$  pathway is one of the most convergent evolutionary adaptations in life with at least 66 independent origins (Sage et al. 2012). Extensive efforts are underway to emulate Mother Nature and transfer the  $C_4$  pathway into rice and other  $C_3$  crops (von Caemmerer et al. 2012).

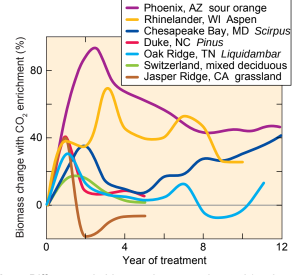
Several observations, however, are inconsistent with the presumption that Rubisco is poorly suited to modern times.

- Earth's atmosphere has contained relatively low  $CO_2$  concentrations (lower than 0.04%) for the past 23 million years (Figure 1). During this period, the plant kingdom experienced major changes including the diversification of modern graminoids, especially grasses and sedges, and the appearance of many new  $C_4$  species, especially when  $CO_2$  concentrations fell below 0.02%, (Sage et al. 2012). In a relatively short period of time (6 or 7 million years) (Osborne and Beerling 2006), the kinetics of Rubisco diverged between  $C_3$  and  $C_4$  plants (Studer et al. 2014). Rubisco in  $C_4$  plants operates under elevated  $CO_2$  conditions, and so the  $C_4$  enzyme has traded a lower specificity for  $CO_2$  relative to  $O_2$  ( $S_{c/o}$ ) for a higher catalytic efficiency ( $k_{cat}$ ) (Galmes et al. 2005; Sage 2002). Surprisingly, the kinetic properties of Rubisco do not differ greatly among higher  $C_3$  plants (Kane et al. 1994; Tcherkez et al. 2006). Thus, the kinetic properties of Rubisco were able to change when a species adopted the  $C_4$  pathway, but such changes were not warranted in  $C_3$  plants because Rubisco may already be "nearly perfectly optimized" for  $C_3$  carbon fixation (Tcherkez et al. 2006).
- Despite 23 million years of low atmospheric CO<sub>2</sub> concentrations, 96% of plant species depend solely on the C<sub>3</sub> carbon fixation pathway (Sage et al. 1999). C<sub>3</sub> species account for over 94% of the Earth's biomass (Still et al. 2003). Species using other carbon fixation pathways are dominant only in hot and dry environments.
- The response of C<sub>3</sub> species to elevated CO<sub>2</sub> atmospheres is highly variable and often depends on plant N status (Cavagnaro et al. 2011; Duval et al. 2012; Finzi et al. 2007; Norby et al. 2010; Reich et al. 2006). Initially, elevated CO<sub>2</sub> stimulates biomass accumulation by about 35% (Figure 2). This stimulation, however, tends to abate upon longer exposures in conjunction with a decline in plant protein concentrations (Cotrufo et al. 1998; Long et al. 2004).

Explanations for the decline in plant protein concentrations at elevated  $CO_2$  include: (*a*) plants under elevated  $CO_2$  grow larger, diluting the protein within their tissues (Ellsworth et al. 2004; Taub and Wang 2008); (*b*) carbohydrates accumulate within leaves, down-regulating the amount of the most prevalent protein Rubisco (Long et al. 2004); (*c*) carbon enrichment of the rhizosphere leads to progressively greater limitations in the soil N available to plants (Reich et al. 2006); and (*d*) elevated  $CO_2$  directly inhibits plant N metabolism, especially the assimilation of  $NO_3^-$  into proteins in shoots of  $C_3$  plants (Bloom et al. 2012b). Recently, several independent meta-analyses conclude that this last explanation is the one most consistent with observations from hundreds of studies (Cheng et al. 2012; Myers et al. 2014; Pleijel and Uddling 2012).



**Fig. 1** A reconstruction of atmospheric  $CO_2$  concentrations based on boron isotope ratios of ancient planktonic foraminifer shells. (Data from Pearson and Palmer 2000)

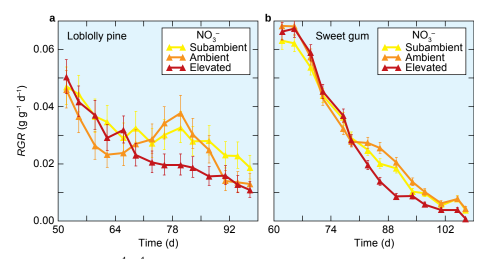


**Fig. 2** Differences in biomass between elevated ( $\approx$  567 ppm) and ambient ( $\approx$  365 ppm) atmospheric CO<sub>2</sub> after years of treatment. Shown are the data from seven different studies using the designated types of plants. (Data from Dukes et al. 2005; Kimball et al. 2007; Korner 2006; Norby et al. 2010; Rasse et al. 2005; Talhelm et al. 2014).

## CO<sub>2</sub> inhibits NO<sub>3</sub>- Assimilation

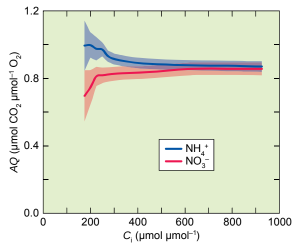
Many independent methods for estimating  $NO_3^-$  assimilation confirm that elevated  $CO_2$  inhibits shoot  $NO_3^-$  assimilation in  $C_3$  plants. These methods include:

- 1. <sup>15</sup>N-labeling. Plants grown on  $NO_3^-$  containing N isotopes at natural abundance levels ( $\approx 0.366\%$  <sup>15</sup>N) were exposed to a pulse of  $NO_3^-$  that was heavily enriched in <sup>15</sup>N. The difference between the <sup>15</sup>N enrichment of total N and that of free  $NO_3^-$  provided an estimate of <sup>15</sup>N-NO<sub>3</sub><sup>-</sup> assimilation, which decreased under CO<sub>2</sub> enrichment (Bloom et al. 2010).
- 2. <sup>14</sup>N-labeling. Plants grown on 99.9% enriched <sup>15</sup>N-NO<sub>3</sub><sup>-</sup> were exposed to a pulse of NO<sub>3</sub><sup>-</sup> containing N isotopes at natural abundance levels ( $\approx 0.366\%$  <sup>15</sup>N); the difference between the <sup>14</sup>N enrichment of total N and that of free NO<sub>3</sub><sup>-</sup> provided an estimate of <sup>14</sup>N-NO<sub>3</sub><sup>-</sup> assimilation, which decreased under CO<sub>2</sub> enrichment (Bloom et al. 2010).
- 3. **Organic N accumulation.** Accumulation of organic N was followed in plants receiving NO<sub>3</sub><sup>-</sup> as a sole N source, and this accumulation decreased under CO<sub>2</sub> enrichment (Aranjuelo et al. 2013; Bloom et al. 2010; Lekshmy et al. 2013; Pleijel and Uddling 2012; Rachmilevitch et al. 2004).
- **4. NO**<sub>3</sub><sup>-</sup> **depletion from a medium.** The decline of NO<sub>3</sub><sup>-</sup> concentrations in a nutrient solution was monitored to calculate net plant NO<sub>3</sub><sup>-</sup> absorption. The difference between this NO<sub>3</sub><sup>-</sup> absorption and the accumulation of free NO<sub>3</sub><sup>-</sup> within plant tissues estimated plant NO<sub>3</sub><sup>-</sup> assimilation, which decreased under CO<sub>2</sub> enrichment (Bloom et al. 2010; Rachmilevitch et al. 2004).
- **Plant growth.** C<sub>3</sub> species received either NO<sub>3</sub> or NH<sub>4</sub> as their sole N source. CO<sub>2</sub> enrichment decreased growth of plants receiving NO<sub>3</sub> (Figure 3) but increased growth of those receiving NH<sub>4</sub> (Bloom et al. 2012b; Bloom et al. 2002; Carlisle et al. 2012).

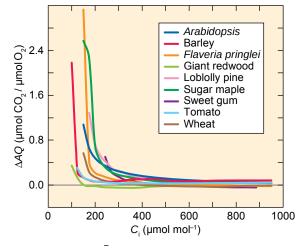


**Fig. 3** Relative growth rate in g g<sup>-1</sup> d<sup>-1</sup> of **(a)** loblolly pine *Pinus taeda* and **(b)** sweet gum *Liquidambar styraciflua* receiving  $NO_3^-$  nutrition in controlled environment chambers at subambient  $CO_2$  (310 µmol mol<sup>-1</sup>, the level of about 50 years ago), ambient  $CO_2$  (400 µmol mol<sup>-1</sup>, current level), or elevated  $CO_2$  (720 µmol mol<sup>-1</sup>, the level anticipated in about 50 years).  $CO_2$  concentration had no significant effect on the growth of plants receiving  $NH_4^+$  nutrition (data not shown). Time is in days after transplanting to a hydroponic solution. Shown are the predicted values and standard errors from mixed linear models with repeated measures on 6 to 10 individual plants. (Bloom et al. 2012b)

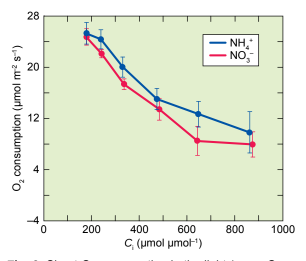
- **Isotopic discrimination by NO**<sub>3</sub><sup>-</sup> **reductase.** Plants were grown under NO<sub>3</sub><sup>-</sup> containing N isotopes at natural abundance levels ( $\approx 0.366\%^{15}$ N). Under CO<sub>2</sub> enrichment, plant tissues became less enriched in <sup>15</sup>N-organic N compounds presumably because (*a*) CO<sub>2</sub> inhibited shoot NO<sub>3</sub><sup>-</sup> assimilation, (*b*) NO<sub>3</sub><sup>-</sup> availability became less limiting to assimilation, (*c*) NO<sub>3</sub><sup>-</sup> reductase discriminated more against <sup>15</sup>N-NO<sub>3</sub><sup>-</sup>, and (*d*) shoots assimilated relatively less <sup>15</sup>N-NO<sub>3</sub><sup>-</sup> (Bloom et al. 2010; Bloom et al. 2014).
- 7.  $\Delta AQ$ . Assimilatory quotient (AQ), the ratio of net  $CO_2$  consumption to net  $O_2$  evolution from shoots was measured in a plant receiving  $NH_4^+$  or  $NO_3^-$  as its sole N source (Figure 4); AQ decreased as  $NO_3^-$  assimilation increased because additional electrons generated from the light-dependent reactions of photosynthesis were transferred first to  $NO_3^-$  and then to  $NO_2^-$ . This stimulated net  $O_2$  evolution, but had



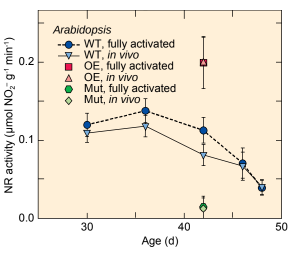
**Fig. 4** Shoot AQ (net  $CO_2$  consumed/net  $O_2$  evolved) as a function of internal  $CO_2$  concentrations ( $C_i$ ) for the 9  $C_3$  species in Figure 4 when they received  $NH_4^+$  or  $NO_3^-$  as a sole N source (mean  $\pm$  SE; solid  $\pm$  shaded area). (Bloom, unpublished data)



**Fig. 5** Shoot  $NO_3^-$  assimilation as a function of shoot internal  $CO_2$  concentration ( $C_i$ ) for 9  $C_3$  species. Shoot  $NO_3^-$  assimilation is assessed by  $\Delta AQ$  (change in the ratio of shoot  $CO_2$  consumption to  $O_2$  evolution with a shift from  $NO_3^-$  to  $NH_4^+$  nutrition). (Bloom et al. 2012b; Searles and Bloom 2003)



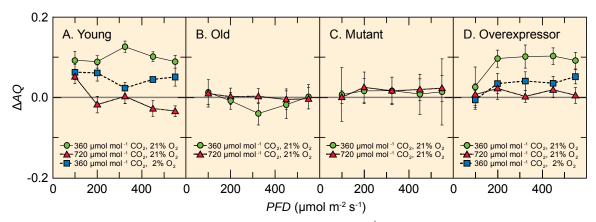
**Fig. 6** Shoot  $O_2$  consumption in the light (gross  $O_2$  – net  $O_2$ ) as a function of  $C_i$  for wheat receiving  $NH_4^+$  or  $NO_3^-$  as a sole N source. Shown are the means  $\pm$  SE for 5–7 replicates per treatment.(Cousins and Bloom 2004)



**Fig. 7** NO $_3$ <sup>-</sup> reductase activity (µmol of NO $_2$ <sup>-</sup> generated per g fresh mass per min) as a function of plant age (d) in leaves of a wild-type *A. thaliana* cv. Columbia (WT), a transgenic line harboring the chimeric gene *Lhch1\*3*::*Nia1\*2* (OE), and a genotype (*nia1 nia2*) with mutations in both structural genes for NO $_3$ <sup>-</sup> reductase (Mut). Because NO $_3$ <sup>-</sup> reductase is regulated through phosphorylation, leaf tissue was assayed under conditions that either dephosphorylated the enzyme (fully activated) or did not change its phosphorylation (*in vivo*). Shown are the mean  $\pm$  SE (n = 5–8 plants). (Rachmilevitch et al. 2004)

little effect on  $CO_2$  consumption; therefore, the change in AQ when a plant received  $NH_4^+$  instead of  $NO_3^-$  ( $\Delta AQ$ ) provided an estimate of shoot  $NO_3^-$  assimilation (Bloom et al. 1989; Bloom et al. 2002; Cen et al. 2001; Cramer and Myers 1948; Rachmilevitch et al. 2004; Van Niel et al. 1953; Warburg and Negelein 1920). In nine taxonomically diverse  $C_3$  species,  $\Delta AQ$  decreased as shoot internal  $CO_2$  increased (Figure 5).

- 8. **O**<sub>2</sub> **consumption.** Shoot O<sub>2</sub> consumption in the light was estimated from the difference between gross O<sub>2</sub> evolution via chlorophyll fluorescence and net O<sub>2</sub> evolution via an O<sub>2</sub> analyzer (Figure 6). At ambient CO<sub>2</sub>, O<sub>2</sub> consumption was lower when wheat plants received NO<sub>3</sub><sup>-</sup> rather than NH<sub>4</sub><sup>+</sup> because NO<sub>3</sub><sup>-</sup> and NO<sub>2</sub><sup>-</sup> were serving as electron acceptors. At elevated CO<sub>2</sub>, O<sub>2</sub> consumption was not significantly different under the two N sources presumably because NO<sub>3</sub><sup>-</sup> assimilation was negligible.
- 9. Altered NO<sub>3</sub> reductase capacity. Shoot CO<sub>2</sub> and O<sub>2</sub> fluxes at ambient and elevated CO<sub>2</sub> were contrasted between stages of plant development or genotypes that have greatly different NO<sub>3</sub> reductase activities *in situ*. In particular, we contrasted 36- vs. 48-d old wild-type Arabidopsis, Arabidopsis NO<sub>3</sub> reductase knockout mutants vs. transgenic Arabidopsis overexpressing NO<sub>3</sub> reductase (Figure 7), and NO<sub>3</sub> reductase-deficient barley mutants vs. wild-type barley. Δ*AQ* (change in the ratio of net CO<sub>2</sub> consumption to net O<sub>2</sub> evolution when a plant received NH<sub>4</sub> instead of NO<sub>3</sub> differed between these stages of development and genotypes under ambient CO<sub>2</sub>, but not under elevated CO<sub>2</sub> (Figure 8). This indicates that none of the stages of development or genotypes were assimilating NO<sub>3</sub> under elevated CO<sub>2</sub> (Bloom et al. 1989; Rachmilevitch et al. 2004).



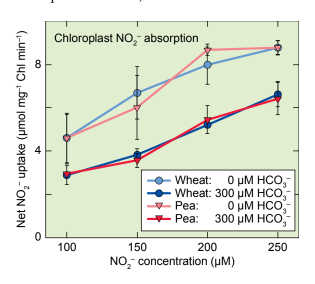
**Fig. 8** Changes in assimilatory quotient with the shift from NO<sub>3</sub><sup>-</sup> to NH<sub>4</sub><sup>+</sup> ( $\Delta AQ$ ) as a function of photosynthetic *PFD* (photon flux density) from shoots of *A. thaliana* cv. Columbia. (A) 36-day-old wild-type plants, (B) 48-d-old wild-type plants (C), genotype with null mutations, and (D) overexpressing line. The plants were grown under ambient CO<sub>2</sub> (360 μmol mol<sup>-1</sup>) and measured under ambient CO<sub>2</sub> and O<sub>2</sub> (360 μmol mol<sup>-1</sup> CO<sub>2</sub> and 21% O<sub>2</sub>; circles), elevated CO<sub>2</sub> (720 μmol mol<sup>-1</sup> CO<sub>2</sub> and 21% O<sub>2</sub>; triangles), or low O<sub>2</sub> (360 μmol mol<sup>-1</sup> CO<sub>2</sub> and 2% O<sub>2</sub>; squares). Shown are the mean ± SE, n = 5 - 8 plants. (Rachmilevitch et al. 2004)

**10.** NO<sub>3</sub> reductase activity. Maximum *in vitro* NO<sub>3</sub> reductase activity generally declined under CO<sub>2</sub> enrichment (Lekshmy et al. 2013; Matt et al. 2001). Presumably, this reflected slower NO<sub>3</sub> assimilation under CO<sub>2</sub> enrichment.

# **Physiological Mechanisms**

Three physiological mechanisms may be responsible for CO<sub>2</sub> inhibition of shoot NO<sub>3</sub> assimilation (Bloom et al. 2010).

• One mechanism is that elevated CO<sub>2</sub> inhibits nitrite (NO<sub>2</sub><sup>-</sup>) transport into chloroplasts (Figure 9). A chloroplast NO<sub>2</sub><sup>-</sup> transporter from higher plants has only recently been identified (Maeda et al. 2014), and so the nature of this inhibition has yet to be determined. Nevertheless, this mechanism can be independent of photosynthesis and, thus, is probably responsible for CO<sub>2</sub> inhibition of shoot NO<sub>3</sub><sup>-</sup> assimilation in Arabidopsis and wheat during the nighttime (Rubio-Asensio, Rachmilevitch, and Bloom, unpublished data).



**Fig. 9** Net  $NO_2^-$  uptake (µmol mg $^{-1}$  chlorophyll min $^{-1}$ ) by isolated chloroplasts as a function of  $NO_2^-$  concentration when the medium contained 0 (light symbols) or 0.3 (dark symbols) µM HCO $_3^-$ . Shown are the mean  $\pm$  SE (n = 3) for wheat (circles) and pea (inverted triangles). (Bloom et al. 2002)

- Another mechanism is that processes in the chloroplast stroma compete for reduced ferredoxin (Fd<sub>r</sub>). FNR (ferredoxin-NADP reductase) has a higher affinity for Fd<sub>r</sub> than NiR (nitrite reductase) (Knaff 1996), and so NO<sub>3</sub><sup>-</sup> assimilation proceeds only if the availability of Fd<sub>r</sub> exceeds that needed for NADPH formation (Backhausen et al. 2000; Robinson 1987). For most plants, this occurs when CO<sub>2</sub> availability limits C<sub>3</sub> carbon fixation (Bloom et al. 2010).
- A third mechanism involves photorespiration. Multiple lines of evidence link photorespiration with shoot NO<sub>3</sub><sup>-</sup> assimilation in C<sub>3</sub> plants. (*a*) Photorespiration stimulates the export of malate from chloroplasts (Backhausen et al. 1998; Taniguchi and Miyake 2012; Voss et al. 2013); this malate in the cytoplasm generates NADH (Igamberdiev et al. 2001; Taniguchi and Miyake 2012) that powers the first step of NO<sub>3</sub><sup>-</sup> assimilation, the reduction of NO<sub>3</sub><sup>-</sup> to NO<sub>2</sub><sup>-</sup> (Quesada et al. 2000; Rathnam 1978; Robinson 1987). (*b*) Conditions that decrease photorespiration—namely, elevated CO<sub>2</sub> and low O<sub>2</sub>—decrease shoot NO<sub>3</sub><sup>-</sup> reduction (Bloom et al. 2010; Rachmilevitch et al. 2004). (*c*) Mutants that alter malate transport or metabolism also alter both photorespiration and NO<sub>3</sub><sup>-</sup> assimilation (Dutilleul et al. 2005; Schneidereit et al. 2006).

The first carboxylation reaction in the  $C_4$  carbon fixation pathway, by contrast, generates ample amounts of malate and NADH in the cytoplasm of mesophyll cells. This explains the  $CO_2$  independence of shoot  $NO_3^-$  assimilation in  $C_4$  plants (Bloom et al. 2010; Bloom et al. 2012b).

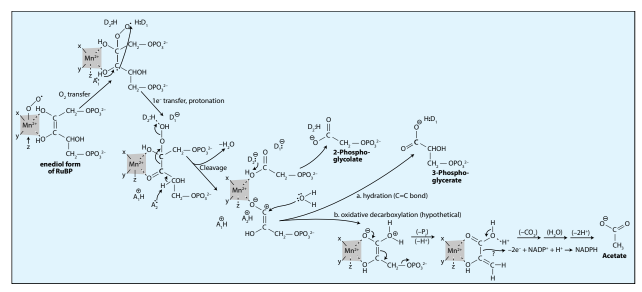
# The Rubisco Complex

Information about the biochemistry of RuBP oxygenation is limited. The stroma of the chloroplast contains similar amounts of Mg<sup>2+</sup> (2 mM, Ishijima et al. 2003) and Mn<sup>2+</sup> (2 mM, Burnell 1988; Robinson and Gibbs 1982). Rubisco may form a complex with either Mg<sup>2+</sup> or Mn<sup>2+</sup> (Pierce and Reddy 1986), but the affinity of Rubisco for Mn<sup>2+</sup> is more than five time greater than that for Mg<sup>2+</sup> (Christeller 1981). The stoichiometry of CO<sub>2</sub> trapping (Miziorko and Sealy 1980) and <sup>31</sup>P and <sup>13</sup>C NMR measurements (Pierce and Reddy 1986) indicate that Mn<sup>2+</sup> and Mg<sup>2+</sup> share a common binding site in the large subunit of Rubisco. Nearly all of the biochemistry of Rubisco has been conducted in the presence of Mg<sup>2+</sup> and in the absence of Mn<sup>2+</sup> because Rubisco when associated with Mn<sup>2+</sup> strongly favors RuBP oxygenation, whereas Rubisco when associated with Mg<sup>2+</sup> favors RuBP carboxylation (Chen and Spreitzer 1992; Christeller and Laing 1979; Houtz et al. 1988; Jordan and Ogren 1981; Raghavendra et al. 1981; Wildner and Henkel 1979).

 ${
m Mg}^{2+}$  has a pair of electrons in its outer shell, whereas  ${
m Mn}^{2+}$  has up to five unpaired electrons and thus participates more readily in redox reactions. In specific,  ${
m Mn}^{2+}$  participates in the catalytic process of RuBP oxygenation (Miziorko and Sealy 1984) during which it becomes excited and transfers an electron with every turnover (Lilley et al. 2003). One possibility is that  ${
m Mn}^{2+}$  transfers electrons to NADP+ (Figure 11). The resultant NADPH activates Rubisco (Laing and Christeller 1976) and then converts OAA to malate for export to the cytoplasm. This malate in the cytoplasm generates NADH to convert  ${
m NO}_3^-$  to  ${
m NO}_2^-$ .

Several additional observations are consistent with this hypothesis. RuBP oxygenation releases 76.3 kcal mol<sup>-1</sup> (Frank et al. 2000), substantially more than the 52 kcal mol<sup>-1</sup> required to reduce NADP<sup>+</sup> to NADPH (Taiz and Zeiger 2010). NADPH complexes strongly with Rubisco and activates the enzyme, but only when CO<sub>2</sub> and Mg<sup>2+</sup> are present in suboptimal concentrations (Chollet and Anderson 1976; Chu and Bassham 1974; Matsumura et al. 2012; McCurry et al. 1981). NADPH binds to the catalytic site of Rubisco through metal-coordinated water molecules (Matsumura et al. 2012).

If Rubisco generates NADPH during RuBP oxygenation,  $C_3$  carbon fixation is more efficient than previously thought, and both  $C_3$  and  $C_4$  carbon fixation at moderate temperatures will expend the equivalent



**Fig. 10** One possible scenario for the intermediates formed during RuBP oxygenation (Chen and Spreitzer 1992; Cleland et al. 1998; Lilley et al. 2003; Oliva et al. 2001; Tapia and Andrés 1992; Tcherkez et al. 2006)

of about 11 ATPs per  $CO_2$  fixed. Indeed, the quantum yield of photosynthesis in an ambient  $CO_2$  and  $O_2$  atmosphere does not differ significantly between  $C_3$  and  $C_4$  species at temperatures between 25° and 30°C (Skillman 2008). Only under hotter and drier conditions does  $C_4$  carbon fixation become more efficient than  $C_3$  fixation. Therefore,  $C_3$  species continue to dominate in most locations.

# Why is photorespiration still prevalent?

Several phenomena are responsible for the persistence of photorespiration through 23 million years of low atmospheric CO<sub>2</sub> concentrations.

- Rubisco oxygenation is inseparable from Rubisco carboxylation (Moroney et al. 2013; Tcherkez et al. 2006). Rubisco catalyzes the carboxylation reaction through stabilizing the formation of the enediol conformation of RuBP (Figure 10). This conformation, however, can react with either CO<sub>2</sub> or O<sub>2</sub>. The specificity of Rubisco for CO<sub>2</sub> over O<sub>2</sub> derives from stabilizing the six carbon intermediate before it is cleaved to form two molecules of PGA. Consequently, any mutation that increases the specificity of Rubisco for CO<sub>2</sub> over O<sub>2</sub> slows the carboxylation reaction.
- Photorespiration maintains redox homeostasis within plant cells (Scheibe and Dietz 2012). Photosynthesis
  generates highly reactive compounds as it captures solar energy and converts it into energy-rich, but
  stable compounds such as carbohydrates. Metabolic pathways, especially under stressful conditions, may
  become unbalanced, and dangerous compounds such as reactive oxygen species (ROS) may accumulate
  (Voss et al. 2013). Photorespiration can dissipate many of these potentially dangerous compounds.
- Photorespiration produces H<sub>2</sub>O<sub>2</sub> in the peroxisome and thus serves as a mechanism for rapidly transferring a signal of photosynthesis to the entire plant cell (Foyer et al. 2009). This signal is involved in photoperiod detection and pathogen defense as well as responses to abiotic stress.
- Photorespiration serves as a mechanism for plants to use NO<sub>3</sub> as a nitrogen source without diverting energy from CO<sub>2</sub> fixation. The following provides details about this phenomenon.

## Nitrate as a nitrogen source

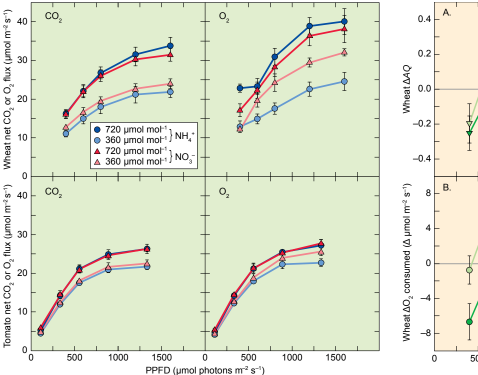
The element nitrogen is a constituent of many organic compounds including all amino acids and nucleic acids. As such, plants require a greater amount of nitrogen than any other mineral element, and its availability generally limits the productivity of natural and agricultural ecosystems (Epstein and Bloom 2005). Conversions among various nitrogen compounds are among the most energy-intensive reactions in life. Consider that plants are generally between 1 and 2% organic nitrogen on a percentage dry weight basis, but that the conversion of NO<sub>3</sub><sup>-</sup> into organic nitrogen expends about 25% of the total energy in shoots (Bloom et al. 1989) and roots (Bloom et al. 1992). These processes expend the energy equivalent of 12 ATP per NO<sub>3</sub><sup>-</sup> assimilated, whereas most biochemical reactions expend the energy equivalent of one or perhaps two ATP.

Most organisms prefer higher energy forms of nitrogen such as  $NH_4^+$  or amino acids. Phytoplankton (Dortch 1990), fungi (Hodge et al. 2010), cyanobacteria (Ohashi et al. 2011), and bacteria (Luque-Almagro et al. 2011) absorb and assimilate  $NO_3^-$  only in the absence of  $NH_4^+$ . In many soils, microorganisms quickly absorb  $NH_4^+$  and either assimilate it into amino acids or nitrify it to  $NO_3^-$ .  $NH_4^+$  also becomes adsorbed on the soil cation exchange matrix. Because soil microorganisms often ignore  $NO_3^-$  and because  $NO_3^-$  as an anion moves relatively freely through the soil,  $NO_3^-$  is often the predominant form of nitrogen available to plants (Epstein and Bloom 2005).

Nitrogen nutrition,  $NH_4^+vs. NO_3^-$ , neither influences net  $CO_2$  consumption (Figure 11) nor cyclic electron flow around photosystem I at low light levels (Walker et al. 2014). This is consistent with the lack of competition for reductant between  $CO_2$  fixation and  $NO_3^-$  assimilation (Robinson 1988) because, as discussed previously, FNR has a higher affinity for  $Fd_r$  than NiR. At high light levels and ambient  $CO_2$  and  $O_2$  concentrations, net  $O_2$  evolution is faster (Figures 11 and 12) and cyclic electron flow around photosystem I is higher (Walker et al. 2014) when plants receive  $NO_3^-$  rather than  $NH_4^+$  as a nitrogen source. Presumably, plants use reductant generated from the light dependent reactions rather than mitochondrial respiration to assimilate  $NO_3^-$  when  $CO_2$  concentration limits  $CO_2$  fixation.

When factors other than  $CO_2$  limit  $CO_2$  fixation, plants may delay assimilating the  $NO_3^-$  that they have absorbed. Free  $NO_3^-$  may comprise as much as 60% of the total nitrogen in a plant (Maynard et al. 1976). This  $NO_3^-$  serves as a metabolically benign osmoticant that balances other ions such as potassium in plant tissues and helps to maintain a favorable cellular water status (Bloom et al. 2012a; Burns et al. 2010; Hanson and Hitz 1983; McIntyre 1997; Veen and Kleinendorst 1986).

In summary, the linkage between photorespiration and  $NO_3^-$  assimilation provides higher plants with a relatively abundant nitrogen source that other organisms cannot afford to use, but that  $C_3$  plants can use with little additional cost. Yes, photorespiration may sacrifice 20% to 35% of  $CO_2$  fixation, but plants that are dependent on  $NO_3^-$  as a nitrogen source are spared the expense of either devoting 25% of their photosynthate to  $NO_3^-$  assimilation or suffering protein deprivation. Apparently, over the last 23 million years, 96% of higher plant species have adapted to this tradeoff.



**Fig. 11** Response of net  $CO_2$  consumption (left panels) and net  $O_2$  evolution (right panels) to photosynthetic photon flux density (PPFD) in wheat (upper panels) and tomato (lower panels) leaves when the plants received  $NH_4^+$  (blue) or  $NO_3^-$  (red) nutrition and were exposed to an atmosphere containing 720 (dark colors) or 360 (light colors)  $\mu$ mol  $mol^{-1}$   $CO_2$ . Shown are the means  $\pm$  SE for 6 wheat plants and 6 to 9 tomato plants per treatment. Notice that in both species,  $CO_2$  fluxes do not differ with N source, and that  $O_2$  fluxes are faster under  $NO_3^-$  nutrition than  $NH_4^+$  nutrition, but only at higher light levels and 360  $\mu$ mol  $mol^{-1}$   $CO_2$ . (Cousins and Bloom 2004; Searles and Bloom 2003)

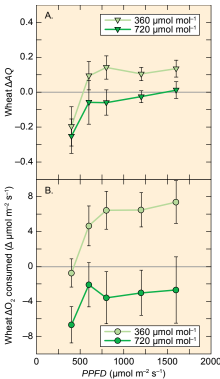


Fig. 12 Responses of wheat shoots (mean  $\pm$  SE, n = 6) to photosynthetic photon flux density (PPFD). (A) Changes in assimilatory quotient ( $AQ = \text{net } CO_2 \text{ consumed } / \text{ net } O_2$ evolved) with the shift from NO<sub>3</sub><sup>-</sup> to NH<sub>4</sub><sup>+</sup> as a N source. (B) Changes in the gross O2 consumed (gross O2 evolved minus net O2 evolved) with the shift from NO<sub>3</sub><sup>-</sup> to NH<sub>4</sub><sup>+</sup> as a N source. As light levels increased and 360 µmol mol<sup>-1</sup> CO<sub>2</sub> limited carbon fixation, exposure to NO<sub>3</sub> stimulated the light dependent reactions of photosynthesis to split water, evolve oxygen, and transfer electrons to NO<sub>3</sub> and NO<sub>2</sub> rather than to CO<sub>2</sub>, and decreased gross consumption (Cousins and Bloom 2004).

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#### **Conflicts of Interest**

The author has no conflicts of interest with regards to this research.

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