Modeling Grain Nitrogen Accumulation and Protein Composition to Understand the Sink/Source Regulations of Nitrogen Remobilization for Wheat

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A functional explanation for the regulation of grain nitrogen (N) accumulation in cereal by environmental and genetic factors remains elusive. Here, new mechanistic hypotheses of grain N accumulation are proposed and tested for wheat (Triticum aestivum). First, we tested experimentally the hypothesis that grain N accumulation is mostly source regulated. Four contrasting cultivars, in terms of their grain N concentrations and yield potentials, were grown with non-limiting N supply. Grain number per ear was reduced by removing the top part of the ear at anthesis. Reduction in grain number gave a significant increase in N content per grain for all cultivars, showing that grain N accumulation was source regulated. However, on a per ear basis, cultivars with a high grain number fully compensated their N accumulation for reduced grain number at anthesis. Cultivars with a lower grain number did not compensate completely, and grain N per ear was decreased by 16%. Second, new mechanistic hypotheses of the origins of grain N source regulation and its response to environment were tested by simulation. The hypotheses were: (a) The regulation by N sources of grain N accumulation applies only for the storage proteins (i.e. gliadin and glutenin fractions); (b) accumulation of structural and metabolic proteins (i.e. albumin-globulin and amphiphilic fractions) is sink-regulated; and (c) N partitioning between gliadins and glutenins is constant during grain development and unmodified by growing conditions. Comparison of experimental and simulation results of the accumulation of grain protein fractions under wide ranges of N fertilization, temperatures, and irrigation supported these hypotheses.

One challenge for global nutrition in the next decade is to increase food yield per unit ground area in a sustainable manner while maintaining its end use value (Cassman, 1999; Tilman, 1999; Tilman et al., 2002). Grain protein concentration and composition are major determinants of grain nutritional value (Feil, 1997). The concentration of Lys in grain, the most limiting amino acid in cereals for human and monogastric animals, increases with increasing grain protein concentration (Feil, 1997) despite the decrease of its concentration in total protein (Mossé et al., 1985). Grain protein concentration and composition are also the major determinants of flour functional properties (Weegels et al., 1996; Shewry and Halford, 2002). However, the inverse relationship between grain yield and protein concentration, reported for several species, may prevent breeders from improving these two traits simultaneously (Stewart and Dwyer, 1990; Delzer et al., 1995; Feil, 1997; Brancourt-Hulmel et al., 2003). To break this inverse relationship, genetic increments in grain protein yield must keep pace with those in grain yield.

Therefore, efforts to overcome the inverse relationship between grain yield and protein concentration must concentrate on improving grain protein accumulation per square meter and per grain (Feil, 1997; Triboï and Triboï-Blondel, 2002).

An increase in grain protein content may come from either improved capacity of the grain to accumulate nitrogen (N) or through greater N supply to the grains (Triboï and Triboï-Blondel, 2002). Several studies have shown some degree of control over grain N by intrinsic grain characteristics for wheat (Triticum aestivum; Borghi et al., 1986), barley (Hordeum vulgare; Mattsson et al., 1993), and maize (Zea mays; Wyss et al., 1991). Although others have shown control of grain N accumulation by the level of N supply for wheat (Barlow et al., 1983; Barneix and Guitman, 1993; Ma et al., 1995, 1996), barley (Dreccher et al., 1997; Voltas and Araus, 1997), maize (Wyss et al., 1991), pea (Pisum sativum; Lhuillier-Soundele et al., 1999a, 1999b), and soybean (Glycine max; Saravitz and Raper, 1995; Nakasathien et al., 2000). Comparison of the capacity of in vitro-cultured grains or seeds from low- and high-protein genotypes of wheat (Donovan et al., 1977), maize (Wyss et al., 1991), and soybean (Hayati et al., 1996) to accumulate N has led to the conclusion that genetic differences in grain or seed N content and concentration are caused, at least
in part, by differences in protein synthetic capacity. The opposite conclusion was reached for barley when comparing a high-protein accession of wild barley (*Hordeum spontaneum* Koch), with low-protein barley cv Ruth, which were able to accumulate 300 and 350 g proteins kg⁻¹ dry mass, respectively (Corke and Atsmon, 1988). Hence, it is still not clear if environmental and genetic differences in grain protein accumulation are regulated by process within the grains or by the N supply from the vegetative organs or are colimitcd by both (Feil, 1997; Tribol and Tribol-Blondel, 2002). Nevertheless, most crop and plant simulation models assume grain N accumulation to be sink regulated (e.g. Porter, 1993; for full reference, see Jamieson and Semenov, 2000). Most studies of the regulation of grain N accumulation for cereals have not considered the partitioning of N in the grain. We believe that to make progress in our understanding of the regulation of grain N accumulation, we should consider the physiological function of N in the grain.

Grain proteins can be divided into structural/metabolic (N<sub>str</sub>) and storage (N<sub>stora</sub>) proteins (Shewry and Halford, 2002). Structural/metabolic proteins consist of albumin, globulin, and amphiphilic proteins. In wheat, storage proteins are divided into two broad fractions. These are gliadins (N<sub>gli</sub>), which are present as monomers, and glutenins (N<sub>glu</sub>), which form polymers. Structural/metabolic protein fractions accumulate mainly during the early phase of grain growth, when most endosperm cells are still dividing, whereas the accumulation of storage proteins fractions occurs later when cell division as stopped and grain growth is only due to cell expansion (Stone and Nicolas, 1996; Tribol et al., 2003). Although grain protein composition depends primarily on genotype, it is significantly affected by environmental factors and their interactions (Graybosch et al., 1996; Huebner et al., 1997; Tribol et al., 2000; Zhu and Khan, 2001). However, the mechanism by which genotype and environmental factors modified the accumulation of the protein fractions are unknown, and, to date, no attempt has been made to model the partitioning of grain or seed N to different protein fractions.

In this study, we manipulated the sink to source ratio of four contrasted wheat cultivars to show that, overall, grain N is regulated by the supply of N to the grain. This was further confirmed by a simulation study using the wheat simulation model Sirius (Jamieson and Semenov, 2000), in which grain N accumulation is driven by N availability in the sources. We were able to explain a wide variation in observed grain N concentration at the canopy level, induced by N fertilization and postanthesis high and low air temperatures and water deficit. In this paper, we extended the concepts of grain N dynamic in Sirius to include a functional explanation for the regulation of the source regulation of grain N accumulation in a series of new mechanistic hypotheses formalized as a simulation model of the accumulation of grain protein fractions. The main hypotheses were: (a) The apparent overall source regulation of grain N accumulation is due to the synthesis of storage proteins, (b) the synthesis of structural and metabolic proteins is sink regulated, and (c) the allocation of N between the storage protein fractions gliadin and glutenin is constant during grain filling and is not modified by growing conditions. Comparison of experimental and simulated results for a wide range of environmental conditions provided a strong support to these functional hypotheses.

**RESULTS**

**Grain N Accumulation Is Source Regulated for Both High- and Low-Yielding Cultivars**

First, we analyzed the level of supply limitation of grain N accumulation in four cultivars with different potential grain numbers per square meter, an increase in which has been one of the major factors contributing to grain yield increases over the last 40 years (Reynolds et al., 1999; Brancourt-Hulmel et al., 2003). Sink to source ratio was modified by removing the top part of the ear on the main stems at anthesis or 250 degree-days (°Cd) later. The experiment was done in the field under non-limiting soil N supply.

Grain yield, yield components, and N content and concentration for the four cultivars and the different treatments show that, under normal conditions, grain number per ear was highest for the cultivars Arche and Réctal, intermediate for Renan, and lowest for Tamaro (Table I). Grain yield was not significantly different for the cultivars Arche, Réctal, and Renan but was 52% to 60% lower for Tamaro compared with the three other cultivars. The four cultivars analyzed could be separated as low (Arche and Réctal) and high (Renan and Tamaro) protein cultivars (Table I).

The ear halving treatment at anthesis reduced the number of grains per ear (i.e. per square meter) by 26% to 32%. This treatment leaded to an increase in N content per grain of 37%, 43%, 25%, and 14% for Arche, Réctal, Renan, and Tamaro, respectively (Table I). However, not all cultivars fully compensated for the reduced grain number. Grain N per ear decreased by 16% for the two cultivars with the lower grain number per square meter (i.e. Renan and Tamaro; Fig. 1). In contrast, grain N per ear was not modified by the reduction in grain number per ear at anthesis for the two cultivars with the higher grain number per square meter (i.e. Arche and Réctal).

Ear halving at 250 °Cd after anthesis reduced the sink size by 37% to 40%, leading to an increase of N content per grain of 29%, 24%, 12%, and 9% for Arche, Réctal, Renan, and Tamaro, respectively (Table I), whereas grain N per ear decreased by 22% to 33% for all four cultivars (Fig. 1).
Variations of Grain N Can Be Predicted Based on the Level of N Supply from the Plant

Regulation of grain N accumulation was further analyzed by simulating N uptake and redistribution for wheat crops grown in the field with a combination of rates and timings of N fertilization and in controlled environments, where different postanthesis temperatures and watering regimes were applied postanthesis. The wheat simulation model Sirius was used to simulate dry matter and N accumulation in the different organs of the crops for these experiments. Simulated and observed kinetics of grain N accumulation for the different experimental treatments agreed well (data not shown), and simulated and observed final grain N were well correlated ($r^2 = 0.83$, 16 degrees of freedom [d.f.]; Fig. 2). The square
Sink/Source Regulation of Grain N Accumulation Revisited

Consideration of the protein fractions in the grain gives a new perspective to the supply regulation of grain N accumulation. An example of the kinetics of accumulation of $N_{\text{stru}}$, $N_{\text{gli}}$, and $N_{\text{gln}}$ obtained for crops grown in the field with a combination of two rates and timings of N fertilization is shown in Figure 3. Pre-anthesis N fertilization increased only slightly (7%) the final quantity of $N_{\text{stru}}$ but increased the final quantities of $N_{\text{gli}}$ and $N_{\text{gln}}$ by 33% and 22%, respectively (Fig. 3, A and B). Under conditions of normal pre-anthesis N fertilization, postanthesis N fertilization increased the final quantity of $N_{\text{stru}}$ by 25% but that of $N_{\text{gli}}$ and $N_{\text{gln}}$ by 95% and 49%, respectively (Fig. 3, C and D). Thus, the accumulation of $N_{\text{gli}}$ and $N_{\text{gln}}$ are significantly enhanced by N fertilization, whereas $N_{\text{stru}}$ is little affected.

The model of accumulation of grain protein fractions described here gave accurate simulations of the accumulation of $N_{\text{stru}}$, $N_{\text{gli}}$, and $N_{\text{gln}}$, even for conditions of non-limiting soil N supply, such as the treatment H15 (Fig. 3). Similar agreement was observed for the 14 other treatments of Figure 2 (data not shown). Simulated and observed $N_{\text{gli}}$ ($r^2 = 0.86$, 16 d.f.) and $N_{\text{gln}}$ ($r^2 = 0.96$, 16 d.f.) at harvest ripeness were well correlated (Fig. 4). The square root of the mean square error of prediction was 26 $\mu$g N grain$^{-1}$ over a range of 84 to 315 $\mu$g N grain$^{-1}$ for $N_{\text{gli}}$ and 31 $\mu$g N grain$^{-1}$ over a range of 215 to 508 $\mu$g N grain$^{-1}$ for $N_{\text{gln}}$.

The supply limitation of grain N uptake may apply predominantly at a particular stage of the development of the grain. Hence, we used our model of grain N partitioning to analyze the joint evolution of the
and supply of total N \((N_{\text{tot}})\) during the development of wheat grains grown in the field with different rates and timing of N fertilization. Crops received either 0 (treatment L) or 10 (H) g N m\(^{-2}\) at the beginning of stem elongation, followed by either 3 (L3 or H3) or 15 (L15 or H15) g N m\(^{-2}\) at anthesis. A, Treatment L3; B, L15; C, H3; D, H15.

![Figure 5. Time course of simulated demand for structural N (—) and supply of total N (—) during the development of wheat grains grown in the field with different rates and timing of N fertilization. Crops received either 0 (treatment L) or 10 (H) g N m\(^{-2}\) at the beginning of stem elongation, followed by either 3 (L3 or H3) or 15 (L15 or H15) g N m\(^{-2}\) at anthesis. A, Treatment L3; B, L15; C, H3; D, H15.](Image)

**DISCUSSION**

The experiments and simulations reported here were designed to analyze the source/sink regulation of grain N accumulation and to assess its genetic variability. Several lines of evidence from studies on detached ears cultured in vitro (Barlow et al., 1983; Corke and Atsmon, 1988) and isolated plants cultivated in pots under controlled environments (Barneix and Guitman, 1993; Drecceer et al., 1997) have suggested that grain N accumulation for many cereal species is source regulated, but apparently this never has been investigated at the canopy level under field or controlled environment conditions, and most studies have been limited to one genotype. Moreover, no functional hypothesis has been proposed to account for the source regulation of grain N accumulation. We modified the sink to source ratio of four contrasted genotypes of wheat grown in the field. The results show that the level of source regulation of grain N accumulation depends on the genotypes, but none of the genotypes were sink limited. The hypotheses framed above have been formulated in a simulation model that predicted the dynamic changes of grain protein composition, an important nutritional and economic trait for cereals. The simulation results presented here support these hypotheses over a broad range of environmental conditions.

Ear halving increases the availability of N to the remaining grains either at anthesis or 250 °Cd later, when cell division has ended and grain growth is solely due to cell expansion (Gleadow et al., 1982; Singh and Jenner, 1982). Ear halving has been shown to increase the final number of cells per grain for wheat (Brookehurst, 1977). For both ear halving treatments, N content per grain increased for all four cultivars, indicating that the storage capacities of the grains were not reached for the control treatments; thus, the capacity of the sink to synthesize proteins did not regulate grain N accumulation for the four cultivars. If the N sources were the major regulators, one would expect grain N per ear to be constant, i.e. independent of grain number. In the experiments presented here, the two cultivars with the lower grain number per square meter and per ear (Renan and Tamaro) were unable to compensate completely for the reduced grain number per ear, so grain N accumulation for these cultivars became sink regulated. In contrast, the two cultivars with the higher grain number per square meter and per ear (Arche and Récal) were able to fully compensate for the reduced grain number per ear. Thus, grain N accumulation for these cultivars was still supply regulated. The only way we could introduce a sink limitation of grain N accumulation in these two cultivars was to artificially reduce the total sink number too late for compensation to occur. This latter result suggests that the compensation observed for Arche and Récal when ablated at anthesis was due to an increased cell number per endosperm.

We were able accurately to predict total grain N accumulation over a large range of grain N for Thésee, a high-yield potential and grain number cultivar, by assuming grain N accumulation to be determined by the size of the source of N, defined as the total nonstructural crop N at anthesis. This gives further support to the previous conclusion that, overall, the accumulation of grain N, at least for high-yielding cultivars with high grain number, is regulated by the source of N and not by the activity of the grain.

Using the model of accumulation of protein fractions described here, the comparison of the simulated demand and supply of grain N suggested that grain N accumulation was sink limited or colimited by both source and sink for the first 10 to 15 d after anthesis. This emphasizes the importance of the early stage of grain development, characterized by active cell division in the endosperm, in setting the potential grain size and \(N_{\text{stru}}\). In contrast with the early phase of grain development, grain N accumulation was always source limited during the grain filling phase of grain development, characterized by active cell division in the endosperm, in setting the potential grain size and \(N_{\text{stru}}\).
period, even when soil N was non-limiting. Moreover, simulations and observations of the accumulation of grain protein fractions for developing grains obtained from plants grown in the field with different rates and timings of N fertilization and in the controlled environment chambers with different postanthesis temperatures and watering regimes agreed well, verifying the hypothesis that the supply limitation of grain N accumulation results from the accumulation of storage proteins and not from that of structural proteins. Thus, the sink/source limitation of grain protein accumulation is related to differences in the timing of deposition between structural/metabolic proteins versus storage proteins, as postulated earlier for barley (Dreccer et al., 1997).

The source regulation of the accumulation of storage proteins gives a mechanistic explanation of the effect of overexpressing glutenin genes on protein composition and concentration where the transformation of wheat with high-M_{s} glutenin subunit genes results in increased quantities and proportions of the high-M_{s} glutenin subunits (Alt peter et al., 1996; Blechl and Anderson, 1996; Barro et al., 1997; Alvarez et al., 2000) but with no difference in total protein quantity and concentration (Rooke et al., 1999). The source regulation of the accumulation of storage proteins is also in good agreement with the presence of two regulatory elements in the promoter region of the genes of several grain storage proteins: the “endosperm motif,” which act as a positive element under high-N conditions; and the “GNC4-like motif,” which act as a negative element under low-N conditions for grains of barley (Hammond-Kosack et al., 1993; Müller and Knudsen, 1993).

The hypothesis introduced in our simulation model of grain protein accumulation that the partitioning coefficient for N_{str} and N_{glc} is constant during grain development and is not modified by the growth conditions was verified. This implies that any modification of the gliadins to glutenins ratio is only the result of modification of total N content per grain and that the processes leading to the synthesis of storage proteins in the grain are not affected by the concentration of N. We observed similar result for the albumin-globulin and the amphi philic proteins, the constituent of N_{str} (data not shown). Preliminary data indicate that this is true for the other cultivars studied here, i.e. Arche, Récital, Renan, and Tamaro (V. Samoil, P. Martre, and E. Tribó, unpublished data), and, importantly, the same partitioning parameters applied. These results imply that the protein fractions and amino acids composition of wheat grains from widely different cultivars can be deduced directly from the total quantity of N per grain. Furthermore, we suggest that the genotype-environment interactions for the composition of protein fractions reported earlier (Graybosch et al., 1996; Tribó et al., 2000) act only via variations of total grain N and, thus, N availability and not via the allocation of N between the different protein fractions.

Functional genomics and proteomics studies aiming at understanding the regulation of grain protein level and composition for cereals, especially wheat (Clarke et al., 2001; Lagudah et al., 2001; Shewry et al., 2001) and rice (Oryza sativa; Tyagi and Mohanty, 2000), have focused on the “protein warehouse” (i.e. the grain). These studies are valuable to better understand the development of the grain and to genetically modify grain protein composition and increase the sink demand. However, the supply limitation of grain N accumulation, as shown here, means that to increase grain yield while maintaining high nutritional and processing values requires understanding of the functioning of the “protein factory” (i.e. the vegetative organs) and its interaction with the grain.

**MATERIALS AND METHODS**

All experiments were at Clermont-Ferrand, France (45°47’ N, 3°10’ E, 329-m elevation) with winter wheat (Triticum aestivum) cv Thésée, Arche, Récital, Renan, and Tamaro.

**Ear Halving Experiments**

Source/sink regulation of grain N accumulation was studied in the field for four cultivars (Arche, Récital, Renan, and Tamaro) with contrasting potential grain number, grain yield, and grain protein concentration. One main plot of 202 m² was sown for each cultivar on November 7, 2001 at a density of 300 grains m⁻². The crops were rain fed. Accumulated rainfall from sowing to anthesis and from anthesis to grain maturity was 124 to 174 and 115 to 135 mm depending on the cultivar, respectively. Average air temperature from sowing to anthesis and from anthesis to grain maturity was 6.8°C to 7.3°C and 17.8°C to 18.7°C depending on the cultivar, respectively. The source to sink ratio was artificially manipulated by removing the top one-half or so of the ears from the main stems on three 0.5-m² subplots per cultivar. At the same time, three 0.5-m² control subplots were identified for each cultivar. Ear halving was performed either at anthesis or 249, 277, 254, and 244°C later for Arche, Récital, Renan, and Tamaro, respectively. Samples of 0.5 m² were taken in each subplot at the time when ears were halved and at grain maturity. Three replicates were used per N treatment.

**Temperature and Drought Experiments**

To study the effects of postanthesis temperature and drought at the canopy level, crops of wheat cv Thésée were grown in 2-m² containers in controlled environment closed-top chambers under natural light (Tribó et al., 2003). From 5 d after anthesis to grain maturity, five air temperatures relative to ambient air temperature were applied in the chambers: −5°C (treatment termed −5, average temperature of 14.9°C); 0°C (0, average temperature of 19.5°C); +5°C (+5, average temperature of 22.3°C); +5°C until 300°C, base 0°C after anthesis, then +10°C until harvest maturity (+5/+10, average temperature of 24.7°C); and +10°C until 300°C after anthesis then +5°C until harvest maturity (+10/+5, average temperature of 23.7°C).

Interactions between postanthesis temperature and drought were studied in a 2nd year of experiments where two air temperatures relative to −5°C and +5°C, average temperature of 12°C and 19.9°C, respectively) were applied from 5 d after anthesis to grain maturity. The crops were rain fed from sowing to anthesis and received 226 mm of rainfall during that period. One container for each temperature treatment received 25 to 50 mm of water every 4 to 7 d until harvest maturity to replace measured crop evapotranspiration (treatments −5W and +5W), whereas the other container received 5% to 15% of

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Martre et al.
the measured crop evapotranspiration from anthesis to harvest maturity (treatments − SD and + SD). Crop evapotranspiration and, thus, crop water requirements under the controlled environment chambers were computed from measurements of the volume of water condensed on the cold exchanger of the chambers and the difference of air vapor pressure between the outlet and inlet of the chambers.

One controlled environment chamber/container was used per treatment. To study the dynamic accumulation of total N and protein fractions, three replicates each of 20 plants (approximately 0.25 m²) were collected every 50 to 135°C day from anthesis to grain maturity. Plants were sampled from the northern side of the containers through their southern side. To minimize the border effects, for each sampling date, the northernmost raw was discarded, and after each sampling, a net of the high of the crop was placed in place of the last raw removed.

N Experiments

The effect of N availability at anthesis in relation to the level of N nutrition before anthesis was studied in a field experiment for crops of wheat cv Thessée sown at a density of 300 seeds m⁻² (Tribo et al., 2003). Crops were sown in plots that had not received N fertilizer since 1948. Three rates of N were supplied at the beginning of stem elongation: 0, 5, and 10 g N m⁻² (treatments L, M, and H), respectively. The H treatments were on plots where leaves of sugar beet (Beta vulgaris) from a previous cultivation and a cut of alfalfa (Medicago sativa) had been buried. At anthesis, each plot was split into three subplots to which 0 (treatments L0, M0, and H0), 3 (L3, M3, and H3), or 15 (L15, M15, and H15) g N m⁻² were applied. Crops were rain fed. Accumulated rainfall from sowing to anthesis and from anthesis to grain maturity was 344 and 61 mm, respectively. Average air temperature from sowing to anthesis and from anthesis to grain maturity was 7.8°C and 19.6°C, respectively. Samples of 0.2 m² were taken in each subplot at anthesis and 290, 505, 712, and 900°C day later. Three replicates were used per N treatment.

Plant Sampling, Protein Extraction, and Total N Content Determination

Grains were separated, and their dry mass was determined on subsamples after oven drying at 70°C to constant mass. The remaining grains were frozen in liquid N, freeze dried, and stored at 4°C before analysis.

The protein fractions albumin-globulin, amphiphilic, gliadin, and glutenin were sequentially extracted from whole meal flour (Tribo et al., 2003). The residue fraction, which represents 1.5% to 9% of the total N content, was frozen in liquid N, freeze dried, and stored at 4°C. Albums were reconstituted with water and fractionated with aqueous propan-2-ol. The residue fraction, which represents 4% to 7% of the total N content, was used for the Kjeldahl method using a Kjeltec 2300 analyzer (Foss Tecator AB, Höganäs, Sweden). The glutenin fraction followed. The residue fraction, which represents 1.5% to 9% of the total N content, was used for the Kjeldahl method using a Kjeltec 2300 analyzer (Foss Tecator AB, Höganäs, Sweden). The glutenin fraction followed.

The Sirius Wheat Simulation Model

We used the wheat simulation model Sirius V99 (Jamieson and Semenov, 2000) to analyze the regulation of grain N accumulation of the crop from the experimental treatments described above. Detailed description of Sirius is given elsewhere (Jamieson et al., 1998; Jamieson and Semenov, 2000). In short, Sirius from emergence to anthesis, N demand is set in proportion to the increment of green area index and structure each day, whereas extra nonstructural N can be stored in proportion to stem biomass. The major assumptions are that specific leaf N concentration is constant at 1.35 g m⁻² of leaf, structural N is 0.5% of biomass accumulated until anthesis, and the crop can store N equivalent to 1% of the stem biomass. At anthesis, all nonstructural starch N (i.e. both stored nonstructural N and N in green tissue) is considered to be available for transfer to the grain. Grain N accumulates at a constant rate, in thermal time, from 100°C day after anthesis until either the total senescence of the canopy or the unconstrained end of grain filling, whichever occurs first. The unconstrained duration of grain filling is assumed to be under genetic control and constant in thermal time. The flux rate of N to the grain in the Sirius model is set at anthesis such as all the nonstructural N would be transferred by the end of unconstraint grain filling. Grain N is supplied from three different sources, accessed in sequence. The first is excess stem N and N released by natural leaf senescence. If these are insufficient, soil N is taken. Should these combined sources be insufficient, then the required N is obtained by accelerating leaf senescence.

The initial quantity of soil organic N at sowing was adjusted in Sirius to match the observed crop N content at anthesis using the treatments 0 and L0 for the controlled environment closed-top chamber and field experiments, respectively. Phenological development was not part of this study. Thus, the phylochron in Sirius was adjusted so that the simulated and observed anthesis dates matched. A phylochron value of 93°C day was used for the controlled environment chamber experiments, and 112°C day was used for the field experiments. Where appropriate, others genetic parameters in Sirius were set as for wheat cv Claire.

Modeling the Accumulation of Grain Protein Fractions

Although grain yield and protein content are regulated at the square meter scale (Jamieson and Semenov, 2000), grain N partitioning appeared to be regulated at the grain scale (Stone and Nicolas, 1996; Tribo et al., 2003). Thus, accumulation of grain protein fractions were modeled at the grain scale. The total grain N (Ngrain) was divided into structural (Nstr) and storage (Nstor) N:

\[ N_{str}(t) = N_{ord}(t) + N_{sto}(t) \]  
\[ N_{sto}(t) = a_{gi}N_{str}(t) \]  
\[ N_{gi}(t) = (1 - a_{gi})N_{str}(t) \]  

where Nstr is composed of the albumin-globulin and amphiphilic protein fractions, and Nsto is composed of the gliadin (Ngli) and glutenin (Ngln) protein fractions. From the analysis of treatment 0 of the experiment in the controlled environment chambers described above, we assumed a constant partitioning of Nsto between Ngli and Ngln:

\[ \frac{dN_{str}}{dt} = k_{str}N_{str}T_c, \quad T_c < D_{ord} \]  
\[ \frac{dN_{demand}}{dt} = N_{ord}(T_c = D_{ord})T_c, \quad D_{ord} \leq T_c \leq D_{str} \]  
\[ \frac{dN_{demand}}{dt} = 0, \quad T_c > D_{str} \]  

where kstr is the initial relative rate of accumulation of Nstr, Tc is the average daily temperature, Tc is the thermal time after anthesis, base 0°C, and Dord and Dstr are the durations in thermal time of the cell division and DNA endoreduplication phases, respectively.

The daily flux of Nstr was expressed as the minimum of the daily demand for Nstr (Nstr_demand) and the daily supply of total N (Nstr_supply). Based on previous work (Stone and Nicolas, 1996), we made several hypotheses to model Nstr_supply. During the initial cell division phase, accumulation of Nstr is exponential. During the cell expansion phase, the flux of Nstr is determined by the size of the pool of Nstr at the end of the cell division phase. We also assumed that the end of accumulation of Nstr coincides with the end of the DNA endoreduplication phase.

Using treatment 0 of the experiment in the controlled environment chambers, kstr and astr were estimated using regression analysis as 8.44 × 10⁻³ (Mr = 1) and 0.38 (dimensionless), respectively, and Nstr (Tc = 0) and Nstr (Tc = 25°C) as 27.27 and 3.50 μg N grain⁻¹, respectively. For Dord and Dstr, we used the values of 25°C found for wheat (Gleadlow et al., 1982; Singh and
Jenner, 1982) and maize grains (Engelen-Eigles et al., 2000), and for $D_{pr}$, we used the values of 450 °C found for maize grains (Engelen-Eigles et al., 2000).

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