REVIEW

Food security through translational biology between wheat and rice

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Comparative biology, rice, translational biology, wheat, yield potential

Abstract
Wheat and rice are the most important food crops in agriculture providing around 50% of all calories consumed in the human diet. While both are C₃ species, the evolution and domestication of wheat and rice occurred in very different environments, resulting in diverse anatomical and metabolic adaptation. This review focuses on the current understanding of their adaptation in an agronomic context. The similarities and differences between wheat and rice are discussed, focusing on traits related to phenology, photosynthesis, assimilate partitioning, and lodging resistance, these being the main abiotic drivers of yield expression in most agro-ecosystems. Currently, there are significant knowledge gaps in the major biological processes that account not only for differential adaption among cultivars within each species, but even between the two species. By addressing what is known as well as where gaps exist in a comparative context, this review aims to highlight translational research approaches that could provide insights into the genetic improvement of both crops.

Introduction
Wheat and rice are important cereal crops providing more than 50% of daily caloric intake worldwide. Taxonomically, wheat and rice represent different subfamilies of the Poaceae: Pooideae and Bambusoideae, respectively, and represent important models for comparative studies of the grasses because various aspects of their biology have been well studied (Sorrells et al. 2003; Shingaki-Wells et al. 2011; Cantu et al. 2013). Both wheat and rice have been domesticated in different climates, wheat is almost exclusively a dry-land winter crop, whereas rice is typically grown under tropical climates where most of the production comes from anaerobic/flooded soils (Nagai and Makino 2009; Shingaki-Wells et al. 2011). Such distinct evolutionary backgrounds triggered differential anatomical and developmental aspects resulting in various optimal growth conditions, yet they operate similar central metabolic processes (C₃ metabolism).

In the recent decades, the comparative biology between wheat and rice has been the subject of many studies (Sorrells et al. 2003; Shingaki-Wells et al. 2011; Cantu et al. 2013; for review, see Valluru et al. 2014). The study of two species may be useful in highlighting mechanisms of yield potential and stress adaptation in view of their domestication. Such distinct knowledge bases can lead to new knowledge, hypotheses, and predictions about related species and contribute to our knowledge of crop species evolution and adaptation to environment. Crop improvement programs can use comparative biology to transfer information on “traits” between relatively close species as well as more distant taxonomic groups. The genetic and molecular bases of yield-related traits were recently presented as a useful comparative biology approach between wheat and rice for application in translational research (Valluru et al. 2014). This review compares agronomic and physiological traits in two crop species.
Grain Yields of Wheat and Rice: Genetic Gains and Impacts

Genetic gains for wheat yields in the post Green Revolution period are widely reported to be in between 0.5% and 0.8% per year and globally the average annual absolute genetic gain of spring wheat reached about 40 kg ha\(^{-1}\) year\(^{-1}\) (Morgounov et al. 2010) despite the similar relative gains in irrigated and rainfed environments (Rajaram and Braun, 2008). Genetic gains in spring wheat continue to date (1995–2010) in both irrigated (Sharma et al. 2012) and rainfed environments (Manès et al. 2012) at an average of 0.6% and 1.0% p.a., respectively. Genetic gains in winter wheat appear to be stagnating in many parts of the world including the great plains of North America and the Europe (Brisson et al. 2010; Graybosch and Peterson 2010).

In rice, absolute yield gain in the range of 16–80 kg ha\(^{-1}\) year\(^{-1}\) were reported, being greater for Philippines (75–80 kg ha\(^{-1}\) year\(^{-1}\)) and smaller for Brazil (16–45 kg ha\(^{-1}\) year\(^{-1}\)) during the period 1966–1981 (Peng et al. 2000; Bresheghello et al. 2011). The hybrid super rice shows 10–20% yield gain in China (Peng et al. 2008), while at IRRI tropical hybrids have showed 11–14% yield gain in Philippines (Fischer and Edmeades 2010). A recent comparison between hybrids and inbreds found 14–18% yield advantage for the hybrids (Bueno and Lafarge 2009). Overall, improvements in yield gain have been achieved through an ideotype approach (e.g., shorter plants, more fertile tillers, high early vigor, and CGR [crop growth rate], larger panicles) and the development of hybrid rice (which itself benefited from the ideotype approach).

Agronomic and Physiological Trait Associated With Yield Potential

With recent emphasis on food security, two major avenues have been highlighted to improve the genetic potential of yield in wheat: (1) increasing photosynthetic capacity and efficiency to raise the biological threshold in terms of total plant mass available for yield formation; and (2) optimizing partitioning to grain while maintaining lodging resistance such that increased genetic potential is realized as agronomic gains (Reynolds et al. 2009; Foulkes et al. 2011; Parry et al. 2011). Prebreeding to combine favorable expression of yield potential traits – focused on achieving a more optimal balance between assimilation capacity (source) and grain sites (sink) – has been successful in increasing yield levels in international trials (Reynolds et al. 2015).

In rice, trait-based selection was used to add specific traits into new plant types (e.g., NPTs) whereby NPTs served as a basis for the generation of high-yielding cultivars/hybrids with larger panicles and higher tillering fertility (Peng et al. 2008). However, yield-based selection has shown to be more effective in increasing grain yields in rice than trait-based selection; although some researchers question the relevance of the traits used (Yuan et al. 2011) like the higher tillering fertility that may be inversely correlated to the tillering capacity (Lafarge and Bueno 2009). Here, we briefly summarize important agronomic and physiological traits contributing to yield potential.

Crop phenology

For cereals, reproductive growth is clearly the most critical physiological function to understand and modify thereby permitting accumulation of biomass to be translated into maximal expression of yield. While wheat exhibits a great plasticity for phenology (Sadras et al. 2009), it is yet to achieve its theoretical maximum harvest index of 0.6, and typically expresses values between 0.4 and 0.5, suggesting major opportunities for yield improvement without even changing RUE (radiation use efficiency) or biomass (Foulkes et al. 2011). Breeding efforts during the 20th century have reduced sowing to anthesis duration in durum wheat in Spain (8 days) and Italy (2 days) (Álvaro et al. 2008; Isidro et al. 2011), although this trend has reversed in some recent CIMMYT cultivars (Fischer 2001; Underdahl et al. 2008). The duration between sowing to terminal spikelet decreased while the duration between booting to anthesis increased, which diminished floret abortion (24%), increased grain number (23%) and grain set (13%) from old to modern cultivars (Isidro et al. 2011). Furthermore, the stem elongation period has been shown to exhibit the greatest sensitivity to environment, independent of the major phenological genes (Garcia 2011). It is known that earliness per se affects the duration of preanthesis phases and spike development in cereals (Faricelli et al. 2009; Borrás-Gelonch et al. 2011, 2012). These studies suggest that earliness per se could be important to understand phenological phase plasticity to the environment, and it remains to be determined to what degree environmental effects confound or interact with genetic expression of phenological phases.

In rice, the genetic variation in postflowering period (from flowering to maturity) is relatively small for a given environment (Peng et al. 2000), while the preflowering phase (from sowing to flowering) varies greatly (Vergara and Chang 1985; Yin et al. 1997a). The cultivars released at IRRI since 1980s show slightly decreased crop duration (6 days) than the cultivars released before 1980s (Peng et al. 2000) due to shortened vegetative period, while the reproductive and grain-filling periods were largely unaffected. One of the objectives of the reduced crop cycle in the tropics is to grow three crops a year. In Texas,
rice cultivars (1944–1992) showed a 12–18 days decrease in heading time (Tabien et al. 2008). The Brazilian cultivars (1984–2009) showed an average genetic loss of −0.25 day years$^{-1}$ in days to heading (Breseghello et al. 2011). Taken all together, both wheat and rice breeding programs have affected phenological pattern with crop duration and the length of the preanthesis period slightly decreased in both crops.

**Photosynthesis and related traits**

**Light interception**

LAI (leaf area index) beyond 3 is generally considered not an option for increasing light interception in wheat (Parry et al. 2011) because no apparent improvement in percent of light interception was observed in historical cultivars (Acreche et al. 2009). In contrast, LAI reaches 9 in rice depending on the developmental stage (Lü et al. 2007; Zheng et al. 2008; Zhang et al. 2009). Recent 3D models suggest that mean leaf inclination angles of rice cultivars range between 65° and 75° (70–90 in the upper canopy, while it is 30–60 in the lower canopy) (Zheng et al. 2008).

Modern rice and wheat cultivars have more erect and longer leaves in the upper canopy (flag-leaf). On average, rice plants receive 700–800 μmol photons m$^{-2}$ sec$^{-1}$ at the canopy top, with a total light interception of 80% of incident (Zheng et al. 2008). It has been estimated that leaves, panicles, and stems contribute about 80%, 15%, and 0.5% to the total light interception in rice (Saitoh et al. 2002). Reducing the panicle height to below top leaves or removing panicles (this allowed top leaves to access more light) have shown to increase leaf gas exchange rate about 15–50% (Setter et al. 1995; Saitoh et al. 2002). While more erect leaves/smaller leaves may allow light distribution to the lower canopy (Horton 2000), the genetic manipulation of leaf position, size, and thickness/density still needs to be explored since its manipulation is not straightforward.

**Leaf photosynthesis**

The relationship between the $A_{\text{max}}$ (light-saturated only) and grain yield is not straightforward (Long et al. 2006) although the differences in $A_{\text{max}}$ and its genetic variation can moderately contribute to variations in yields (Nelson 1988). Since modern hexaploid wheat species have lower $A_{\text{max}}$ (postulated to be due to an inhibitory effect of D genome on gene expression in the A and/or B genomes and larger mesophyll cell sizes) than diploid and tetraploid species (Koç et al. 2003), these genetic pools may serve as sources for increasing $A_{\text{max}}$. For example, emmer wheat ($Triticum dicoccoides$) shows a large genetic diversity for $A_{\text{max}}$, which is independent of leaf anatomy (Carver and Nevo 1990). Genetic advances in wheat yield does not appear to be related to changes in photosynthesis rate on an area basis when measured in the flag-leaf or the spike, but only to a higher, whole spike photosynthesis rate (Zhou et al. 2014), suggesting that under potential yield conditions, the spike with respect to its size, may contribute more than the flag-leaf to yield formation. The awns of wheat are an important photosynthetic and transpiration organ on the spike. It has been reported that the pathway for assimilation movement from awns to the kernels is minimal (Evans et al. 1972). Furthermore, the activity of PEP carboxylase is much higher in awns than in flag leaves throughout ontogeny, and is particularly high at the late stages of grain filling (Li et al. 2002, 2006).

In rice, increasing $A_{\text{max}}$ has been proposed to be a more effective strategy for increasing RUE than improving canopy architecture as modern rice varieties are believed to have an optimal canopy architecture in terms of light and N distribution (Peng and Ismail 2004). A large genotypic variation exists for $A_{\text{max}}$ in rice and its relatives (Cook and Evans 1983) particularly in diploid wild species (Yeo et al. 1994). Such large variation in $A_{\text{max}}$ is associated with variation in stomatal conductance ($g_s$) at mid-day conditions, Rubisco content, specific leaf area, and total leaf protein (Hubbart et al. 2007). $A_{\text{max}}$ at each developmental stage was better explained by $g_s$, especially under mid-day conditions while N content explains changes in $A_{\text{max}}$ associated with crop development (Ohsumi et al. 2007). Interestingly, since genotypic variation in N is unrelated to $g_s$, these two traits could be modified independently to increase $A_{\text{max}}$. Nevertheless, under light-saturated conditions, $g_s$ generally decreases due to decreased leaf water potential, therefore increasing leaf hydraulic conductance under saturated light should be targeted in breeding (Taylaran et al. 2011).

The physiological regulation of two parameters, TPU (triose phosphate utilization) involved in inorganic phosphate (Pi) recycling, and $g_{m}$, the mesophyll conductance, has received far less attention. Limited sink strength results in the accumulation of photoassimilates in the leaf, as a signal for downregulation of photosynthesis, reducing TPU (Zhu et al. 2010; Nebauer et al. 2011). This points to the value of exploring genetic variability in TPU regulation. Also, the reduction in $g_{m}$, standing for the diffusive conductance of CO$_2$ from the stomata to the chloroplast, significantly limits photosynthesis (Flexas et al. 2008; Griffiths and Helliker 2013) with variation in leaf anatomy assumed to be a major factor (Tomas et al. 2013). However, despite substantial evidence for large genotypic variability of $g_{m}$, exactly how leaf anatomy interacts with $g_{m}$ to
determine leaf [CO₂] status remains unclear. Overall, these studies suggest that while both crops exhibit different limitations to increase net photosynthesis, manipulating specific trait dynamics may improve photosynthesis rate in both crops (Fig. 1).

**Canopy photosynthesis and N distribution**

Canopy photosynthesis is largely a function of the vertical distribution of light and nitrogen (light-harvesting proteins and Rubisco), as well as sink activity (Dreccer et al. 2000; Bertheloot et al. 2008). The relationship between plant height and photosynthetic capacity is always not consistent (Gent and Kiyomoto 1984; Gent 1995; Bishop and Bugbee 1998). Semi-dwarf wheat often shows higher canopy photosynthesis at anthesis, which may be linked to stronger sinks associated with more efficient assimilate partitioning at anthesis (Bishop and Bugbee 1998), suggesting that in tall genotypes, the rate of photosynthesis might be partly constrained by assimilate export from leaves. In addition, cultivars with greater capacity to store N in nonphotosynthetic tissues may improve canopy N distribution (Dreccer et al. 1998) particularly if it follows the light gradient (Dreccer et al. 2000; Bertheloot et al. 2008). Recently, the contribution of ear photosynthesis to grain filling was highlighted, especially in situations of source limitation (Maydup et al. 2012).

In rice, an improved canopy structure has increased canopy photosynthesis particularly during the grain-filling period (Zhang and Kokubun 2004). However, the relationships between canopy net photosynthetic rate and biomass, and yield were inconsistent (Peng 2000) probably due to a high genetic variation in canopy photosynthesis during postanthesis (Zhang and Kokubun 2004). With LAI of 5, erect leaves allow higher canopy photosynthesis, while with LAI lower than 3, droopy leaves can allow higher canopy photosynthesis (Peng and Ismail 2004). Therefore, an ideal phenotype should have a prostrate-leaf canopy in the very early growth stage to intercept PAR effectively, while more erect leaves at the later stages would allow light to penetrate more efficiently. Moreover, rice leaves can efficiently maintain lower canopy temperature via efficient transpiration cooling given sufficient water availability (Wassmann et al. 2009; Gao et al. 2011). Depending on the N concentration, either linear ([Makino et al. 1988]) or curvilinear (Cook and Evans 1983) relationships with photosynthesis was observed (Shimoda 2012). Modeling studies suggest that if leaf N content can be increased above 1.8 g N m⁻² in upper leaves, daily canopy photosynthesis rate can be increased regardless of plant density and light condition (Shiratsuchi et al. 2006). Such steeper gradient of leaf N in the canopy increases daily canopy photosynthesis by 20–25%. Both leaf area and tillering capacity should be large at early stages (Laforge and Bueno 2009), but restricted during grain filling to avoid strong mutual shading and to allow radiation to reach deeper leaf layers to realize gains in canopy photosynthesis with increasing leaf N content. In this regard, it is particularly interesting that new plant types have very erect leaf orientation and relatively small number of productive tillers that could be useful to increase canopy photosynthesis (Horton 2000).

**Radiation use efficiency**

Until last century, RUE during preanthesis has not been significantly improved in modern wheat cultivars (Gutierrez-Rodriguez 2000). In irrigated environments, no
differences in RUE or light interception between old and modern cultivars were found, while modern lines were shown to have higher RUE after anthesis (Calderini et al. 1997) largely driven by high sink. Cultivars with cooler canopy temperature exhibit an enhanced gas exchange rate (Ayeneh et al. 2002); hence, cooler canopy can increase crop yields especially under mild stress (Ayeneh et al. 2002). A 2.5°C range in canopy temperature depression at anthesis has been reported for Iranian wheat cultivars (Karimizadeh and Mohannadi 2011). Although diverse mechanisms may theoretically explain genetic variation in canopy temperature (Rebetzke et al. 2012), so far only root depth and capacity has been implicated through direct measurement (Lopes and Reynolds, 2010) and genetic studies in dry residual moisture and hot, irrigated environments (Pinto et al. 2010; Pinto et al. 2014). These studies suggest that RUE can still be improved in both crops and the underlying mechanisms of CTD differing between genotypes need to be explored.

In rice, RUE has reported to increase from old cultivars to modern cultivars. New indica varieties expressed more RUE (2.14 g MJ\(^{-1}\)) than traditional (1.8 g MJ\(^{-1}\)) and japonica varieties (1.8 g MJ\(^{-1}\)) (Boschetti et al. 2006) due to differences in morphophysiological characters (Kiniry et al. 2001). However, RUE in rice is considered to be lower (under aerobic conditions, ~1 g MJ\(^{-1}\); in flooding, ~2.2 g MJ\(^{-1}\)) as compared to the average quantum requirement of C\(_3\) species (2.9 g MJ\(^{-1}\))(Katsura et al. 2010). This may underlie the poor association between RUE and yield superiority of super hybrid rice varieties (Katsura et al. 2008, 2010). Nevertheless, a higher RUE especially during postanthesis can increase thousand-grain weight (Sadras and Lawson 2011). In wheat, photosynthetic capacity can be improved by increasing green leaf area duration and chlorophyll content of tissue, while stomatal conductance and a cooler canopy temperature are indicative of improved RUE (Sadras and Lawson 2011; Xiao et al. 2012).

**Leaf senescence**

In wheat, leaves contribute about 40% to the daily rate of grain N accumulation at mid grain filling (Simpson et al. 1983). Hence, a delayed leaf senescence or stay green, which maintains active photosynthesis for longer period may increase grain yields (Gregersen et al. 2008; Bogard et al. 2011). In contrast, accelerated senescence leads to low C but high N remobilization indicating the plasticity of senescence linked to carbon and N remobilization. Such senescence regulation of C and N could be a viable strategy to alter both C and N remobilization to the actively developing grains. While several patterns of senescence were proposed (Thomas and Howarth 2000), an ideal senescence phenotype in wheat, in general in cereals, still needs to be identified (Gregersen et al. 2008; Bogard et al. 2011). The advent of high-throughput spectral tools such as the Greenseeker has made it more feasible to study total canopy N remobilization dynamics (Lopes and Reynolds 2012).

In rice, a functional stay-green phenotype exhibits higher light conversion efficiency and net photosynthetic rate at saturated light throughout grain filling (Fu and Lee 2012). It also maintains mesophyll conductance due to high chlorophyll content and delays the degradation of photosystem II. Interestingly, the stay-green trait is associated with high root activity so higher capacity to accumulate and maintain high N (Fu et al. 2009) and with higher grain-filling rate under limited radiation conditions (Bueno and Lafarge, unpublished data). These studies suggest that the stay-green trait is beneficial to increase root activity and N accumulation during grain-filling period under favorable environments.

**Nitrogen use efficiency and biological nitrification inhibition**

Genetic gains in NUE under low N supply were associated with the improvements in NUpE (uptake efficiency) in spring wheat and to NUtE (utilization efficiency) in winter wheat (Brancourt-Hulmel 2003), while under high N, both components contribute approximately equally to NUE (Ortiz-Monasterio et al. 1997; Muurinen et al. 2006; Wang et al. 2011). These studies indicate that NUpE may be influenced by G × N interaction, while G × E interaction may influence NUtE. Although leaf senescence kinetics could explain the genetic variability associated with yield variation under low N (Gaju et al. 2011), the strong negative association between NUpE and NUtE highlights the necessity of understanding mechanisms influencing the uptake and utilization of N by crops. Modified root architecture associated with dwarfing genes is one parameter influencing NUE (Wojciechowski et al. 2009; Gooding et al. 2012). Alternatively, BNI (biological nitrification inhibition) may improve NUE. The wild wheat, *Leymus racemosus*, shows higher BNI release rates ranging from 20 to 30 allyl thiourea g\(^{-1}\) root dry wt. day\(^{-1}\) (Subbarao et al. 2007). A locus for high BNI was transferred to cultivated wheat using chromosome addition lines of wild wheat, however, these lines carry many undesirable traits compromising yield potential.

In rice, a large genotypic variation in NUE has been reported, ranging from 38 to 84 kg kg\(^{-1}\) for grain and 100 to 160 kg kg\(^{-1}\) for biomass, probably under aerobic and rainfed management (Korouzas and Ntanos 2003). However, improving NUE for grain in favorable (irrigated) environments has been limited in breeding (Samonte
et al. 2006). Indica cultivars generally have higher NUE for grain than japonica cultivars (Koutroubas and Ntanos 2003). Such large variations could be related to yield potential because high-yielding cultivars show less genetic variation in NUE for grain (Samonte et al. 2006). Moreover, NUE for grain is also influenced by plant height, crop growing duration (short and late maturing cultivars have high NUE for grain than tall and early maturing varieties; Koutroubas and Ntanos 2003), percentage of straw N, grain N concentration, and HI. At HI of 0.5, both straw and grain N equally contribute to NUE for grain; therefore, increasing HI would be associated with NUE for grain. However, genotypic variation in grain N concentration is low (0.95–1.2; Tirol- Padre et al. 1996), therefore, it may not be feasible to improve NUE for grain by reducing grain N concentration using conventional breeding as grain N concentration is affected more by environments than by genotypes (Ladh et al. 1998). Genetic variability for nitrogen metabolism in the developing ear of maize has been suggested to improve yields (Cañas et al. 2012). BNI improves N uptake and NUE due to its inhibitory effects on nitrification (Subbarao et al. 2006). A significant genotypic variation in BNI (<10% to >50%) was reported (Pariasca Tanaka et al. 2010). However, traditional and upland varieties exhibit high BNI activity than the modern irrigated lowland varieties. While nitrification rates in paddy soils are lower compared to typical upland soils, application of synthetic nitrification inhibitors has been shown to increase yields (Li et al. 2009) suggesting that BNI has the potential to improve yields in rice.

Source: sink regulation and lodging resistance

An approach to increase spike dry matter is to increase assimilate partitioning to spike at anthesis (Demotes-Mainard and Jeuffroy 2004). To some extent, dwarf and semi-dwarf cultivars have achieved this by increasing assimilates partitioning to developing spikes (Álvaro et al. 2008; Foulkes et al. 2011). For example, the dry matter partitioning to spikes at anthesis improved from 9% in old cultivars to 27% in modern semi-dwarf durum wheat in Italy and Spain (Álvaro et al. 2008). The genetic range for partitioning to the respective plant organs at anthesis (spikes, 0.12–0.29; leaf lamina, 0.19–0.31; stems and leaf sheath, 0.48–0.63; as a proportion of above-ground biomass; Reynolds et al. 2009) suggests that there is a possibility to increase assimilate partitioning to spikes (Foulkes et al. 2011). However, decreasing plant height further by developing double dwarfs may not provide any additional benefits because of reduced light interception and total biomass (Butler et al. 2005). Moreover, reduced cell elongation and smaller cells have reduced seedling vigor in resource-poor environments (Botwright et al. 2005).

AGB (above-ground biomass) was not systematically increased from old to modern cultivars until relatively recently (China [Zheng et al. 2011]; U.K. [Shearman et al. 2005]; Australia [Sadras and Lawson 2011]; and Mexico [Aisawi et al. 2015]). Increased AGB shows two common trends with yield components: thousand kernel weight or grain weight in cultivars of Australia (Sadras and Lawson 2011) and Henan Province of China (Zheng et al. 2011), and grains per unit area or spikes per unit area in cultivars of U.K. (Shearman et al. 2005) and Shandong of China (Xiao et al. 2012). This suggests that biomass partitioning to yield components has been greatly modified resulting in distinct wheat ideotypes (Álvaro et al. 2008). An increased AGB especially during postanthesis shows strong correlation with yield (Reynolds et al. 2009). Nevertheless, the association between yield and biomass depends heavily on environment. Under stress, there is usually a strong association (Reynolds et al. 2007) but less so often when conditions are favorable. In addition, higher WSC (water-soluble carbohydrates) at anthesis are associated with higher grain weight and lower WSC storage is associated with increased grain number (Rebetzke et al. 2008; Sadras and Lawson 2011). While this trend is consistent with most modern cultivars, often an opposite trend with higher WSC storage and grain number was reported (Shearman et al. 2005; Xiao et al. 2012). These studies did not report any changes in grain weight, hence the associations between WSCs and yield components are unclear.

In rice, there has been a marked increase in HI and grain production by modification of the reproductive structures in the modern cultivars (Evans et al. 1984; Mohapatra et al. 2011) and hybrids (Yang et al. 2007), suggesting an increased partitioning of assimilates to reproductive organs (Lafarge and Bueno 2009). Modern cultivars show increased biomass production over old cultivars. Yield improvement in cultivars released before 1980 was associated with increases in HI, while the yield improvement in the cultivars released after 1980 was associated with increases in biomass (Peng et al. 2000) although it largely depends on comparisons between either semi-dwarf and traditional or among semi-dwarfs (Evans et al. 1984; Yamauchi 1994; Peng et al. 1998). Also, higher yields of hybrids were attributed to both high HI (Bueno and Lafarge 2009) and biomass (Bueno and Lafarge 2009; Zhang et al. 2009) mainly through higher growth rate of priority sinks and higher sink regulation, that is the timing and magnitude of the successive sink activities over crop growth (Bueno et al. 2010), to the favor of stem biomass at the end of the vegetative phase, panicle biomass at the end of the reproductive phase, and remobilization
from stem to panicle at the beginning of the filling phase (Lafarge and Bueno 2009; Bueno et al. 2010). Yet, it was reported that higher yield was attributed to higher biomass accumulation under favorable conditions and to higher HI under suboptimal conditions in the tropics (Peng et al. 2003). A recent study conducted on 32 high-yielding contrasted genotypes (hybrids and inbreds) from diverse geographical origin, and respective of their maturity groups, highlighted, however, that relevant traits were those favoring light capture in the favorable (dry) season and short plant height and delayed leaf senescence in the suboptimal (wet) season (Bueno and Lafarge submitted).

It has been estimated that during heading to maturity, an inbred rice allocates dry matter about 39.5% to panicles, 4.3% to leaves, 33.5% to stems, 6.5% to late tillers, and 6.7% to roots (Kato et al. 2006), suggesting that panicles have high demand for dry matter partitioning (Zhao et al. 2008). Indeed, in favorable conditions, hybrid rice shows a higher partitioning coefficient to culms in the late vegetative and early reproductive while lower partitioning coefficient to culms and higher partitioning coefficient to panicles in the later reproductive phases as compared to inbreds (Lafarge and Bueno 2009). This suggests a strong sink regulation and remobilization capacity with hybrids over the crop cycle (Bueno et al. 2010) through larger biomass partitioning to culm at late reproductive phase and HI. Indeed, in high-yielding recently bred varieties, equipped with large panicles (Fujita et al. 2013), grain filling of late flowering basal spikelets is particularly poor (Panda et al. 2009). Both NSC accumulation in the stems at heading (Fu et al. 2011) and reduction in ethylene concentration in the basal spikelets (Wang et al. 2012) enhances sink strength of the basal spikelets at early grain filling by increasing number of endosperm cells and activities of sucrose synthase and AGPase (adenosine diphosphoglucose pyrophosphorylase). These studies suggest that increasing NSC accumulation in the stems and reducing ethylene production in the basal spikelets will induce fast-synchronized filling within the panicle and higher yield potential. In addition, there is growing evidence that the T6P (trehalose-6-phosphate) plays a central role in regulating carbohydrate and starch metabolism in plants as it indirectly reflects sucrose concentration and has been widely accepted as an indicator of sucrose status (O’Hara et al. 2013; Yadav et al. 2014). By regulating sucrose export to the grain, T6P could then increase sink strength in case of its higher concentration within the grain and has been shown to increase grain yields (Nuccio et al. 2015).

### Lodging resistance

Strategies to raise HI should consider assimilate partitioning to structural organs to avoid any stem and root lodging. Models describing lodging resistance in wheat have highlighted stem thickness, stem wall material strength, and area of root plate in the stem and root lodging resistance (Berry et al. 2003b, 2007) (Fig. 2). Interestingly, lodged plants exhibited stronger source limitation during grain filling than unlodged ones (Acreche and Slafer 2011), highlighting the importance of photosynthetic activity...
during grain filling to avoid any stem or root lodging. Reducing unproductive tiller number could also reduce lodging problems and increase grains per spike (9%) but decrease grain yields (8%) (Duggan et al. 2005; Gaju et al. 2011), as observed in China (Zheng et al. 2011) and Spain (Royo et al. 2006; Álvaro et al. 2008). The net redistribution of N and C to surviving shoots after their death is inconsistent being negative (Berry et al. 2003a) or positive (Thorne and Wood 1987). Although the efficiency of such transport is unknown, a high, early tiller production may enhance grain yields in cereals (Pasuquin et al. 2008).

Rice exhibits a large genetic variability in lodging resistance. Several lodging-related traits were identified (Islam et al. 2007). Lodging resistance was strongly correlated with the culm diameter and wall thickness of the basal internodes (Chuanren et al. 2004). A stem diameter of 5.3–5.5 mm (plant height of 114 cm under typhoon conditions) (Chuanren et al. 2004; Ishimaru et al. 2005) and stem wall thickness of 0.49 (Chuanren et al. 2004) were associated with lodging resistance. While plant height was not necessarily an important factor (Ishimaru et al. 2007), leaf sheath and the extent of wrapping (breaking strength of the shoot was increased by 30–60%) will be an important trait for selection against lodging at all developmental stages as shown in rice (Setter et al. 1994). In addition, mechanical properties such as tensile breaking point and the tensile elastic modulus are associated with lodging resistance (Chuanren et al. 2004). Clearly, while plant height was not necessarily being reduced, increasing stem wall thickness and mechanical properties can contribute to enhance lodging resistance.

**Trait Differences and Similarities Between Wheat and Rice**

Over the last two decades, the linear rate of yield change for wheat has been 25 kg ha⁻¹ year⁻¹, while it is 38 kg ha⁻¹ year⁻¹ for rice. However, the relative rates of yield increase seem to be the same (0.9%) for both crops (Fischer and Edmeades 2010). While reduction in plant height, increased biomass, and HI commonly resulted in yield gain in both crops, increased kernel number per unit area in wheat and increased CGR in rice are largely associated with yield gains (Fischer and Edmeades 2010; Pozo et al. 2014). The CGR between stem elongation and anthesis is increased in wheat particularly in the cultivars of Australia (Sadras and Lawson 2011), while CGR and early vigor (higher LAI) have been significantly increased in rice particularly in hybrids, which increased biomass accumulation and grain yield (Peng et al. 1998; Bueno and Lafarge 2009). In addition, net photosynthesis and RUE have been slightly increased in modern wheat while hybrid rice has a lower net photosynthetic rate during vegetative and grain-filling phases because of lower leaf N concentration (Peng et al. 1998).

Both wheat and rice crop phenophases are responsive to the environment. Breeding efforts have altered vegetative phase durations in both crops. The genetic factors such as earliness per se loci are absent in rice (Faricelli et al. 2009, 2010). Both crops were domesticated in different climates and so differ significantly in their growth conditions; wheat shows appreciably faster growth rate and allocates more biomass to leaves than rice; however, the optimal temperature for biomass production is higher for rice than wheat (Nagai and Makino 2009). While both species significantly contrast in photoperiod sensitivity, the response of developmental processes to optimum temperature is similar in both species and is not affected by either natural selection or artificial breeding (Parent and Tardieu 2012). This would suggest a strong interaction between developmental events and the environment in particular temperature per se.

Both species differ in LAI. Since both crops differ for leaf N content per unit area, they also differ for radiation conversion factor (Mitchell et al. 1998). Interestingly, the correlation between leaf N content and grain number vary with genotype and growth conditions in wheat, while it is reported to be independent in rice (Wada and Matsushima 1962; Makino 2011). Since the N concentration in the straw and grain in rice is lower than wheat, the NUE for biomass and grain is higher in rice (Ladha et al. 1998). Such differential N levels in the biomass and its allocation influence leaf senescence in both species. Furthermore, awns are absent or short in cultivated rice (Luo et al. 2013; Hua et al. 2015), while awns have been shown to contribute to net ear photosynthesis in wheat (Evans et al. 1972; Sanchez-Bragado et al. 2014).

The source–sink regulation is different between two crops. In rice (hybrids), the source capture is relatively higher than that of wheat, while the sink strength is more efficient during grain-filling period. For example, biomass production before stem elongation was higher in rice than in wheat due to high early vigor and CGR (Bueno and Lafarge 2009; Bueno et al. 2010). While assimilate partitioning to panicles over other organs is larger in rice after heading (Kato et al. 2006; Kumar et al. 2006), this has been shown to be limited in wheat. Although the remobilized carbon reserve percentage and its contribution to grain appears higher in wheat than rice, the HI is lower in wheat than in rice probably due to the existence of alternative sinks (Yang and Zhang 2006). Moreover, stem structural differences exist; the width of the stem wall in wheat (0.65 mm) is greater than that in rice (0.45 mm), while stem diameter in wheat (4.7–4.9) is less than the stem diameter of rice (5.3–5.5 mm) (Chuanren et al. 2004; Reynolds et al. 2009; Wu et al. 2011).
By virtue of the growth conditions, the crop management practices and environmental influences on crop growth and development differ between two species. While the conventional crop management practices in wheat have largely been unmodified, in rice, sound management practices have been in practice. Among others, planting patterns, planting density, and spacing have resulted in more open canopies, which increased the overall performance of the rice crop canopy in terms of leaf angle, tiller dynamics, light interception, and photosynthetic rate (Table 1).

### A Research Gap Analysis for Future Research Priorities

Both, wheat and rice, crops have been explored in different ways. However, the genetic gains in yields of wheat and rice across the globe are less than 1% per annum. The following opportunities, but not limited to, may be promising to further increase in genetic gains in yields of both crops.

### Yield components

Recent gains in crop yields has been achieved through larger panicle size with higher grain number per panicle in rice (Ying et al. 1998), while increased grain number predominantly enhanced grain yields in wheat (Pozo et al. 2014; Zhou et al. 2014). Historical data, however, support that grain number (either through extra panicles/spikes or extra grain per panicle/spike) needs to be increased in both crops. In addition, a number of genes important for yield formation have been identified in both crops (see review, Valluru et al. 2014, table 3). For example, genes controlling leaf and panicle size (SPIKE) and grain assimilate accumulation (GIF1 and Rg5) have been identified in rice (Fujita et al. 2013; Ishimaru et al. 2005). In wheat, structural modification to AGPase gene has been proposed to increase starch accumulation in grain (Meyer et al. 2007). Furthermore, GS3 and Gnl controlling grain number and weight have been identified in rice. These genes have largely not been explored in both crops (Valluru et al. 2014).

### Harvest index

Increasing HI is not a strong candidate, as it is already high, above 0.5 in high-yielding varieties and sink regulation is also high and effectively expressed in hybrid rice (Ying et al. 1998; Lafarge et al. 2010). In wheat, an average HI reported is 0.45 (Zheng et al. 2011; Xiao et al. 2012). Increasing HI above the current values may lead to high risk of lodging unless plants have a lower center of gravity (i.e., are shorter), but then this would reduce light interception, consequently reducing biomass. More rigid stems involving anatomical changes like increasing concentrations of structural carbohydrates like lignin could be a long-term breeding objective.

### Biomass

Increasing biomass accumulation may be a promising avenue in both crops. In rice, hybrids have significantly

| Trait          | Wheat References                                                                 | Rice References                                                                 | References |
|----------------|---------------------------------------------------------------------------------|---------------------------------------------------------------------------------|------------|
| Preanthesis period | Preanthesis period reduced in modern genotypes                                 | Preanthesis period reduced in modern genotypes                                 | Peng et al. (2000) |
| Earliness per se | Earliness per se locus are present in wheat                                     | Earliness per se locus are reported to be absent in rice                       | Faricelli et al. (2009); (Faricelli et al. 2010) |
| LAI             | LAI is lower                                                                     | LAI is higher                                                                   | Zheng et al. (2008); Zhang et al. (2009) |
| RUE             | RUE increased in modern cultivars particularly during postanthesis period         | RUE increased in modern cultivars                                              | Boschetti et al. (2006) |
| Awns            | Awns are reported to contribute to net ear photosynthesis                        | Awns are absent or short in the cultivated rice                                | Luo et al. (2013); Hua et al. (2015) |
| Leaf senescence | Leaf senescence is slightly delayed in modern cultivars due to the presence of nonfunctional grain protein content gene | Indica and Japonica exhibit different leaf senescence patterns, early and late, respectively | Abdelkhalik et al. (2005) |

LAI, leaf area index; RUE, radiation use efficiency.
higher aboveground biomass (10–12%) than inbred cultivars (Peng et al. 2008); (Bueno and Lafarge 2009). In wheat, biomass has, on average, a lower genetic gain (0.3), suggesting that biomass production may still be the main way forward to increase yields (Fischer and Edmeades 2010; Sylvester-Bradley et al. 2012). However, the key question that needs to be addressed would be whether the key driver of yield is, biomass accumulation or sink strength? Would higher biomass accumulation induce better sink strength or better sink strength support higher biomass accumulation? A crossing program at CIMMYT that combines high source with high sink traits is showing genetic gains for yield and biomass in international trials, suggesting that source and sink balance is important (Reynolds et al. 2015).

**Leaf and crop photosynthesis**

Improving TPU is promising in both crops. If sink strength is higher, photosynthate could be exported quicker from the source leaves to the sink to reduce downregulation of photosynthesis. Similarly, improving mesophyll conductance could be promising. Changes in leaf anatomy could facilitate CO$_2$ diffusion between the leaf compartments to also reduce downregulation of leaf photosynthesis. Further reducing night respiration could be important although few studies report limited diversity for this trait and limited effect on the overall performance (Peraudeau et al. 2015a,b). If the plant could uptake more N (via higher fertilization and/or better root system and/or improved water management for better uptake), with N being more available to fill grain, then so leaf senescence can be delayed and crop photosynthesis maintain for a longer duration. Numerous proposed avenues for enhancing photosynthesis rate in C$_3$ species are given in Figure 1.

**Sink strength**

Increasing sink strength through improving panicle/spike architecture could be important. Reducing hormonal (especially ethylene) regulation that delays the filling of basal spikelets could be promising for efficient grain filling. In addition, increasing the role of T6P, as sugar regulator, could facilitate the export of carbohydrates to the panicles/spikes.

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**Conflict of Interest**

None declared.

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