Introducing the metabolic engineering of bread wheat to improve grain iron concentration and bioavailability

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Summary

Bread wheat (Triticum aestivum L.) is cultivated on more land than any other crop and produces a fifth of the calories consumed by humans. Wheat endosperm is rich in starch yet contains low concentrations of dietary iron (Fe) and zinc (Zn). Biofortification is a micronutrient intervention aimed at increasing the density and bioavailability of essential vitamins and minerals in staple crops; Fe biofortification of wheat has proved challenging. In this study we employed constitutive expression (CE) of the rice (Oryza sativa L.) nicotianamine synthase 2 (OsNAS2) gene in bread wheat to up-regulate biosynthesis of two low molecular weight metal chelators – nicotianamine (NA) and 2’-deoxymugineic acid (DMA) – that play key roles in metal transport and nutrition. The CE-OsNAS2 plants accumulated higher concentrations of grain Fe, Zn, NA and DMA and synchrotron X-ray fluorescence microscopy (XFM) revealed enhanced localization of Fe and Zn in endosperm and crease tissues, respectively. Iron bioavailability was increased in white flour milled from field-grown CE-OsNAS2 grain and positively correlated with NA and DMA concentrations.

Keywords: nicotianamine, 2’-deoxymugineic acid, biofortification, Zn, Caco-2, X-ray fluorescence microscopy.

Introduction

Micronutrient mineral deficiencies affect over two billion people worldwide with women and children most acutely at risk (Beal et al., 2017; Lopez et al., 2016). Iron (Fe) deficiency is the leading cause of anaemia, a condition that impairs cognitive development and work productivity and increases maternal and child mortality (Kassebaum et al., 2017; Lopez et al., 2016). Zinc (Zn) deficiency causes a variety of disorders including stunted growth in children. Human Fe and Zn deficiencies are most prevalent in SEA (Southeast Asia) and MENA (Middle East and North Africa), in part due to high consumption of micronutrient-poor staple crops such as wheat (Triticum aestivum L.), rice (Oryza sativa L.) and maize (Zea mays L.) (Beal et al., 2017). Rising atmospheric CO2 concentrations will likely decrease Fe and Zn concentrations in C3 grains and could further exacerbate micronutrient deficiencies in SEA and MENA (Myers et al., 2014; Smith et al., 2017).

The wheat grain is comprised primarily of starch (60%–75%) and protein (12%–14%) with only low concentrations of Fe, Zn and other micronutrients (Shewry, 2009; Velu et al., 2014). Iron and Zn are remobilized from senescing vegetative tissues and/or translocated from roots during grain filling (Maillard et al., 2015; Pottier et al., 2014; Waters et al., 2009) and transported into developing grain via the vascular bundle in the crease region. In mature grain, the highest concentrations of Fe and Zn are found in the aleurone cells in conjunction with compounds such as phytic acid, polyphenols and fibre that inhibit bioavailability, while only low Fe and Zn concentrations are localized to endosperm tissues (Brouns et al., 2012; De Brier et al., 2016; Moore et al., 2012; Schiemmer et al., 2009; Singh et al., 2013; Stomph et al., 2011). Grain milling removes inhibitors of micronutrient bioavailability but also a significant proportion of the grain Fe and Zn (Hourston et al., 2017; Zhang et al., 2010).

Wheat biofortification aims to complement micronutrient supplementation and fortification programs by improving grain micronutrient density and/or bioavailability (Bouis et al., 2011; Prentice et al., 2017). Generation of Fe biofortified wheat through conventional breeding is hindered in modern wheat cultivars by inherently low grain Fe concentrations and a lack of genomic resources for this trait (Borrill et al., 2014; Velu et al., 2014), indicating that genetic engineering strategies may be required to generate novel genetic variation (Connon et al., 2017; Singh et al., 2017).

Nicotianamine (NA) is a non-protein amino acid formed by trimerization of S-adenosylmethionine (SAM) in a process mediated by NA synthase (NAS) enzymes (Clemens et al., 2013). Nicotianamine plays key roles in the chelation and transport of metals such as Fe, Zn, manganese (Mn) and copper (Cu) in all higher plants (Takahashi et al., 2003; von Wieren et al., 1999). In graminaceous plant species, NA serves an additional purpose as the precursor to biosynthesis of 2’-deoxymugineic acid (DMA), a root-secreted phytosiderophore that chelates ferric Fe (Fe3+) in the rhizosphere and is subsequently re-absorbed. Nicotianamine and/or DMA are also the main chelators of Fe in white wheat
flour (Eagling et al., 2014a), and increased NA is known to increase Fe bioavailability in polished rice (Lee et al., 2012; Trijatmiko et al., 2016; Zheng et al., 2010). For these reasons, increased NA biosynthesis has emerged as a popular strategy for Fe biofortification of cereals (Johnson et al., 2011; Lee et al., 2012; Singh et al., 2017; Trijatmiko et al., 2016).

In this study of glasshouse- and field-grown plants, we demonstrate that constitutive expression of the rice OsNAS2 gene (CE-OsNAS2) in bread wheat causes not only significant Fe and Zn enrichment of whole grain and flour fractions but also increased concentrations of NA and DMA that are positively correlated with Fe bioavailability in CE-OsNAS2 white flour. Interestingly, our results indicate that Fe bioavailability is higher in NA and DMA-enriched white flour regardless of Fe concentration and that white flour NA and DMA concentrations are more critical than absolute Fe concentration in determining Fe bioavailability. These results provide unique insights into cereal Fe biofortification and highlight the importance of metabolic engineering strategies that enhance not only micronutrient density but also promoters of micronutrient bioavailability.

Results

Generation of independent transformation events with constitutive OsNAS2 expression and selection of biofortified material for advanced nutritional analysis

Bread wheat cultivar (cv.) Bobwhite transformatants with constitutive expression (CE) of the rice nicotianamine synthase 2 (OsNAS2) gene were generated through biolistic transformation of a cassette containing the OsNAS2 gene under regulatory control of the maize ubiquitin 1 promoter (UBI-1) (Figure 1a). Six independent CE-OsNAS2 transformation events termed CE-1, CE-5, CE-7, CE-8, CE-13 and CE-15 were regenerated from tissue culture and Southern blot analysis showed that insert copy number ranged from 1 to 7 among the six events (Figure 1b). Automated imaging facilities at The Plant Accelerator (Adelaide, Australia) were used to phenotype T1 progeny of the six events; two events (CE-1 and CE-5) showed no phenotypic differences from a null segregant (NS) line derived from CE-1 nor wild-type (WT) wheat with respect to shoot area, plant height, total grain number and thousand grain weight (Figure 1c–g). Elemental analysis showed that four of the CE-OsNAS2 events (CE-1, CE-8, CE-13, CE-15) produced T2 grain with significantly increased Fe and Zn concentrations relative to NS and WT and one CE-OsNAS2 event (CE-5) produced T2 grain with significantly increased Zn concentration relative to NS and WT (Figure 1h,i). Wild-type plants did not differ from NS plants for any trait measured and had mean values of 1465 cm² shoot area, 65 cm plant height; 375 total grain number, 38 g thousand grain weight, 40 µg/g DW grain Fe and 78 µg/g DW grain Zn.

Based on low insert copy number, no difference in plant phenotype, and increased grain Fe and Zn concentrations from the T1 to T2 generations, homoygous progeny of the double-insert event CE-1 and corresponding NS line were selected for a range of additional analyses from the T1 to T6 generations as depicted in Figure S1. Glasshouse-grown CE-1 seedlings displayed high OsNAS2 expression in roots and shoots (Figure 1j,k). Expression of a range of endogenous TaNAS, TaNAAAT and TaDMAS genes involved in NA and DMA biosynthesis was not significantly different between CE-1 and NS seedlings, however, a trend towards slightly reduced expression was detected in CE-1 (Figure S2). The CE-1 and NS seedlings did not differ with respect to shoot Fe, Zn and DMA concentration while shoot NA concentration was 1.3-fold higher in CE-1 seedlings (Figure S3). Nicotianamine concentration was 15-fold higher in CE-1 mature grain relative to NS (Figure 1l).

Constitutive OsNAS2 expression alters Fe and Zn accumulation in grain, bracts, rachis and flag leaf at multiple points during grain filling

The main stem flag leaf, rachis, bracts and grain of glasshouse-grown CE-1 and NS plants were harvested at five timepoints post anthesis to determine Fe and Zn accumulation patterns in vegetative and grain tissues during grain filling. The CE-1 grain had significantly higher Fe content at 5–8 days after anthesis (DAA) and 19–21 DAA relative to NS grain (P = 0.006 and P = 0.046; respectively) and showed non-significant trends towards higher grain Fe and Zn content at maturity (Figure 2a, b). Bracts of CE-1 plants had significantly higher Fe and Zn contents at 12–15 DAA relative to NS (P = 0.013 and P ≤ 0.001; respectively) but did not differ from NS bracts at maturity (Figure 2c,d). Rachis of CE-1 plants had significantly higher Fe content at 5–8 DAA (P = 0.012), and significantly higher Zn contents at 12–15 DAA and 19–21 DAA (P = 0.014 and P = 0.020, respectively) but did not differ from NS rachis at maturity (Figure 2e,f). Flag leaf of CE-1 and NS plants contained similar Fe content throughout grain filling. Conversely, CE-1 flag leaf contained significantly lower Zn content at 19–21 DAA and 26–29 DAA relative to NS (P = 0.009 and P = 0.038, respectively) but did not differ from NS flag leaf at maturity (Figure 2g,h).

Iron and Zn concentration data followed similar trends to the content data although more significant differences were detected (Table S1). Vegetative and grain tissue biomass was largely unchanged between CE-1 and NS plants throughout anthesis and was not significantly different for any tissue at maturity (Table S2).

Constitutive OsNAS2 expression increases Fe accumulation in grain endosperm tissue and Fe and Zn accumulation within the crease region

Elemental maps of Fe and Zn in transverse cross-sections of two representative CE-1 and NS grain showed that CE-1 grain had higher Fe signal intensities in all grain tissue types relative to NS with the largest difference detected in endosperm tissues (Figure 3c,d). The Zn signal in both CE-1 and NS grain was localized to the aleurone and crease regions and was not detectable in the endosperm. Line scans across the mid-section demonstrated that CE-1 grain had higher Fe signal intensities relative to NS in all regions of the grain, particularly in the endosperm, while CE-1 grain had slightly higher Zn signal intensities in the aleurone cells relative to NS (Figure 3e,f). The line scans also revealed different Fe and Zn distribution patterns within the crease region of NS and CE-1 grain, with Zn signals appearing as two distinct peaks on either side of the crease while Fe signals clustered into one central peak. Tri-colour elemental maps of Fe, Zn and Cu distribution within the crease region demonstrated that these distinct peaks result from extensive Zn localization in modified aleurone cells bordering the crease in contrast to prominent Fe localization in the centrally located nucellar projection (Figure S4). While CE-1 and NS grain showed similar Zn signal intensities in the modified aleurone cells of the crease, the CE-1 grain had slightly higher Zn signal intensity in the
nucellar projection relative to NS (Figure 3f). The CE-1 and NS grain had similar distribution and signal intensities for Cu and Mn (Figure S5).

Tri-colour elemental maps of Fe, Zn and phosphorus (P) near the grain edge of one representative CE-1 and NS grain demonstrated that the Fe signal in CE-1 grain was enhanced within pericarp, aleurone and endosperm regions relative to NS grain (Figure 4c,d). The Zn signal in CE-1 grain was enhanced primarily within the pericarp and aleurone cells, relative to the NS, and co-localized with Fe in that region. The P signal in both NS and CE-1 grain did not differ in intensity nor distribution and was localized to the aleurone cells. The trends observed in CE-1 and NS tri-colour maps were further confirmed by line scans (Figure 4e,f). In both CE-1 and NS grain, P was localized exclusively to the aleurone cells with equal signal intensity, indicating that the significant enrichment of Fe in CE-1

**Figure 2** Fe and Zn content in vegetative and grain tissues during grain filling of CE-OsNAS2 and NS wheat lines. Fe and Zn content (µg) in NS (open circles) and CE-1 (closed circles) plant tissues between 5–8, 12–15, 19–21 and 26–29 days after anthesis (DAA) as well as at maturity. (a, b) grain; (c, d) bracts; (e, f) rachis; and (g, h) flag leaf tissues were sampled for Fe and Zn content, respectively. Each symbol represents mean ± SEM of at least 3 biological replicates. Asterisks denote the significance between NS and CE-1 for $P < 0.05$ (*), $P \leq 0.01$ (**), $P \leq 0.001$ (***) as determined by student’s t-test.

**Figure 1** Generation and characterization of independent bread wheat transformation events constitutively expressing the rice nicotianamine synthase 2 (OsNAS2) gene. (a) Schematic representation of the T-DNA construct. RB and LB: right and left borders, respectively; UBI-1: maize ubiquitin 1 promoter; OsNAS2: rice nicotianamine synthase 2 gene (LOC_Os03 g19420); nos-ter: nopaline synthase terminator; 35S-pro: dual promoter of 35S cauliflower mosaic virus gene; hyg: hygromycin phosphotransferase gene; 35S-ter: terminator of 35S cauliflower mosaic virus gene. (b) Southern blot analysis of T1 wheat events to determine T-DNA insertion number. Dral: restriction endonuclease; + indicates positive control. (c) Representative plants of null segregant (NS) and the 6 transformation events (CE-OsNAS2) 100 days after sowing. (d–g) Projected shoot area (cm²), plant height (cm), total grain number and thousand grain weight (TGW) of NS (white), leading CE-OsNAS2 event (CE-1, orange) and other CE-OsNAS2 events (grey) at the T1 generation. Bars represent mean ± SEM of at least 7 biological replicates. (h, i) Iron and zinc concentration (µg/g DW) in T2 whole grain of NS, CE-1 and other CE-OsNAS2 events. Bars represent mean ± SEM of at least seven biological replicates. (j, k) Relative quantification of OsNAS2 transcript levels in NS and CE-1 shoots and roots. Bars represent mean ± SEM of three bulked biological replicates, each with three technical replicates of quantitative RT-PCR. (l) Nicotianamine concentration (µmol/mg) in whole grain of NS and CE-1 plants at the T2 generation. Bars represent mean ± SEM of three biological replicates. Asterisks denote the significance between NS and CE-OsNAS2 events for $P < 0.05$ (*), $P \leq 0.01$ (**), $P \leq 0.001$ (***) as determined by student’s t-test. Wild-type plants did not differ from NS plants for any trait measured and therefore only NS data is presented.

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endosperm was not associated with phytic acid (a P containing compound). Sulphur (S) distribution and intensity (a proxy for protein) appeared slightly reduced in the CE-1 aleurone, relative to NS, but similar to NS in the endosperm (Figure 4e,f).

Constitutive *OsNAS2* expression does not alter the phenotype of field-grown plants and increases Fe, Zn, NA and DMA concentrations in grain and white flour

Replicated plots of three CE-1 sibling lines designated CE-1.1, CE-1.2 and CE-1.3 and the NS were evaluated in confined field trials at Katanning and Merredin, Western Australia in 2015. With the exception of CE-1.1 at Merredin, phenotype of the CE-1 sibling lines and NS did not differ with respect to plant height, spike number, biomass, thousand grain weight (TGW) and grain yield (Table S3). Whole grain Fe concentrations were significantly higher for CE-1.1 at both Katanning \((P = 0.042)\) and Merredin \((P = 0.002)\) relative to NS (Figure 5a). Whole grain Zn and P concentrations did not differ between the three CE-1 sibling lines and NS at both Merredin and Katanning. All lines (CE-1 and NS) had higher whole grain Zn and P concentrations at Katanning relative to Merredin, possibly due to lower TGW at Katanning (Figure 5b,c, Table S3). Analysis of NA and DMA showed that whole grain NA concentrations were significantly higher for CE-1.1 and CE-1.2 at Katanning \((P = 0.005\) and \(P \leq 0.001\), respectively), and for all three CE-1 sibling lines at Merredin \((P = 0.004, P = 0.008\) and \(P = 0.016\), respectively), relative to NS. Whole grain DMA concentrations were significantly higher for CE-1.1 and CE-1.2 at Katanning \((P = 0.014\) and \(P = 0.045\), respectively), and CE-1.2 and CE-1.3 at Merredin \((P = 0.022\) and \(P \leq 0.001\), respectively), relative to NS (Figure 5d,e).

White flour Fe concentrations were significantly higher for CE-1.1 and CE-1.2 at Katanning \((P \leq 0.001\) and \(P = 0.012\), respectively) relative to NS (Figure 6a) but did not differ between any line at Merredin. White flour Zn concentrations were significantly higher for all three CE-1 sibling lines at Katanning \((P = 0.002, P = 0.010\) and \(P = 0.036\), respectively) relative to NS (Figure 6b) but did not differ between any line at Merredin.

Figure 3 Distribution of Fe and Zn in CE-*OsNAS2* and NS wheat grain. (a) Grain position where transverse cross-sections were made. (b) Location of major tissue types in transverse section of wheat grain. (c) Bright field images of two representative NS grain sections and corresponding XFM elemental maps of Fe and Zn. (d) Bright field images of two representative CE-1 grain sections and corresponding XFM elemental maps of Fe and Zn. Colour bar represents high (white) and low (black) elemental concentration. White boxes in the bright field images represent areas in two grain sections (one each for NS and CE-1) used to generate line scans. (e) Line scans showing the distribution and signal intensity of Fe and Zn across NS grain. (f) Line scans showing the distribution and signal intensity of Fe and Zn across CE-1 grain. Units for the y-axis are elemental counts per pixel.
and Katanning (Figure 6a–c). White flour NA concentrations were significantly higher for CE-1.1 and CE-1.2 at Katanning (P = 0.004 and P ≤ 0.001, respectively) and for all three CE-1 sibling lines at Merredin (P ≤ 0.001, P = 0.002 and P = 0.002, respectively), relative to NS (Figure 6d). White flour DMA concentrations were significantly higher for all three CE-1 sibling lines at both Katanning (P = 0.010, P = 0.020 and P = 0.035, respectively) and Merredin (P = 0.006, P ≤ 0.001 and P = 0.002, respectively) relative to NS (Figure 6e).

Iron bioavailability is increased in white flour containing increased concentrations of NA and DMA

Iron bioavailability in CE-1.1, CE-1.2, CE-1.3 and NS whole grain and white flour produced from field-grown grain was determined through ferritin assays of in vitro Caco-2 cells incubated with flour digests. The levels of bioavailable Fe in CE-1.1, CE-1.2, CE-1.3 and NS whole grain flour were negligible and did not differ significantly, a result likely due to phytic acid and other inhibitory compounds in the outer layers of wheat grain (Figure 5f). By contrast, levels of bioavailable Fe were significantly increased in CE-1.1, CE-1.2, CE 1.3 white flour digests from Katanning (P = 0.005, P = 0.009 and P = 0.045, respectively), and CE-1.1 and CE-1.2 white flour digests from Merredin (P = 0.050 and P = 0.010, respectively) relative to NS (Figure 6f). White flour Fe bioavailability was significantly correlated with NA concentration (r = 0.711, P = 0.048) and DMA concentration (r = 0.812, P = 0.014). White flour Fe bioavailability was not significantly correlated with concentrations of Fe (r = 0.242, P = 0.563), Zn (r = -0.132, P = 0.755) and P (r = -0.433, P = 0.284) (Table 1).

**Discussion**

S-adenosylmethionine (SAM) is an ubiquitous cosubstrate involved in the production of polyamines, ethylene, biotin, NA and DMA in plants (Mao et al., 2015; Roeder et al., 2009). Because increased NAS gene activity diverts more endogenous SAM towards NA and DMA production, it is important to carefully phenotype Fe and Zn biofortified plants with constitutive NAS expression to identify any pleiotropic effects that may negatively impact on plant phenotype. In our study we analyzed multiple CE-OsNAS2 transformation events in several glasshouse experiments, and the lead CE-1 event in multi-location field trials.
Figure 5  Whole grain nutrition of field grown CE-1 and NS wheat lines. Nutrient and metabolite concentrations in whole grain samples of NS (white) and three CE-1 sibling lines (CE-1.1, 1.2 and 1.3, grey) at the T6 generation. (a–c) Whole grain Fe, Zn and P concentrations (µg/g) of NS and CE-1 plants grown at Katanning and Merredin field sites. (d, e) Whole grain NA and DMA concentrations (µmol/mg) of NS and CE-1 plants grown at Katanning and Merredin field sites. (f) Whole grain Fe bioavailability of NS and CE-1 plants grown at Merredin field site. Bars represent mean ± SEM of 3 biological replicates. Asterisks denote the significance between NS and each CE-1 wheat line for $P < 0.05$ (*), $P \leq 0.01$ (**), $P \leq 0.001$ (***).
**Figure 6** White flour nutrition of field grown CE-1 and NS wheat lines. Nutrient and metabolite concentrations in white flour samples of NS (white) and three CE-1 sibling lines (CE-1.1, 1.2 and 1.3, grey) at the T6 generation. (a–c) White flour Fe, Zn and P concentrations (µg/g) of NS and CE-1 plants grown at Katanning and Merredin field sites. (d, e) White flour NA and DMA concentrations (µmol/mg) of NS and CE-1 plants grown at Katanning and Merredin field sites. (f) White flour Fe bioavailability of NS and CE-1 plants grown at Merredin and Katanning field sites. Bars represent mean ± SEM of 3 biological replicates. Asterisks denote the significance between NS and each CE-1 wheat line for $P < 0.05$ (*), $P \leq 0.01$ (**), $P \leq 0.001$ (***).
barrier to endosperm loading (Johnson et al., 2011; Kyriacou et al., 2014). Determining whether differences in transport mechanisms or sink strength account for the Zn distribution differences between wheat and rice grain merits further investigation.

While flour milled from CE-1 grain contained significantly higher concentrations of NA and DMA at both Merredin and Katanning field sites, relative to NS, while white flour Fe and Zn concentrations only differed from NS at Katanning (Figure 6a–e). The NS and CE-1 wheat plants were larger and higher yielding at the Merredin field site (Table S3), which may have had a dilution effect on grain micronutrient concentrations and thereby minimized differences between the genotypes. This result demonstrates the importance of multi-location field trials for accurate assessment of wheat Fe and Zn biofortification traits, yet also uniquely enabled us to examine the effects of NA and DMA as promoters of Fe bioavailability (Eagling et al., 2014b; Glahn et al., 1998; Lee et al., 2012; Tako et al., 2016) under differing Fe concentrations. We used the in vitro Caco-2 cell line model to measure Fe bioavailability in NS and CE-1 white flour produced at both field sites and found that white flour NA and DMA concentrations were significantly and positively correlated with Fe bioavailability at both sites while white flour Fe concentration was not significantly correlated with Fe bioavailability (Figure 6, Table 1). This result was most evident at the Merredin field trial, where white flour Fe concentrations of two CE-1 sibling lines (CE-1.2 and CE-1.3) were virtually identical to that of NS, however, CE-1.2 white flour contained 1.7-fold more bioavailable Fe than NS (Figure 6a and f). In addition to Fe, white flour P concentrations were unchanged between NS and CE-1 sibling lines, indicating that altered white flour phytate/Fe ratios are not likely to be responsible for the increase in Fe bioavailability (Figure 6c). These results reinforce the finding that NA and/or DMA bind Fe in a readily bioavailable form within wheat white flour (Eagling et al., 2014a) and suggests that NA and/or DMA mediate increased uptake of Fe into human cells independent of endosperm Fe concentration. As such, NA and DMA can be viewed as important phytonutrients that increase Fe availability in plant foods (Martin and Li, 2017; Nozoye, 2018), and metabolic engineering strategies focused on increasing their concentrations could help offset the effects of decreased mineral concentrations forecast for C3 grains by mid-century.

### Methods

#### Vector construction and generation of wheat transformation events

The full-length coding sequence of OsNAS2 (LOC_Os03g19420) was PCR amplified from rice (Oryza sativa L.) cv. Nipponbare genomic DNA (Johnson et al., 2011). Recombination into a modified pMDC32 vector (Curtis and Grossniklaus, 2003) with the hygromycin phosphotransferase plant-selectable marker gene placed OsNAS2 under transcriptional control of the maize (Zea mays L.) ubiquitin 1 promoter. Particle bombardment of the construct into immature wheat (Triticum aestivum L.) cv. Bobwhite embryos (1.0–1.5 mm in length) was performed at the University of Adelaide (Adelaide, Australia) using established protocols (Kovalchuk et al., 2009). Plants were grown in glasshouse conditions (12 h photoperiod, 23 °C day/12 °C night, 50% humidity) in soil (coconut peat and sand mixture) with complete fertilizers.

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| Table 1 Pearson’s correlation analysis of NA, DMA, Fe, Zn and P concentrations in NS and CE-1 white flour and ferritin production in Caco-2 cells |
|-----------------|--------|--------|--------|--------|--------|
|                  | DMA (µmol/mg) | Fe (µg/g) | Zn (µg/g) | P (µg/g) | Ferritin (ng/mg protein) |
| NA (µmol/mg)    | 0.912**    | 0.440   | 0.239   | −0.180  | 0.711*   |
| DMA (µmol/mg)   | 0.002   | 0.276   | 0.569   | 0.670   | 0.048   |
| Zn (µg/g)       | 0.355   | 0.132   | 0.145   | 0.812*  |
| Fe (µg/g)       | 0.389   | 0.755   | 0.732   | 0.014   |
| Zn (µg/g)       | 0.893**  | 0.627   | 0.242   |
| P (µg/g)        | 0.003   | 0.096   | 0.563   |
| **P < 0.05, **P ≤ 0.01, ***P ≤ 0.001** |

In Western Australia, and found that important agronomical traits such as shoot area, plant height, total grain number and TGW did not differ from NS in most cases (Figure 1, Tables S2 and S3). These results are similar to those reported for glasshouse and field-grown 35S::OsNAS2 rice (Johnson et al., 2011; Trijatmiko et al., 2016), as well as glasshouse grown UBI-1::OsNAS2 wheat (Singh et al., 2017), and demonstrate that constitutive expression of the OsNAS2 gene to biofortify bread wheat with Fe and Zn does not negatively impact on plant phenotype and yield. Determining whether the changes observed in Fe and Zn accumulation patterns during grain filling (Figure 2) relate to increased remobilization of Fe and Zn from vegetative to grain tissues in CE-OsNAS2 plants will require further analysis using radioactive or stable Fe and Zn isotopes (Pottier et al., 2014).

The most striking result from our synchrotron X-ray fluorescence microscopy (XFM) analyses was the increased Fe signal in endosperm tissue of CE-1 grain relative to NS (Figures 3d and 4d). Line scans showed that the increased Fe signal extended throughout CE-1 endosperm and did not co-localize with P (Figures 3f and 4f), indicating that Fe is not associated with phytic acid in this tissue nor the inhibitory effects that phytic acid has on mineral absorption in humans (Hurrell, 2003). By contrast, both CE-1 and NS grain showed high Zn signal co-localized with P in aleurone cells and essentially no Zn signal in the endosperm (Figure 3f and 4f). The observed differences in grain Fe and Zn localization demonstrate that wheat white flour contains Zn that is derived almost entirely from outside of the endosperm. We postulate that the wheat crease, which is difficult to remove through milling (Campbell, 2007), represents the major source of Zn in white flour. Line scans across the crease region showed similar Zn signal intensities in CE-1 and NS modified aleurone cells yet higher Zn signal intensity in the CE-1 nucellar projection (Figure 3e–f; Figure 5g), a tissue which has been shown to contain NA-bound Fe (De Brier et al., 2016), and this difference likely accounts for the observed Zn concentration increase in CE-1 white flour (Figure 6b). Interestingly, synchrotron XFM studies of rice grain reveal a radically different Zn distribution pattern that extends throughout the endosperm and shows no obvious
Genomic DNA (10 μg) was isolated from CE-OsNAS2 leaf tissue and digested with Dral and HindIII restriction enzymes. Restriction fragments were separated by gel electrophoresis (0.8% agarose) alongside a positive barley control and blotted to a nylon membrane. Two independent hybridizations of a 32P-labelled probe to both nopaline synthase terminator and dual 35S promoter were performed using established protocols (Pallotta et al., 2014).

Automated phenotyping analysis

Grain was sown in white plastic pots (14 × 19 cm) containing 2.5 kg of soil mixture (equal parts clay-loam soil and coconut peat) and Osmocote® fertilizer. Plants were maintained under glasshouse conditions (12 h photoperiod, 24 °C day/18 °C night, 50%-90% humidity) in the phenotyping platform of The Plant Accelerator (Adelaide, Australia). Projected shoot area and plant height were measured 100 days after sowing using the conveyor automated imaging system (Berger et al., 2012). Grain number and TGW were manually determined at harvest.

Inductively coupled plasma optical emission spectrometry (ICP-OES)

Plant tissues were submerged in 0.1% Tween 20 solution, rinsed with dH2O and oven dried for 48 h at 60°C before grinding to a powder using an IKA tube mill (www.ika.com). Inductively coupled plasma optical emission spectrometry (ICP-OES) analysis was conducted at Waite Analytical Services (Adelaide, SA, Australia), the Robert W. Holley Centre for Agriculture and Health (USDA-ARS, Ithaca, NY) and the CSBP Soil and Plant Analysis Laboratory (Perth, WA, Australia).

Quantitative reverse transcription PCR (qRT-PCR)

Shoot and root tissues (without the crown) of 4-week-old plants were separated, cleaned with dH2O and snap frozen. Three plants of each genotype (representing one biological replicate) were combined and total RNA was extracted from pulverized frozen plant tissue (100–150 mg) using TRIzol Reagent (Life Technologies, Carlsbad, CA) and a commercial kit (Direct-zol®; ZymoResearch, Irvine, CA). Genomic DNA was removed from RNA (2 μg) using a DNase I treatment (Promega, Madison, WI) and reverse transcription was performed using a commercial kit (Bioline).

Consensus primers were designed to amplify homologous groups of TaNAS, TaNAAT and TaDMAS gene families using Primer3 (http://bioinfo.ut.ee/primer3-0.4.0) software. Each biological replicate was analyzed in triplicate and transcripts were quantified against four replicates of 10-fold serial dilutions (10²–10⁸) for each purified PCR template (DNA Clean & Concentrator™-5; ZymoResearch). Expression levels of OsNAS2, TaNAS, TaNAAT and TaDMAS were measured in root and shoot tissues using qRT-PCR analysis (CFX384- BioRad). The geometric mean expression of three housekeeping genes: TaCyclophilin, TaGAPDH and TaELF and TaGAPDH, TaActin and TaELF, was used to normalize OsNAS2, TaNAS, TaNAAT and TaDMAS gene expression within shoot and root tissues, respectively (Schreiber et al., 2009; Vandesonompel et al., 2002). All primers had annealing temperatures between 61–65 °C and primer sequences and efficiencies are provided (Table S4).

Quantification of NA and DMA

Quantification of 6-aminoquinolyl-N-hydroxysuccinimidyl carbamate (AQC)-derivatized NA in whole grain was performed via liquid chromatography-mass spectrometry (LC-MS) using established protocols (Callahan et al., 2007; Johnson et al., 2011). Quantification of 9-fluorenylmethoxycarbonyl chloride (FMOCl)-derivatized NA and DMA in whole grain (Figure 5d,e) and white flour (Figure 6d,e) was performed via RP LC-MS on an 1290 Infinity II and 6490 Triple Quadrupole LC/MS system (Agilent Technologies Inc., Santa Clara, CA) using established protocols (Selby-Pham et al., 2017). Briefly, sequential methanol (100%) and deionized H2O (18MΩ) extractions of pulverized wheat grain or white flour (25 mg) were combined and added (5 μL) to sodium borate buffer (pH = 8, 1 M, 10 μL), EDTA buffer (pH = 8, 50 mM, 10 μL) and fresh FMOCl-KCl solutions (50 mM, 40 μL). After incubation (60 °C, 700 rpm, 15 min), the derivatization reaction was quenched via the addition of formic acid (FA; pH = 4, 5%, 8.9 μL). A Zorbas Eclipse XDB-C18 Rapid Resolution H5 2.1x100 mm, 1.8 μm particle size column (Agilent Technologies Inc.) was used during chromatography with aqueous (0.1% v/v FA in dH2O) and organic (0.1% v/v FA in acetonitrile) mobile phases. For quantification, a stock aqueous solution of NA and DMA (Toro Research Chemicals, Toronto, ON, Canada) was prepared at 750 μM and a calibration set was generated in the range of 0.005 to 75 μM.

Analysis of Fe and Zn accumulation post anthesis

Plants were grown in glasshouse conditions (12 h photoperiod, 18 °C day/13 °C night, 40%-80% humidity) in Hortico® potting mix with Osmocote® fertilizer at The University of Melbourne (Victoria, Australia). The main stem flag leaf, rachis, bracts and grain were harvested at 5–8 DAA (days after anthesis), 12–15 DAA, 19–21 DAA, 26–29 DAA and maturity. Samples were washed, oven dried for 48 h at 60°C and ground to a powder before analysis by inductively coupled plasma mass spectrometry (ICP-MS) at the Environmental Analysis Laboratory (Lismore, NSW, Australia).

Synchrotron X-ray fluorescence microscopy (XFM)

Elemental X-ray fluorescence (XRF) maps of Fe, Zn, Cu and Mn in transverse cross-sections of two representative CE-1 and NS grain (four sets of maps total) were collected at the XFM beamline at the Australian Synchrotron (Melbourne, Australia) as previously described (Van Malderen et al., 2017). Briefly, the beam energy was set at 15.6 keV and the beam focused to approximately 2 × μm² using Kirkpatrick-Baez mirrors. Samples were analyzed continuously in the horizontal direction with a sampling interval of 4 μm and a step size of 4 μm in the vertical direction (pixel transit time was set at 5.2 ms). The XRF signal from the 80 μm transverse grain sections was collected using a 384-element Maia detector system. Tri-colour elemental maps showing the distribution of Fe, Zn and P near the grain edge of one representative CE-1 and NS grain (two sets of maps total; different grain from those used with the Maia detector) were collected using a separate Vortex-EM detector. The tri-colour maps were used as guides to select rectangular areas of approximately 14 × 140 μm near the grain edge for the generation of Fe, Zn, P and S line scans. Elemental maps were generated using GeoPIXE (http://nmp.csiro.au/GeoPIXE.html) software. The NS grain contained 38 μg/g DW Fe, 71 μg/g DW Zn, 4700 μg/g DW P and 1630 μg/g DW Mn.

<ref>Callahan et al., 2007</ref>
DW S while CE-1 grain contained 69 µg/g DW Fe, 122 µg/g DW Zn, 5300 µg/g DW P and 2100 µg/g DW S (Table S5).

Confined field trials

Confined field trials were conducted in Western Australia from June to December 2015 at the New Genes for New Environment facilities located in Merredin (31.483° S, 118.277° E) and Katanning (33.689° S, 117.555° E). Grain were sown in 2 m² plots with three replicate plots per genotype and arranged in a randomized block design at each site. Rows were spaced at 30 cm and grain were sown at a rate of 60 kg/ha. At maturity, average plant height was determined from three representative measurements per plot and spike number, total biomass and TGW were determined from 0.15 m² subsamples per plot (Table S3). Grain yield was calculated from the amount of grain harvested per 2 m² plot and extrapolated to kg/ha. Soil properties of both field sites are provided in Table S6.

Production of white flour

Whole grain samples harvested at Merredin and Katanning were conditioned to 13% moisture content for 24 h prior to milling. Each sample was milled using a Quadrumat Junior laboratory mill (Brabender, Duisburg, Germany) at constant temperature and run through a 280 μm sieve to isolate the white flour fraction. Average flour extraction for all lines from Merredin and Katanning was 71.5 ± 0.2%.

Caco-2 cell culture assessment of Fe bioavailability

Whole grain and white flour samples were digested for Caco-2 cell Fe-bioavailability analysis as previously described (Glahn et al., 1998; Trijatmiko et al., 2016). The Caco-2 cells were maintained in supplemented Dulbecco’s modified Eagle medium (DMEM) for 11 days post-seeding and then replaced with supplemented minimum essential media (MEM) solution 48 h prior to the experiment. On the experiment day, gastric-digested samples (1.5 mL) were added to cylindrical Transwell inserts (Corning Life Sciences, Corning, NY) fitted with a semipermeable (15 000 Da MWCO) basolateral membrane (Spectra/Por 2.1, Spectrum Medical, Gardena, CA). The inserts were placed within wells containing Caco-2 cell monolayers and incubated for 2 h (37 °C), after which the inserts were removed and additional MEM (1 mL) added to the cells before incubation for 22 h (37 °C). After incubation, growth medium was removed by aspiration and the Caco-2 cells were washed twice with a solution (pH = 7.0) containing NaCl (140 mmol/L), KCl (5 mmol/L) and PIPES (10 mmol/L) and harvested with the addition of dH₂O (1.5 mL) and brief sonication (140 mmol/L), KCl (5 mmol/L) and PIPES (10 mmol/L) and harvested with the addition of dH₂O (1.5 mL) and brief sonication (140 mmol/L), KCl (140 mmol/L), KCl (5 mmol/L) and PIPES (10 mmol/L) and harvested with the addition of dH₂O (1.5 mL) and brief sonication (140 mmol/L), KCl (140 mmol/L), KCl (5 mmol/L) and PIPES (10 mmol/L) and harvested with the addition of dH₂O (1.5 mL) and brief sonication

Statistical analysis

All graphs and statistical analyzes were generated using Minitab® 17 Statistical Software, Minitab, State College, PA and SigmaPlot v13, Systat Software Inc., San Jose. Data are presented as mean ± SEM with biological replicate numbers noted in table and figure legends. Student’s t-test was used to determine significant differences between means. Data are available upon request to the corresponding author of this paper.

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Conflict of interest

The authors declare no conflict of interest.

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Recent advancements in iron biofortification have demonstrated significant progress in enhancing iron levels in staple crops. The project has achieved notable success in various regions around the world, with multiple countries reporting increased iron levels in their crops. However, the effectiveness of iron biofortification strategies in different environments and crop species remains a critical area for further investigation.

To optimize the performance of iron biofortification approaches, it is essential to consider the specific needs of different regions and crops. This involves understanding the agronomic and biological factors that influence iron availability and uptake. The project is actively exploring the integration of diverse techniques, such as genetic improvement, agronomic practices, and innovative development strategies, to maximize the impact of iron biofortification.

In conclusion, the iron biofortification project has made significant strides in enhancing iron levels in staple crops. By continuing to refine and adapt the strategies employed, we can expect to make further advancements in this field, contributing to improved nutrition and health outcomes in vulnerable populations worldwide.
Table S1 Iron and zinc concentrations (µg/g DW) in NS and CE-1 plant tissues at 5–8, 12–15, 19–21 and 26–29 days after anthesis (DAA) as well as maturity.

Table S2 Biomass (mg DW) of NS and CE-1 plant tissues at 5–8, 12–15, 19–21 and 26–29 days after anthesis (DAA) as well as maturity.

Table S3 Agronomic performance of NS and CE-1 sibling lines grown at the Katanning and Merredin field sites.

Table S4 Wheat genes and primers used for quantitative reverse transcription PCR (qRT-PCR) analysis of CE-1 and NS seedling shoot and root tissue.

Table S5 Elemental concentrations (µg/g DW) of T₃ grain harvested from 10 NS and 9 CE-1 plants including the two batches used for synchrotron XFM analysis (in bold).

Table S6 Soil properties of Katanning (K) and Merredin (M) field sites.
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