Discovery and validation of candidate genes for grain iron and zinc metabolism in pearl millet *Pennisetum glaucum* (L.) R. Br.

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Pearl millet is an important crop for alleviating micronutrient malnutrition through genomics-assisted breeding for grain Fe (GFeC) and Zn (GZnC) content. In this study, we identified candidate genes related to iron (Fe) and zinc (Zn) metabolism through gene expression analysis and correlated it with known QTL regions for GFeC/GZnC. From a total of 114 Fe and Zn metabolism-related genes that were selected from the related crop species, we studied 29 genes. Different developmental stages exhibited tissue and stage-specific expressions for Fe and Zn metabolism genes in parents contrasting for GFeC and GZnC. Results revealed that *PglZIP*, *PglNRAMP* and *PglFER* gene families were candidates for GFeC and GZnC. Ferritin-like gene, *PglFER1* may be the potential candidate gene for GFeC. Promoter analysis revealed Fe and Zn deficiency, hormone, metal-responsive, and salt-regulated elements. Genomic regions underlying GFeC and GZnC were validated by annotating major QTL regions for grain Fe and Zn. Interestingly, *PglZIP* and *PglNRAMP* gene families were found common with a previously reported linkage group 7 major QTL region for GFeC and GZnC. The study provides insights into the foundation for functional dissection of different Fe and Zn metabolism genes homologs and their subsequent use in pearl millet molecular breeding programs globally.

Nutritional security is the key component to improve the health status of the world’s population as they are primarily dependent on plant-based diets. Plants are the major source of essential nutrients and nutraceuticals for the normal growth and development of humans. However, half of the global population, especially people from Asia and Africa suffer from a deficiency of micronutrients such as Fe, Zn, and selenium as they depend upon cereal crops for food. Biofortification is an approach to overcome the micronutrient deficiency/starvation by delivering nutrient-dense crops at the doorsteps of resource-poor people. Biofortification under HarvestPlus-Consultative Group for International Agricultural Research (CGIAR) micronutrients project has focused primarily on seven major staple crops (rice, beans, cassava, maize, sweet potato, pearl millet, and wheat) targeting important micronutrients like Fe and Zn. It is reported that 79% of pre-school and school-going children and 56% of women in India suffer from anemia due to iron deficiency. Likewise, 50% of the world population suffers from diarrhea, impaired physical growth, and suppressed immune function due to zinc deficiency. Fe supplementation program in India started in 1970 but failed to address the problem of iron deficiency. To address malnutrition, high grain Fe and Zn containing pearl millet lines were developed using the biofortification approach of conventional breeding (HarvestPlus, 2009). Earlier reports have been revealed no significant correlation in grain yield and seed size with that of grain Fe and Zn content. Hence, while breeding for high mineral content in pearl millet, the large grain size was recommended for selection as an associated trait. The progenies derived from AIMP92901, a high-yielding OPV, exhibited large intra-population variability with the highest levels of Fe (~110 mg/kg) and Zn (~65 mg/kg) than their parents.

An alternative approach to address the issue of genetic enhancement of grain Fe, Zn content is by modulating the metal transporters that facilitate their uptake, translocation, and storage. Study of a gene family of regulated

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transporter-like protein (ZIP), Zn-regulated transporter and metabolism responsible for Fe and Zn homeostasis by either uptake or remobilization in intracellular compartments are crucial\(^9\). The ZIP transporters enhance Zn uptake in plant species like Solanum lycopersicum\(^9\), Glycine max\(^10\), Arabidopsis thaliana\(^11\), Hordeum vulgare\(^11\), and Oryza sativa\(^12\). ZIP is differentially regulated in various tissues under mineral deprivation and abundance in soils\(^12\). Millets and other cereals with high Fe and Zn content in seeds can be engineered by seed-specific expression of ZIP transporter genes. Fe and Zn are vital for plant growth, development and transport to the grain filling site\(^13\). As a structural motif, Zn plays an important role in many proteins, including DNA-binding Zn-finger proteins\(^13\), RING finger proteins, and LIM domain-containing proteins\(^13\), and participates in governing cellular processes such as growth, development, and differentiation. While sufficient Zn content enhances crop productivity\(^14\), Fe serves an important role in electron transfer during respiration and photosynthesis, though high concentrations of intracellular Fe may undergo Fe\(^3+\)/Fe\(^2+\) redox reactions and cause damage to the plants\(^15\). Mineral deficiency hampers growth, yield and grain quality in cereals\(^16\), but excess Fe and Zn may cause significant toxicity to biological systems\(^17\). Therefore, plants developed a tightly regulated system to balance the uptake, utilization and storage of these metal ions\(^18\). Since Zn cannot diffuse across the cell membrane, specific Zn transporter is required to transport Zn into the cytosol and also into the stelar cells\(^19\). Metal transporters have been identified in plants, including the P\(_1\)p-ATPase family, zinc-regulated transporter (ZRT), iron-regulated transporter (IRT)-like protein (ZIP), natural resistance-associated macrophage protein (NRAMP) family, and cation diffusion facilitator (CDF) family where these two metals play crucial roles\(^20\).

Several factors affect the uptake of both Fe and Zn from soil to roots, transport to leaves and stems and subsequently remobilization to developing grains\(^21\). Flag leaves are the major source of the remobilization of metals for developing seeds\(^21\). The gene(s) involved in Fe movement from soil to seed are not highly selective for Fe, and Zn, but other metals are also transported and allocated to different organs and extents\(^21\). Currently, it is not clear how coordinately Fe and Zn metabolism genes are expressed at developmental stages of pearl millet. This information is critical to understand the genetics and molecular mechanisms that help to identify candidate genes associated with their accumulation, toxicity, and homeostasis. Deployment of such information has been used recently in rice for developing lines with high grain Fe and Zn\(^22\). Several genes/ gene families involved in Fe and Zn homeostasis like ZRT, ZIP, yellow stripe-like (YSL), NRAMP, nicotianamine synthase (NAS), nicotianamine aminotransferase (NAAT), heavy metal ATPases (HMA), metal tolerant protein (MTP), zinc-induced facilitator-like (ZIFL) and others have been characterized in cereal crops. But a thorough understanding of the regulatory events leading to the roles and functions of Fe and Zn metabolism genes of pearl millet should help us to gain new insights into micronutrient homeostasis and ultimately mitigating malnutrition. Also, identifying their potential in metal transport, homeostasis, and their expression levels to regulate the ions is the prime aim of such an endeavor. The present study involves the identification of genes associated with Fe and Zn metabolism, gene structures, characterization, chromosomal localizations, grain Fe and Zn quantification and tissue-specific expression of these genes in P. glaucum. Unraveling the intricate mechanisms associated with Fe and Zn metabolism and the gene expression analysis should also help us in understanding better grain micronutrient nutrition in cereal crops\(^23\).

Pearl millet is a nutritionally important cereal not only for the rural poor but also in arid and semi-arid tropics. Scientific research is still limited in comparison with other cereals\(^24\) in this regard. To date, no genes associated with Fe and Zn metabolism genes have been characterized in P. glaucum and also their transport capabilities at the time of grain filling. But, such findings help us in developing better lines with high grain Fe and Zn content and subsequently reducing malnutrition among resource-poor populations. As yet, a comprehensive analysis of all the Fe and Zn metabolism gene family members is not available in P. glaucum. An effort has been made to identify those genes that get expressed to mobilize nutrients from root to flag leaves and then into developing grains in pearl millet.

**Results**

**In silico identification of Fe and Zn metabolism genes.** A total of 114 Fe and Zn-responsive gene sequences from different plant species including Setaria italica (11), Zea mays (11), Oryza sativa (39), Sorghum bicolor (3) and Arabidopsis thaliana (49) were downloaded from NCBI. BLAST was performed against pearl millet sequences which resulted in the identification of 29 putative Fe and Zn metabolism genes. The reliability of these porters was checked by confirming the presence of Fe and Zn conserved domains by at what developmental stages of pearl millet. This information is critical to understand the genetics and molecular mechanisms that help to identify candidate genes associated with their accumulation, toxicity, and homeostasis. Deployment of such information has been used recently in rice for developing lines with high grain Fe and Zn22. Several genes/ gene families involved in Fe and Zn homeostasis like ZRT, ZIP, yellow stripe-like (YSL), NRAMP, nicotianamine synthase (NAS), nicotianamine aminotransferase (NAAT), heavy metal ATPases (HMA), metal tolerant protein (MTP), zinc-induced facilitator-like (ZIFL) and others have been characterized in cereal crops. But a thorough understanding of the regulatory events leading to the roles and functions of Fe and Zn metabolism genes of pearl millet should help us to gain new insights into micronutrient homeostasis and ultimately mitigating malnutrition. Also, identifying their potential in metal transport, homeostasis, and their expression levels to regulate the ions is the prime aim of such an endeavor. The present study involves the identification of genes associated with Fe and Zn metabolism, gene structures, characterization, chromosomal localizations, grain Fe and Zn quantification and tissue-specific expression of these genes in P. glaucum. Unraveling the intricate mechanisms associated with Fe and Zn metabolism and the gene expression analysis should also help us in understanding better grain micronutrient nutrition in cereal crops23.

**Chromosomal location and structure of Fe and Zn metabolism genes.** The 29 genes identified for Fe and Zn transporters in the present study were distributed on all 7 chromosomes of pearl millet and their homologs were found in Setaria italica, Sorghum bicolor, Oryza sativa and Zea mays which are shown in CIRCOS5 Fig. 1. Of the 29 genes, PglZIP1 was found on chromosome 1; 5 genes on chromosome 2; 13 on 3, 2 on 4, 3 on 5, 2 on 6 and 3 on chromosome 7. Results indicated that chromosome 3 is the hot spot for these genes as it is homing maximum genes (Fig. 2). The gene structures analysis revealed that three genes are intron less while remaining displayed many introns. The exons ranged between 1 (Pgl_GLEAN_10002987) and 13 (Pgl_GLEAN_10000860) (Supplementary Table S1).
Sequence analysis and conserved motif analysis of Fe and Zn metabolism genes. The number of amino acids in the proteins ranged between 65 (PglYSL2, not considered as full-length gene) and 883 (PglNAS1), and the pI ranged from 5.32 (PglFER1) to 10.02 (PglYSL2). Out of 29 proteins, 10 were found acidic and the remaining basic. Molecular weights varied between 7146.53 (PglYSL2) and 96605.68 (PglNAS1) daltons. Except for PglFER1 and PglNAS1, all Fe and Zn transporter proteins were found hydrophobic. They showed the highest aliphatic index that varied from 132.59 (PglNRAMP5) to 82.52 (PglFER1). Transmembrane helices ranged from 1 to 15, but 2 proteins (PglFER1 and PglNAS1) exhibited no helices. Subcellular localization revealed that of 29, 21 were in the plastid, 4 in the vacuole, 2 in the chloroplast, 1 each in the cytosol and extracellular region. The phosphorylation sites of these proteins (Supplementary Table S2) indicated that they are phosphorylated at serine residues, followed by threonine. But, the YSL subgroups were found phosphorylated at tyrosine residues. The PKC, CKII, PKA, UNSP, and CDC2 were common kinases for Fe and Zn protein phosphorylations. The identified Fe and Zn transporter proteins belonging to various subgroups did not display inter-subgroup similarity, hence each family was submitted to MEME separately for conserved motif analysis.

Figure 1. Circular genome visualization (CIRCOS) of pearl millet. Fe and Zn metabolism genes and their homologous map with other cereals (pink: P. glaucum chromosomes (Pg1–7); red: O. sativa chromosome (Os 1–12); brown: S. bicolor chromosomes (Sb 1–10); green: Z. mays chromosomes (Zm 1–10); light green: S. italica chromosomes (Si1–10).
The MEME analysis revealed ten conserved motifs, which showed various compositions and distribution patterns among the family members. All the YSL group proteins exhibited ten conserved motifs except YSL2, which showed 2 motifs. Motifs 6 and 10 were conserved at N terminus, while 5 and 8 at C terminus. Out of 29 identified genes, 10 showed conserved motifs among the YSL members. Motifs 6 and 3 are the most conserved at N terminus, while motifs 7 and 8 are commonly present at C terminus. In the ZIP family, motif 3 was found as the most conserved at N terminus, followed by motif 5, while 3 and 4 represent NES motifs. In the Nramp family, 10 were identified as the conserved motifs. While motif 3 and 9 were the most conserved at N terminus, motif 6 and 7 conserved at C terminus. Among all, motif 1 and 3 appeared as the utmost conserved (Fig. 3).

Phylogenetic analysis. A phylogenetic tree was constructed for 29 identified Fe and Zn metabolism genes, belonging to 5 groups in pearl millet. The analysis revealed four clades, of which five subfamilies clustered and showed group-specific distribution based on the conserved domains (Fig. 4a). While FER1 and NAS1 were clustered into one group, YSL2 fell into the ZIP family, ZIP3 and ZIP6 into the NRAMP subgroup. A total of 9 paralogs, 4 tandem duplications (PglYSL1/PglYSL4, PglYSL5/PglYSL6, and PglYSL10/PglYSL11) on chromosome 3, one PglNRAMP5/PglNRAMP6 on chromosome 7, and 5 segmental duplication events were noticed. The phylogenetic tree was constructed to identify the evolutionary relationships of 5 gene families of Fe and Zn metabolism genes in pearl millet with other crop plants (Fig. 4b–f). The trees showed species-specific, group-specific and class-specific clades. Pearl millet showed 4 orthologous events, of which one with Oryza (Os11G0138400/Pgl_GLEAN_10003023), while remaining with Setaria (Si024505/Pgl_GLEAN_10026644, Si010411/Pgl_GLEAN_10012759 and Si03616/Pgl_GLEAN_1005377). All four genes belonged to the ZIP family (Fig. 4d).

Promoter analysis. Promoter analysis revealed that the majority of them contain Zn deficiency-responsive (ZDRE), iron-responsive (IRO2) and iron deficiency elements (IDE), which are responsible for iron and zinc deficiencies. Interestingly, two IRO2 elements were found in PglFER1 gene which is a potential candidate gene for GFeC. In PglZIP2 and PglZIP4, single ZDRE element was found, whereas for PglZIP2 two, PglZIP4 one, PglNRAMP5 two, and PglZIP9 three IDEs were found. They also contained dehydration-responsive elements (DRE) for drought, low temperature-responsive elements (LTE), salt-regulated elements (SRE, GT1GM) and abscisic acid-responsive elements (ABRE). They exhibited the highest number of ABRE along with Cu-responsive elements. Among all, genes 3, 7, 11, 22, and 27 contained large amounts of stress-responsive elements (Supplementary Table S3).

Grain Fe and Zn content. The grain Fe and Zn content in bi-parental RIL mapping population parents AIMP 92901-S1-183-2-2-B-08 (hereafter referred to as AIMP 92901) and ICMS 8511-S1-17-2-1-1-B-P03 (hereafter referred to as ICMS 8511) pearl millet lines were 110.70 ± 2.44 ppm, 79.11 ± 22.03 ppm, 31.26 ± 3.26 ppm and 26.63 ± 0.68 ppm, respectively. On the other hand, in control line MRC-HS-130-2-1-B-B-3-B-B-B-1-3-1 (hereafter called MRC), Fe and Zn content was 107 ± 14.22 ppm and 63 ± 13.03 ppm, respectively (Supplementary Table S4).
Gene expression profiles in different tissues in contrasting genotypes. To investigate the role of Fe and Zn metabolism genes in the accumulation of grain Fe and Zn content, the expression levels of genes were quantified by qRT-PCR analysis. The expression of these genes was analyzed among different tissue types.
Figure 4. (a) Phylogenetic tree of Fe and Zn metabolic gene families in Pearl millet. N-J phylogenetic tree of Fe and Zn metabolism genes in pearl millet grouped into 4 showing ZIP, YSL, Nramp and FER. NAS is seen in a single group. A total of seven paralogs were observed out of which 4 segmental and 3 regional duplications were noticed. (b) Phylogenetic tree showing the relationship between Pearl millet, other cereals Ferritin (FER) family genes. (c) Phylogenetic tree showing the relationship between Pearl millet, other cereals NAS family genes. (d) Phylogenetic tree showing the relationship between Pearl millet, other cereals ZIP family genes. (e) Phylogenetic tree showing the relationship between Pearl millet, other cereals NRAMP family genes. (f) Phylogenetic tree showing the relationship between Pearl millet, other cereals YSL family genes.
Figure 4. (continued)
and developmental stages to unravel their precise involvement in the transport, remobilization and grain loading of micronutrients. The expression values of different genes recorded during different developmental stages of three genotypes AIMP, ICMS and MRC are illustrated (Fig. 5a) by MultiExperiment viewer (MeV) v4.1.0. It has been noticed that the majority of the genes were up-regulated in root tissues at the reproductive stage.
compared to vegetative stages. In ICMS also, the up-regulation of genes was observed in roots and leaves at the vegetative stage. The PglFER1, PglZIP2, PglZIP8, PglNRAMP2, PglYSL1 were found to be the candidate genes with high expressions in seeds showed in CIRCOS (Fig. 5b). They showed moderate expressions in the roots at the vegetative stage, indicating their Fe and Zn restoration nature in seeds. In ICMS, the majority of them were expressed at the vegetative stage compared to the reproductive stage, while in AIMP they were expressed in roots at the reproductive stage similar to MRC, the check.

**Correlation of differentially expressed genes with previously reported QTLs.** The earlier identified LG1 and LG7 high grain QTL regions for GFeC and GZnC were scanned for presence of PglZIP, PglNRAMP, PglYSL and PglFER family genes that were differentially expressed between the two parents. Upon sequence comparison between these genes and the QTL regions defined by the set of linked markers, we could...
not locate any common set of genes for LG1 QTL interval. However, interestingly, for LG7 QTL interval two gene families, PglZIP and PglNRAMP were found to be common.

**miR159 associated with Fe.** A host of plant miRNAs were screened in addition to checking the miRNAs from miRBASE. miR159 is the only miRNA that appeared to be associated with Fe, but, no miRNAs could be found for Zn. Surprisingly, careful dissemination and search from PMRD yielded only one miRNA in pearl millet. Before this, we have considered all the miRNAs in *Arabidopsis* and made a Smith-Waterman/Blast search to find any potential miRNAs, but in vain except miR159.

**Protein–protein interaction analysis predictions.** The predicted protein–protein interaction network revealed interactions of Fe and Zn accumulation associated proteins with several other transporter proteins of pearl millet (found ontology with *Sorghum* and *Seteria*). Fe and Zn transporter proteins have been noticed as centers for interactions and displayed interactions with various other proteins associated with the metabolism of fructose, galactose, and also with ATP binding protein, hexokinase and protein kinase (Supplementary Table S5; Fig. 6).

**Discussion**

The deficiencies and improper consumption of micronutrients like minerals and vitamins lead to multiple disorders and diseases in humans and are collectively stated as hidden hunger or malnutrition. The connecting link between health and food has been well documented as human beings require several organic and mineral nutrients for their proper growth and development. Hyper and hypo levels of iron and zinc in the soil cause many physiological and metabolism-related complications in plants. But plants have developed a tightly regulated cellular iron and zinc homeostasis, including the uptake, distribution within different tissues and their utilization. Since both iron and zinc are the vital nutrients, their deficiencies can lead to a variety of problems like retarded growth, skeletal abnormalities, learning capacity and wellbeing of humans. Their deficiencies cause nutritional disorders in human beings, especially in developing world. Edible parts of many cereal cultivars have inherently low iron and zinc content. Millets are nutritionally rich and can serve potentially as crops of nutritional security in the developing and underdeveloped countries. Therefore, biofortification of cereals or increasing their bioavailability in the edible parts like endosperm can cost-effectively solve the problem of malnutrition and save millions of lives across the globe. Among the Fe and Zn superfamily, genes such as YSL, ZIP, NRAMP, FER, NAS have been identified in many plants like *Arabidopsis thaliana*, *Setaria italica* and *Oryza sativa*. These genes were responsible for Fe and Zn translocation from soil to the grain filling sight. Genome-wide studies of different families of Fe and Zn have been characterized individually in cereals, but there were no reports on the identification and gene expressions of Fe and Zn in pearl millet, though it has high micronutrients.
the present study, 12 PglYSL, 9 PglZIP, 6 PglNRAMP and a single gene for PglFER and PglNAS were identified and characterized from *P. glaucum*. However, form *PglYSL* family, one gene PglYSL2 was found to be of much shorter length (just 65 amino acids). This may be because of differential divergence of this gene in pearl millet, or because of lack of capture of complete sequence during the genome assembly process. Hence, *PglYSL* was not considered in further studies.

Palmer et al. identified 520 genes that belong to 40 different transporter classes in the *Panicum virgatum* (*C4* switchgrass) genome. There are five sub-families for Fe and Zn transport based on their conserved domains. Attempts were made earlier for in silico identification and expression analysis of either YSL, ZIP or NAS gene family or NRAMP members in a wide array of plants. The YSL or YSL is a membrane-bound Fe (III)-phytosiderophores (PS) transporter and is distantly related to the Oligopeptide Transporter (OPT) family proteins. Though non-grass species do not synthesize PS, many YSL genes were detected in monocots, dicots, and lower groups of plants. Meena et al. identified 8 and 18 YSL family members in *Arabidopsis thaliana* and *Oryza sativa*, respectively. Similarly, in the present study, 12 genes belonging to the YSL family were detected suggesting that they are highly conserved across plant species. Species-specific variations in the gene numbers were also known to exist for many gene families in cereals. The ZIP family members have been characterized in higher plants and are involved in metal uptake, transport, detoxification and storage of Fe and Zn in plant cells. In *Oryza sativa*, 17 ZIP genes, 15 in *Arabidopsis thaliana*, 12 in *Poncirus trifoliata* and 9 in *Zea mays* were reported. Alagarasan et al. on the other hand, identified 36 ZIP family members in Setaria italicca. In contrast, in the present study, 8 ZIP family members (out of a total of 29 iron and zinc transporter genes) were noticed in the genome of *P. glaucum*. This number is close to the published data in maize that belongs to the same family. Members of the NRAMP family genes are also widespread from bacteria to plants and transport Fe and Mn (manganese) across cellular membranes. In the present study, 6 NRAMP members were identified in pearl millet like that of *Arabidopsis*, while Qin et al. reported 13 in soybean. In *Arabidopsis*, 4 ZIP genes were detected, and 3 in *Lupinus luteus*. Gross et al. identified 2 FER genes in maize and rice, respectively, but only one is detected in the present study. A systematic analysis for NAS members could identify 3 and 4 NAS members in *Oryza sativa* and *Arabidopsis thaliana*, respectively, while in the current study merely one NAS member was identified in pearl millet. But in maize and barley, 9 NAS genes were detected. This suggests that NAS members in pearl millet are encoded by a few genes instead of a large gene family members. Zhou et al. pointed out that large NAS gene members in maize and barley may be due to gene duplication events that evolved during the evolution of new species.

YSL members are acidic with a large number of glutamic and aspartic acid residues. It was observed that these proteins are highly hydrophobic (27 out of 29) and are membrane-bound transporters. Except for two, the rest of the Fe and Zn porters have transmembrane helices in pearl millet. Such variability can occur between transmembrane domains 7 and 8 and also at the N-terminal portion of the YSLs. Chu et al. reported that ysl1/ysl2 double mutants result in strong interveinal chlorosis and also reproductive defects implying that these two YSL members are needed for loading metals into the reproductive tissues. Meena et al. found that the intron/exon structure and domains of YSL members from rice and *Arabidopsis* are conserved during evolution. They also noticed less variation in sequence lengths of amino acid residues of rice (554–728) and *Arabidopsis* (664–724) YSL members. Though they are transport proteins, plasma membrane localization of these proteins could not be detected among the 29. But, others could detect plasma membrane-bound porters/proteins as the probable sub-cellular localization. This difference could be due to the diverse bioinformatics tool(s) that we use for this purpose. Guerinot pointed out that a metal-binding domain is seen between transmembrane 3 and 4. A phylogenetic tree was constructed to find out the evolutionary relationships of these porters. The *Arabidopsis YSL* family members share high sequence similarity. Based on sequence similarity, the YSL family can be divided into three conserved groups, while we noticed only two groups (Fig. 3). It was noticed that amino acid sequences of PglZIPs were related to ZIPS identified in other plant species. This infers that PglZIPs share a common evolutionary ancestor along with other cereals. Two subfamilies appear for the members of Nram that are common both in dicots and monocots as noticed in the present study and also in soybean. Phylogenetic analysis revealed that NASs from the members of Poaceae were divided into two classes. Further, more members existed in class I in maize and barley than in rice. Schnable et al. suggested that nearly 25% of the genes in maize possess closely related paralogs due to genome duplication events. Many NAS genes in some of the monocots indicate that there may be functional redundancy as also noticed by Mizuno et al. Nevertheless, the evidence is lacking at present to exclude or include the possibility of functional redundancy and if they are pseudogenes or not. Strozycki et al. working with *Lupinus* suggested that eudicot ferritins are both structurally and functionally diverse. They found out that 14-bp-long iron-dependent regulatory sequence (IDRS) is the characteristic feature for all higher plant ferritin promoters. They also found that a highly conserved IDRS can be extended (extIDRS) up to 22 bp. Such an analysis of ferritin gene promoters for specific or conservative elements and their validation only can resolve such a complexity. In the present study, the analysis of promoter regions was carried out for all the 29 genes. The presence of Fe and Zn deficiency elements, hormone-responsive and salt-regulated elements indicate that Fe and Zn transporters are associated with diverse stresses. However, the promoters need to be validated further in the lab. The grain Fe and Zn content were higher in AIMP compared to ICMS indicating that AIMP should be utilized for future breeding programs. In pearl millet, QTLs for high GFeC and GZnC revealed a co-localization of QTLs on chromosome 3. Major QTLs were noticed earlier in other populations also, and LG3 and LG1 served as co-localizations for high grain Fe and Zn. Current work thus forms a perfect computational framework for in silico characterization of five gene families that can be utilized in marker-assisted selection (MAS) of important cereal crops for enriching grain Fe and Zn.

The present work reveals that major differential gene expressions were recorded in PglYSL2, PglZIP2, PglZIP8, PglNRAMP2, PglFER1 in the two contrasting parents. Such Fe and Zn transporter gene expressions in diverse tissues under various developmental stages provide evidence in the acquisition of iron and zinc, and tight regulation
of metal ion homeostasis in the grains of cereals. Chu et al. demonstrated that Arabidopsis YSL1 and YSL3 are vital in iron transport and responsible for loading Fe, Cu, and Zn from source to sink i.e. grain filling site. On the other hand, YSL4 and YSL6 appear to be involved in iron transport and metal mobilization into the seeds of Arabidopsis. Menna et al. reported a preferential pattern of YSL gene expressions in rice and Arabidopsis which suggest spatiotemporal regulation of YSL genes during plant development. Interestingly, both AtYSL1, 3 and 6 were highly expressed in senescent leaves indicating that they are implicated in the remobilization of metals. It is known that some ZIP genes in Arabidopsis and rice play crucial roles in transporting Zn or Fe. ZIP genes exhibit different expression profiles like tissue specificity and response to the availability of Fe and Zn in the environment. The gene expression patterns of PglZIPs indicate diverse functions carried out by ZIPs. While expression of MizZIP1 was reported in Zn-deficient roots and leaves, MizZIP2 was stimulated in roots and stems by Zn deficiency. Yang et al. observed that OsZIP7a was induced mostly in roots where Fe-deficiency is common, but OsZIP8 was activated in Zn-deficient shoots as well as roots. In Arabidopsis, bZIP19 and bZIP23 regulated the adaptation to Zn deficiency by increasing the transcription of ZIPs. In maize, ZmZIP4, 5, 7 and 8 were upregulated in shoots and roots after excess Fe was applied. They pointed out that ZmZIP4, 5, 7 and 8 may be associated with excessive Fe detoxification and storage. Also, ZmZIP4 and ZmZIP5 were stimulated during embryo development and they were repressed in later stages. Such an expression suggests that these two ZIPs are essential for the growth of plumule and radicle in maize. Qin et al. studied the soybean NRAMP gene expressions and suggested these genes function in many tissues and developmental stages. GmNRAMP genes were found to be differentially regulated upon N, P, K, Fe and S deficiencies as well as toxicities of Fe, Cu, Cd, and Mn. These results infer that NRAMPs are functionally associated with many nutrient stress-responsive pathways. Zhou et al. found that class I NAS genes were induced under Fe deficiency, but suppressed under excess Fe conditions in Zea mays. In contrast, they noticed that the expression pattern of class II NAS genes was a reversal of class I genes, thus complementing each other. While ZmNAS1; 1/1; 2 was mainly expressed in cortex and stele of root tissues under sufficient Fe conditions, but under deficit conditions, its expression was noticed also in the epidermis, and shoot apices. The expression patterns suggest that the two classes may be regulated at the transcriptional level and respond to the demand for iron uptake and transport. Fobis-Loisy et al. noticed in maize differential accumulation of FM1 and FM2 (two sub-classes) mRNAs in response to iron, ABA and water deficit conditions which could be due to differential transcription of ZmFer1 and ZmFer2. Bournier et al. found out that a molecular link exists between the control of iron and phosphate homeostasis in Arabidopsis. Further, genome-wide analysis of gene expression profiling revealed that an intact COP9 signalosome is essential for correct expression of Fe homeostasis genes in Arabidopsis. Such a possibility cannot be ruled out in pearl millet. Our in silico analysis has further validated key genes that are differentially expressed like ZIP, and NRAMP family members which have been found significantly on LG 7 QTL by annotation study and found to be a candidate gene.

Besides, miRBASE, other plant miRNAs were screened in the present study for identifying miRNAs associated with the regulation of Fe, Zn transporters. We found that pgl-miR159 is known to be the only matured miRNA sequence that was identified from the Plant miRNA database (PMRD). With the plant miRNAs predominantly targeting transcription factors, we aimed to identify functional targeted genes responsible for various physiological processes using psRNAtarget. For the lome miRNA, we found that 1-amino-cyclopropane-1-carboxylate synthase 8 (ACS8) is the probable target. While this gene was found in Arabidopsis, it was not yet annotated in pearl millet. We believe that ACS transcripts are responsible for Fe deficiency, which is invariably associated with increased abundance of ACS transcripts in plant leaves. Our study revealed that among the 29 proteins, 6 proteins (PglFER1, PglZIP2, PglZIP3, PglZIP4, PglZIP5, and PglZIP9) formed tight networking with other proteins (Fig. 6). The protein Pgl_GLEAN_10005377 (PglZIP2) displayed the highest interactions. (Supplementary Table S5). PglFER and PglZIP might play a key role in mineral transportation and accumulation in seeds.

Conclusions Pearl millet is a climate-resilient nutricereal with substantial genetic variability for grain Fe and Zn content. Our results from the gene expression studies, corroborated by QTL interval annotation revealed that the gene families PglYSL, PglZIP, PglNRAMP, PglNAS, and PglFER play an important role in Fe and Zn homeostasis in pearl millet. Ferritin-like gene PglFER1 along with other key candidate genes should be further investigated in diverse pearl millet germplasm. This study provides insights into the foundation for functional dissection of different Fe, Zn metabolism genes homologs. The genomic resources developed in this study may be used for the development of breeder-friendly marker systems for improvement of grain Fe and Zn content in the pearl millet breeding programs globally.

Materials and methods
In silico identification of Fe and Zn metabolism genes. The Fe and Zn transporter gene sequences of Setaria italica, Zea mays, Oryza sativa, Brachypodium distachyon, and Triticum aestivum were retrieved from NCBI (https://www.ncbi.nlm.nih.gov/) database and searched against Cenchrus americanus (pearl millet) genome to find out their homologs. GENSCAN (http://genes.mit.edu/GENSCAN.html) was used to retrieve the gene and corresponding protein sequences. All the identified putative protein sequences were subjected to Motif Search (https://www.genome.jp/tools/motif/) to check their reliability and identification of conserved domains.

Chromosomal locations and gene structure. The identified Fe and Zn metabolism genes are mapped to their respective chromosomes based on the information provided in the pearl millet genome database (http://https://www.gigadb.org/dataset/100192). The gene structure display server (https://gds.cbi.pku.edu.cn) soft-
In silico protein analysis. The molecular weight (MW), isoelectric point (pI), and grand average of hydropathy (GRAVY) of Fe and Zn metabolism genes were identified using ProtParam of Expasy tools74 (https://web.expasy.org/protparam), while the phosphorylation sites were predicted by employing NetPhosK3 software of Expasy tools75. The putative transmembrane helices within genes were identified using TMHMM software76. For finding out the subcellular localization of transporters, the WOLFSORT program (https://wolfsort.org/) was used77.

In silico prediction of cis-regulatory elements and phylogenetic analysis. Genomic sequences measuring 2000 bp of upstream of the start codon of Fe and Zn metabolism genes were extracted for the identification of cis-acting elements. The elements were analyzed by employing PLANTCARE software78. The Neighbor-Joining (NJ) phylogenetic tree was constructed for the Fe and Zn protein sequences of pearl millet and other cereals like Setaria, Oryza, etc., by using MEGA 6.2 software79 by employing the Poisson correction, pairwise deletion, and bootstrap value (1000 replicates) permutation.

Plant material. In the present investigation, two contrasting parents of a bi-parental RIL mapping population AIMP 92,901 (high grain Fe and Zn), and ICMS 8511 (low grain Fe and Zn) were used77. Another line MRC was used as a check which records very high grain Fe and Zn content. The experiment was conducted under controlled environmental research facility (CERF), ICRISAT greenhouse conditions at the geographical location of longitude 17.51017° N, latitude 78.27540° E and altitude 623.14 m. Since the impact of the geographical location of plants remains a potential aspect while considering for nutrient accumulation and biological activities, the precise location of crop grown area is mentioned here. Following plant tissues were used for qRT-PCR analysis: flag leaf 1 (during vegetative stage), flag leaf 2 (during reproductive stage), root 1 (during active vegetative growth), root 2 (during the grain filling stage) and mature seeds.

Element analysis for grain Fe and Zn content. Grain Fe and Zn content were analyzed at the Charles Renard Analytical Laboratory, ICRISAT, Patancheru, Hyderabad, India following the method described by Wheat et al.80. The ground samples were digested in closed tubes, and Fe and Zn in the digests were analyzed using Inductively Coupled Plasma Optical Emission Spectrometry (ICP-OES). RNA was isolated at the vegetative and reproductive stages. The flag leaf and root were harvested first at the vegetative stage once and again the same was collected at the mature embryo (reproductive) stage and frozen in liquid nitrogen and stored at – 80 °C until further use. The values are represented as means ± standard errors of the replicated data.

RNA isolation, cDNA synthesis and qRT-PCR analysis. Total RNA was extracted from different tissues of pearl millet using the Invitrogen RNA isolation kit according to the manufacturer’s instructions. A total of 2.5 μl RNA with 5 μg concentration was taken for conversion to cDNA using a first-strand cDNA synthesis kit (Thermo Scientific) for qRT-PCR analysis of pearl millet Fe and Zn transporters. The template was diluted with nuclease-free water (1:10). Gene expression analysis was performed for 29 gene-specific primers, along with three reference genes (Supplementary Table S6), namely elongation factor 1-alpha (EF-1α), eukaryotic initiation factor 4A (EIF4A) and an acyl carrier protein (ACP)81 for gene normalization in 96-well optical PCR plates. The SYBR Green PCR master mix (Applied Biosystems, CA, USA) was used for gene expression analysis according to the manufacturer’s instructions. The ABI 7500 real-time PCR system (Applied Biosystems, USA) was used with the following thermal cycling conditions; the cycle 1 at 95 °C for 10 min, followed by 40 cycles at 95 °C for 15 s and 60 °C for 1 min (variable based on primers gradient temperatures) (Supplementary Table S6). The amplicon dissociation curves were recorded with a fluorescent lamp after the 40th cycle by heating from 58 to 95 °C within 20 min. Three biological replicates were taken in addition to two technical replicates each time. The relative expression values of 29 genes in five tissues as mentioned earlier at two developmental stages (vegetative and reproductive) of 3 contrasting genotypes (low and high Fe and Zn content) were calculated using qbase+ software82.

Correlation of differentially expressed genes with previously reported QTLs. We correlated differentially expressed genes in the plant materials (ICMS 8511 and AIMP 92901) which also happen to be parents of a bi-parental RIL mapping population. This RIL population was earlier used to identity LG1 and LG7 high grain QTL regions for GFeC and GZnC. In silico approaches, the LG1 and LG7 QTL interval was scanned for presence of common genes that were differentially expressed and were also present in the QTL region.

In silico identification of miRNA(s). For the in-silico identification of miRNA associated with Fe and Zn we have used 29 putative gene sequences as input to find perfect or complementary miRNA binding to its target sequences. The pearl millet miRNA targets were identified through pairwise homolog search by subjecting mature miRNA sequences as query against manually curated reference mRNA sequences (RefSeq, NCBI) and assembled EST sequences of pearl millet using Plant small RNA Target67 (psRNATarget). Analysis server was used for obtaining the Fe and Zn gene structures—exons, introns, and untranslated sequence regions (UTRs) based on the alignments of their coding sequences. MEME software72 was employed to analyze the new sequence patterns, their distributions, significance7 and identify the motifs by setting different default parameters like the number of motifs from 1–10, with a motif width of 5–50, and the number of motif sites from 5–10.
and Plant miRNA database (PMRD): https://bioinformatics.cau.edu.cn/PMRD/. To further confirm this, we have also done a UVA Fasta/Smith-Watermann algorithm search using the Pearl millet and Arabidopsis whole genomic sequences taking the matured sequences as query from miRNA databases. The brief methodology representing in pictorial form in supplementary Fig. 1.

**Protein–protein interaction.** The protein–protein interaction (PPI) was prepared using 29 identified genes. As the pearl millet PPI network details are not available in the STRING database, initially all the orthologs of *P. glaucum* in the *S. italica* were listed out. For each gene (29), corresponding and interacting proteins were listed by searching in the STRING database and redundant proteins were discarded. Network analysis and functional enrichment in the network were performed by Cytoscape.

**Data availability**

All the data presented in the manuscript are publicly available. The gene expression data generated during this study are available from the corresponding author(s) on request. All authors have read and accepted the MS.

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**References**

1. Zhao, F. J., Ma, J. F., Meharg, A. A. & McGrath, S. P. Arsenic uptake and metabolism in plants. *New Phytol.* **181**, 777–794 (2009).
2. Bouis, H. E. et al. Biofortification: a new tool to reduce micronutrient malnutrition. *Food Nutri. Bulletin.* **32** (1-suppl1), S31–40 (2011).
3. Welch, R. M. & Graham, R. D. Breeding for micronutrients in staple food crops from a human nutrition perspective. *J. Exp. Bot.* **55**, 353–364 (2004).
4. Krishnaswamy, K. The problem and consequences of the double burden: A brief overview. *Programme and abstracts. In Symposium on Nutritional Security for India–Issues and Way Forward, Ind. Natl. Sci Acad.* New Delhi 3–4 (2009).
5. Gibson, R. S. Zinc deficiency and human health: etiology, health consequences, and future solutions. *Plant Soil* **361**, 291–299 (2012).
6. Anand, T., Rahi, M., Sharma, P. & Ingle, G. K. Issues in prevention of iron deficiency anemia in India. *Nutrition* **30**, 764–770 (2014).
7. Kumar, S. et al. Mapping grain iron and zinc content quantitative trait loci in an inbred-derived immortal population of pearl millet. *Genes* **9**, 248 (2018).
8. Connorton, J. M. & Balk, J. Iron biofortification of staple crops: lessons and challenges in plant genetics. *Plant Cell Physiol.* **60**, 1447–1456 (2019).
9. Guerinot, M. L. The ZIP family of metal-transporting zinc/ cadmium transporters. *Biochim. Biophys. Acta.* **1465**, 190–198 (2000).
10. Mäser, P. Phylogenetic relationships within cation transporter families of Arabidopsis. *Plant Physiol.* **126**, 1646–1667 (2001).
11. Colangelo, E. P. & Guerinot, M. L. Put the metal to the petal: metal uptake and transport throughout plants. *Curr. Opin. Plant Biol.* **9**, 322–330 (2006).
12. Eckhardt, U., Mas Marques, A. & Buckhout, T. J. Iron-regulated cation metabolism from tomato complement metal uptake deficient yeast mutants. *Plant Mol. Biol.* **45**, 437–448 (2001).
13. Moreau, S. et al. GmZIP1 encodes aminobis-specific zinc transporter in soybean. *J. Biol. Chem.* **15**, 4738–4746 (2002).
14. Plaza, S. et al. Expression and functional analysis of metal transporter genes in two contrasting ecotypes of the hyperaccumulator *Thlaspi caerulescens*. *J. Exp. Bot.* **58**, 1717–1728 (2007).
15. Pedas, P. et al. Manganese efficiency in barley: identification and characterization of the major iron transporter HvIRT1. *Plant Physiol.* **148**, 455–466 (2008).
16. Bashir, K., Ishimaru, Y. & Nishizawa, N. K. Molecular mechanisms of zinc uptake and translocation in rice. *Plant Soil* **361**, 189–201 (2012).
17. Ramesh, S. A., Shin, R., Eide, D. J. & Schachtman, D. P. Deferential metal selectivity and gene expression of two zinc metabolism from rice. *Plant Physiol.* **133**, 126–134 (2003).
18. Haydon, M. J. & Cobbett, C. S. Metalloregulation of ligands for essential metal ions in plants. *New Phytol.* **174**, 499–506 (2007).
19. Valle, B. L. & Auld, D. S. Zinc coordination, function, and structure of zinc enzymes and other proteins. *Biochem. 29*, 5647–5659 (1990).
20. Cakmak, I. Enrichment of cereal grains with zinc: agronomic or genetic biofortification?. *Plant Soil* **302**, 1–2 (2008).
21. Briat, J. F. & Lebrun, M. Plant responses to metal toxicity. *C R Acad. Sci. III*(322), 43–54 (1999).
22. Niigababa, E. et al. Winter wheat grain quality, zinc and iron concentration affected by a combined foliar spray of zinc and iron fertilizers. *Agronomy* **9**, 250 (2019).
23. Price, A. H. & Hendry, G. A. F. Iron-catalysed oxygen radical formation and its possible contribution to drought damage in nine native grasses and three cereals. *Plant Cell Environ.* **14**, 477–484 (1991).
24. Palmgren, M. G. et al. Zinc biofortification of cereals: problems and solutions. *Trends Plant Sci.* **13**, 464–473 (2008).
25. Kambe, T., Yamaguchi-Iwai, Y., Sasaki, R. & Nagao, M. Overview of mammalian zinc metabolism. *Cell. Mol. Life Sci.* **61**, 49–68 (2004).
26. Sperotto, R. A. et al. Identification of up-regulated genes in flag leaves during rice grain filling and characterization of OsNAC5, a new ABA-dependent transcription factor. *Planta* **230**, 985–1002 (2009).
27. Sperotto, R. A. et al. Iron biofortification in rice: it’s a long way to the top. *Plant Sci.* **190**, 24–39 (2012).
28. Neeraja, C. N. et al. Transporter genes identified in landraces associated with high zinc in polished rice through panicle transcriptome for biofortification. *PLoS ONE* **13**, e0192362 (2018).
29. Alagarasan, G., Dubey, M., Aswathyl, K. S. & Chandell, G. Genome wide identification of orthologous ZIP genes associated with zinc and iron translocation in *Setaria italica*. *Front. Plant Sci.* **8**, 775 (2017).
30. Manwarthing, H. R., Bleugh, H. P. J. & Yadav, R. The challenges and opportunities associated with biofortification of pearl millet (*Pennisetum glaucum*) with elevated levels of grain iron and zinc. *Front. Plant Sci.* **7**, 1944 (2016).
31. Krzywinska, M. et al. Circos: an information aesthetic for comparative genomics. *Genome Res.* **19**(9), 1639–1645 (2009).
32. Kabir, A. H. et al. Regulation of phytosiderophore release and antioxidant defense in roots driven by shoot-based auxin signalling confers tolerance to excess iron in wheat. *Front. Plant Sci.* **7**, 1684 (2016).
33. Salgueiro, M. J. et al. Zinc status and immune system relationship: a review. *Biol. Trace Elem. Res.* **76**, 193–205. https://doi.org/10.1385/BTRE:76:3:193 (2000).
34. De Benoist, B., McLean, E., Andersson, M. & Rogers, L. Iodine deficiency in 2007: global progress since 2003. *Food Nutr. Bull.* **29**, 195–202 (2008).
35. Shahzad, Z., Rouached, H. & Rakha, A. Combating mineral malnutrition through iron and zinc biofortification of cereals. *Comprehensive Rev. Food Sci. Food Saf.* 13, 329–346 (2014).
36. Weber, M. et al. Comparative microarray analysis of Arabidopsis thaliana and *Arabidopsis halleri* roots identifies nicotianamine synthase, a ZIP transporter and other genes as potential metal hyperaccumulation factors. *Plant J.* 37, 269–281 (2004).
37. Bouis, H. E. & Welch, R. M. Biofortification—a sustainable agricultural strategy for reducing micronutrient malnutrition in the global south. *Crop Sci.* 50, S20 (2010).
38. Palmer, C. M. et al. MYB10 and MYB72 are required for growth under iron-limiting conditions. *PLoS Genet.* 9(11), e1003953 (2013).
39. Chu, H. H. et al. Successful reproduction requires the function of Arabidopsis YELLOW STRIPE-LIKE1 and YELLOW STRIPE-LIKE3 metal-nicotianamine transporters in both vegetative and reproductive structures. *Plant Physiol.* 154, 197–210. https://doi.org/10.1104/pp.115.259103 (2010).
40. Varanesev, R., Ozyigit, I. I. & Filiz, E. Comparative and phylogenetic analysis of zinc transporter genes/proteins in plants. *Turk. J. Biol.* 40, 600–611 (2016).
41. Zhou, X. et al. Genome-wide identification, classification and expression profiling of nicotianamine synthase (NAS) gene family in maize. *BMC Genom.* 14, 238. https://doi.org/10.1186/1471-2164-14-238 (2013).
42. Qin, L. et al. Genome-wide identification and expression analysis of NRAMP family genes in soybean (*Glycine max* L.). *Front. Plant Sci.* 8, 1436 (2017).
43. Curie, C. et al. Maize yellow stripe1 encodes a membrane protein directly involved in Fe (III) uptake. *Nature* 409, 346–349 (2001).
44. Menna, E., Fossati, G., Scita, G. & Matteoli, M. From filopodia to synapses: the role of actin-capping and anti-capping proteins. *Euro. J. Neurosci.* 34, 1655–1662 (2011).
45. Nagaraju, M. et al. Genome-wide in silico analysis of dehydrins in *Sorghum bicolor*, *Setaria italica* and *Zea mays* and quantitative analysis of dehydrin gene expressions under abiotic stresses in *Sorghum bicolor*. *Plant Gene* 13, 64–75 (2018).
46. Li, S. et al. Identification and characterization of the zinc-regulated transporters, iron-regulated transporter-like protein (ZIP) gene family in maize. *BMC Plant Biol.* 13, 114. https://doi.org/10.1186/1471-2229-13-114 (2013).
47. Chen, W. et al. Genome-wide analysis and expression pattern of OsZIP1, OsZIP3, and OsZIP4 in two rice (*Oryza sativa* L.) genotypes with different zinc efficiency. *Russ. J. Plant Physiol.* 55, 400–409 (2008).
48. Fu, X. Z. et al. Genome-wide identification, cloning and functional analysis of the zinc-Iron-regulated transporter-like protein (ZIP) gene family in *Trifoliate Orange* (*Poncirus trifoliata* L. Raf.), *Front. Plant Sci.* 8, 588. https://doi.org/10.3389/fpls.2017.00588 (2017).
49. Nevo, Y. & Nelson, N. The NRAMP family of metal-ion transporters. *Biochim. Biophys. Acta Mol. Cell Res.* 1763, 699–620 (2006).
50. Tarantino, D. et al. Differential involvement of the IDRS cis-element in the developmental and environmental regulation of the AtFer1 ferritin gene from *Arabidopsis thaliana*. *Plant Biol.* 217, 709–716 (2003).
51. Strzyzcki, P. M. et al. A new family of ferritin genes from *Lupinus luteus*-comparative analysis of plant ferritins, their gene structure, and evolution. *Mol. Bio. Evol.* 27, 91–101 (2010).
52. Gross, I. et al. Iron homeostasis related genes in rice. *Genet. Mol. Biol.* 26, 477–497 (2003).
53. Klätte, M. et al. The analysis of Arabidopsis nicotianamine synthase mutants reveals functions for nicotianamine in seed iron loading and iron deficiency responses. *Plant Physiol.* 150, 257–271 (2009).
54. Schnable, P. S. et al. The B73 maize genome: complexity, diversity, and dynamics. *Science* 326, 1112–1115 (2009).
55. Mizuno, D. et al. Three nicotianamine synthase genes isolated from maize are differentially regulated by iron nutritional status. *Plant Physiol.* 132, 1989–1997 (2003).
56. Kumar, S. et al. Mapping quantitative trait loci controlling high iron and zinc content in self and open pollinated grains of pearl millet (*Pennisetum glaucum* (L.) R. Br.). *Front. Plant Sci.* 7, 1636 (2016). eCollection 2016.
57. Lee, S. et al. Zinc deficiency-inducible OsZIP8 encodes a plasma membrane-localized zinc transporter in rice. *Mol. Cells* 29, 551–558 (2010).
58. Lopez-Millan, A. F., Ellis, D. R. & Grusak, M. A. Identification and characterization of several new members of the ZIP family of metal ion transporters in *Medicago truncatula*. *Plant Physiol.* 54, 358–396 (2004).
59. Burleigh, S. H., Kristensen, B. K. & Rechmann, I. E. A plasma membrane zinc transporter from *Medicago truncatula* is up-regulated in roots by Zn fertilization, yet down-regulated by arbuscular mycorrhizal colonization. *Plant Mol. Biol.* 52, 1077–1088 (2003).
60. Yang, X., Huang, J., Jiang, Y. & Zhang, H. S. Cloning and functional identification of two members of the ZIP (Zrt, Irt-like protein) gene family in rice (*Oryza sativa* L.). *Mol. Biol. Rep.* 36, 281–287 (2009).
61. Assunção, A. G. et al. Arabidopsis thaliana transcription factors bZIP19 and bZIP23 regulate the adaptation to zinc deficiency. *Proc. Natl. Acad. Sci. U.S.A.* 107, 10296–10301 (2010).
62. Jebri-Loisy, I. et al. Structure and differential expression of two maize ferritin genes in response to iron and abscisic acid. *Euro. J. Biochem.* 231, 609–619 (1995).
63. Bournier, M. et al. Arabidopsis ferritin 1 (AtFer1) gene regulation by the phosphate starvation response 1 (AtPHR1) transcription factor reveals a direct molecular link between iron and phosphate homeostasis. *J. Biol. Chem.* 288, 22670–22680 (2013).
64. Erogulu, S. & Aksoy, E. Genome-wide analysis of gene expression profiling revealed that COP9 signalosome is essential for correct expression of Fe homeostasis genes in Arabidopsis. *Bio Met.* 30, 685–698 (2017).
65. Kozomara, A. & Griffiths-Jones, S. miRBase: Integrating microRNA annotation and deep-sequencing data. *Nucl. Acids Res.* 39, D152–D157 (2011).
66. Zhang, Z. et al. PMRD: plant microRNA database. *Nucl. Acids Res.* 38, D806–813 (2010).
67. Dai, X., Zhuang, Z. & Zhao, X. P. psRNATarget: a plant small RNA target analysis server (2017 release). *Nucl. Acids Res.* 46(W1), W42–W42. https://doi.org/10.1093/nar/gkx516 (2018).
68. Ye, L. et al. MPK3/MPK6 are iron-induced deficiency ethylene production in *Arabidopsis*. *Front. Plant Sci.* 6, 953. https://doi.org/10.3389/fpls.2015.00953 (2015).
69. Descalsota-Empleo, G. I. et al. Genetic mapping of QTL for agronomic traits and grain mineral elements in rice. *Crop J.* 7, 560–572 (2019).
70. Varshney, R. K. et al. Pearl millet genome sequence provides a resource for improving agronomic traits in arid environments. *Nat. Biotechnol.* 35, 969–976. https://doi.org/10.1038/nbt.3943 (2017).
71. Letunic, I. et al. SMART 4.0: towards genomic data integration. *Nucl. Acids Res.* 32, D142–D144 (2004).
72. Bailey, T. L. & Gribskov, M. Methods and statistics for combining motif match scores. *J. Comput. Biol.* 5, 211–221 (1998).
73. Bailey, T. L., Williams, N., Misleh, C. & Li, W. W. M. MEME: discovering and analyzing DNA and protein sequence motifs. *Nucl. Acids Res.* 34, W369–W373 (2006).
74. Gasteiger, E. et al. Protein identification and analysis tools on the ExPASy server 571–607 (In The proteomics protocols. Handbook, Humana Press, 2005).
75. Blom, N. et al. Prediction of post-translational glycosylation and phosphorylation of proteins from the amino acid sequence. *Proteomics* 4, 1633–1649 (2004).
76. Müller, S., Croning, M. D. & Appweiler, R. Evaluation of methods for the prediction of membrane spanning regions. *Bioinformatics* 17, 646–653 (2001).
77. Horton, P. et al. WoLF PSORT: protein localization predictor. *Nucl. Acids Res.* 35, W585–W587 (2007).
78. Lescot, M. et al. PlantCARE, a database of plant cis-acting regulatory elements and a portal to tools for in silico analysis of promoter sequences. Nucl. Acids Res. 30, 325–327 (2002).
79. Tamura, K., Dudley, J., Nei, M. & Kumar, S. MEGA4: molecular evolutionary genetics analysis (MEGA) software version 4.0. Mol. Biol. Evol. 24, 1596–1599 (2007).
80. Wheal, M. S., Fowles, T. O. & Palmer, L. T. A cost-effective acid digestion method using closed polypropylene tubes for inductively coupled plasma optical emission spectrometry (ICP-OES) analysis of plant essential elements. Analyst. Met. 3, 2854–2863 (2011).
81. Reddy, P. S. et al. Cloning and validation of reference genes for normalization of gene expression studies in pearl millet [Pennisetum glaucum (L.) R. Br.] by quantitative real-time PCR. Plant Genet. 1, 35–42 (2015).
82. Helleman, J. et al. qBase relative quantification framework and software for analysis and automated analysis of real-time quantitative PCR data. Genome Biol. 8, R19 (2007).
83. Dai, X., & Zhao, P. X. psRNATarget: a plant small RNA target analysis server. Nucleic Acids Res. 39(suppl_2), W155–W159 (2011).
84. Vishwakarma, N. P. & Jadeja, V. J. Identification of miRNA encoded by Jatropha curcas from EST and GSS. Plant Signal. Behav. 8(2), e23152 (2013).
85. Vlasblom, J. et al. GenePro: a Cytoscape plug-in for advanced visualization and analysis of interaction networks. Bioinformatics 22, 2178–2179 (2006).

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Author contributions
R.K.S. and P.B.K. planned the experiments and coordinated this study. M.D.M. generated lab and field data. M.D.M. and M.P. generated gene expression data at the Osmania University. M.D.M. and P.B.K. performed bioinformatics experiments. R.K.S., P.B.K., M.P. provided technical guidance during the conduct of research work. M.D.M., P.B.K., R.K.S. drafted the manuscript. R.K.S. edited the manuscript for submission to the Journal.

Competing interests
The authors declare no competing interests.

Additional information
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