Structure, Function, Regulation and Phylogenetic Relationship of ZIP Family Transporters of Plants

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Zinc (Zn) is an essential micronutrient for plants and humans. Nearly 50% of the agriculture soils of world are Zn-deficient. The low availability of Zn reduces the yield and quality of the crops. The zinc-regulated, iron-regulated transporter-like proteins (ZIP) family and iron-regulated transporters (IRTs) are involved in cellular uptake of Zn, its intracellular trafficking and detoxification in plants. In addition to Zn, ZIP family transporters also transport other divalent metal cations (such as Cd²⁺, Fe²⁺, and Cu²⁺). ZIP transporters play a crucial role in biofortification of grains with Zn. Only a very limited information is available on structural features and mechanism of Zn transport of plant ZIP family transporters. In this article, we present a detailed account on structure, function, regulations and phylogenetic relationships of plant ZIP transporters. We give an insight to structure of plant ZIPS through homology modeling and multiple sequence alignment with Bordetella bronchiseptica ZIP (BbZIP) protein whose crystal structure has been solved recently. We also provide details on ZIP transporter genes identified and characterized in rice and other plants till date. Functional characterization of plant ZIP transporters will help for the better crop yield and human health in future.

Keywords: ZIP transporters, homology modeling, transcription factor, functional characterization, genetic modification

INTRODUCTION

In agriculture, the low availability of nutrients has reduced the crop production. The optimal supply of micro-nutrients in soil solutions is vital for normal agriculture production. Zinc (Zn) is one of the most essential micronutrients; it is irreplaceable for plant growth and metabolism (Marschner, 1995, 2011). Both deficient and excess Zn has impaired the physiological and biochemical process of the plant (Cakmak, 2000; Marichali et al., 2014). Zn deficiency is one of the most serious problems worldwide reducing quality of crops (Ruel and Bouis, 1998; Sadeghzadeh, 2013), apart from yield loss and the reduced Zn content in grains (Krithika and Balachandar, 2016). Soil Zn deficiency and human Zn deficiency are often closely associated (Cakmak, 2008). Zn deficiency has been a major cause of death for children in many countries (Black et al., 2008). Regmi et al. (2010) reported that the lack of Zn was a major nutritional problem in humans; more than 3 billion world population were affected by various health problems due to low supply of Zn in their food. For example, approximately 50% of the paddy field is Zn deficient and rice grown on these soils usually produces very less yield with poor nutritional quality (Krithika and Balachandar, 2016). It was estimated that
the Zn sufficient rice plant had 40 mg/kg of Zn in its grains, whereas under Zn deficient condition it showed only 10 mg/kg of Zn (Wissuwa et al., 2008). All over the world, 50% of agriculture soils are Zn deficient (FAO, 2000). Zn deficiency was observed in a wide range of soil types such as high pH calcareous soils, sandy soils, and high phosphorus (P) fertilized soils (Marschner, 1995). In 1972, the United States considered Zn deficiency as the most common micronutrient deficiency in crops (Lindsay, 1972). Zn deficiency is one of the major limiting factors affecting crop production badly. So, improving crop varieties with Zn efficiency is helpful to overcome the Zn deficiency problem.

To overcome low Zn availability, plants have evolved a complex array of tightly controlled adaptive mechanisms. The Zinc-regulated, Iron-regulated transporter-like Protein (ZIP) family has been identified and characterized in prokaryotes, eukaryotes, and archaeotes and has been validated to be involved in metal uptake and transport including Zn$^{2+}$ (Grotz and Guerinot, 2006; Kavitha et al., 2015). The presence of ZIP transporters in such a diverse organism indicates their pivotal role in Zn homeostasis. The ZIP transporters are also located in various cell organelles and their role in transport is to balance the uptake, utilization, and storage of Zn$^{2+}$ under Zn stress condition (Ramesh et al., 2003; Palmgren et al., 2008). The ZIP transporters are also located in various cell organelles and they are actively involved in Zn homeostasis and plant’s adaptation to low and high Zn soils (Tiong et al., 2015). In crops, most of the ZIP transporters are poorly understood with only ZIPs of a few plants have been functionally characterized (Kavitha et al., 2015). The detailed analysis of the structure of the plant ZIP transporter is still lacking. Recently, the crystal structure of the ZIP protein was deduced from a bacterium Bordetella bronchiseptica and its metal transport mechanism was also predicted (Zhang et al., 2017). So, in this article, we present the details on the structure, mechanism and phylogenetic relationships of plant ZIP transporters. We provided insight into the structure of plant ZIPs through homology modeling and multiple sequence alignment using B. bronchiseptica ZIP (BbZIP) protein as a template. These findings would be a valuable theoretical knowledge for future studies on Zn transporters in crops. We also present details on ZIP transporter genes identified and characterized till date in various plants.

**ROLE OF Zn IN CROPS**

Zinc is one of the eight essential micronutrients in plants (Hänisch and Mendel, 2009). In crops, Zn deficiency was first identified in rice on calcareous soils in India (Nene, 1966; Yoshida and Tanaka, 1969). Plant cells require optimum levels of Zn for normal physiological functions. Zn is involved in the maintenance of the structural and functional integrity of biological membrane and facilitation of protein synthesis, gene expression and regulation and defense against disease (Cakmak, 2000; Andreini et al., 2006; Sadeghzadeh, 2013). Also, it is involved as a structural, catalytic, and intracellular and intercellular signaling component and Zn is the only metal required for the activity of all six classes of the enzymes (Sadeghzadeh, 2013). Especially Zn is essential for the activity of metallo-enzymes that are involved in protein and nucleic acid metabolism (Eide, 2006).

Zinc is needed by a small quantity (0.5–2 μM) from the soil for the normal plant function (Krishna et al., 2017), but yet crucial for many physiological and metabolic pathways (Chen et al., 2008). In rice, 1.5 μM Zn is an optimum level for growth on agar nutrient solution (Impa et al., 2013). Zn content less than 15–20 μg per gram of dry leaf tissues of the plant is a sign of Zn deficiency (Mitra, 2015). Zn is actively involved in specific reactions of metabolic pathways, such as tryptophan biosynthesis, which in turn is the precursor of indole-3-acetic acid (IAA) and other phytohormones. In plants, Zn$^{2+}$ ions are directly involved in the synthesis of tryptophan and auxin (Horák and Trčka, 1976; Marschner, 2011). Zn deficiency decreases the level of phytohormones such as auxin, abscisic acid, gibberellins, and cytokinin’s (Kumar et al., 2016). It significantly affects the plant cell division, cell enlargement, and differentiation (Skøog, 1940; Tsui, 1948). Cakmak et al. (1989) observed that under Zn deficient condition, the level of IAA in the shoot tips and young leaves of bean was reduced to 50% of that in Zn sufficient condition. Zn deficiency reduces the plant growth, yield, and quality in crops such as rice (Chen et al., 2008), mungbean (Samreen et al., 2017), and maize (Wang and Jin, 2005). Zn deficiency decreases the activity of key photosynthetic enzymes, namely carbonic anhydrase, (Brown et al., 1993) which is crucial for crop production (Ali et al., 2008; Mousavi, 2011; Xi-wen et al., 2011). The Zn deficiency induces male sterility in maize (Sharma et al., 1987) and wheat (Sharma et al., 1979). Ekiz et al. (1998) noticed that the crops such as wheat, oat, barley, and triticale show significant decrease in growth and grain yield under Zn-deficient conditions. Zn is a not replaceable micro-nutrient for crops. Zn deficiency is a common problem in all parts of the globe. Therefore, much more research is necessary for improving Zn use efficiency in crops for growth and higher yields under Zn deficient soils.

**ROLE OF ZIP TRANSPORTERS IN Zn TRANSPORT**

In plants, Zn is acquired and transported predominantly as Zn$^{2+}$ (divalent). Zn ions can also be bound with root exudates like
malate, citrate, oxalate and other low molecular weight organic acids which aided to move toward the root surface area (Suzuki et al., 2006). The charged Zn\(^{2+}\) ion do not freely diffuse across the lipid bilayer membranes (Eide, 2005). Zn first enters the root cell wall’s free space by a diffusion process from the soil solutions (Hacisalihoglu and Kochian, 2003). Transport of Zn\(^{2+}\) into the cortex takes place via symplastic or apoplastic pathway (Kumar et al., 2016). The Zn transporter proteins are required to carry Zn\(^{2+}\) into the cells and transport out of intracellular compartments. They include a low-affinity (Km = 2–5 \(\mu\)M) and high-affinity (Km = 0.6–2 nM) membrane transporter systems (Hacisalihoglu et al., 2001; Kumar et al., 2016). The high-affinity transporter system is dominantly active under low Zn soil solutions (Hacisalihoglu and Kochian, 2003). Transport of Zn\(^{2+}\) into the lipid bilayer membranes (Eide, 2005). Zn first enters the parenchymatous cells which subsequently loaded into the xylem. The heavy metal ATPase 2 (HMA2) and HMA4 transporters of P-type ATPase family are involved in xylem loading of Zn from the xylem parenchymatous cells (Hussain et al., 2004; Hanikenne et al., 2008). The ZIP family transporters are mainly involved in up take, transport and distribution of Zn in the whole plant. The presence of ZIP transporters in a diverse organism indicates their importance in Zn transport and homeostasis. Therefore, understanding of their expression levels, localization, and function in crops is essential. The functional genomics and biotechnological approaches can be used to develop Zn deficiency tolerant crops in the future. It assists in improving the quality and quantity of crops especially enrichment of Zn content in grains and to overcome Zn deficiency problems worldwide. The ZIP transporter families can be used for genetic modification in crops to improve the fortification of Zn.

**STRUCTURE OF THE ZIP TRANSPORTER**

Only a very limited information is available on structural features and mechanism of Zn transport of plant ZIP family transporters. Plant ZIP family transporters are predicted to have 8 transmembrane (TM) domains (\(\alpha\)-helices) with 8 being the most prevalent form (Guerinot, 2000). The molecular weight of the Zn transporters ranged from 33.1 to 51.4 kDa and protein sequence ranged from 322 to 478 amino acids (Vatansever et al., 2016). Understanding the mechanism of ZIP transporters requires their high-resolution crystal structures. The crystal structure of the plant ZIP protein is not yet available. However, recently, a high-resolution crystal structure of prokaryotic ZIP protein was deduced from a bacterium B. bronchiseptica and its metal (Cd\(^{2+}\) and Zn\(^{2+}\)) transport mechanism was also predicted (Zhang et al., 2017). The crystal structure of BbZIP transporter protein showed eight TM domains (TM1–TM8). The structure of BbZIP was deduced with an inward-open confirmation and occluded at the extracellular side with a binuclear metal center located in the center. The eight TMs formed a closely associated \(\alpha\)-helix bundle. The first three TMs (TM1–TM3) can be superimposed on the last three TMs (TM6–TM8) by rotating 180\(^\circ\), and TM4 and TM5 are symmetrically related and sandwiched by the two 3 TM repeats (Zhang et al., 2017). The binuclear metal center is formed by TM4 and TM5 with several conserved amino acid residues (His177, Asn178, Pro180, Glu181, and Gly182) and metal-binding motifs (Gln207, Asp/Asn208, Pro210, Glu211, and Gly212). The structure of the BbZIP showed four Cd\(^{2+}\) and seven Zn\(^{2+}\) metal-binding sites (Zhang et al., 2017). The topology of BbZIP shows that both the N and the C termini are exposed to the extracellular space and the putative metal transport pathway found at the extracellular side is blocked by hydrophobic residues of the TM2 (Met99 and Ala102), TM5 (Leu200 and Iso204), and TM7 (Met269) (Zhang et al., 2017). The invariant Ser106 on TM2 is situated on the bottom of the shallow and negatively charged entrance cavity, it is important for guiding metals into the transport pathway. Two invariant metal-chelating residues Asp113 and Asp305 present in the entrance cavity have been predicted to be crucial for recruiting metal substrates. The BbZIP structure shows multiple conserved metal-binding sites near the metal exit cavity, indicating that these constitute a route of metal release to the cytoplasm. The bound metal released into the cytoplasm through a chain of metal-chelating residues such as His177, Glu276, His275, Pro180, Pro210, and Asp144 (Zhang et al., 2017). The Zn\(^{2+}\) binding is penta coordinated by Glu181, Glu207, and Glu211 and two molecules of water (Zhang et al., 2017). This structure provides a model to analyze the plant ZIP transporters. So we have used this as a template to model and analyze the residues in plant ZIP transporters.

**Comparison of Metal Binding Residues in BbZIP and Plant ZIPS**

We have analyzed the BbZIP protein sequence with plant ZIP transporters such as Arabidopsis (12 ZIPS), rice (12 ZIPS), and maize (10 ZIPS) through a ClustalW alignment (Tamura et al., 2013). The functional residues of the BbZIP and their corresponding plant ZIP residues are presented in Figure 1. We have also included the complete alignment file (Supplementary Figure S1). Except a few, most of the functional residues of Arabidopsis, rice and maize ZIP protein sequences are not conserved/homologous with BbZIP. But it is interesting to see that all the plant ZIP protein sequences (except AtZIP10 and OsZIP16) have conserved His117 residue found in BbZIP (Figure 1). In AtZIP10 (Gln222) and OsZIP13 (Gln140) BbZIP’s His177 is replaced by Gln. Gly182 of BbZIP is conserved in all plant ZIPS (Figure 1). The His177 and Gly182 are involved in the metal release from the metal-binding site of the BbZIP. Similarly, Glu211 and Gly212 are metal-binding residues in BbZIP. Glu211 of BbZIP is conserved in all plant ZIP proteins expect for AtZIP2 (Ala243), AtZIP11 (Ala215), OsZIP1 (Ala242), and OsZIP2 (Ala247) where Glu is replaced by Ala. Similarly, expect AtZIP2 (Ala224), AtZIP11 (Ala216), OsZIP1 (Ala243), and OsZIP2 (Ala248), other plants ZIP transporter proteins possess conserved Gly212 of BbZIP. Another interesting observation is that metal-binding site residues such as Asn178, Pro180, and Pro210 are conserved only with OsZIP13,
OsZIP16, and ZmZIP6 of plants (Figure 1). Metal binding site residues Gln207 and Asp208 and metal-chelating residue Asp305 are not conserved with any other plant ZIPS. The Gln207 is replaced by His residue in all plant ZIPS (Figure 1). An invariable Ser106 is an important residue for guiding metals into the transport pathway of the BbZIP. In plants, Ser is completely replaced by His or Asp residue with His being a dominant residue. Overall, the metal binding and transport residues of OsZIP13, OsZIP16, and ZmZIP6 are more conserved with BbZIP compared to other plant ZIPS. These variations indicated that plant ZIPS transporters may have partially overlapping but distinct metal transport mechanism compared to BbZIP as some of the residues involved in metal binding and transport are not conserved in plant ZIPS. We have also done homology modeling of plant ZIPS whose details are discussed below.

Homology Modeling of Plant ZIP Transporters

In order to gain more insights to the structure of plant ZIP transporters, we have modeled some of the plant ZIP transporters by homology modeling using BbZIP (PDB Id: 5TSA) as a template with Zn$^{2+}$ as a ligand. The models
were then compared with the structure of BbZIP. In BbZIP, Zn$^{2+}$ is coordinated by Glu181, Gln207, and Gln211 and water molecular at metal binding site as per our modeling (Figure 2; Zhang et al., 2017). The Zn$^{2+}$ binding site is also covered by Glu276, His177, Met269, and Met99 which are involved in Zn transport (Figure 2). However, the plant ZIP transporters showed greater variation at Zn$^{2+}$ binding site except for Glu181/211 (equivalent to BbZIP) based on the homology models (Figure 2 and Table 1). Zn$^{2+}$ binding is coordinated by Glu141 and Glu170 in ZmZIP6, Glu252 and Asp190 in ZmZIP11, Glu173 and Glu240 in OsZIP16, Glu253 and Asn30 in OsZIP13. Even Zn$^{2+}$ is co-ordinated only by a single Glu residue in AtZIP1 and AtZIP2 and not co-ordinated by any residues in AtZIP8 (Figure 2 and Table 1). The residues surrounding the Zn$^{2+}$ binding site include Glu276, Met269, His177, and Met99 in BbZIP. These residues are partially conserved in some plant ZIPs and not at all conserved in other ZIPs modeled (Table 1). For e.g., His177 is conserved in OsZIP6 (His169) and ZmZIP6 (His166); shifting in the positioning of His is seen in AtZIP1 (His238), AtZIP2 (His239), OsZIP13 (His249), and ZmZIP11 (His278) as indicated in the alignment above (Figures 1, 2 and Table 1). Similar variation was seen for residues involved in Zn transport in the plant ZIPs analyzed (Table 1). It might be due to partially overlapping but distinct mechanism for binding and transport of Zn$^{2+}$ by plant ZIPs which is also reflected in alignment (Supplementary Figure S1). This requires further studies to confirm the roles of functional residues especially by site directed mutagenesis with yeast mutants to gain more knowledge on residues involved in Zn$^{2+}$ binding and transport in plant ZIPS. Also, the high-resolution crystal structure of plant ZIP is needed for understanding any specific Zn$^{2+}$ transport mechanism in plants.

### Phylogenetic Analysis of Plant ZIP Proteins

The phylogenetic tree was constructed from 113 ZIP protein sequences collected from 14 plant species. These include eight monocot plants and six dicot plants (Supplementary File S1). The phylogenetic tree consisted of eight major clusters (MCs) (MC1–MC8) and sub-divided into 1–2 sub-clusters (SCs) and each SC was again divided into small clusters (Figure 3). More ZIP transporters are clustered in MC-8. The MC-8 had 21 ZIP transporters and it is followed by MC-5 (20 ZIPs) and MC-2 (18 ZIPs). The ZIP proteins of *Brassica rapa* (BrZIP1), *Cucumis sativus* (CsZIP1), *Medicago truncatula* (MtZIP1), and *Glycine max* (GmZIP1) were clustered with AtZIP1; BrZIP3 is closely clustered with AtZIP3 (Figure 3). The AtZIP1 and AtZIP3 proteins are clustered closely which are characterized as low-affinity transporters (Grotz et al., 1998). In *Arabidopsis*, AtZIP1, and AtZIP3 are highly expressed in roots in response to Zn deficiency condition suggesting their role in Zn uptake from soil (Grotz et al., 1998). OsZIP1 and OsZIP3 are characterized as low-affinity Zn transporters in rice plants (Ramesh et al., 2003) and these are placed distinctly in the phylogenetic tree. The OsZIP1 is clustered with monocot ZIP transporters such as TaZIP1, *Panicum hallii* ZIP1 (PhZIP1) and SbZIP1. The expression analysis showed the OsZIP1 gene is induced in roots under Zn
starvation (Ramesh et al., 2003). Similarly, the OsZIP3 is closely clustered with HvZIP3, TaZIP3, BdZIP3, and SbZIP3. HvZIP3 was reported as a low-affinity transporter of barely (Pedas et al., 2009). Both OsZIP3 and HvZIP3 are highly expressed in both shoot and root under Zn deficient condition (Chen et al., 2008). AtZIP2 is a high-affinity transporter in Arabidopsis (Grotz et al., 1998), it is closely clustered with BrZIP2, SIZIP2 MtZIP2, TaZIP1, OsZIP1, PhZIP1, and SbZIP1 proteins. It is interesting to see that both low-affinity (OsZIP1) and high-affinity (AtZIP2) ZIPs are closely clustered. The monocot and dicot plant ZIPs are clustered in separate clusters. ZIP family member are mostly clustered together as per their numbers (Figure 3), for e.g., ZIP1s of Arabidopsis, mustard, barley clover, soybean and sweet orange are closely clustered. The identification and characterization of the ZIP gene family are still lacking for many crops. More molecular and functional genomic studies will aid for the characterization of ZIP transporter in other plants in future.

**REGULATION OF ZIP TRANSPORTERS IN PLANTS**

All living organisms need to maintain optimum concentrations of nutrients including Zn to sustain cellular functions (Shaul et al., 1999). Plants possess the homeostatic networks to maintain Zn levels in a narrow concentration to avoid either deficiency or toxicity. Zn homeostasis needs a complex network of cellular functions such as Zn uptake, accumulation, trafficking, sequestration, remobilization, and detoxification (Clemens, 2001; Krishna et al., 2017). ZIP transporters are regulated to provide a suitable quantity of Zn into all cell types and at all stages of development (Clemens, 2001). The Zn homeostasis equilibrium was considered for external and internal requirements of Zn (Claus and Chavarría-Krauser, 2012). Unfortunately, very little information is available on the regulation of plant ZIP transporters and mechanism of Zn homeostasis. Transport of divalent metal ions like Zn$^{2+}$ by different transporters may pose some difficulty for high resolution studies to understand the role of plant ZIPs in Zn homeostasis. Expression levels of genes are increased for some ZIP transporters under Zn deficiency to facilitate the higher uptake of Zn from the soil. For e.g., in barley, HvZIP3, HvZIP5, and HvZIP8 genes are highly induced in roots and are involved in Zn uptake under Zn deficient condition (Pedas et al., 2009).

Till date plant’s sensing and transferring the signal of Zn deficiency remains poorly understood. However, based on the available data, some transcription factors (TFs) are found to be essential for regulation of target genes and maintaining Zn homeostasis. Two TFs identified in model plant Arabidopsis are shown to be crucial in the adaptation response to Zn deficiency. The basic-region leucine zipper (bZIP) is a TF involved in the regulation of many physiological processes including abiotic and biotic stress responses (Corrêa et al., 2008). These TFs belong to the F group of bZIP and showed histidine-rich motifs at the basic N-terminal region (Nijhawan et al., 2008; Assunção et al., 2010). bZIP TFs are also involved in the up-regulation of ZIP transporters in Arabidopsis during Zn deficiency (Assunção et al., 2013). The TFs bZIP19 and bZIP23 are considered to be the essential regulators of candidate genes including ZIPs under Zn deficiency. In plants, bZIP19 and bZIP23 proteins exist as monomers under Zn sufficient condition. The Zn deficiency leads to the activation (binding) of bZIP9 and bZIP23 (dimerization) which induces the expression of ZIP genes. The bZIP19 and bZIP23 dimer bind to 10bp Zn-deficiency responsive elements (RTGTCGACAY) present in the promoter region of target genes (Assunção et al., 2010; Lilay et al., 2018). The histidine-rich motif is conserved among group F bZIP TFs (Jakoby et al., 2002; Castro et al., 2017) and suggested to play key role as a Zn-sensor (Assunção et al., 2013; Henriques et al., 2017). Totally, 33 bZIP19 genes are identified and characterized in cereal crops using in silico approaches and histidine-rich motifs are found in most of these bZIP19 TFs (Henriques et al., 2017).

In Arabidopsis, the TFs AtbZIP19 and AtbZIP23 induce the expression of AtZIP4 gene under Zn deficiency (Assunção et al., 2010). These TFs are also involved in the activation (up-regulation) of a specific subset of genes such as AtZIP1, AtZIP2, AtZIP4, AtZIP5, AtZIP9, AtZIP10, and AtZIP12 (Assunção et al., 2010; Inaba et al., 2015; Lilay et al., 2018). Similarly, three bZIP TFs such as bZIP1, bZIP2, and bZIP3 were identified in common bean which are similar to Arabidopsis TFs bZIP19, bZIP23, and bZIP24 (Astdil et al., 2013). In whole-genome transcriptome analysis, 3 bZIP TFs (PvBZIP1, PvBZIP2, and PvBZIP3) are found to be expressed under Zn deficiency condition in common bean (Astdil-Reyes et al., 2015). But

**TABLE 1 |** Details on residues identified to be involved in Zn$^{2+}$ transport through homology modeling using BbZIP as a template.

| Name of the protein | UniProt Accession/Phytozome ID | Length (total amino acids) | Zn$^{2+}$ coordinating residues | Zn$^{2+}$ transporting residues |
|---------------------|--------------------------------|---------------------------|--------------------------------|-------------------------------|
| BbZIP               | BB2405                         | 309                       | Glu181; Glu211 Gin207          | Glu276; Met 269 His 177; Met 99 |
| AtZIP1              | Q8I123                         | 355                       | Glu242                          | His238; Lys 89 Ile 206; Met 312 |
| AtZIP2              | Q9LTH9                         | 353                       | Glu216                          | His239; Leu 77 Phe 73          |
| AtZIP8              | Q8S3W4                         | 347                       | –                               | Glu248; Asn 251 Ile 209; Ala 247 Phe 71 |
| OsZIP13             | LOC_Os07g12890 (Chen et al., 2008) | 276                     | Asn 30; Glu 253                | Ile 59; Met 321 His 249; Ala 85 Cys 28 |
| OsZIP16             | LOC_Os08g01030 (Chen et al., 2008) | 282                     | Glu 173; Glu 240               | His 169; Met 48 Val 144; Ala 49 |
| ZmZIP6              | GRMZM2G0050484_T01 (Mondal et al., 2013) | 297                     | Glu 141; Glu 170               | Glu 237; Met 51 His 166; Ala 47 |
| ZmZIP11             | GRMZM2G034551 (Mondal et al., 2013) | 396                     | Glu 282 Asp 190               | His 278; Ile 74 Lys 198; Met 159 Ala 70 |
no information is available on target genes of these TFs. Similarly, seven bZIP TFs were identified in the wheat genome. Seven out of four TabZIP TFs such as TabZIPF1-7DL, TabZIPF3b-7BL, TabZIPF4-7AL, and TabZIPF4-7DL of wheat were used for functional complementation assay (Evens et al., 2017). The role of these four TabZIP TFs in the Zn homeostatic mechanism was determined by expression in the Arabidopsis double mutant line bzip19-4 bzip23-2 under Zn deficient condition. The TabZIPF1-7DL and TabZIPF4-7AL could partially complement the double mutant line (bzip19-4 bzip23-2) under Zn-deficient condition. TabZIPF3b-7BL have slightly increased the growth of the double mutant line under Zn-deficient condition whereas TabZIPF4-7DL did not complement the double mutant lines (Evens et al., 2017). Similarly, seven bZIP TFs HvbZIP1, HvbZIP10, HvbZIP55, HvbZIP56, HvbZIP57, HvbZIP58, and HvbZIP62 were identified.

FIGURE 3 | Phylogenetic tree of ZIP transporter family proteins of plants. The ZIP phylogenetic tree was constructed from 113 ZIP transporter protein sequences collected from 14 plant species. These included 8 monocot (Blue) and 6 dicot plants (Black). Each major cluster (MC1-MC8) is highlighted with different color. The low-affinity and high-affinity ZIP transporters are highlighted in red and green, respectively. The protein sequences of ZIP transporter family members were collected from Phytozome (www.phytozome.net) website. The phylogenetic tree was constructed by MEGA version 6 software with the maximum likelihood method based on the Jones-Taylor-Thornton matrix-based model. The bootstrap values are from 1000 replicates. The phylogeny tree was visualized by iTOL (https://itol.embl.de/).
in barely (Nazri et al., 2017). Among these, the TFs HvZIP56 and HvZIP62 could restore the growth of bzip19 and bzip23 double mutant of Arabidopsis under Zn deficiency condition (Nazri et al., 2017). Recently, Lilay et al. (2020) identified and characterized three bZIP TFs in rice such as OsbZIP48, OsbZIP49, and OsbZIP50. The OsbZIP48 and OsbZIP50 complemented the Arabidopsis double mutant (bzip19/bzip23) under Zn deficient condition but the OsbZIP49 does not complement the Arabidopsis double mutant (bzip19/bzip23) (Lilay et al., 2020).

In plants, concentrations of both the macro- and micro-nutrients influence each other through molecular cross-talks. Signals of P nutrition interact with those of the micronutrients like Zn and Fe. The TFs such as Phosphate starvation response 1 (PHR1) (Rubio et al., 2001), ZAT6 (Devai et al., 2007b), WRKY75 (Devai et al., 2007a), and MYB62 (Devai et al., 2009) are involved in P deficiency response. The PHR1 acts as a positive regulator of inorganic phosphate (Pi) starvation responsive genes and PHR1 along with Phosphate 1 (PHO1) are involved in Zn and Fe homeostasis in Arabidopsis (Khan et al., 2014; Rai et al., 2015). Zn deficiency also induced the expression of P stress-responsive genes through PHR1 and increased the uptake of Pi (Bouain et al., 2014). PHR1 seems to be a positive regulator of the ZIP transporters (AtZIP2 and AtZIP4) under Pi deficiency in Arabidopsis (Briot et al., 2015). The cross-talk of signals between P, Zn, and Fe were highlighted by many articles (Briot et al., 2015; Xie et al., 2019). Only a very little research is done on the characterization Zn responsive and ZIP related TFs in crops and most of the reports are also confined only to plants like Arabidopsis. Therefore, more studies are needed on the identification and characterization of Zn responsive TFs in crops. This could help to understand the molecular mechanism of Zn deficiency tolerance and may aid in developing Zn deficiency tolerant crops.

ZIP FAMILY TRANSPORTERS IDENTIFIED IN VARIOUS CROPS

We have enlisted the ZIP family members identified so far in crops (Supplementary Table S1), and the details are discussed below. ZIP family transporter genes have been identified and functions characterized majorly in model plants like Arabidopsis and rice. Identification and characterization of the ZIP family genes are still lacking for many crops. Expression levels of ZIP family genes showed dynamic pattern in crops. For e.g., a few ZIP transporters are expressed only under deplete Zn conditions, so their expression is declined within 2 h when Zn is added to the medium (Van de Mortel et al., 2006). The ZIP transporters are localized to plasma membrane and membranes of many intracellular organelles (Figure 4). It is proved that the plant ZIP transporters are involved in Zn uptake at the cellular level when tested in yeast complementation test (Fu et al., 2017). In model plant Arabidopsis, the ZIP transporters such as AtZIP1, AtZIP2, AtZIP3, and AtZIP4 have been functionally characterized to be the transporters with different affinities (Grotz et al., 1998; Mäser et al., 2001; Assunção et al., 2010). Evidence also shows the direct involvement of ZIP transporter in enhancing Zn accumulation at edible parts of the plants (Ramegowda et al., 2013; Gaitán-Solís et al., 2015). Rice has been the only cereal model crop whose ZIP family transporters have been studied to a reasonable extend when compared to other crops. So, the details on rice ZIP and IRT transporters are discussed in detail in the following section which could serve as a model to study the ZIP and IRT transporters of other crops especially cereals. We also discussed the details on ZIP family transporters identified in other crops.

Rice

Rice is a staple food for more than 560 million people in the world; it is one of the highly sensitive crops to Zn deficiency (Krithika and Balachandar, 2016). Many works were conducted on rice to understand the Zn transport mechanism and rice seems to be a good model for other crops to study the Zn transport. Totally 16 ZIP transporter members were identified in rice (Ramesh et al., 2003; Ishimaru et al., 2005; Chen et al., 2008). Some OsZIP genes are expressed in both roots and shoots and others are expressed in whole parts of the plant such as root, culms, leaves, and spikelets under Zn deficiency as reported by different authors (Figure 5A). In rice, several ZIPs such as OsZIP1, OsZIP3, OsZIP4, OsZIP5, OsZIP7, and OsZIP8 were reported to be responsible for Zn uptake from soil, translocation from root to shoot as well as for grain filling (Ramesh et al., 2003; Ishimaru et al., 2005, 2007; Lee et al., 2010b; Meng et al., 2018). In rice, understanding the molecular mechanisms of Zn transport is important for improving the Zn content in the edible part. Ishimaru et al. (2011) highlighted the mechanism of Zn uptake and translocation in rice. The OsZIP1 and OsZIP3 seem to be important for Zn uptake from the soil, OsZIP4, OsZIP5, and OsZIP8 for root to shoot translocation, while OsZIP4 and OsZIP8 played a role in grain filling (Bashir et al., 2012). The expression levels of OsZIP3 and OsZIP4 are higher in the roots of Zn-efficient genotype (IR8192) than those of in-efficient genotype (Erjiufeng) suggesting that these genes could contribute to high Zn efficiency (Chen et al., 2008).

Under Zn deficient condition, OsZIP1, OsZIP3, and OsZIP4 are up-regulated in the roots and OsZIP4 over-expressed in the shoot of both genotypes (Chen et al., 2008). These transporters are localized to the vascular bundles in shoots and in the vascular bundles and epidermal cells in roots of rice (Ramesh et al., 2003; Ishimaru et al., 2006). The expression analysis shows that OsZIP4 is highly expressed under Zn deficient conditions in roots and shoots in rice (Ishimaru et al., 2005). It is expressed in the meristem of the roots and shoots, and also in vascular bundles of the roots and shoots under Zn deficient condition (Ishimaru et al., 2005; Ishimaru et al., 2011). The expression of OsZIP1 gene is seen only in root tissues under Zn starvation, but OsZIP3 is expressed in both roots and shoots under both Zn sufficient and deficient conditions (Ramesh et al., 2003). OsZIP7 and OsZIP8 are expressed in roots and shoots under Zn deficient condition (Yang et al., 2009; Tan et al., 2019). The OsZIP3 and OsZIP4 are highly expressed in the nodal regions under Zn deficiency condition (Sasaki et al., 2015). OsZIP1 is expressed in the epidermis and vascular tissues of roots and leaves of rice (Ramesh et al., 2003; Bashir et al., 2012). The OsZIP4, OsZIP5,
FIGURE 4 | Localization of ZIP transporter family proteins in the plant cell. The ZIP transporter family proteins are actively involved in uptake, transport, detoxification and homeostasis of Zn within plant cells (Li et al., 2013). Under low Zn condition, many ZIP transporters (ZIP1-ZIP12) are expressed and are localized in plasma membrane of the cell in rice, (Ramesh et al., 2003; Ishimaru et al., 2005; Yang et al., 2009; Lee et al., 2010a,b) hardy orange (Fu et al., 2017), maize (Li et al., 2013; Mondal et al., 2013), Arabidopsis (Milner et al., 2013), and barley (Fedas et al., 2009; Tiong et al., 2015). Some of the ZIP transporters are also localized to the membranes of intracellular organelles such as chloroplast (ZmZIP5, ZmZIP7 (Li et al., 2013)), vacuole (HvZIP5 (Tiong et al., 2015) and AtZIP1 (Milner et al., 2013)), and endoplasmic reticulum (ZmZIP1-ZmZIP8 (Li et al., 2013)).

and OsZIP8 transporters are localized to plasma membrane and are involved in Zn influx (Ishimaru et al., 2005; Lee et al., 2010a,b). The OsZIP3 gene was expressed in the nodal region which is responsible for unloading Zn from xylem (Sasaki et al., 2015). OsZIP7 is located in the parenchyma cells of vascular bundles in nodal region, and in the stele in the roots of rice (Tan et al., 2019). OsZIPs might be actively involved in the uptake and transport of Zn in rice. Similarly, Ishimaru et al. (2011) suggest that OsZIP4 may be responsible for Zn translocation to aerial parts.

The OsIRTs are similar to OsZIP metal transporters; these are the members of ZIP family transporters (Guerinot, 2000). Especially OsZIP4, OsZIP5, OsZIP6, and OsZIP7 showed sequence similarity to OsIRT1 (Ishimaru et al., 2005). The expression analysis revealed that OsIRT1 and OsIRT2 are mainly expressed in roots of rice under low Fe conditions (Bugnio et al., 2002; Ishimaru et al., 2006; Lee et al., 2009). The expression level OsIRT1 was much higher than OsIRT2 in the Fe deficient root of rice (Ishimaru et al., 2006). OsIRT1 and OsIRT2 have been cloned and characterized in yeast (Bugnio et al., 2002; Ishimaru et al., 2006). The OsIRT1 and OsIRT2 proteins localize to the plasma membrane and have been shown to complement the growth defect of a yeast Fe uptake mutant, confirming that they are functional Fe$^{2+}$ transporters under low Fe condition (Bugnio et al., 2002; Ishimaru et al., 2006). The OsIRT transporters also capable of transporting other divalent metal cations such as Cd$^{2+}$, Zn$^{2+}$, Cu$^{2+}$, and Mn$^{2+}$ (Bugnio et al., 2002; Nakanishi et al., 2006). Cellular and sub-cellular metal homeostasis is very crucial for maintaining the optimal metabolic process and cellular functioning (Bashir et al., 2016). Therefore, identification and characterization of IRT transporters are essential for understanding the mechanism of Fe homeostasis in rice and other plants.

The information available on Zn transporters of rice plants are high when compared to other crops such as maize, wheat, barley, foxtail millet, orange, and common bean, etc. This large information is helpful for understanding the expression pattern of Zn transporters, their localization and metal homeostasis in other crops. Apart from these reports we need further studies on the biofortification of rice with Zn in order to increase the
The ZIP transporters are involved in Zn biofortification in other crops. For example, the OsZIP4 and OsZIP8 are actively involved in the grain filling of Zn in rice (Ishimaru et al., 2007; Lee et al., 2008). The expression analysis indicated that ZmZIP4, ZmZIP5, ZmZIP7, and ZmZIP8 are sensitive to Zn status during the seedlings stage of maize (Li et al., 2013). Under Zn deficient condition, ZmZIP5 and ZmZIP8 are induced in shoot and ZmZIP3 is up-regulated in both root and shoot (Li et al., 2013). The expression of ZmZIP4, ZmZIP5, ZmZIP7, and ZmZIP8 decreased in shoots and ZmZIP3 is down-regulated in roots of maize under excess Zn supply (Li et al., 2013). Similarly, 12 ZmZIP genes (ZmZIP1–ZmZIP12) were identified by Mondal et al. (2013) in maize. Ten ZmZIP genes (ZmZIP1, ZmZIP2, and ZmZIP4–ZmZIP11) are tissue-specific and are highly expressed in flag leaf except ZmZIP3 and ZmZIP12, under Zn deficient condition (Mondal et al., 2013). The ZmZIP2, ZmZIP5, ZmZIP6, ZmZIP8, and ZmZIP11 are expressed in the kernel under Zn deficient condition (Mondal et al., 2013). ZmZIP4, ZmZIP5, ZmZIP7, and ZmZIP9 proteins are located in cytoplasm and chloroplast (Mondal et al., 2013; Figure 4). It is well known that some ZIP transporters are involved in Zn biofortification in other crops. For example, the OsZIP4 and OsZIP8 are actively involved in the grain filling of Zn in rice (Ishimaru et al., 2007; Lee et al., 2010b). The ZmZIP5 and ZmZIP11 are highly expressed in the flag leaf, which might play a vital role in the mobilization of Zn from flag leaf to developing kernel for the accumulation of the large amount of Zn in the kernel (Mondal et al., 2013; Figure 5B). These ZmZIPs (ZmZIP5 and ZmZIP11) might contribute to the biofortification of maize with Zn to improve the quality (Mondal et al., 2013). Since maize is highly sensitive to Zn deficiency, more high resolution studies are needed to improve the quality of maize grown under low Zn soils.

**Maize**

Maize is one of the most widely cultivated and important cereal crops for human and animal food (Shiferaw et al., 2011). Zn is the most common limiting micronutrient in maize yield worldwide (Alloway, 2009). The maize genome has been sequenced and assembled (Palmer et al., 2003). Li et al. (2013) identified eight ZIP transporters (ZmZIP1–ZmZIP8) in maize genome. All eight ZmZIP proteins are localized to the plasma membrane (Li et al., 2013). The expression analysis indicated that ZmZIP3, ZmZIP4, ZmZIP5, ZmZIP7, and ZmZIP8 are tissue-specific and are highly expressed in shoot and ZmZIP3 is up-regulated in both root and shoot (Li et al., 2013). The expression level of ZmZIP4, ZmZIP5, ZmZIP7, and Zm ZIP8 decreased in shoots and ZmZIP3 is down-regulated in roots of maize under excess Zn supply (Li et al., 2013). Similarly, 12 ZmZIP genes (ZmZIP1–ZmZIP12) were identified by Mondal et al. (2013) in maize. Ten ZmZIP genes (ZmZIP1, ZmZIP2, and ZmZIP4–ZmZIP11) are tissue-specific and are highly expressed in flag leaf except ZmZIP3 and ZmZIP12, under Zn deficient condition (Mondal et al., 2013). The ZmZIP2, ZmZIP5, ZmZIP6, ZmZIP8, and ZmZIP11 are expressed in the kernel under Zn deficient condition (Mondal et al., 2013). ZmZIP4, ZmZIP5, ZmZIP7, and ZmZIP9 proteins are located in cytoplasm and chloroplast (Mondal et al., 2013; Figure 4). It is well known that some ZIP transporters are involved in Zn biofortification in other crops. For example, the OsZIP4 and OsZIP8 are actively involved in the grain filling of Zn in rice (Ishimaru et al., 2007; Lee et al., 2010b). The ZmZIP5 and ZmZIP11 are highly expressed in the flag leaf, which might play a vital role in the mobilization of Zn from flag leaf to developing kernel for the accumulation of the large amount of Zn in the kernel (Mondal et al., 2013; Figure 5B). These ZmZIPs (ZmZIP5 and ZmZIP11) might contribute to the biofortification of maize with Zn to improve the quality (Mondal et al., 2013). Since maize is highly sensitive to Zn deficiency, more high resolution studies are needed to improve the quality of maize grown under low Zn soils.

**Wheat**

Wheat is a major cereal crop for whole world and it provides about one-half of humans’ food calories and a large part of their nutrient requirements (Shiferaw et al., 2013). Complete assembly of the hexaploid bread wheat genome is available now (Zimin et al., 2017). Previously, very little was known about ZIP family transporters except for TdZIP1 from wild emmer wheat (Triticum turgidum ssp. dicoccoides), a Zn transporter with higher expression under Zn deficiency (Durma z et al., 2011). Recently, 14 TaZIP genes were identified in bread wheat (Triticum aestivum) (Evens et al., 2017). Out of 14 ZIPs, five (TaZIP3, TaZIP5, TaZIP6, TaZIP7, and TaZIP13) were analyzed for the expression level in shoot and root under Zn starvation. In shoot, expression of all five TaZIPs increased under low Zn conditions but the timing varied between individual genes. In roots, TaZIP3, TaZIP5, TaZIP7, and TaZIP13 showed increased expression under Zn starvation with expression of only TaZIP6 gene remains fairly stable (Evens et al., 2017). Similarly, Deshpande et al. (2018) analyzed the relative expression of five selective ZIP genes in durum wheat (Triticum durum) genotypes UC 1114 and MACKS 3125 under three different foliar application of Zn (Deshpande et al., 2018). In flag leaves, expression level of three TaZIP genes (TaZIP1, TaZIP3, and TaZIP7) decreased and two genes

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**Figure 5** | Expression of ZIP transporter genes in different parts of rice (A), maize (B), and wheat (C). The ZIP transporters are expressed in root, nodes, clumps, shoot/stem, leaf, flag leaf, kernel, and spikelets under Zn deficient levels. Under low Zn condition, root tissues show the expression of ZIP genes in rice (Farnesh et al., 2003; Ishimaru et al., 2005; Chen et al., 2008; Yang et al., 2009; Suzuki et al., 2012), maize (Li et al., 2013; Mondal et al., 2013) and wheat (Durma z et al., 2011; Evens et al., 2017; Deshpande et al., 2018), clumps in rice (Yang et al., 2009), shoot/stem in rice (Farnesh et al., 2003; Ishimaru et al., 2005; Chen et al., 2008; Yang et al., 2009; Suzuki et al., 2012), maize (Li et al., 2013; Mondal et al., 2013), and wheat (Durma z et al., 2011; Evens et al., 2017). Leaf tissue also showed the over expression of ZIP genes under low Zn soil in rice (Farnesh et al., 2003; Ishimaru et al., 2005; Chen et al., 2008; Li et al., 2009) and wheat (Deshpande et al., 2018), kernel tissue in maize (Li et al., 2013) and spikelets tissue in rice (Yang et al., 2009) and wheat (Deshpande et al., 2018).
AMF-mediated Zn transport and signals and TFs involved in this symbiotic Zn uptake.

**Foxtail Millet**

Foxtail millet is one of the food security minor cereal crops in low input regions. It is a domesticated diploid C4 crop having a small genome (∼515 Mb) with short life cycle (Doust et al., 2009). So, it is considered as a model crop for genetic and genomic studies of monocots (Lata et al., 2013; Ceasar et al., 2014). The whole-genome sequence is available for two different genotypes of foxtail millet (Benetzen et al., 2012; Zhang et al., 2012). But, the information about ZIP genes is limited. Alagarasan et al. (2017) identified seven SiZIP genes (SiZIP1–SiZIP7) in foxtail millet. The expression pattern of seven SiZIP genes was analyzed in root, leaf, stem and spike tissues of foxtail millet grown under drought stress without any additional nutrient supplementation. All seven SiZIP genes were induced in all four tissues (root, leaf, stem, and spike) with various levels of expression. The SiZIP2, SiZIP3, SiZIP4, and SiZIP5 showed relatively higher expression and low level of expression was observed with SiZIP6, in all tissues. SiZIP1 gene is moderately expressed in root, shoot and spike and very least expression was seen in leaf (Alagarasan et al., 2017). The highly induced SiZIP genes (SiZIP2, SiZIP3, SiZIP4, and SiZIP5) could be used for the bio-fortification process for the enrichment of Zn into the seeds of foxtail millet. The functional characterization of ZIP genes in foxtail millet could aid to improve the Zn uptake in foxtail millet and other minor millets.

**Orange**

Orange is an important fruit crop with 60 million metric tonnes of annual production worldwide and has high nutritional values, vitamins and other nutrients (Etebu and Nwauzoma, 2014). Till date, orange is the only fruit crop in which ZIP transporters are identified. In navel orange (Citrus sinensis), four ZIP genes such as CsZIP1, CsZIP2, CsZIP3, and CsZIP4 were identified and expression analysis show that CsZIP3 and CsZIP4 are highly expressed in mild, moderate and severely affected Zn deficient leaf. CsZIP1 gene is down-regulated under mild Zn deficient leaf and up-regulated in moderate and severely Zn depleted leaf. There is no change in the expression of CsZIP2 in all tissues when compared with control (Fei et al., 2016). Recently, 13 ZIP (PtZIP1–PtZIP3 and PtZIP5–PtZIP14) genes were identified in trifoliate orange (Poncirus trifoliata), another variety of orange fruit crop. PtZIP1, PtZIP2, PtZIP3, PtZIP5, PtZIP6, and PtZIP9 are highly induced in roots, whereas PtZIP1, PtZIP2, PtZIP5, PtZIP6, and PtZIP7 are highly expressed in leaves, under Zn-deficient condition (Fu et al., 2017). In future, functional characterization of ZIP genes in orange is very important to improve the Zn content in the edible parts of orange. Therefore, it will help to reduce the Zn deficiency problems in humans. So, more studies are needed to improve the quality of orange in terms of biofortification.

**Common Bean**

Common bean is an important food crop for human as it contains a rich source of nutrients (Gepts et al., 2008), especially a source of dietary Zn (Astudillo-Reyes et al., 2015). Totally 19...
PvZIP genes (PvZIP1–PvZIP19) were identified in common bean. But, only seven PvZIP genes such as PvZIP2, PvZIP6, PvZIP7, PvZIP12, PvZIP13, PvZIP16, and PvZIP18 were analyzed for their expression dynamics. The selection of PvZIP genes is based on their location in the genome relative to the presence of QTLs for seed Zn content (Astudillo et al., 2013). PvZIP12, PvZIP13, and PvZIP16 genes are expressed in root, leaf and pod of the common bean under Zn deficient condition (Astudillo et al., 2013). Other PvZIP genes are not expressed in any tissue types in Zn deficient condition. The PvZIP12 is highly expressed in leaves at vegetative stage and PvZIP13 is highly expressed in leaves at flowering stage. Hence, the PvZIP12 gene could be involved in mobilizing the Zn to seeds of bean. The identification and characterization of the candidate genes related to PvZIP genes are not expressed in any tissue types in Zn deficient condition. The expression dynamics of the candidate genes related to PvZIP transporters in crops may help to improve Zn uptake and nutrient content of seeds.

**FUNCTIONAL CHARACTERIZATION OF ZIP TRANSPORTERS IN CROPS**

Till date, only a little information is available on functional characterization of ZIP transporters in crops (Milner et al., 2013). Mostly, the investigations on the functions of plant ZIP transporters have been conducted by yeast complementation assays which revealed that ZIP transporters are capable of transporting various divalent cations (Guerinot, 2000). For functional characterization, Zn sensitive yeast mutant (zrt1/zrt2) which is defective in both the ZRT1 high—affinity and the ZRT2 low—affinity uptake transporters and is susceptible to Zn deficient conditions, was employed (Zhao and Eide, 1996a,b). In most of these studies, genes of model plant *A. thaliana* are used. The *AtZIP* genes such as *AtZIP1*, *AtZIP2*, and *AtZIP3*

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**Table 2** | Details on the ZIP family transporter genes identified in plants.

| Name of the plant | Common name | Name of ZIP genes | Method of characterization | Affinity References | References |
|-------------------|-------------|-------------------|----------------------------|---------------------|------------|
| Arabidopsis thaliana | Arabidopsis | AtZIP1-AtZIP4 | Yeast complementation assay | AtZIP1 (Km = 13 µM) and AtZIP3 (Km = 14 µM) | Grotz et al., 1998 |
| Poncirus trifoliate | Trifoliate orange | PzZIP1-PzZIP3, PzZIP5-PzZIP7, PzZIP9 and PzZIP10-PzZIP14 | qRT-PCR and yeast complementation assay | – | Fu et al., 2017 |
| Setaria italica | Foxtail millet | SzZIP1-Sz ZIP7 | RT-PCR | – | Alagarasan et al., 2017 |
| Glycine max | Soybean | GmZIP1, GmZIP4, GmZIP4, GmZIP6, GmZIP10 and GmZIP11 | Yeast complementation assay | GmZIP1 (Km = 13.8 µM) | Moreau et al., 2002 |
| Oryza sativa | Rice | OsZIP1-OsZIP16 | qRT-PCR and yeast complementation assay | OsZIP1 (Km = 16.3 µM) and OsZIP3 (Km = 18.5 µM) | – |
| Phaseolus vulgaris | Common bean | PvZIP1-PvZIP18 | qRT-PCR | – | Astudillo et al., 2013 |
| Zea mays | Maize | ZmZIP1-ZmZIP12 | qRT-PCR and yeast complementation assay | ZmZIP3, ZmZIP4, ZmZIP5, ZmZIP7 and ZmZIP8 | – |
| Triticum aestivum | Wheat | TaZIP1-TaZIP3, TaZIP5-TaZIP7, TaZIP9-TaZIP11, TaZIP13, TaZIP14 and TaZIP16 | qRT-PCR and yeast complementation assay | – | Evens et al., 2017 |
| Citrus sinensis | Navel orange | CsZIP1-CsZIP4 | qRT-PCR | – | Fei et al., 2016 |
| Hordeum vulgare | Barley | HvZIP1-HvZIP3, HvZIP5-HvZIP8, HvZIP10, HvZIP11, HvZIP13, HvZIP14 and HvZIP16 | qRT-PCR and yeast complementation assay | HvZIP3 and HvZIP8 | Fei et al., 2016; Pedas et al., 2009; Tiong et al., 2015 |
| Triticum durum | Durum wheat | TdZIP1, TdZIP3, TdZIP7 TdZIP10 and TdZIP15 | qRT-PCR | – | Deshpande et al., 2018 |
| Triticum dicoccoides | Emmer wheat | TdZIP1 | qRT-PCR and yeast complementation assay | – | Durmaz et al., 2011 |

*The details on name of the plant, name of ZIP genes, affinity of ZIP transporter and mode of characterization are provided.*
enabled Zn transport based on the yeast complementation assay (Eide, 1998; Grotz et al., 1998; Guerinot, 2000). In another study, six ZIP genes such as AtZIP1, AtZIP2, AtZIP3, AtZIP7, AtZIP11, and AtZIP12, are able to complement the zrt1/zrt2Δ yeast mutant fully or partially under Zn deficient conditions (Milner et al., 2013). Recently, Fu et al. (2017) characterized the Zn transport specificities of the PtZIP genes and PtZIP1, PtZIP2, PtZIP3, and PtZIP12 are also able to complement zrt1zrt2 mutant (Fu et al., 2017).

In plants, the high-affinity ZIP transporter system was highly active under low Zn conditions. Based on a yeast complementation assay, the affinity values of AtZIP1 ($K_m = 13 \, \mu M$) and AtZIP3 ($K_m = 14 \, \mu M$) confirmed to be low-affinity transporters and AtZIP2 ($K_m = 2 \, \mu M$) as high-affinity transporter (Grotz et al., 1998). In rice, OsZIP1 ($K_m = 16.3 \, \mu M$) and OsZIP3 ($K_m = 18.5 \, \mu M$) are characterized as low-affinity Zn transporters (Kam et al., 2003). The GmZIP1 of soybean was identified to be a low-affinity transporter ($K_m = 13.8 \, \mu M$) (Moreau et al., 2002). The high affinity transporters ZmZIP3, ZmZIP4, ZmZIP5, ZmZIP7, and ZmZIP8 were identified in maize (Li et al., 2013; Mondal et al., 2013). Similarly, Pedas et al. (2009) found that HvZIP3 and HvZIP8 are low-affinity transporters and HvZIP5 is a high-affinity Zn transporter in barely based on yeast complementation assay. However, additional kinetic assay is essential for confirming this hypothesis. Till now, very little information is available on the affinities of ZIP transporters in crops (Table 2). Further, heterologous expression of plant ZIP transporters in yeast may not replicate the same function and may yield false results due to being a completely different system and lack of key signals involved in the regulation of Zn concentration. Testing the function of plant ZIPs in the same plant by knock-out studies may yield confirmative results. To this end, newly adopted genome editing tool CRISPR/Cas9 may help for the efficient generation of double and multiple mutants for plant ZIPs to test their function in the same system. Characterization of other ZIP transporters could help to understand the crucial roles of ZIP transporter family genes in crops.

### MANIPULATION OF EXPRESSION LEVELS OF ZIP GENES THROUGH TRANSGENIC MODIFICATION

The plant ZIP genes were over expressed through transgenic modification in some studies. Till now, only a few genes such as AtZIP1, OsZIP1, ZmZIP3, OsZIP4, OsZIP5, and HvZIP7 were used for over-expression analysis in crops (Table 3). For example, OsZIP1 gene was transferred into finger millet using pGreen0179 vector under the control of Bx17 promoter through Agrobacterium-mediated transformation; the transgenic plants showed significantly improved accumulation of Zn in seeds (Ramegowda et al., 2013). Similarly, over-expression of AtZIP1 gene in cassava showed an increase of 25% Zn content in the edible part of the plant (Gaitán-Solís et al., 2015). Over expression of OsZIP4, OsZIP5, and OsZIP8 in transgenic rice decreased the Zn content in shoot and seed; but significantly increased the root Zn content (Ishimaru et al., 2007; Lee et al., 2010a,b). It clearly revealed that these OsZIP transporters play a crucial role in Zn uptake from the soil. Over-expression of OsZIP4 in rice plants showed the reduction of plant growth under Zn deficient conditions (Ishimaru et al., 2007). Approximately, 50% reduction observed in plant height and root length in OsZIP4 over-expressing transgenic rice plants (Ishimaru et al., 2007). At the flowering stage, OsZIP5 and OsZIP8 over-expressing transgenic rice plants are shorter and had fewer tillers (Lee et al., 2010a,b). Also, these transgenic plants produce very fewer grains (Lee et al., 2010a,b). Similarly, over-expression of OsIRT1 in rice altered the plant architecture and increased the Fe and Zn contents in mature seeds and vegetative parts of the plant under Zn deficient condition (Lee and An, 2009). These studies indicate that the ZIP transporters are responsible for improving

### TABLE 3 | Details on genetic manipulation of ZIP transporter genes reported in various plants.

| Name of the ZIP gene | Source | Host | Name of vector used | Mode of transformation | Observation | References |
|----------------------|--------|------|---------------------|------------------------|-------------|------------|
| ZmZIP3 Zea mays      | Arabidopsis thaliana | pBI121 | Agrobacterium tumefaciens | Improved Zn accumulation in the roots | Li et al., 2015 |
| AtZIP1 Arabidopsis thaliana | Manihot esculenta | pCAMBIA2301 | Agrobacterium tumefaciens | Higher Zn concentrations in the edible portion | Gaitán-Solís et al., 2015 |
| HvZIP7 Hordeum vulgare | Hordeum vulgare | pMDC32 | Agrobacterium tumefaciens | Increases root to shoot translocation of Zn | Tiong et al., 2014 |
| OsZIP1 Oryza sativa | Eleusine coracana and Nicotiana tabacum | pGreen0179 | Agrobacterium tumefaciens | Enhanced Zn concentration in seed and plant tissue | Ramegowda et al., 2013 |
| OsZIP5 Oryza sativa | Oryza sativa | pCAMBIA1302 | Agrobacterium tumefaciens | Decreased Zn concentration in shoot and increased in root | Lee et al., 2010a |
| OsZIP8 Oryza sativa | Oryza sativa | pCAMBIA1302 | Agrobacterium tumefaciens | Decreased Zn concentration in shoot and seed; increased in root Zn distribution | Lee et al., 2010b |
| OsZIP4 Oryza sativa | Oryza sativa | pG121Hm | Agrobacterium tumefaciens | Decreased Zn concentration in shoot and seed; increased in root Zn distribution | Ishimaru et al., 2007 |
| AtZIP1 Arabidopsis thaliana | Hordeum vulgare | pWVec8 | Agrobacterium tumefaciens | Increased Zn content in seed | Ramesh et al., 2004 |

The name of the plant, vectors and the role of ZIP genes are mentioned.
the nutritional quality in crops. So, more effort is needed to identify the specific function of all ZIP genes in crops and it could help to develop transgenic plants related to biofortification of edible parts with Zn/Fe in crops.

CONCLUSION AND FUTURE PROSPECTS

Zinc is not only essential for plant growth but also crucial for human health. It is estimated that nearly 50% of the world’s population is at the risk of Zn deficiency problems (Pedas and Husted, 2009). In humans, Zn deficiency impairs growth and development, affects the nervous system, reduces immunity and can cause death (Ekweagwu et al., 2008; Menguer et al., 2018). Researchers need to pay more attention to the biofortification of food crops with priority to Zn. The ZIP transporters play a crucial role in biofortification with Zn. In soil, Zn deficiency problem is usually tackled by adding Zn containing fertilizers. But, it is only a temporary solution. Also, the subsistence farmers cannot afford to buy Zn fertilizers all the time due to high market price. Identification of candidate genes related to ZIP transporters, their characterization and tapping this information to develop crops via transgenic and/or the marker-assisted selection may help for developing Zn efficient crops. Genetic modification of crops with ZIP genes is also helpful in improving the crops related to Zn use efficiency. Therefore, the structural and functional insight of plant ZIP transporter are essential. The bioinformatics study showed that the plant ZIP transporters may have partially overlapping but distinct metal transport mechanism compared to BbZIP as some of the residues involved in metal binding and transport are not conserved in plant ZIPS. The plant ZIP transporters showed greater variation at the Zn$^{2+}$ binding site when compared to BbZIP. The phylogenetic analysis of 113 plant ZIP proteins from 14 plant species revealed that the ZIP family members are mainly clustered together as per their numbers. Monocot and dicot plant ZIPS are clustered in separate clusters and both low-affinity and high-affinity ZIPS are closely clustered. These findings would be a valuable theoretical knowledge for future studies in terms of understanding the gene, protein, functional residues of Zn transporters in crops and might be helpful to overcome the problems associated with Zn deficiency. Many ZIP family genes involved in Zn transport have been characterized in model plants. Identification and functional characterization of certain ZIP genes, their relative expression and localization have been done only in a few crops such as rice and maize. More information on Zn transporters are available for rice plants when compared to other crops so this can serve as a model to study Zn transport in other cereal crops. ZIP family members involved in Zn transport and sequestration represent some of the clearest candidate genes for increased Zn content in crops. But very little information is available on how Zn is transported from leaf xylem to phloem of developing seeds and ultimately unloaded into seeds with the help of ZIP genes. This needs further research in the coming years. Identification and characterization of ZIP proteins and related TFs in many crops would help for the better understanding of Zn homeostasis. A more holistic and high-resolution studies on ZIP transporters in crops will help to overcome the problems associated with low Zn soils and to improve the human health.

DATA AVAILABILITY STATEMENT

All datasets generated for this study are included in the article/Supplementary Material.

AUTHOR CONTRIBUTIONS

TA, TM, and SA conceptualized and wrote the manuscript. TA and SA analyzed the protein sequences with alignment, modeling and phylogeny. GV and SI assisted, and TA, TM, GV, and SA edited, and updated the manuscript. SA and SI contributed critically in revising and improving the manuscript for publication.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2020.00662/full#supplementary-material

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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