Enhanced Gene Expression Rather than Natural Polymorphism in Coding Sequence of the OsbZIP23 Determines Drought Tolerance and Yield Improvement in Rice Genotypes

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Abstract

Drought is one of the major limiting factors for productivity of crops including rice (Oryza sativa L.). Understanding the role of allelic variations of key regulatory genes involved in stress-tolerance is essential for developing an effective strategy to combat drought. The bZIP transcription factors play a crucial role in abiotic-stress adaptation in plants via abscisic acid (ABA) signaling pathway. The present study aimed to search for allelic polymorphism in the OsbZIP23 gene across selected drought-tolerant and drought-sensitive rice genotypes, and to characterize the new allele through overexpression (OE) and gene-silencing (RNAi). Analyses of the coding DNA sequence (CDS) of the cloned OsbZIP23 gene revealed single nucleotide polymorphism at four places and a 15-nucleotide deletion at one place. The single-copy OsbZIP23 gene is expressed at relatively higher level in leaf tissues of drought-tolerant genotypes, and its abundance is more in reproductive stage. Cloning and sequence analyses of the OsbZIP23-promoter from drought-tolerant O. rufipogon and drought-sensitive IR20 cultivar showed variation in the number of stress-responsive cis-elements and a 35-nucleotide deletion at 5'-UTR in IR20. Analysis of the GFP reporter gene function revealed that the promoter activity of O. rufipogon and drought-sensitive IR20 cultivar showed variation in the number of stress-responsive cis-elements and a 35-nucleotide deletion at 5'-UTR in IR20. Analysis of the GFP reporter gene function revealed that the promoter activity of O. rufipogon is comparatively higher than that of IR20. The overexpression of any of the two polymorphic forms (1083 bp and 1068 bp CDS) of OsbZIP23 improved drought tolerance and yield-related traits significantly by retaining higher content of cellular water, soluble sugar and proline; and exhibited decrease in membrane lipid peroxidation in comparison to RNAi lines and non-transgenic plants. The OE lines showed higher expression of target genes-OsRab16B, OsRab21 and OsLEA3-1 and increased ABA sensitivity; indicating that OsbZIP23 is a positive transcriptional-regulator of the ABA-signaling pathway. Taken together, the present study concludes that the enhanced gene expression rather than natural
polymorphism in coding sequence of OsbZIP23 is accountable for improved drought tolerance and yield performance in rice genotypes.

Introduction

Plants are continuously exposed to wide ranges of environmental stresses such as drought, salinity, heat, low temperature and nutrient deficiency. The growth and developmental constraints due to such environmental stresses result in sub-optimal performance and reduced crop production. Drought is one such serious environmental stress which occurs when crop plants are subjected to insufficient soil moisture to meet their demands. Rice (Oryza sativa L.), which feeds more than half of the global population is adversely affected by soil moisture stress resulting in estimated 50% production loss worldwide [1]. Drought severely affects rice plants at morphological, physiological and molecular level. It is of vital importance to understand how plants respond to drought stress and what molecular mechanisms are being used to tackle this abiotic stress. Subsequently, the knowledge on the genetic factors or genes would help generating stress tolerant crop plants with better performance and improved yield. Allele mining is a promising way to identify naturally existing genetic variants of candidate genes, and thereby enabling direct access to key alleles conferring resistance to environmental stresses and to use such alleles for crop improvement programme.

The phytohormone abscisic acid (ABA), which is synthesized significantly under water deficit condition, plays a regulatory role in diverse physiological processes in plants coordinating a complex regulatory network that enables plant to cope with the decreased water availability [2–3]. The threshold level of ABA in planta is the central switch of drought tolerance, and controls various growth and developmental aspects such as cell division, seed germination, seedling growth, stomatal movement, synthesis of storage proteins and lipids and seed maturation. The ABA-dependent stress responses are regulated by the expression of stress-responsive genes [4–6]. The hierarchically upstream genetic regulators in this ABA stress-responsive pathway, mainly the transcription factors (TF) directly controls the expression of multiple other stress response related genes by binding to the cis-regulatory elements in their promoter region. The products from the later sets of genes are actually involved in biochemical and physiological functions to protect the plants against stresses. Thus, the TF genes are master regulators of stress responses. The basic leucine zipper (bZIP) TF family is one of the most diverse families in higher plants, whose members have different roles, particularly in stress responses and hormone signaling [7]. The members of the bZIP family contain the characteristic bZIP domain which is highly conserved and composed of a basic and a leucine zipper region [8]. They have been identified for their functions relating to the activation of hierarchically downstream gene expression by binding to the ACGT-motifs of ABA responsive elements (ABREs) in the promoter regions of these downstream genes [9]. Thus, these bZIP TFs were termed as ABRE-binding factors (ABFs) or ABRE-binding proteins (AREBs) [10].

In Arabidopsis, 75 identified bZIP TFs are subdivided into 10 groups. Group A bZIPS of Arabidopsis contains 13 members, many of them have been well studied and found to play central role in ABA-dependent stress signaling [11]. The group A subfamily III bZIPS in Arabidopsis were given special attentions (also known as ABF/AREB/ABI5) due to their biotechnological potentials. Among the seven members of these bZIP ABF genes, AREB1/ABF2, AREB2/ABF4 and ABF3 have been found to be expressed under abiotic stress condition, mainly in vegetative tissues [12–13]; and plants overexpressing these transcription factors
showed enhanced drought tolerance [14–15]. The triple mutant of areb1 areb2 abf3 showed highest level of decreased ABA sensitivity and reduced drought tolerance compared to single or double mutant, indicating that these TFs cooperatively regulate ABA stress signaling and require ABA for their complete activation [16]. The triple mutant of areb1 areb2 abf3 displayed remarkably impaired stress responsive gene expression leading to the identification of novel downstream factors, including late embryogenesis abundant (LEA) genes, group A type 2C phosphatase (PP2C) and numerous transcription factors [17]. In conclusion, the AREB/ABF genes are the major regulators of ABA-dependent gene expression in Arabidopsis.

Rice genome contains 89 putative bZIP TFs clustered into 11 groups based on their amino acid sequence similarity and DNA-binding specificity [18]. Several members have been studied for their functions related to stress responses including ABRE-binding factor OSBZ8, whose expression is strongly induced by ABA and positively correlated with salinity tolerance in rice [19]. The expression of LIP19, another member of rice bZIP TFs have been observed to be induced by low temperature, and it acts as a molecular switch in low temperature signaling pathway in rice [20]. Further, RF2a and RF2b bZIP TFs in rice have been found to be associated with biotic stress tolerance [21–22]. In comparison to comprehensive research on group A subfamily III bZIPs of Arabidopsis, relatively few orthologs in rice have been studied functionally for their roles in ABA-mediated stress responses, which include TRAB1, OsABI5, OsbZIP23 and OsbZIP46. The TRAB1 has been found to be activated by ABA-dependent phosphorylation and it interacts with VP1 transcription factor [23]. The OsABI5 has been suggested to be involved in ABA signaling, regulating fertility and stress responses [24]. The overexpression of OsbZIP23 has conferred increased sensitivity to ABA and enhanced stress tolerance in rice [25]. Overexpression of intact OsbZIP46 has exhibited no effect on drought tolerance; however a constitutively active form of OsbZIP46 improved drought tolerance in rice upon overexpression [26]. Among the above-mentioned four members of group A subfamily III bZIPs of rice, the OsbZIP23 is certainly the key player, which functions as a transcriptional activator and positively regulate the ABA-dependent stress responsive pathway, and hence increases the stress tolerance in rice [27].

Therefore, our present study aimed to search for natural allelic variations of the OsbZIP23 gene across the selected drought-tolerant and drought-sensitive rice genotypes and few other members of the Poaceae family. After identifying the natural polymorphisms in the coding DNA sequence (CDS) and promoter regions of the OsbZIP23 gene between drought-tolerant and drought-sensitive rice genotypes, we have functionally characterized the two promoter segments and two CDSs. The promoter activity has been investigated through GFP reporter gene expression analysis to find out the effect of natural polymorphism on OsbZIP23 gene expression. On the other hand, the two allelic variants of OsbZIP23 CDS have been studied through transgenic overexpression and RNAi-mediated endogenous gene silencing. The present study reveals that the OsbZIP23-mediated drought tolerance and yield improvement in rice genotypes depends on the enhanced gene expression due to increased promoter activity rather than the natural polymorphism in coding sequence.

**Materials and Methods**

**Plant materials and growth conditions**

Nine cultivated indica rice (Oryza sativa L.) genotypes, which include drought tolerant- Vandana, Manipuri, Nagina22 and drought sensitive- IR20, IR36, IR64. IR72, Swarna and HRC300 were chosen. Two wild progenitors of rice viz., Oryza rufipogon, Oryza nivara and four other representative members of the Poaceae family viz., Zea mays (maize), Sorghum bicolor (sorghum), Pennisetum glaucum (bajra) and a grass genotype Brachypodium was also included. All
the selected genotypes were used for allelic polymorphism analysis. Rice genotypes were grown in the glass house at 25/28°C with a 16/8 h photoperiod and 70% relative humidity for all experimental works. For all phenotypic assays, seeds from the same harvest and same storage conditions were used.

**Drought stress treatment**

To detect transcript level of OsbZIP23 gene at vegetative and reproductive (grain filling) stages, leaf samples were collected from rice genotypes growing under three different conditions: (i) under normal watered condition that is referred as before stress (BS), (ii) under water stress (8 days) condition, which is referred as after stress (AS) and (iii) the rewatering (3 days) condition that is referred as after recovery (AR). Drought stress treatment in T1 transgenic lines and non-transgenic (NT) plants were performed under vegetative and early reproductive (panicle initiation) stages following the method reported earlier [28] with certain modification. For this, seeds of transgenic lines were germinated on MS agar medium containing 50mg/L hygromycin. The germinated transgenic and NT (germinated on MS agar only) seedlings were transferred into liquid ½ MS medium under 16h light/8h dark cycle at 25°C for one week. After one week, plants were transplanted into pots and grown for 20 days under standard growth conditions. Upon attaining vegetative and panicle initiation stages (about 35 days and 60 days of seeding, respectively) plants were subjected to drought stress by withholding water supply for a period of 8 days until a lethal effect of dehydration was observed on RNAi lines and NT plants. In each cases, rewatering was done for 3 days after the stress period and survival rate (%) of the transgenic lines and NT plants were determined as the number of visible green plants. For analysis of grain yield under drought stress in transgenic lines and NT plants, drought treatment at flowering stage was imposed in the PVC pipes as described previously [29]. Plants from each line were planted individually in the PVC pipe, and allowed to grow normally until they reach the flowering stage. Drought stress was initiated at flowering stage and after 10 days of water deprivation (when serious drought stress was visualized on RNAi lines and NT plants), plants were irrigated to allow recovery at the flowering and seed maturation stage. After the seed maturation, grain yield was calculated in transgenic lines and NT plants.

**Isolation of total cellular RNA and synthesis of 1st strand cDNA**

Total RNA from leaf tissues of rice plants was isolated using RNeasy Mini Kit (Qiagen) following the manufacturer’s instruction. First strand cDNA was synthesized from 2 μg of total RNA with gene-specific reverse primer using Transcriptor 1st strand cDNA synthesis kit version 6.0 (Roche molecular biochemical) following manufacturer’s protocol.

**Isolation, cloning and sequence analysis of OsbZIP23 coding DNA sequence**

Isolation of the coding DNA sequence (CDS) of OsbZIP23 gene was carried out by RT-PCR from leaf RNAs of drought-tolerant and drought-sensitive rice plants and non-rice (Poaceae family) plants mentioned in plant material section. The PCR was performed using 2 μl of the 1st strand cDNA with gene-specific primer pairs bZ23F-bZ23R (S1 Table) by the following thermal profile: initial denaturation at 98°C for 2 min, followed by 30 cycles of 98°C/15 s, 60°C/15 s, 72°C/1 min and a final extension at 72°C for 8 min in a thermocycler (Applied Biosystems). The PCR products (~1083 bp size) were separately cloned in pUC18 plasmid. Nucleotide polymorphism of all the CDSs was analyzed after sequencing of positive clones. All the CDS-derived polypeptides were used for multiple amino acid sequence alignment through Jalview 2 [30].
Real-time PCR analysis
Real-time PCR (qRT-PCR) for specific transcript was performed in Eppendorf Realplex² Master Cycler using SYBR green based relative quantification method using 5 prime kits (Eppendorf). Three replicates were routinely used for each sample. Relative gene expression levels were determined as reported earlier [31]. Rice polyubiquitin1 gene (OsUbi1) was used as the internal control for normalization in each case. The primers used for real-time PCR are listed in S1 Table.

Southern hybridization
The genomic DNA was isolated from leaf tissues of rice plants following the standard protocol [32]. For each sample, 15 µg of genomic DNA was digested with HindIII, run on 1% agarose gel overnight and transferred onto nylon membrane (Hybond-N⁺). A 739 bp DNA of Osb-ZIP23 CDS (from O. rufipogon) was radiolabelled with P³²-dCTP (3500 Ci/mmol) by random priming using rediprime II DNA labeling system (GE Healthcare, USA) following manufacturer’s instructions. Southern hybridization was performed [33] at 65°C in Church buffer (0.5 M phosphate buffer (pH-7.2), 1mM EDTA, 1% BSA and 7% SDS) using OsbZIP23 gene probe. Similar procedure was used for Southern hybridization of transgenic rice lines developed in this study. The developed multi-sensitive X-ray film (Perkin Elmer) was scanned with a Cylone® Plus phosphor system (Perkin Elmer).

Isolation, cloning and in silico analysis of predicted OsbZIP23 promoter
The predicted promoter sequence of ~1600 bp was PCR amplified from the genomic DNA isolated from leaves of O. rufipogon (drought-tolerant) and IR20 (drought-sensitive) rice genotypes using gene-specific primers bZ23PF-bZ23PR (S1 Table) by the following thermal profile: initial denaturation at 98°C for 2 min, followed by 30 cycles of 98°C/10 s, 55°C/15 s, 72°C/2 min and a final extension at 72°C for 8 min in a thermocycler (Applied Biosystems). The PCR products were then separately cloned in pUC18 plasmid, and positive clones were sequenced. The sequences of the cloned DNA fragments were analyzed for cis-acting elements and promoter core elements using PLACE [34] and Plant Promoter Database 3.0 [35] web tools.

Generation of transgenic rice plants with the OsbZIP23 promoter-GFP reporter gene construct and assay of GFP fluorescence by confocal microscopy
The OsbZIP23 promoter-GFP reporter gene construct was prepared by fusing the putative OsbZIP23 promoter DNA fragment of O. rufipogon (1612 bp) and IR20 (1578 bp) in HindIII and BamHI restriction sites with GFP reporter gene followed by NOS transcription terminator in pCAMBIA1300 plasmid. The individual construct was introduced into drought-tolerant Vanda rice genotype through Agrobacterium-mediated transformation [36] and transgenic lines were confirmed by Southern hybridization using GFP reporter gene as hybridization probe. GFP fluorescence in leaves of transgenic rice lines was visualized with finely cross-sectioned leaf samples under the confocal microscope (Olympus FV1200). GFP was excited at 490 nm wavelength by 30 mW multi-Argon laser and the emission was at 520 nm. Images were taken at 40X magnification. The relative expression of GFP gene was monitored by qRT-PCR using specific primer set. The primers used are listed in S1 Table.
Generation of transgenic rice plants with the OsbZIP23 gene overexpression and gene silencing construct

For construction of the OsbZIP23 overexpression plasmid, the full length CDS of allelic forms from *O. rufipogon* and *O. nivara* were put separately (in BamHI and KpnI restriction sites) under the rice polyubiquitin 1 (*OsUbi1*) promoter (HindIII and BamHI restriction sites) and NOS transcription terminator (KpnI and EcoRI restriction sites) in pCAMBIA1300 plasmid. Thus, two OsbZIP23 overexpression (OE) constructs - OER for 1068 bp CDS of *O. rufipogon* and OEN for 1083 bp CDS of *O. nivara* were prepared. An RNAi-mediated gene silencing construct of OsbZIP23 gene was also prepared. For this, a DNA fragment of 399 bp from 3’ part of OsbZIP23 CDS in sense (SalI and BamHI restriction sites) and antisense (BamHI and KpnI restriction sites) orientation separated by a linker DNA (BamHI restriction site on both sides) was put under the OsUbi1 promoter (HindIII and SalI restriction site) and NOS transcription terminator (KpnI and EcoRI restriction sites) in pCAMBIA1300 plasmid. The OEN, OER and RNAi constructs were introduced separately into drought-sensitive IR20 rice genotype through *Agrobacterium*-mediated transformation. The relative expression level of OsbZIP23 gene in OEN, OER and RNAi transgenic rice lines was monitored by qRT-PCR using gene-specific primers. The primers used are listed in S1 Table.

Immunoblotting for detection of OsbZIP23 protein in rice plants

The total protein was isolated from leaves of transgenic and NT rice plants and western blot analysis was carried out taking 40 μg of total protein following the previously reported method [37]. Affinity purified polyclonal antibody raised against OsbZIP23 protein in rabbit (catalog no- E580041-A-SE, Enogene Biotech, USA) was used as primary antibody (1: 1,000 dilution) and monoclonal plant anti-actin antibody (Sigma, catalog no.-A0480) was used as loading control (1:500 dilution). Immuno-detection was carried out with the Lumi-LightPLUS western blotting kit (Roche Molecular Biochemicals), following manufacturer’s instruction.

ABA sensitivity test during seed germination and seedling growth

For analysis of ABA sensitivity at seed germination stage, seeds of non-transgenic (NT) and transgenic rice plants were put on MS agar medium plates with different concentration of ABA (0, 1, 3 and 6 μM), and allowed to germinate for 10 days. After 10 days, seed germination rates were calculated as percentage (%). Similarly for seedling stage, seeds of NT and transgenic plants were germinated on ½ MS liquid medium with different concentration of ABA (0, 1, 3 and 6 μM). The seedlings were grown for 14 days in a growth chamber (28 ± 2°C, 16/8 h photoperiod and 70% relative humidity) and shoot height and root length of seedlings were recorded.

Measurement of water loss rate and relative water content in rice plants

Water loss rate (WLR) and relative water contents (RWC) were measured using leaves of transgenic and NT rice plants following reported method [38–39]. Two sets of experiment were performed for transgenic and NT plants and the results are consistent. The result from one set of experiment is presented. For WLR and RWC, each data point represents the average of three replicates.

Measurement of proline and soluble sugar contents in leaf tissues of rice plants

Free proline content was determined using 50 mg of rice leaf tissue of transgenic and NT plants following the previously reported method [40]. Total soluble sugars in NT plants and
transgenic lines were determined from 100 mg of rice tissues using anthrone reagent [41]. Two sets of experiment were performed for transgenic and NT plants and the results are consistent. The result from one set of experiment is presented. For proline and soluble sugars, each data point represents the average of three replicates.

Measurement of malondialdehyde (MDA) contents and *in vivo* localization of reactive oxygen species (ROS) in leaf tissues of rice plants

The MDA content was determined using 100 mg of leaf tissue following the previously described method [42]. The absorbance (A) of the samples was measured at 450, 532 and 600 nm. The amount of MDA was calculated according to the formula: MDA content = 6.45×[(A<sub>532</sub>−A<sub>600</sub>)/0.56×A<sub>450</sub>]. Two sets of experiments were performed for transgenic and NT plants and the results are consistent. The result from one set of experiment is presented. For MDA content, each data point represents the average of three replicates. *In vivo* localization of reactive oxygen species (ROS) was done using the intact leaf samples following the method as reported earlier [43].

Accession Numbers

Sequence data from this article can be accessed in the GenBank database with the following accession numbers: Rice polyubiquitin 1 (*OsUbi1*) gene (AF184279), *OsUbi1* promoter (AY785814), OsbZIP23 promoter from *O. rufipogon* (KP779639), OsbZIP23 promoter from IR20 (KP779640), OsbZIP23 CDS from *O. rufipogon* (KP779637), OsbZIP23 CDS from *O. nivara* (KP779638), OsRab16B (AF333275), OsRab21 (Y00842) and OsLEA3-1 (DQ789359).

Results

Analysis of allelic polymorphism in OsbZIP23 CDS across selected rice genotypes and four other members of the Poaceae family differing in response to drought stress

For detection of allelic polymorphism in OsbZIP23 gene, nine *indica* rice cultivars of drought-tolerant and drought-sensitive genotypes were selected along with two rice wild progenitors and four other representative members of the Poaceae family. The full length coding DNA sequence (CDS) of OsbZIP23 was amplified from leaf RNA samples of selected genotypes through RT-PCR using the gene-specific primers. The amplicons were found to have an expected size of ~1083 bp (Fig 1A). The amplicons from each genotype was cloned in pUC18 plasmid and positive clones were sequenced. Multiple sequence alignment of the CDS and the derived polypeptide of OsbZIP23 across the 11 rice genotypes and four other members of the Poaceae family revealed the presence of natural polymorphism (Fig 1B). Four non-synonymous single nucleotide polymorphisms (SNPs) and one 15-nucleotide deletion were detected. First SNP at position 606<sup>th</sup> (C to A) from the start codon led to 202<sup>nd</sup> amino acid alteration from Phenylalanine (F) to Leucine (L), and the 2<sup>nd</sup> SNP at position 789<sup>th</sup> (A to T) from the start codon led to 263<sup>rd</sup> amino acid alteration from Lysine (K) to Asparagine (N). Phenylalanine and Lysine were present at 202<sup>nd</sup> and 263<sup>rd</sup> amino acid positions, respectively in IR20, IR36, Swarna, Vandana, HRC300, bajra and *Brachypodium*; whereas Leucine and Asparagine were present at the corresponding positions in IR72, Nagina22, Manipuri, *O. rufipogon*, *O. nivara*, maize and sorghum. The 3<sup>rd</sup> SNP was observed in the OsbZIP23 CDS of bajra, where nucleotide change (A to T) at position 44<sup>th</sup> from the start codon altered the 15<sup>th</sup> amino acid from Glutamine (Q) to Leucine (L). The 4<sup>th</sup> SNP was observed in the OsbZIP23 CDS of *O.
Fig 1. Analysis of natural polymorphism in the coding DNA sequence of OsbZIP23 gene. (A) PCR amplification of OsbZIP23 CDS from selected genotypes using gene-specific primers. Lanes 1–15 represents O. rufipogon, O. nivara, Vandana, Swarna, Nagina22, Manipuri, HRC300, IR20, IR36, IR64, IR72, sorghum, maize, Brachypodium and bajra respectively. Lane M– Hinfi digested pUC18 plasmid as a standard molecular weight marker. (B) Allelic polymorphism in the CDS of OsbZIP23 gene across selected rice genotypes is represented by multiple sequence alignment in Jalview software.

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nivara and sorghum, where nucleotide change (C to T) at the position 257th from the start codon led to an altered amino acid at 86th position from Alanine (A) to Valine (V). Interestingly, a major 15-nucleotide deletion resulting in elimination of five amino acids from position 85th to 89th in the N-terminal region of OsbZIP23 was observed in wild progenitor O. rufipogon (Fig 1B). Although the 5-amino acid deletion sequence of O. rufipogon is AAEHA in almost all the genotypes tested, but this sequence is AVEHA in O. nivara and sorghum (Fig 1B).

Analyses of relative expression and copy number of OsbZIP23 gene in the selected rice genotypes

To investigate the OsbZIP23 gene expression patterns in different tissues of the rice plant, we performed qRT-PCR analyses using the total RNA isolated from various tissues of Vandana, a cultivated drought-tolerant indica rice genotype. The OsbZIP23 transcript was found to be present in all tested tissues, including root, shoot, stem, leaf, leaf sheath and panicle, with highest transcript expression levels in leaf and leaf sheath than in other tissues (Fig 2A). The allelic polymorphism in the coding region of OsbZIP23 gene impelled us to check the relative expression level of this gene by qRT-PCR in the selected rice genotypes, which include nine cultivated indica rice genotypes viz., Vandana, Nagina22, Manipuri, Swarna, HRC300, IR20, IR36, IR64, IR72 and two wild progenitors of rice O. rufipogon and O. nivara. For this analysis, rice genotypes under the vegetative and reproductive (grain filling) stages were selected. Total RNA was isolated from leaves of all the rice genotypes grown under three different conditions: (i) before stress (BS), (ii) after stress (AS) and (iii) after recovery (AR). The qRT-PCR analyses showed

![Fig 2. Analysis of the transcript expression level and copy number of the endogenous OsbZIP23 gene. (A) Relative expression level of OsbZIP23 gene in different tissues of an upland indica rice genotype, Vandana. Real time PCR analysis of OsbZIP23 transcript in vegetative (B) and reproductive (grain filling) stage (C), in 11 selected rice genotypes grown under before stress (BS), after stress (AS) and after recovery (AR) condition as described in method section. Rice polyubiquitin1 (OsUbi1) gene was taken as internal reference. Data bars represent the mean ±SD of triplicate measurement. Statistical analysis of Student’s t test indicated significant differences (* P<0.05, **P<0.01). (D) Southern blot showing the single copy of endogenous OsbZIP23 gene in 11 selected rice genotypes. Lanes 1–11 represent O. rufipogon, O. nivara, Vandana, IR20, Swarna, HRC300, IR36, IR64, IR72, Nagina22, Manipuri, HRC300 rice genotypes, respectively.](https://doi.org/10.1371/journal.pone.0150763.g002)
that the relative expression level of OsbZIP23 transcript increased in AS and AR samples of drought-tolerant genotypes *O. rufipogon*, *O. nivara*, Nagina22, Vandana and Manipuri in both vegetative (Fig 2B) and reproductive stages (Fig 2C). However, the relative expression levels of OsbZIP23 transcript in AS and AR samples of drought-sensitive genotypes IR20, IR36, IR64, IR72, HRC300 and Swarna were not much comparable to that of BS samples in both vegetative (Fig 2B) and reproductive stages (Fig 2C). It was also observed that the abundance of OsbZIP23 transcript in all the genotypes were comparatively higher in reproductive stage (Fig 2C) than in the vegetative stage (Fig 2B). The differences in the relative expression level of OsbZIP23 transcript in the selected rice genotypes grown under the same conditions, led us to check the copy number of OsbZIP23 gene in the genome of all the selected rice genotypes through Southern blot hybridization using OsbZIP23 gene-specific probe. Southern blot analysis revealed a single band signal with *Hind*III digested genomic DNA samples in all the selected rice genotypes (Fig 2D), indicating the presence of only one copy of this gene in all the genotypes tested.

**In silico analysis of the predicted promoter sequence of OsbZIP23 gene from drought tolerant and drought sensitive rice genotypes**

The wide variation in the relative expression level of OsbZIP23 transcript between the drought-tolerant wild progenitor *O. rufipogon* and drought-sensitive genotype IR20 prompted us to analyze the promoter region of OsbZIP23 gene from these two rice genotypes. For this, ~1.6 kb DNA fragment upstream of the start codon was PCR amplified from the two genomic DNA samples using the gene-specific primers (Fig 3A), cloned and sequenced. Alignment of the predicted OsbZIP23 promoter sequences from *O. rufipogon* and IR20 showed a major deletion of 35 nucleotide (17 nucleotide upstream of the start codon) in the 5'-UTR segment of IR20 genotype in comparison to *O. rufipogon* (Fig 3B). Searching for the regulatory motifs in the promoter sequence using the PLACE database predicted several putative stress response-related cis-elements, which vary in number within *O. rufipogon* and IR20 genotypes. ABA-responsive element (ABRE) core has 11 hits in *O. rufipogon* and 12 hits in IR20; dehydration responsive element (DRE) has 3 hits in both the genotypes. Among other cis-elements MYC-core is present 10 times in *O. rufipogon* and 11 times in IR20, whereas MYB-core occurs 9 times in *O. rufipogon* and 7 times in IR20. Interestingly, the 35-nucleotide difference causes two extra MYC-cores in *O. rufipogon* (Fig 3B). Apart from these stress responsive cis-elements, some light responsive elements (LRE) were also found to be present in both the promoter sequences. Analysis revealed that the predicted transcription start site (TSS) is present at 216 bp upstream from the ATG codon in case of *O. rufipogon*, compared to 181 bp upstream in case of IR20. The OsbZIP23 promoter was predicted to be TATA-less, but the core promoter elements like CCAAT-box and Y-patches were found to be present around TSS in both the sequences.

**The OsbZIP23 gene promoter from the drought tolerant genotype drives higher level of the reporter gene expression than that of drought sensitive one**

To examine the differences in OsbZIP23 promoter activity, if any, between *O. rufipogon* and IR20 rice genotypes, two GFP reporter gene constructs were prepared (S1 and S2 Figs) using the OsbZIP23 promoter fragment isolated from *O. rufipogon* and IR20, and introduced into drought-tolerant Vandana rice genotype through *Agrobacterium*-mediated transformation. The tissue culture regenerated plantlets were screened in hygromycin and subjected to Southern blot analysis, which revealed single integration transgenic lines (Fig 4A). The transformed lines developed with the OsbZIP23 promoter-GFP-NOS construct in case of *O. rufipogon* and IR20 were designated as RuP and 20P, respectively. The T₀ seeds from the respective lines were
Fig 3. Cloning and in silico analysis of OsbZIP23 promoter sequences. (A) PCR amplification of the OsbZIP23 gene promoter region from two selected rice genotypes using gene-specific primers. Lanes 1–2 represent O. rufipogon and IR20, respectively. Lane M- HinfI digested pUC18 plasmid as a standard molecular weight marker. (B) Alignment of nucleotide sequences and distribution of major cis-regulatory elements in the OsbZIP23 promoter of O. rufipogon and IR20.

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germinated in hygromycin containing media and Southern blot analysis was performed, which showed similar transgene integration pattern in T1 plants (data not shown) as it was observed in T0 plants (Fig 4A). The Southern positive T1 plants were used for further analyses. To check the activity of both the promoters, the GFP fluorescence was observed in cross-sectioned leaf samples collected under drought stress condition from the transgenic RuP and 20P lines using confocal microscopy. It was found that the GFP fluorescence in RuP#2 and RuP#3 lines were comparatively higher than the 20P#1 and 20P#2 lines, whereas the non-transgenic (NT) plants showed no GFP fluorescence (Fig 4B). Further to quantitate the relative expression level of GFP in all the four lines, qRT-PCR analysis was carried out taking 20P#2 as calibrator (the least GFP fluorescence was observed in 20P#2 line). The qRT-PCR analysis revealed that the relative expression level of GFP in RuP#2 and RuP#3 lines were significantly (P<0.01) greater than the 20P#1 line (Fig 4C). These results indicated that the OsbZIP23 promoter isolated from the drought-tolerant O. rufipogon genotype exhibits higher functional activity than the drought-sensitive IR20 genotype.

**Generation of OsbZIP23 overexpression and silencing transgenic rice lines**

To study further the loss-of-function and gain-of-function phenotypes of OsbZIP23 in planta, overexpression (OE) and RNAi-mediated silencing (RNAi) of the OsbZIP23 gene was carried out. For OE, two polymorphic forms of OsbZIP23 CDS from the wild progenitors- O. rufipogon
and *O. nivara* were selected to prepare two different OE constructs, designated as OER and OEN, respectively (S3 and S4 Figs). The polypeptide of *O. rufipogon* has a deletion of 5-amino acid residues between the transactivation domain I and domain II of OsbZIP23 (S6 Fig). Similarly for studying gene silencing, an RNAi construct of *OsbZIP23* was also prepared (S5 Fig).

The individual genetic construct (OER, OEN and RNAi) was then introduced into the drought-sensitive IR20 rice genotype by *Agrobacterium*-mediated transformation. The tissue culture regenerated OE and RNAi plantlets were screened through hygromycin selection and subjected to Southern blot hybridization using *OsbZIP23* gene-specific probe (Fig 5A and 5B).

Three single integration T1 OE lines (OER#5, OER#7 and OEN#9) and two single integration T1 RNAi lines (RNAi#1 and RNAi#4) along with the control non-transgenic (NT) plants were selected for subsequent studies to examine for the transgene expression and their performance under drought stress. The independent transgenic lines along with NT plants were examined for *OsbZIP23* transcript expression level by real-time PCR analysis (qRT-PCR). The qRT-PCR analysis showed that the relative expression of *OsbZIP23* in OE lines increased significantly (*P*<0.01), whereas the relative expression level of endogenous *OsbZIP23* in RNAi lines decreased significantly (*P*<0.01) in comparison to NT plants (Fig 5C). Further, western blot analysis revealed that the abundance of OsbZIP23 protein increased in OE lines and decreased in the RNAi lines compared to NT plants (Fig 5D), supporting the qRT-PCR results.
Overexpression but not the silencing of OsbZIP23 gene improves drought tolerance and grain yield in rice

To analyze the tolerance of transgenic lines against drought stress under vegetative and early reproductive (panicle initiation) stages, two sets of plant samples (vegetative and reproductive stage) were subjected to drought stress. After the stress period, plants were irrigated for 3 days and survival rate (%) of the OE, RNAi and NT plants was determined. Analysis revealed that the survival rate of OE lines was significantly (P < 0.01) higher than the RNAi lines and NT plants in case of samples at vegetative stage (Fig 6A and 6B) as well as at reproductive stage (Fig 6C and 6D). The decreased drought tolerance of RNAi lines and increased drought tolerance of OE lines in both the stages of plant growth and development suggested that the OsbZIP23 has a crucial role to play in rice plants for adaptation to drought stress. The differences in the period of drought tolerance of OE and RNAi lines prompted us to check the grain yield of these lines under drought stress condition. For this, OE, RNAi and NT plants growing under normal condition in PVC pipes were subjected to progressive drought stress upon attaining the flowering stage. After the stress period, plants were irrigated and allowed to recover at the flowering and seed maturation stage (Fig 6E). It was found that the panicle weight and spikelet fertility of OE...
lines were significantly (P < 0.01) higher than the RNAi lines and NT plants (Fig 6F, 6G and 6H). The panicle weights of OE lines were ~2.5-3g per panicle, whereas the RNAi lines and NT plants had panicle weighing ~1–1.5g. Similarly, the spikelet fertility was found to be within 75–90% in case of OE lines. On the contrary, the spikelet fertility of RNAi lines was found to be ~40–50%, which was significantly lower than the OE lines. These results indicated that the Osb-ZIP23 gene has a significant role to play in drought tolerance and grain yield in rice.

Transgenic lines overexpressing OsbZIP23 display improved water retention capacity by accumulating more proline and soluble sugars thereby decreasing the cellular oxidative damage cause by drought stress

The opposite drought tolerance phenotype of OE and RNAi lines further accounted for the analysis of two water retention parameters viz., water loss rate (WLR) and relative water content (RWC). During 0 to 24 hours of dehydration treatment, OE lines exhibited least water loss rate (WLR) and retained the highest relative water content (RWC), while RNAi lines had highest WLR and lowest RWC in comparison to those of NT plants (Fig 7A and 7B). Morphological differences were apparently visible, as the OE leaves largely retained their turgor and the RNAi and NT leaves withered severely. These results suggested that the OsbZIP23 plays a positive role in improving the ability of the plants to retain water under dehydrated conditions. To elucidate the physiological mechanism by which OsbZIP23 improves the water retaining ability of plants under drought stress, proline and soluble sugar contents were determined in the transgenic and NT plants. Under normal growth condition, no significant differences in the contents of proline and soluble sugars were detected amongst OE lines, RNAi lines and NT plants. However, under drought stress condition, the OE lines accumulated dramatically higher contents of proline and soluble sugars, but RNAi lines accumulated smaller amount of these molecules, compared with those in NT plants (Fig 7C and 7D). These findings indicated that the OsbZIP23 helps in water retention by increasing the accumulation of osmolytes, like proline and soluble sugars. Drought stress disrupt the cellular homeostasis of plants by increasing the activity of reactive oxygen species (ROS), which results in high cell toxicity, membrane lipid peroxidation and even cell death. Lipid peroxidation is used as an indicator of oxidative stress. Membrane lipid damage during drought stress was analyzed by measuring the malondialdehyde (MDA) content in OE lines, RNAi lines and NT plants. The results showed that under normal growth condition, there were no significant differences in MDA content amongst OE, RNAi and NT plants. However, under drought stress, lipid peroxidation levels increased in both transgenic lines and NT plants, but the content of MDA in OE lines were significantly (P < 0.01) lower than in the RNAi lines and NT plants (Fig 7E). The decreased level of MDA in the OE lines under drought stress implied that they might be subjected to least serious oxidative damage than the RNAi lines and NT plants. Therefore, it was of interest to detect the accumulation of reactive oxygen species (ROS) in OE, RNAi and NT plants. There were no apparent differences in the level of H₂O₂ and O₂⁻ ions among the leaves of OE, RNAi and NT plants under normal growth conditions after staining with DAB and NBT. However, under drought condition, leaves of RNAi and NT plants exhibited higher level of H₂O₂ and O₂⁻ accumulations than that of OE lines (Fig 7F and 7G), demonstrating that OsbZIP23 helps to minimize the stress-induced oxidative damage in situ. These results suggest that transgenic lines overexpressing OsbZIP23 alleviate the effect of drought stress by increasing the accumulation of proline and soluble sugars, which helps the plant to retain more water and avoid the oxidative damage caused by ROS in dehydrating cells.
The OsbZIP23 regulates expression of ABA-dependent and stress-responsive genes in transgenic plants

To gain deeper understanding of the function of OsbZIP23 under drought stress, we monitored the transcript levels of three drought-inducible representative genes viz., OsRab16B, OsRab21 and OsLEA3-1, working in downstream of OsbZIP23 in the ABA signaling pathway. The OsRab16B and OsRab21 genes encode dehydrin proteins, which act by decreasing cellular osmotic potential; whereas the OsLEA3-1 encodes late embryogenesis abundant (LEA) protein. The relative expression level of OsRab16B, OsRab21 and OsLEA3-1 in vegetative stage were significantly (P < 0.01) increased in OE lines than in the NT plants, as revealed through qRT-PCR analysis (Fig 8A, 8B and 8C). On the contrary, the relative expression levels of these three genes were significantly (P < 0.01) lower in RNAi lines compared to NT plants (Fig 8A, 8B and 8C).
Further, we analysed the relative expression level of these three genes along with OsbZIP23 in the reproductive stage. The result showed similar trend of increased expression in OE lines and decreased expression in RNAi lines in comparison to NT plants (S7 Fig). These findings suggested that the OsbZIP23 overexpression activates but down-regulation affects the transcription of hierarchically downstream genes involved in ABA signaling and stress responses.

ABA sensitivity differs in OsbZIP23 OE and RNAi lines at germination and post-germination stages

To check the ABA sensitivity of rice transgenic lines, seeds of two OE lines (OER#5 and OEN#9), two RNAi lines (RNAi#1 and RNAi#4) and NT plants were germinated on MS agar medium in different concentration of ABA (0, 1, 3 and 6 μM) for 10 days. The germination rate of OE lines had no significant difference with RNAi lines and NT plants at 0 and 1 μM ABA, but it was significantly (P < 0.01) lower in OE lines than that of the RNAi lines and NT plants at 3 and 6 μM ABA (Fig 9A, 9B and 9C), suggesting that the ABA sensitivity was increased in OsbZIP23 OE lines at germination stage. The ABA sensitivity of transgenic plants was investigated at the post-germination stage. It was observed that at 0 μM concentration of ABA, the shoots and roots of OE seedlings (grown for 2 weeks) showed no significant morphological differences with RNAi lines and NT plants. However, the length of shoots and roots of OE lines exhibited significant disparity in comparison to RNAi lines and NT plants at 1, 3 and 6 μM ABA (Fig 9D, 9E, 9F and 9G). These results indicated that the overexpression but not the down-regulation of OsbZIP23 had increased ABA sensitivity at the post-germination stage, similar to the situation at germination stage.

Discussion

Progress in plant breeding has made possible the accumulation of superior alleles from vast plant genetic resources for generation of improved and high yielding varieties of agricultural crops. However, these superior alleles were not utilized properly as these were left behind during selective evolution and domestication carried out by farmers and breeders. These untapped genetic variations existing in wild relatives and land races of crop plants could be explored for
Fig 9. Evaluating ABA sensitivity of OsbZIP23 OE and RNAi lines at germination and post-germination stages. Germination performance of seeds from (A) OsbZIP23 OE lines (OER#5, OEN#9), and (B) OsbZIP23 RNAi lines (RNAi#1, RNAi#4) in comparison to NT plants on MS agar medium containing 0, 1, 3 and 6 μM ABA at 10th day. (C) Calculation of the germination rates (%) of OsbZIP23 OE, RNAi and NT seeds. (D and E) Performance of OE, RNAi and NT seedlings in ½ MS liquid medium containing 0, 1, 3 and 6 μM of ABA. Measurement of (F) shoot length and (G) root length of OE, RNAi and NT seedlings grown on different concentrations of ABA after 14 days. Data bars represent the mean ±SD of triplicate measurement. Statistical analysis by Student's t-test indicated significant differences (*P<0.05, ** P<0.01). All the results were based on three independent experiments.

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development of superior cultivars [44]. Allele mining is a well-established approach to dissect naturally existing allelic variations of candidate genes controlling key agronomic traits, which have potential applications in crop improvement. Nucleotide polymorphism in the promoter region or in the coding DNA sequence can have dramatic effects on the expression of the candidate gene or the functional activity of the corresponding protein. The consequences of these effects account for the attainment of desirable or undesirable agronomic traits. Although a number of genes have been reported to be involved in the adaptation of various abiotic stresses in plants, the allelic diversity of these genes is poorly understood [45]. To cope with drought stress, it is of vital importance to understand the role of naturally occurring allelic variants of known genes working in stress responsive pathways leading to plant’s tolerance for successful growth and development [46].

We investigated the allelic polymorphism of OsbZIP23 gene that encodes a crucial transcription factor in drought stress adaptation pathway in nine selected indica rice genotypes of drought-tolerant and drought-sensitive cultivars along with two wild progenitors of rice and four other representative members of the Poaceae family. Our study revealed the presence of non-synonymous single nucleotide polymorphisms (SNPs) at four places in the coding DNA sequence (CDS) and a deletion of 15 nucleotides at one place in the CDS of this gene (Fig 1B). The first two non-synonymous SNPs result in a non-polar amino acid substitution (F to L) at the 202\textsuperscript{nd} position and a positively charged amino acid substitution (K to N) at the 263\textsuperscript{rd} position in some of the genotypes (Fig 1B). The 3\textsuperscript{rd} non-synonymous SNP was observed in OsbZIP23 CDS of bajra, where a polar amino acid (Q) at the 15\textsuperscript{th} position changes to the non-polar amino acid (L). The 4\textsuperscript{th} non-synonymous SNP was observed in the OsbZIP23 CDS of O. nivara and sorghum, where a non-polar amino acid substitution (A to V) occurs at the 86\textsuperscript{th} position of the protein. In wild rice progenitor O. rufipogon, which is a drought-tolerant genotype, a major 15-nucleotide deletion in the CDS of OsbZIP23 was observed; eliminating the amino acid sequence AAEHA (or AVEHA compared to O. nivara and sorghum) from the position 85\textsuperscript{th} to 89\textsuperscript{th} at the N-terminal region (Fig 1B). This sequence of five amino acids is normally located specifically between the transactivation domain I and domain II of the OsbZIP23 polypeptide (S6 Fig). Out of the deleted five amino acids (AAEHA), three were non-polar (A) residues, one negatively charged (E) residue and one positively charged (H) residue. The 5-amino acid deletion of O. rufipogon is not observed in any other rice genotypes, and this may be correlated to the fact that all are indica sub-types that are derived from O. nivara progenitor. Similar reports of non-synonymous SNPs and deletions leading to alteration of functions of some other genes have been reported previously [45, 47].

Presence of allelic polymorphism in the CDS of OsbZIP23 gene in the selected drought-tolerant and drought-sensitive rice genotypes as well as wild rice progenitors prompted us to further demonstrate the relative expression level of this gene in all the rice genotypes through qRT-PCR analyses. To know which plant part to select for analysis of these 11 rice genotypes, at first we investigated different tissues from one drought-tolerant genotype, Vandana. The expression of OsbZIP23 was found to be tissue specific, with the highest expression level in leaf followed by leaf sheath (Fig 2A). Hence leaf tissues were used for all subsequent analyses of OsbZIP23 gene expression. The relative expression level of OsbZIP23 in the selected rice genotypes was found to be significantly variable during before stress (BS), after stress (AS) and after recovery (AR) conditions in both vegetative and reproductive stages (Fig 2B and 2C). The relative expression levels of OsbZIP23 under vegetative and reproductive stages were found to be much higher in drought-tolerant genotypes, compared to the drought-sensitive ones. It was also observed that the relative expression level of OsbZIP23 in all the rice genotypes was consistently higher in reproductive stage as compared to vegetative stage (Fig 2B and 2C), which signifies the important role of OsbZIP23 in drought adaptation of plants at the reproductive stage.
The variation in the relative expression level of OsbZIP23 in the selected rice genotypes further guided us to verify the copy number of endogenous OsbZIP23 in the rice genotypes through Southern blot analysis, which revealed the single copy of endogenous OsbZIP23 gene in all the rice genotypes tested (Fig 2D). Presence of single copy of the endogenous OsbZIP23 gene in all the rice genotypes suggests that the copy number has no effect on variable relative expression level of OsbZIP23 in drought-tolerant and drought-sensitive rice genotypes.

The findings that all the drought-tolerant rice genotypes express the OsbZIP23 gene at a much higher level than drought-sensitive genotypes and there is no anticipated copy number effect, directed us to dig into the promoter regions of this gene in different rice genotypes. We selected one drought-tolerant (wild rice O. rufipogon) and one drought-sensitive (O. sativa cultivar IR20) rice genotypes to isolate, clone and characterize the promoter region through GFP reporter gene expression. The prediction based analysis of both the isolated promoter sequences revealed the presence of various stress responsive cis-acting elements (Fig 3B). The number of these cis-regulatory elements varies within these two promoter sequences. One striking observation is that the OsbZIP23 promoter of IR20 genotype carries a deletion of a sequence of 35 nucleotides in comparison to O. rufipogon promoter at the 5'-UTR region. Within this 35-nucleotide deletion, two extra MYB-binding elements are found to be present in O. rufipogon. Interestingly, the OsbZIP23 promoter is TATA-less, however Y- patches are found to be present around transcription start site (TSS). Y-patches are the core promoter elements, which are direction-sensitive pyrimidine rich boxes found around TSS [48–49]. The variation in the number of cis-acting elements as well as a deletion of 35-nucleotide sequence prompted us to check the functional activity of both the promoters by expression analysis of the GFP reporter gene using the promoter-probe constructs (S1 and S2 Figs). Analysis of transgenic leaf tissues revealed that both the fluorescence intensity (Fig 4B) and the relative expression level (Fig 4C) of GFP are much higher in RuP#2 and RuP#3 lines than the 20P#1 and 20P#2 lines under drought stress condition. These results suggest that the promoter activity of O. rufipogon is comparatively higher than the IR20 rice genotype, which could be a reason that under similar growth condition, the expression of OsbZIP23 gene varies significantly in drought-tolerant and drought-sensitive genotypes.

We have cloned and sequenced the OsbZIP23 CDS from the six drought-sensitive and five drought-tolerant rice genotypes. The detailed sequence analysis of the derived polypeptides revealed no striking differences, except the deleted 5-amino acid sequence (AAEHA) in O. rufipogon, which is normally present in all other rice genotypes (except O. nivara which has AVEHA in place of AAEHA) between the transactivation domain I and domain II of OsbZIP23 (S6 Fig). Therefore, we wanted to examine the variation on functional activity, if any, between the two natural allelic variants of OsbZIP23 CDS from two wild rice progenitors, i.e., O. nivara (having 1083 bp CDS) and O. rufipogon (having 1068 bp CDS with deleted 15 nucleotides encoding AVEHA in comparison to O. nivara). The respective CDS was transgenically expressed in a drought-sensitive cultivar (IR20) using the constitutive OsUbi1 promoter driven expression cassette (S3 and S4 Figs). Further to compare the overexpression of these two CDS in transgenic rice lines (OEN and OER), RNAi-mediated OsbZIP23 gene silencing transgenic lines (RNAi) were also developed with an appropriate genetic construct (S5 Fig). The overexpression (OE) and RNAi-mediated silencing (RNAi) of OsbZIP23 gene in drought-sensitive indica rice genotype IR20 resulted in expected increase and decrease of OsbZIP23 expression in OE lines (OEN#9, OER#5 and OER#7) and RNAi lines (RNAi#1 and RNAi#4), respectively in comparison to non-transgenic (NT) plants at both transcriptional (Fig 5C) and translational levels (Fig 5D). Under the drought stress condition, the OE and RNAi transgenic lines showed opposite phenotypes. In both vegetative and reproductive stages, the OE lines exhibited increased drought tolerance in terms of higher survival rate than the RNAi lines and NT plants;
whereas the RNAi lines performed significantly poorer than the NT plants under drought stress condition (Fig 6A, 6B, 6C and 6D). Analysis of yield attributing traits revealed that the OE lines had increased panicle weight and spikelet fertility in comparison to RNAi lines and NT plants under drought stress (Fig 6E, 6F, 6G and 6H). These results clearly established that the transgenic overexpression of OsbZIP23 improves drought tolerance and grain yield in rice. On the contrary, down-regulation of the endogenous OsbZIP23 makes the rice plants drought sensitive and poor yielder. Previous works on overexpression or silencing of ABA-responsive genes have documented similar results [39, 50]. The ability to retain water is a crucial factor for plants to combat drought. The water loss rate (WLR) and relative water content (RWC) are the two important parameters that reflect the water status of the plant [39, 51]. Measurement of WLR and RWC in leaf tissues revealed that the OE lines had highest water retention capacity compared to RNAi lines and NT plants (Fig 7A and 7B). This finding strongly indicates that the OsbZIP23 is involved in maintaining the ability of plants to retain water. We further investigated the physiological mechanism by which OsbZIP23 enables the plant to retain water and mitigate the effect of drought stress. It is established that under water-limiting condition, proline and sugars are accumulated in plant cells, resulting decrease in the cellular osmotic potential [52]. Our results documented that the content of both proline and soluble sugars in the leaf tissues of OsbZIP23 OE lines was much higher than the RNAi lines and NT plants under drought stress condition (Fig 7C and 7D). Drought stress can reduce photosynthesis, and results in accumulation of excess reactive oxygen species (ROS), which leads to cell toxicity, membrane peroxidation and even cell death [33]. There is a constant need for efficient mechanisms to avoid oxidative damage to cell. Malondialdehyde (MDA) is a stress marker that measures the degree of damage caused by stress in plants [54]. Our analysis on transgenic lines displaying lower MDA content in OE lines in comparison to RNAi lines and NT plants under drought stress (Fig 7E) implies that the drought stress-induced oxidative damage causes less harm to OE lines than RNAi lines and NT plants. Therefore, accumulation of ROS under drought stress in OE lines, RNAi lines and NT plants were checked. Accumulation of peroxide and superoxide free radicals in leaf tissues of OE lines was negligible in comparison to RNAi lines and NT plants under drought stress condition (Fig 7F and 7G). It can be inferred that the transgenically overexpressed OsbZIP23 has improved drought tolerance by ROS detoxification, and thereby leading to less membrane damage, which indeed resulted from increased water retention through accumulation of compatible osmolytes.

To understand the molecular mechanism behind the OsbZIP23-mediated drought stress adaptation, we analyzed the transcript level of three representative inducible marker genes of ABA signaling pathway viz., OsRab16B, OsRab21 and OsLEA3-1. Rabs are dehydrin family proteins reported to decrease the cellular osmotic potential while protecting the plants under drought stress [55]. LEA or late embryogenesis abundant proteins have been proposed to maintain the cell membrane structure stable under water deficit condition by reducing the damage caused by high concentration of ions in dehydrating cells [56], and are regulated by bZIP transcription factors through ABRE cis-elements [26]. The increased and decreased expression of OsRab16B, OsRab21 and OsLEA3-1 genes in OE and RNAi lines, respectively (Fig 8 and S7 Fig) suggests that the OsbZIP23 is a transcriptional activator of these genes in rice. The high expression level of these genes, in turn, most likely leads to up-regulation of other genes that are controlled by OsRab16B and OsRab21 and OsLEA3-1 in an ABA-dependent manner, and could be explained by the established ABA signaling pathway (S8 Fig) [57]. It is well known that the phytohormone ABA maintains seed dormancy, inhibits seed germination and retards seedling growth [39]. Drought stress induces ABA biosynthesis and triggers ABA-dependent signaling pathway. Thus, we investigated ABA-sensitivity of OE and RNAi lines, which displayed significantly increased and decreased sensitivity, respectively to
exogenous ABA at both germination and post-germination stages (Fig 9). These results are consistent with the previous observations on increased ABA sensitivity of the transgenic rice lines overexpressing OsbZIP23 gene [25], confirming that the ABA-signaling pathway is positively regulated by the OsbZIP23 transcription factor.

Conclusions
Allelic polymorphism analysis in OsbZIP23 coding DNA sequence showed presence of some non-synonymous SNPs in different genotypes as well as a major 5-amino acid deletion in O. rufipogon. The OsbZIP23 transcript expression profiling under vegetative and reproductive stages revealed comparatively higher expression level of OsbZIP23 in drought-tolerant genotypes than the drought-sensitive ones. The OsbZIP23 promoter of drought-tolerant O. rufipogon showed higher promoter activity than the drought-sensitive IR20 rice genotype. Overexpression of either of the two polymorphic forms of OsbZIP23 from wild rice genotypes and endogenous OsbZIP23 gene silencing through RNAi revealed that the enhanced expression of OsbZIP23 improves drought tolerance and grain yield in drought-sensitive rice genotype. Further the overexpression lines exhibited increase in water retention capacity through higher accumulation of compatible osmolytes, leading to the decrease in membrane damage caused by the activity of reactive oxygen species. The drought tolerance and yield improvement in rice genotypes are found to be dependent on the enhanced gene expression of OsbZIP23 due to higher promoter activity rather than the natural polymorphism in coding sequence. Therefore, the present study advances our understanding on developing drought-tolerant elite rice cultivars utilizing highly expressed promoter to drive the OsbZIP23 transgene. The information generated through the present work will help us to develop gene-based molecular marker for selective breeding to improve drought tolerance in rice crop utilizing this type of transcriptional regulator.

Supporting Information
S1 Fig. Schematic representation of OsbZIP23 promoter-GFP reporter gene construct in pCAMBIA1300 plasmid. Promoter sequence of 1612 bp from drought tolerant wild genotype O. rufipogon was fused to GFP reporter gene and the transgenic lines developed with this genetic construct were designated as RuP series. (TIF)

S2 Fig. Schematic representation of OsbZIP23 promoter-GFP reporter gene construct in pCAMBIA1300 plasmid. Promoter sequence of 1578 bp from drought sensitive indica rice genotype IR20 was fused with GFP reporter gene and the transgenic lines developed with this expression cassette were designated as 20P series. (TIF)

S3 Fig. Schematic representation of the overexpression (OE) construct of 1068 bp CDS of OsbZIP23 from O. rufipogon in pCAMBIA1300 plasmid. The transgenic rice lines developed with this genetic construct were designated as OER series. (TIF)

S4 Fig. Schematic representation of the overexpression (OE) construct of 1083 bp CDS of OsbZIP23 from O. nivara in pCAMBIA1300 plasmid. The transgenic rice lines developed with this genetic construct were designated as OER series. (TIF)
S5 Fig. Schematic representation of the RNAi-mediated silencing (RNAi) construct of Osb-ZIP23 gene in pCAMBIA1300 plasmid. The 399 bp of OsbZIP23 CDS (3’-part) from O. rufipogon was cloned in sense and antisense direction containing an arbitrary 200 bp linker. The transgenic rice lines developed with this genetic construct were designated as RNAi series.

(TIF)

S6 Fig. Structural domain prediction of OsbZIP23 protein using ScanProsite web tool (http://prosite.expasy.org/scanprosite). In both drought tolerant genotypes (A) O. rufipogon and (B) O. nivara, three conserved transactivation domains (domain I, II and III) and one basic leucine zipper (bZIP) domain was predicted. Presence of three protein kinase C phosphorylation site (red pointed serine and threonine), five casein kinase II phosphorylation site (underlined threonine residues) and 11 N-myristoylation sites (indicated in blue colour) were also predicted. The 5-amino acid (AVEHA) deletion in O. rufipogon is indicated with gap. Note that this 5-amino acid sequence is AAEHA in all other genotypes tested except O. nivara and sorghum, which have AVEHA (Fig 1B).

(TIF)

S7 Fig. Real time PCR analysis in reproductive stage showing relative expression level of OsbZIP23 and three selected representative drought inducible genes working downstream in the ABA signaling pathway. (A) OsbZIP23 (B) OsRab16B (C) OsRab21, and (D) OsLEA3-1 in OsbZIP23 OE lines, RNAi lines and NT plants, where rice polyubiquitin1 (OsUbi1) gene was taken as internal reference. Data bars represent the mean ±SD of triplicate measurement. Statistical analysis by Student’s t-test indicated significant differences (** P<0.01).

(TIFF)

S8 Fig. Schematic representation of the ABA signaling pathway under drought stress. Drought induces accumulation of ABA, which is regulated by PARP and LOS5. ABA is perceived by the PYR1 receptor and induces the expression of SnRK2 kinases by releasing ABI1 phosphatase. This in turn activates the downstream AREB/bZIP-transcription factors. The activated AREBs (through the interaction with ABRE cis-elements) induce the expression of a series of downstream genes to produce different classes of osmoprotectants, which protects the plant by decreasing cellular osmotic potential. Adopted from Yang et al., 2010.

(TIFF)

S1 Table. List of primers used in this study.

(DOCX)

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Author Contributions

Conceived and designed the experiments: SG SKS MKM AD. Performed the experiments: AD MKS SG. Analyzed the data: AD SKS MKM. Wrote the paper: AD SG MKM.
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