Materials and Methods

Analysis of public microarray data

We collected microarray data sets from NCBI Gene Expression Omnibus (Edgar et al. 2002). The data contain 11 sets of rice drought studies, which are comprised of expression profiles in various genotypes, different growth stages from seedling to reproductive stage and different organs under normal and drought stress conditions (Table S1). The data were categorized into 5 sets according to the microarray platform and data normalization status. For the Affymetrix microarray data (platform: GPL2025), raw data (CEL file format) were downloaded, and global normalization was applied via Robust Multi-array Average (RMA) (Irizarry et al. 2003). Probe sets corresponding to only one gene model from the Rice Genome Annotation Project (Kawahara et al. 2013) were selected for the present study. For the NimbleGen Rice 3'-Tiling Microarray (platform: GPL15594 and GPL7344), the data was summarized by calculating the average of probe signals for each gene per chip. For the Agilent microarray data (platform: GPL7252), the data was subjected to per chip quantile normalization by using the limma R statistical package (Ritchie et al. 2015). Log₂-transformed signals were defined as normalized gene expression.

Genes in the 4 gene families, GAPDH, actin, ubiquitin and cyclophilin, were mined from the functional annotation in the gene model of Rice Genome Annotation Project and the Rice Annotation Project Database (RAP-DB; http://rapdb.dna.affrc.go.jp/). Several genes in these 4 gene families were defined as potential ubiquitous genes in
this study; i.e. 7 of GAPDH, 13 of actin, 31 of ubiquitin and 23 of cyclophilin. (Table S2). As an index of gene expression stability, CV of normalized gene expression in a dataset was calculated for each gene using Microsoft Excel. The rank was assigned to genes according to their CV by using the rank function in Excel, with the lowest CV garnering the highest score in each gene family. The ranks were used to screen stably expressed genes across the 5 different microarray datasets by calculating a weighted rank score (WRS), which reflects variety of samples in each microarray data set. WRS in each data set was given by following the equation

\[ WRS = R \times (n_g + n_e + n_d + n_o) \]

\( R \) denotes the rank in the data set. \( n_g \) (number of genotypes), \( n_e \) (number of environmental conditions), \( n_d \) (number of developmental stages), and \( n_o \) (number of organs) were counted in the dataset.

WRS in the 5 datasets were averaged per each gene (Table S2), and it was compared among each gene family members. We assume that a smaller WRS represents a more stable gene expression across microarray data.

**Plant Materials**

Seeds from *IR64* (*Oryza sativa* L. var. *indica*) were germinated in moist filter paper at 30°C for 5 days. Following germination, these were sown in potted soil and grown in a greenhouse with temperature control under normal lighting conditions in the International Rice Research Institute (Los Baños, Laguna,
14°10'11.81"N, 121°15'39.22"E) for 4 weeks from April to May 2015. Drought stress was applied by stopping irrigation for 10 days until leaf rolling was observed and soil moisture content was reduced to approximately 15%. A control treatment comprising of well-watered plants was kept by maintaining enough water content throughout the drought stress. Volumetric soil water content was monitored using ML3 ThetaProbe Soil Moisture Sensor (Delta-T Devices Ltd.). Three biological replicates each for shoots and roots were collected for each treatment. These were frozen in liquid nitrogen and stored at -80°C.

A tropical japonica cultivar WAB 56-104, a parental line for NERICA (New Rice For Africa), was grown in a greenhouse of the International Rice Research Institute in 2015. Germinating seeds were planted in a soil of 30 cm height plastic plots in January. When plants were at the booting stage in March, irrigation was stopped, and volumetric soil water content was monitored to prepare samples under progressive drought stress. Roots and flag leaves were collected at 0, 3, 5 and 7 days after initiation of stress. The soil moisture content was 50% (the maximum), 30%, 20% and 15%, respectively. Flag leaves and roots were collected with 3 biological replicates for each treatment. The samples were frozen in liquid nitrogen and stored at -80°C.

**RNA preparation**

Total RNA was extracted from frozen samples by using the PureLink Plant RNA Purification Reagent (Invitrogen, Thermo Fisher Scientific Inc., USA) as per the manufacturer’s instructions. Total RNA was subjected to DNase I digestion to
eliminate contamination of DNA by using TURBO DNA-free Kit (Invitrogen, Thermo Fisher Scientific Inc., USA). The extracted RNA samples were analyzed using a Nano Drop 2000c spectrophotometer (NanoDrop Technologies), and the absorbance (A) in 230, 260 and 280 nm were measured. RNA samples with $A_{260}/A_{280}$ ratio between 1.9 to 2.1 and $A_{260}/A_{230}$ ratio greater than 2.0 were used for first strand cDNA synthesis and further qRT-PCR analysis.

**qRT-PCR assay**

For IR64 samples, 4 μg of total RNA was used for 1st strand cDNA synthesis with oligo d(T)$_{15}$ primer using the ImProm-II Reverse transcription system (Promega Corporation, Madison, USA) according to the manufacturer’s instructions. IR64 cDNA samples were diluted 5-fold and used as a template for each qRT-PCR assay. For WAB 56-104, 100 ng and 500 ng of total RNA in roots and leaves, respectively, were employed for 1st strand cDNA synthesis. WAB 56-104 cDNA samples were then diluted 10-fold and used a template for each qRT-PCR assay.

For IR64 samples, a 10 μl reaction volume consisted of 1 μl of diluted cDNA, 5 μl of 2X SYBR Select PCR master mix (Applied Biosystems, Thermo Fisher Scientific Inc., USA) and 0.4 μl of 10mM primer for each primer pair. Four μl of diluted cDNA was used for WAB 56-104. The reaction conditions followed this protocol: 50°C for 2 minutes, 95°C for 4 minutes, then 40 cycles of 95°C for 10 seconds and 60°C for 30 seconds. Dissociation curve analysis was done for the IR64 seedling stage samples by subjecting the PCR samples to 60°C to 95°C at 1% temperature ramp speed.
Each assay was carried out with two technical replicates for each biological replicate. Ct values were determined by the 7500 Fast Software v2.3 (Applied Biosystems, USA). The threshold level was automatically set at 10 times of the standard deviation of the fluorescence value of the baseline. Ct values in the same biological replicate were averaged into the representative Ct value, and these were then used for statistical analysis.

Primer information

Transcript sequences of the reference genes assayed were retrieved from the Rice Genome Annotation Project and then used to design gene-specific primers using Primer-BLAST from the NCBI website (Ye et al. 2012; http://www.ncbi.nlm.nih.gov/tools/primer-blast/) with the following parameters: product length of 75-100 bp, melting temperature of 60°C and GC content of 40 to 60%. Sequence specificity of the primers was tested through a BLAST search against the Oryza sativa var. japonica sequence.

Statistical analysis by the ΔCt approach

ΔCt approach, documented by Silver et al. (2006), was applied to evaluate gene expression stability. In brief, this method employs average of standard deviation of ΔCt among all the reference genes tested as an expression stability index. In this analysis, we tested plural (up to 4 genes) reference genes. Mean SD of ΔCt was calculated by an in-house Perl script. Geometric mean of Ct values was applied to
calculate ΔCt value of each plural reference gene set as following the method of Vandesompele et al. (2002).

**Analysis of microarray data at a seedling stage**

We applied in-house rice microarray data generated from *IR64* seedling stage roots under well watered and drought stressed condition (Dixit et al. 2015). The microarray was designed in 56K Agilent rice microarray platform (60K × 8-plex format), which allows monitoring 52,901 rice genes that are corresponding to rice gene model MSU 7.0. The microarray TIF images were processed to run batch extractions by choosing appropriate grid using Agilent’s Feature Extraction Software version 11.5.1.1. Raw signal intensities were converted into log2 expression values, and per chip quantile normalization was carried out for each organ by using limma R statistical package (Ritchie et al. 2015).

**Reference for Materials and Methods**


expression and hybridization array data repository. Nucleic Acids Res 30: 207-210


