Methodology

A quantitative RT-PCR platform for high-throughput expression profiling of 2500 rice transcription factors

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Abstract

Background: Quantitative reverse transcription – polymerase chain reaction (qRT-PCR) has been demonstrated to be particularly suitable for the analysis of weakly expressed genes, such as those encoding transcription factors. Rice (Oryza sativa L.) is an important crop and the most advanced model for monocotyledonous species; its nuclear genome has been sequenced and molecular tools are being developed for functional analyses. However, high-throughput methods for rice research are still limited and a large-scale qRT-PCR platform for gene expression analyses has not been reported.

Results: We established a qRT-PCR platform enabling the multi-parallel determination of the expression levels of more than 2500 rice transcription factor genes. Additionally, using different rice cultivars, tissues and physiological conditions, we evaluated the expression stability of seven reference genes. We demonstrate this resource allows specific and reliable detection of the expression of transcription factor genes in rice.

Conclusion: Multi-parallel qRT-PCR allows the versatile and sensitive transcriptome profiling of large numbers of rice transcription factor genes. The new platform complements existing microarray-based expression profiling techniques, by allowing the analysis of lowly expressed transcription factor genes to determine their involvement in developmental or physiological processes. We expect that this resource will be of broad utility to the scientific community in the further development of rice as an important model for plant science.
based hybridisation typically allows the detection of one transcript per cell [3,4], qRT-PCR can detect one transcript per 1000 cells [5]. Recent improvements in qRT-PCR methodology have eliminated many of the initial problems that were associated with quantitative gene expression studies, such as those arising from alternative splicing events [6]. Despite such developments qRT-PCR is mostly used to detect relatively small numbers of genes.

Transcription factors (TFs) are proteins (trans-acting factors) that enhance or repress gene expression through their binding to specific DNA sequences (cis-acting elements) in the promoters of their target genes. The functional characterization of TFs is crucial for the reconstruction of transcriptional regulatory networks controlling developmental and physiological processes such as growth, organ formation and the response to hormonal or environmental stimuli [7,8]. Transcription factor genes represent a sizable fraction of the genomes of all eukaryotic organisms, including higher plants [7]. Analysis of the rice genome [9,10] indicated that approximately 2.6% of the identified genes encode TFs [10]. Currently, the functional analysis of TFs in monocotyledonous species lags considerably behind that of the model dicotyledonous species Arabidopsis thaliana.

Microarray expression profiling in rice has not been widely reported with relatively few publicly available data. Studies using qRT-PCR have also not been widely reported, have focused on small groups of genes, and in many cases were only used to confirm expression changes from microarray experiments. Therefore, the utility of qRT-PCR as a high-throughput method in rice has not been investigated.

To facilitate the analysis of rice TFs we have recently established a database [11,12]. The coding sequences of more than 2500 identified rice TFs were used to design primers for a large-scale qRT-PCR platform. The comparative analysis of several rice varieties and tissues described here has confirmed the broad applicability of the platform.

Results and Discussion

Primer design

Analysis of the rice genome indicated that 21% of all genes give rise to alternatively spliced transcripts [13]. In the case of TFs, splice variants can affect the architecture of the DNA-binding domain and often show tissue-specific expression patterns [14]. To distinguish between such variants, splice variant-specific primer pairs were designed for the 5.7% of all TF loci (131 TF loci) where this was possible. In total, primer pairs for 2508 gene models derived from 2306 loci were designed (Additional file 1). The design of primers followed a set of stringent criteria, as generally suggested in qRT-PCR protocols (e.g. Primer Express Software v2.0 Application Manual, Applied Biosystems). To minimize the risk of amplifying contaminating genomic DNA, primers spanning at least one exon-exon junction, or annealing to different exons, were designed where possible (56% of predicted gene models). However, 35% of genes contained no introns. The specificity of each primer was confirmed by comparing its sequence with all predicted rice coding sequences (CDS) using the BLASTN tool at TIGR [15] to ensure that at least one primer of each pair targets a unique site within the set of predicted rice CDS.

RNA sampling and control for genomic DNA contamination

RNA was initially extracted using a phenol-based method from two different tissues (root and shoot) of four rice cultivars, three of which were indica cultivars (Cham, DR2 and Lua man) and the fourth was a japonica cultivar (Niponbare). This protocol (e.g. as described by Czechowski et al. and Jain et al. [5,16]) gave satisfactory total RNA yield, but the RNA quality was too low for the synthesis of high-quality cDNA [data not shown] [17]. We therefore used the guanidinium thiocyanate-based RNeasy Plant Mini Kit (Qiagen, Hilden, Germany) [18]. Total RNA extraction was straightforward and provided RNA with high yield and quality from both root and shoot tissue (45–75 μg total RNA/100 mg fresh weight).

RNA preparations are usually contaminated with low amounts of genomic DNA, which can result in non-specific amplification [19]. The manufacturer’s recommended on-column DNase treatment was not sufficient to remove interfering genomic DNA. Therefore, we performed a second DNase incubation on the isolated RNA to eliminate detectable genomic DNA contamination. Genomic DNA was detected by qRT-PCR using 0.125 μg of isolated RNA as template, and three different primer pairs annealing to intergenic regions of chromosomes 1 (AP003727, positions 7366–7426) and 7 (AP006456, positions 27334624–27334684), and an intron of the gene Os01g01840. It is important to select more than a single genomic region to assess genomic DNA contamination because chromosomal sites can be differentially accessible to DNase I. Omitting the second DNase digestion always resulted in the amplification of some, but not all, of these genomic regions (data not shown). In most cases the C<sub>T</sub> values obtained were >30. Synthesis of cDNA from the isolated RNA was only performed when all three genomic control amplifications scored negative.

We chose a two-step qRT-PCR protocol where reverse transcription and PCR-mediated cDNA amplification are carried out in subsequent steps in separate tubes. The two-step protocol is preferred when SYBR Green is used as a detection dye because it diminishes unwanted primer
dimer formation [20]. The reverse transcriptase reaction was primed with oligo-(dT) instead of random-sequence primers because the latter preferentially selects for more abundant mRNA species and not for transcripts of weakly expressed genes, such as many TF genes. A flowchart of the protocol is shown in Figure 1.

**Reaction specificity**

All 2508 primer pairs were checked by qRT-PCR using cDNAs synthesised from the roots of the rice cultivars, Cham and DR2 kept under control and salt stress (100 mM NaCl) conditions. Additionally, shoot cDNA from DR2 was used to test a smaller set of primer pairs targeting 192 TF genes. Melting curve analysis was performed for all PCR products to confirm the occurrence of specific amplification peaks and the absence of primer-dimer formation. Finally, all 2508 PCR products from the control DR2 root sample were run on 4% agarose gels and photographically documented (available upon request).

Approximately 3% of all TF genes analysed (i.e. 73 out of 2508 genes) did not yield detectable PCR amplicons, indicating no or weak expression under the employed conditions. In most cases (57 TF genes) at least one primer spanned an exon-exon junction thus precluding tests for priming efficiency on genomic DNA template. Only 2.5% of all reactions (61 TF genes) yielded unspecific PCR products as indicated by multiple or incorrectly sized amplicons.

**Primer efficiency**

Although the fluorescence during qRT-PCR is primarily determined by the starting abundance of a given cDNA template, its increase during the run is strongly affected by the amplification efficiency. When this efficiency is 100%, the amount of a cDNA targeted by a given primer pair is doubled in every PCR cycle of the exponential phase. However, the priming site, the sequence specificity of the primer, and the unwanted formation of primer-dimers are factors that can significantly affect the efficiency of individual PCR reactions. We used the LinRegPCR software [21] to determine the PCR efficiency of each primer pair of the TF qRT-PCR platform, taking into account all amplification profiles obtained (27588 in total). Firstly, the correlation coefficients (R) assigned to efficiency values were used to evaluate the amplification curves and all reactions with an R < 0.990, reflecting low-quality amplifications, were excluded from further analyses. Subsequently, average PCR efficiencies were computed for each individual primer pair across all analyzed samples. This showed that 8% of the 2508 primer pairs (200 TF genes) displayed PCR efficiencies greater than 1.90, and 87% of the primer pairs (2182 TF genes) had efficiencies of 1.51–1.90. Only 5% of the primer pairs (125 TF genes) had PCR efficiencies with mean values below 1.4, and these mostly represented reactions that lacked detectable fragment amplification (C<sub>T</sub> > 40) or that generated unspecific PCR products. The efficiencies of reference genes tested in this work ranged from 1.73 to 1.96 (see below; Table 1).

The TF qRT-PCR platform was developed primarily using the nuclear genome sequence of the *japonica* cultivar Nipponbare (TIGR annotation) [15], which might affect its general applicability for experiments involving *indica* cultivars. SNP (single nucleotide polymorphism) variation among the two rice subspecies was found to be low (<0.4%) [22], and these mostly occur in intergenic regions which contain approximately four times more SNPs than occur within genes [23]. SNPs may affect the comparative analysis of gene expression levels in qRT-PCR experiments. Therefore, to investigate variation of primer specif-


Table 1: Selected reference genes for rice and corresponding primer pair information.

| Locus identifier | Gene name       | Primer sequence F/R [5’-3’] | Amplicon length [bp] | Amplicon T_m [°C] | PCR efficiency |
|------------------|-----------------|-----------------------------|----------------------|------------------|---------------|
| Os03g50890       | Actin           | CTCCCCCATGCTATCCTTCG         | 91                   | 81               | 1.9           |
| Os05g36290*      | Actin1          | ATCCCTGTAGTCACCCTCCTG       | 118                  | 80               | 1.9           |
| Os01g59150*      | β-Tubulin       | GAGTCCACATGCTGCTAAGTT       | 64                   | 80               | 1.96          |
| Os06g11070       | Expressed protein | AGGCTGGTGGAGAGCTCCAT | 101                  | 85               | 1.83          |
| Os03g55270       | TIP41-like      | GTTGGATGCCACCCCGCAA         | 62                   | 77.5             | 1.81          |
| Os08g19610       | Cyclophilin     | CCACCATCAGAGATCGGATTT       | 65                   | 84.2             | 1.73          |
| Os03g08020*      | Elongation factor 1α | GTCAATGGCCACGTCGACTC   | 118                  | 83.5             | 1.85          |

* Primer pair recognizes all splicing variants. Location of forward and reverse primer in the same exon [S] or in different exons [D].


icity between the indica varieties Cham and DR2, PCR efficiencies were separately computed in a cultivar-specific manner for all gene models. Although individual primer pairs of the TF primer platform can exhibit slight differences for their target genes in different varieties/cultivars (Figure 2), this did not significantly affect the overall applicability of the platform for expression profiling experiments, a finding we have also validated in independent experiments (Caldana et al., manuscript in preparation).

**Accuracy and precision of real-time PCR**

To determine the sensitivity and accuracy of the rice qRT-PCR platform, as reported for other platforms [5], we performed additional experiments. Firstly, we assessed the linearity, sensitivity and accuracy using three weakly expressed genes with preferential expression in either shoots (Os12g38200) or roots (Os03g55610 and Os08g38220). Shoot- and root-derived cDNAs were mixed in different ratios (see Figure 3) and transcript abundance of the three genes was determined by qRT-PCR. Linearised values were calculated as $2^{(C_T-C_R)}$, where $C_T$ represents the threshold cycle and directly reflects template abundance. A linear relationship was observed in all three cases, suggesting that, as previously reported for Arabidopsis [5], qRT-PCR was sufficiently accurate in rice even for low-abundant transcripts. These relationships were highly significant with $R^2$ values (coefficient of determination) of 0.61, 0.94, and 0.90 for Os03g55610, Os08g38220, and Os12g38200, respectively (Figure 3).

We also tested the experimental reproducibility of the rice qRT-PCR platform. Czechowski et al. [5] demonstrated qRT-PCR precision by analysing intra-assay variation using the same pool of cDNAs and inter-assay variation using two different pools of cDNAs synthesised from the same batch of RNA, obtaining $R^2$ values of 0.99 and 0.95, respectively. Here, we wanted to determine the variation between different biological replicates which encompasses both technical and biological variation. Using cDNA synthesised from DR2 harvested in three independ-
ent experiments we measured the expression of 201 TF genes. The $\Delta C_T (C_{T, \text{gene of interest}} - C_{T, \text{reference gene}})$ was calculated and the precision of the assay was assessed using the coefficient of variation (CV). Despite the fact that the majority of gene models tested (111 genes) displayed an extremely low expression level ($C_T > 35$), the obtained mean CV was 14%. This is in good agreement with published expression data for human keratinocyte subclones, in which a CV of 18% was found for genes with a $C_T > 30$ [24]. In contrast, CV values are generally higher in microarray-based analyses of genes with such low expression, indicating a lower reproducibility [25]. These findings underscore the advantage of qRT-PCR as an alternative and often superior tool for expression profiling studies, especially for the investigation of genes with low expression level.

Selection of reference genes for qRT-PCR in rice

Generally in qRT-PCR, transcripts of stably expressed genes, also called reference genes, are employed for data normalisation. In rice, previous publications have suggested 18S-rRNA, GADPH, UBI5 and EF-1 $\alpha$ as good reference genes [26,16]. To identify the most suitable reference genes in rice we initially selected nine candidates: 18S-rRNA, ubiquitin (UBQ), actin (ACT), actin1 (ACT1), $\beta$-tubulin (TUB), cyclophilin (CYC), elongation factor 1 $\alpha$ (EF-1 $\alpha$), which are commonly used house-keeping genes in plants, and expressed protein (EP) and TIP41-like protein (TIP41) found to be good reference genes in Arabidopsis [27]. UBQ (Os01g45420), considered as a stable house-keeping gene in various plant species [27,16], had unstable expression across three different rice cultivars (data not shown), and was excluded from further studies. Although the abundance of 18S-rRNA remained constant in different rice cultivars and physiological conditions (data not shown), we did not consider it further as a suitable reference for our analyses, primarily because it requires the use of random hexamers instead of oligo(dT) as primers for the reverse transcriptase.

An overview of the remaining seven reference genes is given in Table 1. CYC, EP and TIP41 are expressed at low, TUB, ACT and ACT1 at intermediate, and EF-1 $\alpha$ at high levels, respectively. Their expression stability was measured by qRT-PCR in a set of 11 different cDNA samples (Additional file 2), and calculated using the gene expression stability measure (M) implemented in the geNORM software [28]. This determines the stability of a reference gene, taking into account the average pair-wise variation (V) of that gene in comparison to all other reference genes tested. Assuming that the difference of gene expression level of two ideal control genes is the same in all experi-
values and the best ranked genes were ACT1 and CYC, followed by EF-1α. However, the weakly expressed gene CYC was the least stable when more diverse conditions were used. Therefore, considering all the experimental categories, ACT1 and EF-1α were the most stable reference genes. Additionally, EP, similar to its Arabidopsis orthologue (At4g33380) [27], showed good stability in all experiments. Generally, we recommend including more than a single reference gene in each qRT-PCR experiment. Conveniently, the geNORM software also calculates the pairwise variation (V), which indicates the optimal number of reference genes to be analysed in a given experiment.

**Conclusion**

A qRT-PCR platform for multi-parallel expression profiling of more than 2500 rice TF genes has been established. It complements existing microarray-based profiling technologies, which are generally not well suited for reliable analysis of weakly expressed genes including TFs. This resource is available for the scientific community to use for their own experiments.

**Methods**

**Plant material**

The three rice (*Oryza sativa* L. *sp. indica*) cultivars (Cham, DR2 and Lua man) were obtained from the Institute of Biotechnology (Hanoi, Vietnam). Additionally, a Nipponbare (*Oryza sativa* L. *sp. japonica*) cultivar provided by the International Rice Research Institute (Manila, Philippines) was used. The plants were grown in hydroponic culture [29] under a day-length of 12 h at 26/22°C (day/night), 70% humidity and 700 μmol m⁻² s⁻¹ light intensity. Roots and shoots were harvested three weeks after germination. Salt-stressed plants (Cham, DR2, and Lua man) were harvested 30 min or 3 h after the application of 100 mM NaCl, with control samples harvested in parallel.

**RNA extraction, DNase I digestion and cDNA synthesis**

The roots and shoots of three biological replicates with five plants each were pooled to isolate total RNA using the RNasy Plant Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer’s protocol. During the extraction, the first on-column DNase I (Qiagen) digestion was carried out. A second DNase I (Roche, Mannheim, Germany) digestion was performed on a total of 60 μg of total RNA according to the manufacturer’s instructions. Absence of genomic DNA contamination was subsequently confirmed by qRT-PCR using three different primer pairs. The primers were designed to amplify two intergenic regions (AP003727, positions 7366 – 7426: forward 5'AGAGAAGACCGCCATGTGG-3' and reverse 5'-CTGGGAACCCACAAAGAATGAC-3'; and AP006456, positions 27334624 – 27334684: forward 5'-TATCCACTCGACAGGACCTAGAC-3', and an intron sequence...
of the gene Os01g01840 (forward 5’-TAGAGAGTTCT-GATCTTGCGCG-3’ and reverse 5’-CGGCCCATCAAT-GAAGTCTT-3’). RNA integrity was checked on 1% (w/v) agarose gels and the concentration measured before and after DNase I digestion. cDNA was synthesized from 5 μg of total RNA using Superscript™ III reverse transcriptase (Invitrogen, Karlsruhe, Germany), according to the manufacturer’s instructions. The efficiency of cDNA synthesis was estimated by qRT-PCR using β-tubulin (Os01g59150, forward primer 5’-GGAGTCACATGCTGCTAAGGT-3’ and reverse primer 5’-TCACTGCGCTTACGGAGG-3’).

**Design and validation of qRT-PCR primers**

Transcription factor sequences were extracted from version 1 of the Rice Transcription Factor Database [11,12] and used to establish the rice TF qRT-PCR platform. The set of 2508 gene models corresponding to 2306 loci of confirmed and putative transcription factors was subsequently used to design primers for ca. 500 genes. The remaining 2000 primer pairs were designed by MWG Biotech AG (Ebersberg, Germany). A standard set of reaction conditions and a set of stringent criteria were used as follows: T_{m} of 60°C ± 2°C, PCR amplicon length of 60 to 150 bp, primer length of 20 ± 5 bp, and a guanine-cytosine content of 45 to 55%. If gene structure allowed, at least one primer was designed to cover an exon-exon junction. The specificity of the primer pair sequence was checked against Version 2 of the rice transcripts (CDS) from the TIGR Rice Database [15] using the BLAST programme. The EXPECT threshold (statistical significance) was set to 1000, as suggested when searching for short, nearly exact matches [30]. The specificity of the amplicons was set to 1000, as suggested when searching for short, nearly exact matches [30]. The specificity of the primer pair sequence was estimated using the LinRegPCR software [21] (Additional file 1).

**Selection of reference genes**

Potential reference genes were chosen based on published data for rice and other plant species and rice orthologues were identified using the TBLASTN programme [15]. The gene models used were: actin (Os03g50680), actin1 (Os05g36290), β-tubulin (Os01g59150), expressed protein (Os06g11070), TIP41-like protein (Os03g55270), cyclophilin (Os08g19610) and elongation factor 1α (Os03g08020). Primer design followed the same criteria as described and details are given in Table 1.

**Quantitative RT-PCR conditions and analysis**

PCR reactions were conducted in an ABI PRISM 7900 HT sequence detection system (Applied Biosystems). A 5 μl reaction containing 0.5 μl of cDNA (1.25 ng/μl), 200 nM of each gene-specific primer and 2.5 μl of SYBR Green master mix (Applied Biosystems Applera, Darmstadt, Germany), was used to monitor double-strand DNA synthesis. The qRT-PCR reactions were carried out following the recommended thermal profile: 50°C for 2 min, 95°C for 10 min, followed by 40 cycles of 95°C for 15 s and 60°C for 1 min. After 40 cycles, the specificity of the amplifications was tested by heating from 60°C to 95°C with a ramp speed of 1.9°C min⁻¹, resulting in melting curves. Data analysis was performed using SDS 2.2.1 software (Applied Biosystems). All amplification curves were analysed with a normalized reporter (Rn; the ratio of the fluorescence emission intensity of SYBR Green to the fluorescence signal of the passive reference dye) threshold of 0.2 to obtain the C_{t} values (threshold cycle). The reference control genes were measured with four replicates in each PCR run, and their average C_{t} was used for relative expression analyses. TF expression data were normalized by subtracting the mean reference gene C_{t} value from their C_{t} value (ΔC_{t}). The Fold Change value was calculated using the expression 2^{−ΔΔC_{t}}, where ΔΔC_{t} represents ΔC_{t}, condition of interest - ΔC_{t} control. The obtained results were transformed to log₂ scale.

**Dilution experiments**

Mixtures of root and shoot cDNA were prepared as described by Czechowski et al. [5] to give the ratios indicated in Figure 3. Primer pairs for two root-specific genes (Os03g55610, 5’-TACCCCCACCCAAACCATCAATCTG and 5’-TGAGAAAGAGCAAGGGCAGTGAG; Os08g38220; 5’-GGGGTTTCATCACTACCGATGA and 5’-TCACCATATCCACCGGCAAG) and one shoot-specific gene (Os12g8320, 5’-TGTTTTCTCCCTCGCTACG and 5’-TGCTGCTGGATCTGAGTCCAATT) were used in qRT-PCR experiments.

**Determination of reference gene expression stability**

To analyse the expression stability of the selected reference genes, the geNORM v.3.4 software was used as described by Vandesompele et al. [28]. The measured C_{t} values were transformed so the highest expression of each reference gene is equal to 1 and its expression in all other conditions is relative to this value (as suggested by the geNORM manual). The gene expression stability (M) was calculated and the most stable control genes were determined.

**Competing interests**

The author(s) declare that they have no competing interests.
Authors' contributions
CC performed the experimental work and drafted the manuscript. WRS helped to outline the project. BMR and SR designed and coordinated the project and wrote the manuscript. Overall supervision was provided by BMR.

Additional material

Additional file 1
Complete list of TF genes, primer data and corresponding PCR efficiencies. The TF primer platform was established based on the data for the gene models according to the Version 2.0 of TIGR Rice genome annotation. The corresponding new identifiers (Version 5.0) are also presented. The ranking of primer pairs taking into account experimental data (Cat column) was done as follows: 1, specific amplification; 2, no amplification; 3, non-specific amplification. 'Primer Location' indicates the sites within genes selected for primer design: J, at least one of the primers extends over an exon-exon junction; D, the primers are located in different exons; S, both primers are located within same exon.

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Additional file 2
cDNA samples used for the validation of the reference genes.

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