Quantitative real-time PCR based evaluation and validation of reference genes in *Gossypium arboreum*

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ABSTRACT

Estimation of gene expression levels plays a crucial role in understanding the function of the target gene(s). Inter-sample variance in gene expression can be more precisely measured if transcripts levels are accurately normalized. Normalization is pre-requisite step prior to the determination of candidate gene expression by qPCR. In this study conducted at ICAR-Central Institute for Cotton Research, Nagpur during 2015–16, six candidate reference genes, viz. actin4 (ACT4), actin7(ACT7), RNA Helicase (RNAH), Serine/threonine-protein phosphatase PP2A-1(PP2A1), ubiquitin7 (UBQ7) and α tubulin (αTUB) were systematically analysed for their expression patterns in different tissues pertaining to three development stages of cotton namely seedling, early reproductive and fiber development. The study has identified actin-4/actin-7/ubiquitin-7 as the most ideal reference genes for fiber development stages whereas actin-4/ ubiquitin-7 and actin-7/RNA helicases for seedling and early reproductive development stages, respectively. Validation of identified reference genes for relative expression analysis of Gacobl9, a COBRA-like protein, demonstrated their usefulness in qPCR analysis in *Gossypium arboreum*.

Key words: Cotton, Gene Expression, *Gossypium arboreum*, Reference gene, qPCR

Cotton, popularly known as white Gold, is one of the world's richest source of natural fiber and also one of the most important cash crops belonging to genus *Gossypium* (Abd El-Moghny et al. 2017). Besides being a major source of raw material for textile industry, having a single elongated epidermal cell, it acts as a molecular reference system for studying cellular differentiation and cell patterning in plants (Kim and Triplett 2001). In the era of functional genomics, the study of gene expression profile plays a crucial role in understanding the function of the candidate gene(s) and to decipher complex gene regulatory network in biological systems (Vandesompele et al. 2002). Gene chip array, RNA blot, quantitative real-time PCR (qPCR), semi-qPCR, etc. are being routinely used by molecular biologists for quantitative estimation of transcript levels in biological samples. However, qPCR exhibiting higher specificity and sensitivity with low sample volume requirement and amenability for high throughput applications remains undoubtedly the preferred and most widely used method (Czechowski et al. 2005). qPCR is also the most favoured choice for validation of large biological data generated through microarray, RNA-seq, and genotyping platforms (Jain et al. 2006). Despite being a robust and powerful technique, qPCR mediated gene expression profiling lacks accuracy if reliable and suitable reference genes are not used (Huggett et al. 2005). Normalization improves the precision of gene expression data by circumventing the limitations raised due to poor RNA quality and variability in reverse transcription & PCR efficiencies (Ermei et al. 2012). Ideally, an internal control must show invariant transcript levels both spatially and temporal across the experimental set-up (Marum et al. 2012). Housekeeping genes are generally considered to have constitutive expression as they are playing very crucial role in maintaining basic cells' surviving activities. However, several scientific studies have reported that these genes also showed variation in their expression may be due to their multiple roles in other cell's activities besides housekeeping functions (Czechowski et al. 2005). In addition to this, identified reference genes for one crop species may express inconsistently in another crop species even under identical experimental setup, thus negating their recurrent broad-spectrum use (Jain et al. 2006, Jian et al. 2006, Marum et al. 2012).
Reference genes have been systematically evaluated and validated for target gene expression normalization by using qPCR and statistical programs in many plant species under varied experimental conditions and developmental stages (Jian et al. 2008, Nicot et al. 2005, Reid et al. 2006, Hu et al. 2009). In cotton, scientific reports pertaining to selection and validation of reference genes are very limited and restricted with only teteraploid cotton *Gossypium hirsutum* (Tu et al. 2007, Artico et al. 2010, Wang et al. 2013, Raghavendra et al. 2014, Fausto et al. 2017). Thus, identified stably expressed and validated internal control genes in *Gossypium hirsutum* are generally in use for target gene expression analysis in *Gossypium arboresum*. Owing to the genomic complexity of the *Gossypium* spp. and availability of whole genome sequencing data containing uncharacterized gene sequences; it will be of immense value to systematic evaluation and validation of the reference genes in *G. arboresum*.

Through this study, six potential candidate internal control genes from *Gossypium arboresum* were statistically analyzed for their transcript expression stability to identify and validate the most suitable ones for target gene normalization by qPCR.

**MATERIALS AND METHODS**

**Plant material:** Total 11 samples each with three biological replicates were sourced from three different developmental stages (Seedling development, early reproductive stage and fiber developmental stage) of *G. arboresum* cv. Roja grown at ICAR-CICR, Nagpur during **kharif** 2015–16. Seedling development stage comprises of hypocotyl, cotyledonary leaves, and root tissues from 15 days old seedling raised on MS medium; early reproductive stage; consisted of sepal, petals, anthers and ovary from flowers at 0 Days Post Anthesis (DPA) and square at -3 DPA and samples set from fiber developmental stage comprising ovule with fibers at 3, 10 and 25 DPA, from plants grown in the greenhouse.

**RNA isolation and cDNA synthesis:** Total RNA was extracted using Spectrum™ Plant Total RNA Kit (Sigma-Aldrich St. Louis, MO, USA) by following the manufacturer’s instructions. Traces of genomic DNA were eliminated by On-Column DNasel digestion set (Sigma-Aldrich St. Louis, MO, USA). Total RNA was subjected to quality and quantity analysis using agarose gel electrophoresis and Qubit2.0 fluorometer (Invitrogen, USA) respectively. A total of 1µg of total RNA was used for first-strand cDNA synthesis using Superscript III Reverse Transcriptase kit (Life Technologies, USA).

**Primer design and qPCR analysis:** qPCR primers for selected six candidate housekeeping genes and Gacobl9 were synthesized from Integrated DNA Technologies (IDT) online tool (Supplementary Table 1). qPCR’s were performed using a Mx 3005 Real-Time PCR System (Agilent Technologies). Each qPCR reaction comprised a final volume of 25 µl containing 2 µl of 1:10 diluted cDNA, 0.5µl of each forward and reverse primer (0.4 µM), and 12.5 µl of 2x 2× Brilliant II SYBR Green PCR Master Mix, 0.375 µl of diluted reference dye and 11.125µl nuclease free water (Agilent Technologies). The following thermal cycling profile was used for all qPCRs: initial hot start at 95°C for 10 min, 35 cycles of 95°C for 30 sec, 60°C for 45 sec and 72°C for 30 seconds and one segment of disassociation/melt curve. Reliability of qPCR run was confirmed by including non-template controls along with total six replication comprising three biological and two technical replications per sample.

**Reference genes transcript data analysis:** The reference genes relative expression data were systematically evaluated with geNorm v 3.0, NormFinder and BestKeeper programs (Vandesompele et al. 2002, Andersen et al. 2004, Pfaffl et al. 2004). Each software is programmed with a unique algorithm and calculates gene stability measure by taking threshold cycle values (Cq values) either in raw or in normalized form using a delta Ct method. The geNorm program ranks

Supplementary Table 1 List of candidate reference genes, their primer sequences selected for evaluation of expression stability in cotton (*Gossypium arboresum* cv. Roja)

| Reference gene | Accession number | Primer sequence |
|---------------|-----------------|-----------------|
| RNAH          | XM_016862161.1  | F: GCCTGATCATGATGCGAGGAT R: CAGGAAGGTTTGGGCCATCTGGGA |
| UBQ7          | DQ116441.1      | F: AGACAAGGAGGACATCCACCTGGA R: GCTTGATCTTCTTGAGGGTTTGGA |
| ACT7          | NM_001327049.1  | F: TACAAAGAATCCGATGTGCTTCC C: R: TACCCAGGAAATCCAGCAACAAACTGG |
| αTUB          | EF151302.1      | F: GCTGAGGAAGGGTTACCAATGAGCCA R: GGTACATCAAACAGCAAGCCA |
| PP2A1         | DT545658.1      | F: GATCCTTGTGGAGGAGTGGGA R: GCGAAGACAGGTTGACGAGAT |
| ACT4          | AY305726.1      | F: TTTGACAGCCGATTAGGCAAGA R: ACTATTCGCCATGACACACTG |
| Gacobl9       | Cotton_A_03290* | F: CTGCCACCCATCTGATCTTAC R: GGCTGTCAAACCGGAGAATA |

*Niu et al. 2015*
the reference genes in an order of expression stability index-M value (lower the M value more the stability of expressed gene) derived based on repeated pairwise variance analysis. Besides this, a statistical parameter known as \( \text{Vn/Vn+1} \) variance is also calculated to identify the number of potential genes for more accurate expression normalization. The inclusion of additional gene becomes statistically insignificant if \( \text{Vn/Vn+1} \) variance scores below 0.15 cutoff. NormFinder algorithm workout both inter and intra-group transcript variance using repeated pairwise analysis by eliminating pseudo matched expression of candidate gene pair(s). Therefore, NormFinder analysis not only specifies the best individual internal control but also a pair of genes which may fall short in competence if analyzed individually. On the contrary, BestKeeper algorithm takes raw Ct values as input data and performs repeated correlation and regression analysis to find suitable reference gene(s).

**RESULTS AND DISCUSSION**

**Comparative expression data of reference genes:** The data for (quantification cycle value) Cq values depicting the initial transcript abundance were collected for each reference genes (Supplementary Table 2, Fig 1.) Cq values vary from 20.66 to 24.46, where \( \text{ACT4} \) showed highest transcript abundance (mean Cq value=20.66) across all samples followed by \( \text{g} \text{TUB}, \text{UBQ7}, \text{ACT7}, \text{RNAH}, \text{PP2A1} \) gene showed lowest transcript levels (mean Cq value=24.46). Interestingly, \( \text{ACT7} \) gene showed relatively stable expression profile with SD value 1.18 followed by 1.31, 1.64, 1.66, 1.84 and 2.91 for \( \text{PP2A1} \), \( \text{UBQ7} \), \( \text{ACT4} \), \( \text{RNAH} \), and \( \text{\alpha TUB} \), respectively.

**geNorm analysis:** geNorm analysis identified \( \text{ACT4/UBQ7} \) as most stable internal control gene pair (0.519) during the seedling developmental stage followed by \( \text{ACT7}, \text{PP2A1}, \text{RNAH} \) and \( \text{\alpha TUB} \) (Fig 2a). Similarly, \( \text{ACT4/ACT7} \) (1.37) and \( \text{UBQ7/ACT7} \) (0.64) were the most promising candidates for early reproductive and fiber development stages, respectively (Fig 2b, 2c). The results from the analysis also show that pairwise \( \text{Vn/Vn+1} \) variance is above 0.15 in all samples across the three developmental stages. It indicates that inclusion of next stably expressed genes are found to contribute significantly to the normalization of gene expression (Fig 3). The results from geNorm are summarised in Supplementary Table 3 and Fig 2.

**NormFinder analysis:** Using NormFinder, \( \text{ACT4} \) (0.139) emerge out to be the promising internal control gene in seedling and early reproductive stage samples, whereas \( \text{PP2A1} \) (0.192) followed by \( \text{ACT4} \) (0.312) in fiber development stages respectively (Supplementary Table 4).

However, comparative analysis between geNorm and NormFinder softwares shows minor differences in normalization suitability ranking of reference genes (Supplementary Table 5).

**BestKeeper analysis:** Using BestKeeper software, three (\( \text{UBQ7}, \text{ACT4} \), and \( \text{\alpha TUB} \)), four (\( \text{ACT7}, \text{RNAH}, \text{ACT4} \) and \( \text{\alpha TUB} \)) and two (\( \text{PP2A1} \) and \( \text{\alpha TUB} \)) candidate genes showed significant correlation at \( p \)-value <0.001 with the BestKeeper Index in samples of seedling, early reproductive and fiber development stages respectively (Supplementary Table 6). SD data showed that \( \text{UBQ7} \) (1.31) is the most consistently expressed internal control gene among samples of seedling developmental stages. Similarly, \( \text{ACT7} \) (1.1) and \( \text{ACT4} \) (0.83) were found suitable for early reproductive and fiber development stages respectively.

**Validation of reference genes:** Gacobl9, a cotton COBRA-like gene was chosen as a target gene for the purpose to validate the qPCR normalization potential

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**Supplementary Table 2** Expression levels of six reference genes in ten biological samples across three experimental sets of cotton (\textit{Gossypium} \textit{arboreum} Cv. Roja)

| Reference genes | Seedling stage mean Cq±SD* | Early reproductive stage mean Cq ±SD | Fiber development stage mean Cq ±SD | Across the developmental stages mean Cq ±SD |
|-----------------|-----------------------------|-------------------------------------|-----------------------------------|----------------------------------|
| RNAH            | 22.64±0.74                  | 24.9±1.71                           | 24.7±1.89                         | 24.3±1.84                        |
| \text{ACT7}     | 24.66±0.92                  | 24.5±1.1                            | 22.69±0.92                        | 24.16±1.18                       |
| \text{ATUB}     | 21.38±2.34                  | 21.67±3.4                           | 18.72±1.41                        | 20.84±2.91                       |
| \text{UBQ7}     | 21.34±1.31                  | 22.51±1.59                          | 21.34±1.03                        | 21.8±1.64                        |
| \text{ACT4}     | 20.05±1.39                  | 21.3±1.94                           | 20.04±0.83                        | 20.66±1.66                       |
| \text{PP2A1}    | 24.26±0.71                  | 25.05±1.67                          | 23.26±0.75                        | 24.46±1.31                       |

*Geometric mean of Ct values observed across all biological samples along with Standard Deviation (SD) value
of identified reference genes. Relative expression of the Gacobl9 during secondary cell developmental stages of cotton fibre (25 DPA) in relation to 10 DPA was quantified using ACT4, ACT7, and UBQ7 as normaliser. We found the higher transcript abundance of target gene and it was in descending order for the normaliser ACT4 (76.49), ACT7 (39.05) and UBQ7 (16.32) reference genes (Fig 4).

Allotetraploid Gossypium species including G. hirsutum (AD1) and G. barbadense (AD2) are the descendants of two diploid species G. arboreum (A genome) and G. raimondii (D genome), resulted from genome union between A and D genomes and subsequent chromosome doubling (Yu-xiang et al. 2013). Of late, successful whole genome sequencing of cultivated diploid progenitor species has generated immense genomic data for exploration (Wang et al. 2012, Li et al. 2014). Whole genome sequencing is an important and fundamental step for incisive scientific studies of wide dimensions, provide a valuable insights, vital for linkage relationship studies, identification of candidate genes, development of DNA based markers which can be
used in crop improvement either through plant breeding or through transgenic approach. With the recent and rapid developments in sequencing technologies, the time and cost aspects of sequencing have seen a great retrenchment. Having the sequence information of whole genome is just the beginning of long worthy journey in science. 

*Gossypium arboreum* (Asiatic cotton) is well known for its tolerance to various climatic resilient features like biotic and abiotic stresses. With basic alphabets of *G. arboreum* genome decoded, assigning the function to the genome polymorphisms revealed in sequencing is most imperative and highly challenging task. Until and unless this sequence is converted to knowledge, the complete benefits of genome sequencing cannot be attained. The *G. arboreum* sequence information lays a strong platform for next generation genomics assisted desi cotton improvement. Hence, in present study, a total of six candidate reference genes were selected in *G. arboreum* based on their DNA sequence similarity to previously validated reference genes in *G. hirsutum* (Artico et al. 2010, Niu et al. 2015). A similar trend of candidate gene validation and selection is widely employed in many crop species (Reid et al. 2006, Wang et al. 2013). Three software packages were used for analysis and identification of suitable reference gene. The inherent differences in algorithms of these softwares resulted in variation in the ranking order of reference genes. For example, in our study, the BestKeeper outputs *ubiquitin7* gene, as most stable in seedling samples (Supplementary Table 6) while it ranked second by NormFinder (Supplementary Table 4). Likewise, *PP2A1* emerged as ideal gene during fiber development stage in NormFinder (Supplementary Table 4) while it was ranked four by geNorm (Supplementary Table 3). Similar results were also evidenced in other plant species (Chen et al. 2015, Wan et al. 2017). Our study revealed that *ACT4* and *ACT7* are the best combinations of internal control genes in all samples of three developmental stages (Supplementary Table 4). Actin filaments constitute a major

**Supplementary Table 4** Reference genes expression stability values calculated by the NormFinder software

| Seedling stage | Early reproductive stage | Fiber development stage | Across the developmental stages |
|----------------|--------------------------|-------------------------|-------------------------------|
| Ranking        | Stability value           | Ranking                 | Stability value               | Ranking | Stability value |
| ACT4           | 0.139                    | ACT4                    | 0.127                        | ACT4    | 0.311           |
| UBQ7           | 0.17                     | ACT7                    | 0.276                        | ACT4    | 0.344           |
| PP2A1          | 0.393                    | UBQ7                    | 0.366                        | UBQ7    | 0.469           |
| ACT7           | 0.482                    | PP2A1                   | 0.438                        | ACT7    | 0.75            |
| RNAH           | 0.491                    | RHAH                    | 0.46                         | αTUB    | 1.168           |
| αTUB           | 0.56                     | αTUB                    | 0.474                        | RHAH    | 1.182           |
| Best combination | Stability value           | Best combination       | Stability value               | Best combination       | Stability value |
| ACT4 &         | 0.118                    | RHAH & ACT7             | 0.187                        | ACT7 &  | 0.298           |
| UBQ7           |                          |                         |                               | ACT4    | 0.043           |

Stability values are listed from the most stable to the least stable.

**Supplementary Table 5** Best combination of reference genes based on geNorm and NormFinder programs

| Seedling stage | Early reproductive stage | Fiber development stage | Across the developmental stages |
|----------------|--------------------------|-------------------------|-------------------------------|
| ACT4           | ACT7                     | ACT4                    | ACT4                          |
| UBQ7           | UBQ7                     | ACT7                    | ACT7                          |
| RHAH           | ACT7                     | UBQ7                    |                               |

Stability values are listed from the most stable to the least stable.
component of plant cell’s cytoskeleton and are principle determinants of its shape and movements. Actin gene variants are traditionally being used and also reported as ideal internal control gene for qPCR in various plant species under different experimental conditions (Artico et al. 2010, Wan et al. 2017, Zhang et al. 2015). In addition, ACT7 along with RNAH was found to be best for early reproductive development stage (Supplementary Table 5) which is also supported by BestKeeper analysis (Supplementary Table 6). During data analysis, we have observed that gene RNAH showed relative stable expression in early reproductive stage while poor and varied expression levels were observed in seedling and fiber development stages (Supplementary Table 5) suggesting tissue or stage-specific selection of normalizer gene. Similarly, the weaker expression of most commonly used housekeeping gene such as αTUB was also noticed (Fig 2, Supplementary Table 3 and 4). In congruence with our results, varied transcript abundance of tubulin as normalizer was also reported in crops like potato, grapes, and soybean (Jain et al. 2006, Nicot et al. 2005, Reid et al. 2006). Apart from actin, ubiquitin (UBQ7) was also emerged as one of the suitable internal control gene for samples representing seedling stage of cotton (Supplementary Table 5). Ubiquitin's are classical enzymes play a very important role in post-translation modification of proteins. Stably expressed ubiquitin isoforms have also been identified and widely used (Ermei et al. 2012, Zhang et al. 2015). Similarly, Artico et al. (2010) identified and reported ubq14...
has suitable housekeeping gene for all the tissues including plant organs and flower buds of *Gossypium hirsutum*.

Overall, the study has identified *actin4/actin7/ubiquitin7* as the most suitable reference genes for fiber development stages, whereas *actin4/ ubiquitin7* and *actin7/RNA helicases* for seeding and early reproductive development stages, respectively. To validate the suitability of the identified internal control genes from the above study, tissue specific expression analysis of gene for COBRA like protein (*Gacobl9*) was performed. *Gacobl9* gene codes for a plant-specific glycosylphosphatidyl inositol (GPI) anchored protein and is reported to be expressed predominantly during secondary cell wall deposition stage of cotton fiber development (Niu et al. 2015). COBRA-like protein plays a key regulatory role in cellular microfibrillar orientation (Roudier et al. 2005). The massive accumulation of cellulose during secondary cell wall synthesis stage (25 DPA) of cotton fiber development necessitates the recruitment of the diverse group of proteins for the microfibrillar arrangement.

Relative expression of the *Gacobl9* during 25 DPA in relation to 10 DPA was quantified using *ACT4*, *ACT7*, and *UBQ7* as normaliser. Its expression during 25 DPA was found to be higher in comparison to 10 DPA. However, the relative transcript abundance of the *Gacobl9* gene was observed to be in descending order for *ACT4*, *ACT7*, and *UBQ7* which was in concurrence with their relative expression stability. The results obtained from our study are in line with the validation of identified reference genes reported elsewhere (Hu et al. 2009, Artico et al. 2010, Lin et al. 2014). In conclusion, our study identified and validated *actin4, actin7*, and *ubiquitin7* for their utilization as normalizers for gene expression studies in *Gossypium arboreum*. However, to avoid misleading outputs, a set of normalization genes should always be carefully evaluated and validated under specific experimental conditions.

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