Genome-wide investigation of CCCH zinc finger family in longan (Dimocarpus longan Lour): characteristic identification and expression profiles in longan somatic embryo

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Liyao Su
Fujian Agriculture and Forestry University

Mengqi Jiang
Fujian Agriculture and Forestry University

Shuqi Huang
Fujian Agriculture and Forestry University

Xiaodong Xue
Fujian Agriculture and Forestry University

Xue Li
Fujian Agriculture and Forestry University

Zhongxiong Lai
Fujian Agriculture and Forestry University

Yuling Lin
Fujian Agriculture and Forestry University
Corresponding Author

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Abstract

Background: CCCH Zinc finger (Znf) transcription factors (TF), as a novel type Znf genes, regulate genes expression by binding on their mRNA and play important roles in plant abiotic stress, growth and development. However, no overall genome-wide analysis or expression profiling of CCCH (C3H) gene family in Dimocarpous longan, especially during the early stages of somatic embryo in longan has been studied. Longan is a tropical/subtropical fruit tree of great economic importance in Southeast Asia, and longan embryogenesis is the main factor affecting fruit quality and yield.

Results: In this study, a comprehensive analysis of longan C3H → DIC3H →gene family was carried out. 49 DIC3H genes were identified from longan genome database, which divided into 3 clades. Besides, genes characteristics, phylogenetic tree, gene structure, motif composition were comprehensively analyzed. The analysis of alternative splicing events (AS) suggested that AS events of DIC3H genes were related to longan non-embryonic and embryonic callus transformation. Promoter analysis indicted that most of DIC3H genes included cis-elements associated with hormones and stress response. Quantitative real-time PCR analysis indicated that 26 DIC3Hs, which possess MeJA and ABA responsive cis-elements, showed different expression patterns and may involved into ABA and MeJA signaling pathway. The expression profiles of 17 DIC3Hs were performed in four stages of longan, the results showed that only DIC3H01/07/14/16/38 was consistent with the data in the transcriptome. DIC3H 07/14/16/36/49 were highly expressed in EC and only DIC3H 04/38 was in GE, suggesting that they have different functions in embryonic development. Finally, sRNAs were verified involved into regulating 6 DIC3Hs.

Conclusion: This study provides the first systematic analysis of CCCH protein in longan somatic embryo. Particularly, CCCH genes may be involved in hormone and stress respond, and somatic embryogenesis. Our results presented here may provide a insight
into the characteristics and functions of this family in somatic embryogenesis.

Background

Transcription factors (TFs), as a gene widely distributed in plants, play an important role in the growth and development of plants and morphogenesis[1, 2]. Zinc finger (Znf) transcription factors is one of the largest TF families containing RING-finger[3], LIM[4], WRKY[5] and DOF[6] gene families which regulate gene expression through DNA-binding and protein binding proteins. However, recent evidence suggests that CCCH (C3H) as a novel type Znf transcription factors regulate gene expression by binding on targets genes mRNA[7-9]. To verify its characteristics, based on the genome-data, the comprehensive analysis of C3H Znf family was performed in dicotyledonous Arabidopsis thaliana[9], Medicago truncatula[10], Populus trichocarpa[11], Clementine mandarin[12], Vitis vinifera[13], Cicer arietinum[14], monocotyledonous Musa acuminata[15], Zea mays[16], and Oryza sativa[9], showed that C3H Znf family widely involved in biotic and abiotic pathways. Besides, C3H Znf members have various functions. In Arabidopsis, ATSZF1 and ATSZF2 negatively regulate the expression of salt responsive genes[17]. Cotton GHZFP1 can be induced by salt, drought and SA, and its transgenic tobacco shows resistance to fungal diseases[18]. Rice OsDOS controls leaf senescence through jasmonate (JA) pathway[19]. Arabidopsis HUA1 is confirmed a regulator for flower morphogenesis[20]. Moreover, C3H Znf genes is an essential regulator for plants somatic embryogenesis. Previous studies have reported that PEI1, as an embryo-specific expression C3H Znf gene, directly regulate the heart-shaped embryo development in Arabidopsis[21]. Cucumber CsSEF1 shows the importance of controlling cell polarity, and marks the cotyledon primordia and procambium tissues in later developmental stages[22]. The above studies indicate that C3H Znf family plays a significant role in plant abiotic stress, growth and development and somatic embryo morphogenesis.
Sapindaceae plants are widely distributed in tropical and subtropical areas, including important tropical fruit trees such as longan, lychee, *Nephelium lappaceum*, and well-known oil plants such as *Sapindus mukurossi*, *Xanthoceras sorbilifolium*. To date, many draft genome sequence of plants have been identified, which has greatly promoted the research of corresponding plants. Longan, as the first Sapindaceae plant completing the genome sequencing[23], provides a reference for studying the molecular genetic characteristics of Sapindaceae plants. So far, comprehensive analysis of gene family in Sapindaceae plants is still limited. Only longan WRKY[24], Ubiquitin-conjugating enzymes[25], Laccase[26] families have been comprehensively analyzed. As an important fruit tree, fruit quality of longan is closely related to economic effects. Embryonic development as one of the main factors regulate longan fruit quality. Therefore, understanding the mechanism of longan embryonic development is critical for improving longan fruit quality.

Despite the *CCCH* Znf gene is great significance in plants, the comprehensive analysis of *CCCH* gene family in plants embryo has not been performed. Longan genome sequencing provides an opportunity to reveal the function at the genome-wide level[23]. The 49 *DIC3Hs* were identified from longan genome database. We further analyzed the gene characteristics, phylogenetic tree, gene structure, motif composition, alternative splicing events and promoter cis-elements. Additionally, the expression profiles of 26 *DIC3Hs* were carried out by RT-qPCR to explore their responses to methyl jasmonate (MeJA), abscisic acid (ABA) and their endogenous inhibitor (Salicylhydroxamic acid, SHAM, Sodium Tungstate Dihydrate, STD) treatment. According to the transcriptome data, 17 *DIC3Hs* were selected to analyze the expression levels in longan embryogenic callus[EC], incomplete compact pro-embryogenic cultures[ICpEC], globular embryos[GE], non-embryonic callus[NEC] and their cleavage sites were verified. Our preliminary results might provide valuable clues for researching the function of the *CCCH* Znf gene family in
plant embryonic development.

Results

Analysis the characteristics of longan C3H gene family

According to annotation files of InterPro software, the 68 candidate longan CCCH Znf family members were found in longan genome database. Then, the BLASTP program and CD search were performed. A total 49 non-redundant CCCH Znf genes were confirmed in longan, then we named them DIC3H01 to DIC3H49. Gene characteristics, including the Arabidopsis orthologs locus, number of exons, length of CDS, molecular weight (kD), isoelectric point (PI), number of CCCH motif and subcellular localization were showed in Table 1. Among the 49 DIC3H genes, the DIC3H41 was identified to be the smallest protein with 136 amino acid, whereas the DIC3H27 was largest with 1811 amino acid. The number exons of the genes range from 1 to 14, the kD range from 14.46 (DIC3H41) to 198.20 (DIC3H27), and the PI range from 4.90 (DIC3H33) to 9.50 (DIC3H28). In addition, the number of CCCH motif of DIC3H gene family was the same as that in Arabidopsis and rice, which was range from 1 to 6. Finally, the subcellular location showed that 9 of DIC3H members were located in cytoplasm, 35 DIC3Hs members were located in nucleus, and the rest was secreted protein.

Phylogenetic analysis and conserved motif multiple sequence alignment

A phylogenetic tree of longan and Arabidopsis was constructed by maximum likelihood (ML) method based the full length of protein sequence. The phylogenetic analysis showed that DIC3Hs and AtC3Hs gene family was divided into 3 clades contained 21, 39 and 55 members, respectively (figure 1). DIC3Hs had 9, 19 and 21 members in each of the three clades. In the first clade, four AtC3Hs members were not classified, and all members of the longan were classified. These results indicted that DIC3Hs had three different
evolutionary directions. Such as AtC3H51 (PEI1), as a key protein for plant embryogenesis, was clustered with DIC3H01, speculated that they had similar function. The longan CCCH Znf domains were further multiple aligned according to the phylogenetic tree. The AtC3Hs (AtC3H01, AtC3H51, AtC3H08) in each clades were selected for a representatives. The results showed that longan CCCH Znf domain sequences were highly conserved in each clades with the length range from 19 to 27 amino acids (figure 2). And it basically belonged to C-X\textsubscript{8}C-X\textsubscript{5}C-X\textsubscript{3}H and C-X\textsubscript{7}C-X\textsubscript{5}C-X\textsubscript{3}H types, suggesting that these two types were parallel evolutionary. Besides, the conservation of clade II was the worst. There are three different types domain in clade II belonging to DIC3H15-1 (C-X\textsubscript{9}C-X\textsubscript{5}C-X\textsubscript{3}H), DIC3H21 (C-X\textsubscript{7}C-X\textsubscript{4}C-X\textsubscript{3}H) and DIC3H25 (C-X\textsubscript{9}C-X\textsubscript{4}C-X\textsubscript{3}H).

**Gene structure and motif composition of DIC3Hs**

The introns and exons of all 49 DIC3Hs were identified for better understanding the evolution of DIC3Hs. As shown in figure 3B, among the 49 DIC3Hs, the number of exons were range from 1 to 14 (eight with one exons, seven with two exons, six with three and four exons, two with five exons, one with six exons, nine with seven exons, three with eight exons, one with nine exons, three with ten exons, one with 11 exons, one with 12 exons and one with 14 exons). In the same class, genes usually had the same structure, such as class I, except DIC3H08, they all contained one intron. All class IIe/f members had no intron, except DIC3H26/15. Besides, within the same class, the intron structure were highly consistent. Although the gene structure and the introns phase were similar with phylogenetic relationship, the different between classes were significant.

The conserve motif was identified by CDD. Comparing the previous researches, the motif found in longan C3H family was the most containing 25 types (figure 3C). Only one C3H domain was observed in 16 DIC3Hs. The rest genes all possessed 2 to 5 domains. The
cluster genes (\textit{DIC3H23/27}, \textit{DIC3H13/20}, \textit{DIC3H45/10}) had consistent motif composition indicating functional similarity in longan. In addition, some motifs were unique to one group, for example, motif 6, motif 7 and motif 23 were special to class \textsc{III}d, \textsc{II}a and \textsc{II}f, respectively. The differentiation of motifs between different members reflected the functional diversification of \textit{DIC3Hs}, and the function of motifs needed further verification. Overall, \textit{DIC3Hs} members consisting of the same gene structure and motif composition were clustered into one branch of phylogenetic tree implying it’s highly conserved.

Analysis the AS events of \textit{DIC3Hs} in longan non-embryonic and embryonic cultures

According to the RNA-seq analysis of longan NEC, EC, ICpEC and GE, the alternative splicing events of \textit{DIC3Hs} were identified. A total of 445 AS events, including alternative 3' splice site (A3'S), alternative 5' splice site (A5'S), intron retaintion (IR) and exon skipping (ES), were detected from 29 \textit{DIC3Hs}. The type of AS event and the statistics of AS events in 29 \textit{DIC3Hs} was showed in Table 2. A3'S events (26.17%) were more frequent than A5'S events (18.30%). IR events were the most frequent with 45.17% (Table 2). This result was the same with previous studies which considered IR events were the most frequent events of AS in plants. Furthermore, the number of genes that with A3’S, A5’S and IR events was basically the same (Table 2). In addition, as the figure 4A shown, AS events might play a key role in longan somatic embryo morphological. For example, in EC stage, IR event sharp decrease and A3’S/A5’S marked increase. The ES event slight rise in ICpEC and GE stages. Meantime, we counted the number of AS events in longan NEC, EC, ICpEC and GE. The results shown that the AS events occur most frequently in the NEC stage and least frequently in the EC stage. (figure 4B). This result suggested that the AS events in \textit{DIC3Hs} was related with longan somatic embryogenesis.
Stress and hormone related *cis*-elements in *DIC3Hs* promoter

To further explore the potential regulatory mechanism of *DIC3Hs* during external stress, the promoters regions, which were up-stream 2Kb sequences of *DIC3H* genes translation starts site, were submitted into PlantCARE database to search *cis*-elements. A total of 559 *cis*-elements related to hormone and stress were detected in *DIC3H* genes (Figure 7). Among them, except *DIC3H07, DIC3H40* and *DIC3H49*, the rest genes contained at least 1 anaerobic induction element. Meanwhile, drought and low-temperature related *cis*-elements possessed in 25 and 16 *DIC3Hs*, respectively. This result showed that *DIC3H* family might response these abiotic stress. In addition, 36 *DIC3H* genes contained 164 MeJA responsive *cis*-elements and 31 *DIC3Hs* possessed 88 abscisic acid responsive element indicating that MeJA and ABA play a key role in *DIC3Hs* regulatory. Furthermore, 34 auxin-responsive elements existed in 20 *DIC3Hs* and 38 gibberellin-responsive elements were found in 23 *DIC3Hs*. 29 salicylic acid responsive element was located in 20 *DIC3Hs*. On the whole, the *cis*-element analysis suggested that *DIC3Hs* family could involved into abiotic stress and hormone responsive.

Expression patterns of *DIC3H* genes after ABA, MeJA and theirs endogenous inhibitor treatments

According to the potential *cis*-elements analysis above, 26 *DIC3H* members, which possessed MeJA and ABA responsive *cis*-element, were selected from 49 *DIC3H* genes. The qPCR was performed to analyze their expression patterns after the identical concentration of MeJA, ABA and their endogenous inhibitor treatments. In ABA treatment, among the 26 *DIC3Hs*, 10 were up-regulated, 8 were down regulated and 8 *DIC3Hs* were no changed (Figure 6). STD was the inhibitor of endogenous ABA. In STD treatment, among the 26 *DIC3Hs*, 4 were up-regulated, 13 were down regulated and 9 were no changed (Figure 6).
Some of DIC3Hs showed the opposite trends in ABA and STD treatment, such as DIC3H10/24/28/37/45/46 (Figure 6). Most of DIC3Hs signal significantly up-regulated responded MeJA. However, in SHAM treatment, the expression of DIC3Hs was almost invariant compared the control. In addition, several DIC3Hs (DIC3H09/24/26/28/30/33/37/46) were up-regulated in MeJA treatment, and down-regulated in SHAM treatment (Figure 7). This results implying that DIC3Hs were involved into ABA and MeJA signaling pathway.

Expression profiling of DIC3Hs with RNA-seq and qPCR in longan non-embryonic and embryonic cultures

The expression patterns of longan CCCH family in the longan NEC, EC, ICpEC and GE transcriptomes were investigated in this study (Transcriptome datas of DIC3H02, DIC3H08, DIC3H28, DIC3H29, DIC3H30 and DIC3H32 were absent.). As the figure 8 showed that the expression of 43 DIC3Hs was divided into 2 group. In the group I, they were at a low expression levels in NEC stage, and high expression between EC stage and GE stage indicating that these genes were related to embryonic of longan somatic embryo. Moreover, 3 DIC3H genes were specific in GE stage and 2 DIC3H genes in ICpEC stage. 12 DIC3Hs highly expressed in NEC and EC stage, which were clustered at group II. This results implied that these genes which highly expressed in specific stage might involved into their morphogenesis.

To further confirm whether the specific expression of DIC3Hs could regulate longan somatic embryo morphogenesis of specific stage, 17 DIC3Hs which highly expressed in a special stage were selected to study. Then, the qPCR was carried out to verify the expression patterns of these DIC3Hs in longan early SE. The results showed that only DIC3H01/07/14/16/38 was consistent with the data in the transcriptome. DIC3H05,
DIC3H31, DIC3H39, DIC3H43 and DIC3H47 were down regulated during longan SE, and DIC3H38 and DIC3H41 showed the reverse trend, suggesting that members of the DIC3Hs gene family may have different functions in embryonic development. Whilst, 6 DIC3Hs (DIC3H07/11/14/16/36/49) were highly expressed in EC, and there were lower expression level of most DIC3Hs in ICpEC and GE than NEC and EC (Figure 9).

Small RNA involved into DIC3Hs transcription

Small RNAs played an important role in plant growth and development. These regulatory small RNAs (mainly include miRNAs and ta-siRNAs, sic passim) negatively regulate gene expression at post-transcriptional level by directing the cleavage of target transcript (mRNA)[30]. Li Yiqun reported that the MulZF1 which is a zinc finger protein containing CCCH domain is the target gene of mul-miRn26 in Morus alba L[31]. To understand whether the DIC3Hs were regulated by sRNA in longan, the modified RLM-RACE was carried out to verified the cleavage site of 17 DIC3Hs which highly expressed in a special stage. As the figure 10 shown, among the 17 DIC3Hs, the fragments of 6 DIC3Hs (DIC3H01/03/05/11/19/39) were detected. The 6 DIC3Hs had 1 to 5 cleavage sites.

Meantime, the longan small RNA (sRNA) database was used to predict the potential sRNA that could cleaved the 6 DIC3Hs. As the results shown, the 14 cleavage sites of 6 DIC3Hs were identified as the putative cleavage site for 131 sRNAs (Figure 10, Additional file 2 to 7). This implied that sRNAs could widely involve into DIC3Hs pathway. For example, each of three cleavage sites of DIC3H01 could be combined with 4, 5 and 17 sRNA, respectively. Among these sRNA, 21 sRNA had been registered in miRBase database. It is suggested that miRNA could regulate DIC3Hs in longan somatic embryogenesis. Meantime, a larger number animal origin miRNAs were found in these sRNAs, indicating that the C3H family might conserved between plant and animal in terms of the formation principle of miRNA. Furthermore, the rest 5 sRNA had no similar in miRBase database and their had a reliable
E value (one with 1.5, one with 2.5, three with 3.0). Thus, we speculated that they might be siRNA or piRNA.

Discussion

Evolutionary conservation and divergence of the DIC3H gene family of longan

In plants, there are many studies about CCCH Znf family. Because of lacking draft genome sequence, CCCH Znf family has not been reported in Sapindaceae plants. The publication of longan draft genome sequence provides a solid foundation for our study[23]. We identify 49 CCCH Znf genes from longan genome, the number of is less than that of Arabidopsis (68)[9], rice (69)[9], maize (68)[16], Vitis vinifera (69)[13], Clementine mandarin (62)[12], more than Medicago truncatula (34)[10]. The size of longan genome is 445 Mb[23], which is greater than Arabidopsis (125 Mb)[32], rice (375 Mb)[33], Clementine mandarin (367 Mb)[34], smaller than Vitis vinifera (125 Mb)[35], maize (2300 Mb)[36], Medicago truncatula (500 Mb)[37]. This results show that the number of plants CCCH Znf genes is may independent with the plants genome size, implying that it is related to species evolution. In addition, the C-X$_7$-C-X$_5$-C-X$_3$-H types of CCCH Znf domain is conserved in plants. In longan, it occupies 96.62%, which outclass Populus trichocarpa (72%)[11], Arabidopsis (83%)[9], Vitis vinifera (79.8%)[13], maize (79.4%)[16] and Cicer arietinum (82.3%)[12]. This indicates that longan CCCH Znf domain is highly conserved. Furthermore, there are three nonconservative CCCH Znf domains in longan. Among them, C-X$_9$-C-X$_5$-C-X$_3$-H and C-X$_7$-C-X$_4$-C-X$_3$-H domain have been reported in Musa acuminata[15], maize[16], rice[9], Arabidopsis[9]. The domain of C-X$_9$-C-X$_4$-C-X$_3$-H type is first identified in longan speculated that DIC3H25 has Special biological functions and
regulatory pathways.

We further study the gene structure and motif composition of longan CCCH Znf family. DIC3Hs within the same groups share similar intron/exon composition, intron phase and conserved domain. This suggest that longan CCCH Znf members are functionally conservative during the evolution. In addition, there are a larger number of conserved domain except CCCH Znf domain, which is a functional region of the protein. 15 conserved domains are identified in *Populus trichocarpa*[11], 13 in *Arabidopsis*[9], 8 in *Medicago truncatula*[10], 6 in *Vitis vinifera*[13], 10 in maize[16], 16 in *Clementine mandarin*[12].

There are 25 conserved domains are found in longan implying functions diversification of longan CCCH Znf family. The above plants share many domains such as ANK, RRM, WD-40, KH, etc, which play important roles in various life activities such as particle transport, signal transduction, RNA/DNA recognition, RNA binding and protein interaction[38-42], suggesting the similarity and importance of CCCH genes function among different plants.

Meanwhile, longan CCCH Znf members have a lot specific conserved domains for example Torus super family, PRK12678 super family, DNA_pol3_delta2 super family, etc. The results show that longan CCCH Znf family may involved in a wider range of life activities.

**DIC3H** genes may response to plant stress and hormone response

Promoter, as a non-coding region upstream of coding gene, plays a key role in regulating gene expression. In this study, identification of a large number of cis-acting elements associated with biotic and abiotic stresses, including MeJA (29.34%), ABA (15.20%), SA (5.19%), Auxin (6.10%), GA (6.80%), drought (6.40%), anaerobic induction (27.01%) and low-temperature responsive elements. It is suggested that the longan CCCH Znf family may involve in these signaling pathways. In MeJA and ABA treatment, longan CCCH Znf members can increase or decrease the expression in response to hormone signals. This results are similar with previous research in *Arabidopsis*[9][43], *Populus trichocarpa*[11],
maize[16] and *Medicago truncatula*[10]. Although many studies have shown that CCCH Znf can respond to exogenous hormone signals, the effects of endogenous hormones on its expression remain unknown. In endogenous inhibitor of MeJA and ABA treatment, the effect of endogenous hormone inhibitors on some *DIC3Hs* are more obvious than exogenous hormones. Besides, some *DIC3Hs* have opposite expression trends to hormone and their endogenous inhibitors, such as *DIC3H10/24/28/37/45/46* to ABA and *STD/DIC3H09/24/26/28/30/33/37/46* to MeJA and SHAM. These results demonstrate the importance of the CCCH Znf transcription factor family in plant stress and hormone response.

**Potential roles of DIC3H genes during plants somatic embryo**

Comprehensive analysis of the *CCCH* Znf family has not been reported in plants somatic embryo. Combined with longan transcriptome data and RT-qPCR analysis, only *DIC3H01/07/14/16/38* was consistent with the data in the transcriptome. *DIC3H07/11/14/16/19/36//38/49* were highly expressed in single stage suggesting that this members can participate in specific stage of longan somatic embryo morphogenesis. Most of them are highly expressed in EC, indicating that these *DIC3Hs* play an important role in the formation of longan embryonic cells. Some *DIC3Hs* up-regulated (*DIC3H38/41*) or down-regulated (*DIC3H05/31/39/43/47*) during NEC to GE indicate that up- or down-regulation of these genes can promote the formation and differentiation of longan somatic cells. These result are similar to the function of *AtPEI1*[21] and *CsSEF1*[22] in somatic embryogenesis. Moreover, sRNA is an endogenous non-coding small molecule regulator, and many studies have shown that sRNA is of importance role to longan somatic embryogenesis[44-46]. While *DIC3Hs* is involved in plant growth and stress response, it is also regulated by sRNA. 14 cleavage sites of 6 *DIC3Hs* may be regulated by 131 presumed sRNA. Moreover, a large number of animal-derived sRNA also reflects the conservatism of
CCCH Znf family in the animals and plants.

Conclusions
In conclusion, 49 DIC3H genes were identified in the longan genome, which divided into 3 clades. The results of a comprehensive analysis demonstrate the importance of CCCH zinc finger genes in the regulation of plant somatic differentiation and in response to hormone and stresses. A systematic and comprehensive analysis of longan CCCH Znf family is conducive to further screening DIC3Hs for functional identification, as well as to improving longan fruit quality and enhancing genetic improvement against stress.

Methods

Plant materials and treatments
The ‘HHZ’ longan friable-embryogenic callus (EC) that preserved by Institute of Horticultural Biotechnology, fujian agriculture and forestry university was used in this experiment. For hormone treatments, the EC was cultivated in liquid MS with 20 g/L sucrose and 50 mg/L MeJA, ABA and theirs endogenous inhibitor (Salicylhydroxamic acid, SHAM, Sodium Tungstate Dihydrate, STD) for 24h. The EC, ICpEC (incomplete compact pro-embryogenic cultures), GE (globular embryos) were induced in solid MS with 1.0 mg/L, 0.5mg/L and 0.1mg/L 2,4-D, respectively. The NEC (non-embryonic callus) was induced from longan mature embryo in solid MS medium.

Identification the longan CCCH Znf family
The longan protein-coding DNA sequences and protein sequences were download from GigaScience Database (http://dx.doi.org/10.5524/100276). The annotation files of InterPro software of all longan genes downloaded in longan genome database. The 68 genes corresponding to IPR000571 (Zinc finger, CCCH-type) was obtained. Then, all obtained
genes was further verified by NCBI BLASTP (Basic Local Alignment Search Tool Protein) and CD Search (Conserved Domain Search Service). So far, 49 DIC3Hs were identified from longan genome database, and named from DIC3H01 to DIC3H49 according to their position on pseudo molecules.

**Gene sequence characteristics analyze**

The ExPasy website (https://web.expasy.org/protparam/) was used to identify the length of sequence, molecular weight and isoelectric points of longan CCCH protein members. The local BLAST was performed to prediction homologous gene of longan CCCH genes in Arabidopsis genome database. In addition, subcellular location of longan CCCH protein members was predicted by LocTrees3 (https://rostlab.org/services/loctree3/). The number of exon and CCCH Znf domain were obtain from longan genome database and NCBI conserved domain search (https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi). The data of Alternative splicing events of 49 DIC3Hs were extracted from longan non-embryonic callus, embryonic callus, incomplete compact pro-embryogenic cultures and globular embryo transcriptome SRA050205.

**Phylogenetic analysis and multiple alignment of CCCH domain**

The CCCH protein sequences of Arabidopsis were downloaded from PlantTF database (http://planttfdb.cbi.pku.edu.cn/). All acquired protein sequences were aligned and constructed ML (maximum likelihood) phylogenetic tree by BioEdit software with default parameters and 1000 bootstrap. The conserved domain amino sequence of longan CCCH members and selected Arabidopsis member were aligned by GeneDoc.

**The cis-elements analysis of DIC3Hs' promoters**

The up-stream sequences (2K) of DIC3Hs CDS (coding sequences) were obtained by TBtools Gtf/Gff3 sequence extractor. Then we deleted the base N found in the promoter of
DIC3H01, DIC3H12, DIC3H28 and DIC3H30. Next, the sequences were submitted to PlantCARE database to predict cis-elements.

RNA extraction and expression levels analyses of DIC3Hs

Total RNA was extracted by Trzol Reagent kit according to the protocol. The cDNA for quantitative PCR was synthesized by using PrimerScript RT Reagen Kit (Takara). Quantitative PCR was preformed with Roche LightCyclers 480 instrument using SYBR Prumix EX Taq™ II (Takara). The 20 µL qPCR reaction was carried out containing 10 µL SYBR Prumix EX Taq™ II, ddH₂O 6.4 µL, 1 µL each primer (10 µM), cDNA 2 µL. To acquire reliable results, three biological repeats and three technical repeats were performed. The reference genes FSD, EF-1α and EIF-4α[27, 28] were used as the internal control. We obtained the relative expression of DIC3Hs according to the 2^ΔCt method, and results were shown as mean and standard deviation (SD). All the primers used in this study were listed in additional file 1.

Small RNA cleaved verification of a part of DIC3Hs

Small RNAs can regulate gene expression by directing the cleavage of target transcript. To understand whether the DIC3Hs were regulated by sRNA in longan, 17 DIC3Hs which highly expressed in a special stage were choosed to verified the cleavage site. The mixture of longan EC, ICpEC and GE total RNA was used to synthesized the cDNA for modified RLM-RACE followed the GeneRacer™ Kit instruction. Using DNAMAN, two gene special primers were designed for modified RLM-RACE. Then, the potential cleavage sites of a part of DIC3Hs were predicted by psRNAtarget software against longan sRNA database[29] with default parameters and a maximum expectation value of 3.5 (except DIC3H36 ). All the primers used in this study were listed in additional file 1.

Abbreviations
C3H-Znf: CCCH Zinc finger transcription factors; Longan: *Dimocarpus longan* Lour.; *DIC3H*: longan CCCH Zinc finger gene; AS: alternative splicing events; RT-PCR: real-time PCR; qPCR: quantitative real-time PCR; MeJA: methyl jasmonate; SHAM: Salicylhydroxamic acid.; STD: Sodium Tungstate Dihydrate; ABA: abscisic acid; EC: embryogenic callus.; ICpEC: incomplete compact pro-embryogenic cultures; GE: globular embryos; NEC: non-embryonic callus.

**Declarations**

**Ethics approval and consent to participate**

Experimental materials provided by the Institute of Horticultural Biotechnology, Fujian Agriculture and Forestry University, Fuzhou, Fujian, China. The 'HHZ' cultivar used in this study were planted and grown in Fujian Agriculture and Forestry University, Fujian Province, China. No specific permits were required for plant collection. The study did not require ethical approval or consent as no endangered or protected plant species were involved.

**Consent for publication**

Not applicable.

**Availability of data and materials**

All data presented in this study are provided either in the manuscript or additional files.

**Competing interests**

The authors declare that they have no competing interests.

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study and collection, analysis, and interpretation of data and in writing the manuscript.

**Authors’ contributions**

ZXL and YLL designed and coordinated the research, and helped to draft the manuscript. YLS participated in its design, carried out the experimental work and wrote the manuscript. MQJ helped to draft the manuscript. SQH, XDX and XL prepared the materials. YLL revised the paper. All authors read and approved the final version of the manuscript. All authors agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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**Authors' information**

1Institute of Horticultural Biotechnology, Fujian Agriculture and Forestry University, Fuzhou 350002, Fujian, China

**Additional Files**

Additional file 1 :The primers used in this study.

Additional file 2 :The potential sRNAs cleave DIC3H01.

Additional file 3 :The potential sRNAs cleave DIC3H04.

Additional file 4 :The potential sRNAs cleave DIC3H05.

Additional file 5 :The potential sRNAs cleave DIC3H11.

Additional file 6 :The potential sRNAs cleave DIC3H19.

Additional file 7 :The potential sRNAs cleave DIC3H36.
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Tables

Table 1 The characteristics of longan ZF_C3H gene family

| Gene name | Locus name      | Arabidopsis orthologs locus | Exons | Protein | Number of CCCH motif | Subcellular local |
|-----------|-----------------|-----------------------------|-------|---------|----------------------|-------------------|
| DIC3H01   | Dlo_001632.     | AT3G02830.1                 | 8     | 440     | 48.10                | 8.86              | nucl              |
| DIC3H02   | Dlo_001685.     | AT3G02830.1                 | 2     | 256     | 29.32                | 7.99              | cytop             |
| DIC3H03   | Dlo_002620.     | AT5G12440.1                 | 7     | 639     | 70.50                | 5.85              | nucl              |
| DIC3H04   | Dlo_003675.     | AT5G12440.1                 | 7     | 183     | 20.35                | 8.64              | secret            |
| DIC3H05   | Dlo_005201.     | ATG41900.1                  | 1     | 739     | 80.41                | 5.73              | nucl              |
| DIC3H06   | Dlo_006391.     | AT5G12850.1                 | 1     | 730     | 79.29                | 6.10              | nucl              |
| DIC3H07   | Dlo_007322.     | AT5G51980.1                 | 9     | 405     | 44.32                | 8.12              | nucl              |
| DIC3H08   | Dlo_008757.     | AT3G12130.1                 | 3     | 293     | 30.64                | 9.36              | secret            |
| DIC3H09   | Dlo_009034.     | AT3G12130.1                 | 4     | 499     | 56.30                | 8.15              | nucl              |
| DIC3H10   | Dlo_009361.     | AT2G29580.1                 | 3     | 274     | 30.81                | 7.12              | nucl              |
| DIC3H11   | Dlo_010735.     | AT2G32930.1/AT2G32930.2     | 7     | 448     | 48.40                | 7.19              | nucl              |
| DIC3H15   | Dlo_014492.     | AT2G02160.1                 | 3     | 721     | 79.80                | 5.30              | nucl              |
| DIC3H16   | Dlo_015528.     | AT2G24830.1                 | 4     | 504     | 56.79                | 5.22              | nucl              |
| DIC3H17   | Dlo_015573.     | AT2G29580.1                 | 5     | 413     | 46.12                | 8.26              | nucl              |
| DIC3H18   | Dlo_016726.     | AT1G32360.1                 | 2     | 379     | 40.01                | 7.12              | secret            |
| DIC3H19   | Dlo_017697.     | AT2G47850.1/AT2G47850.2     | 7     | 473     | 50.84                | 9.17              | nucl              |
| DIC3H20   | Dlo_017983.     | AT3G52980.1                 | 7     | 570     | 64.02                | 5.97              | nucl              |
| DIC3H21   | Dlo_018202.     | AT2G24830.1                 | 11    | 415     | 46.13                | 7.23              | nucl              |
| DIC3H22   | Dlo_018738.     | AT3G27700.1/AT3G27700.2     | 4     | 931     | 101.64               | 6.52              | cytop             |
| DIC3H23   | Dlo_019035.     | AT4G29190.1                 | 1     | 388     | 42.08                | 7.86              | nucl              |
| DIC3H24   | Dlo_019408.     | AT3G19360.1                 | 2     | 333     | 37.37                | 8.68              | secret            |
| Type       | Structure                                      | Events   | Genes  |
|------------|-----------------------------------------------|----------|--------|
| A3'S       | ![A3'S Structure](image)                      | 84 (26.17%) | 22 (30.14%) |
| A5'S       | ![A5'S Structure](image)                      | 59 (18.30%) | 18 (24.66%) |
| IR         | ![IR Structure](image)                       | 145 (45.17%) | 21 (28.77%) |
| ES         | ![ES Structure](image)                       | 33 (10.28%) | 12 (16.44%) |
| Total      |                                               | 321      | 73     |

Table 2: Classification of AS events in longan somatic embryo
Figure 1

The phylogenetic tree of C3H genes from longan and Arabidopsis.

AtC3H01: YQPDKKDYKET.GYCYGDSRFLFD.
AtC3H02: KTELKNKMET.GTIPYGENQFARGL.
AtC3H03: WKTENNMT.EGCPYGENQFARGL.
AtC3H14: FKTENQKET.GAPYIAPRQLAG.
AtC3H18-1: FPKLCCFRN.GTIPYITCFAAG.
AtC3H18-2: YKGRHKRFYTE.EGCPYGENQFARGL.
AtC3H18-3: WKTREIKNKWL.EGCPFNGKNAFAAG.
AtC3H24-1: FKTRIAGKQK.GTCRNGFCAAG.
AtC3H24-2: WKTREIKKWT.EGCPFNGKNAFAAG.
Figure 2

The multiple alignment of DIC3Hs and selected AtC3Hs ZF-CCCH domain amino acid sequences.

Figure 3
The evolution relationship, gene structure and conserve motif in longan C3H gene family. A The phylogenetic tree of DlC3Hs. B Gene structure of longan C3H genes. Yellow box indicate the coding sequence; Blue box is the untranslation 3’ and 5’ region. Black line represent intron; Red box is the ZF-CCCH conserved domain. The numbers of 0, 1, 2 were the phase of corresponding introns. The sequence length can be inferred by bottom scale. C Distribution of conserved domain in longan C3H genes. The 25 motifs are display with different color and the length of protein can be estimated by bottom scale. Motif1: ZF-CCCH super family; Motif 2: zf-U1; Motif3: YTH1 super family; Motif4: WD40 super family; Motif5: TrmA; Motif6: Torus super family; Motif7: SWIB; Motif8: SMC_N super family; Motif9: RRM super family; Motif10: Plus3; Motif11: PHD5_NSD; Motif12: OST-HTH; Motif13: MPP_CWF19_N; Motif14: KH_1 super family; Motif15: HIT_like; Motif16: GYF; Motif17: G-patch; Motif18: FtsK super family; Motif19: DNA_pol3_delta2 super family; Motif20: DFRP_C; Motif21: CwfC_2; Motif22: Atrophin-1 super family; Motif23: ANK; Motif24: Amino_oxidase super family; Motif25: PRK12678 super family
The AS event distribution of DIC3H genes. A The percentage of four AS events find in NEC, EC, ICpEC and GE stages of longan somatic. A3’S: alternative 3’ splice site; A5’S: alternative 5’ splice site; IR: intron retention; ES: exon skipping. B The number distribution of AS event in DIC3Hs in four stages of longan somatic embryo.
Figure 5

The potential cis-elements of DIC3H promoters. The promoter sequences (upstream 2000Kb) of 49 DIC3H genes are analyzed by PlantCARE database. A part of promoter region of DIC3H01, DIC3H12, DIC3H28 and DIC3H30 are absent. The length of sequences can be inferred by the bottom scale.
Expression levels of 26 selected DIC3Hs after identical concentration of ABA and STD treatment.
Figure 7

Expression levels of 26 selected DIC3Hs after identical concentration of MeJA and SHAM treatment.
Figure 8

Expression profile of DIC3Hs in different stages of longan somatic embryo.
Expression patterns of 17 DIC3Hs in longan early somatic embryo. NEC: non-embryonic callus; EC: embryonic callus; ICpEC: incomplete compact pro-embryogenic cultures; GE: globular embryos
Figure 10

Analysis of the sRNAs cleavage site in longan C3Hs. The yellow area is the sRNA binding site, the number above is the cleavage position; the number in the green area is the number of cleavage.

Supplementary Files

This is a list of supplementary files associated with the primary manuscript. Click to download.

Additional file 6  The potential sRNAs cleave DIC3H19.xls
Additional file 7  The potential sRNAs cleave DIC3H36.xls
Additional file 3  The potential sRNAs cleave DIC3H04.xls
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