Gain-of-function in *Arabidopsis* (GAINA) for identifying functional genes in *Hevea brasiliensis*

Han Cheng¹,²*, Jing Gao¹, Haibin Cai¹, Jianshun Zhu¹ and Huasun Huang¹,²*

**Abstract**

**Background:** Forward genetics approaches are not popularly applied in non-model plants due to their complex genomes, long life cycles, backward genetic studies etc. Researchers have to adopt reverse genetic methods to characterize gene functions in non-model plants individually, the efficiency of which is usually low.

**Results:** In this study, we report a gain-of-function in *Arabidopsis* (GAINA) strategy which can be used for batch identification of functional genes in a plant species. This strategy aims to obtain the gain-of-function of rubber tree genes through overexpressing transformation ready full-length cDNA libraries in *Arabidopsis*. An initial transformation test produced about two thousand independent transgenic *Arabidopsis* lines, in which multiple obvious aberrant phenotypes were observed, suggesting the gain-of-function of rubber tree genes. The transferred genes were further isolated and identified. One gene identified to be metallothionein-like protein type 3 gene was further transferred into *Arabidopsis* and reproduced a similar aberrant phenotype.

**Conclusion:** The GAINA system proves to be an efficient tool for batch identification of functional genes in *Hevea brasiliensis*, and also applicable in other non-model plants.

**Keywords:** Gain-of-function, Rubber tree, *Arabidopsis*, Full-length cDNA library, Mutation, Forward genetics

**Background**

Mutants play important roles for identifying new genes with specific functions. Many methods have been applied to generate loss-of-function mutations, including the use of ethyl methanesulfonate, fast-neutron treatment, antisense and RNA interference technology, and insertion mutations by a transposable element or T-DNA (Bolle et al. 2011). These methods have produced a large number of mutant pools in *Arabidopsis* (Alonso et al. 2003; Berardini et al. 2004; Lamesch et al. 2012), rice (Zhang et al. 2006; Krishnan et al. 2009) and maize (Andorf et al. 2015). However, these approaches do not likely discover the genes that are redundant or essential for early embryo development, as these mutations may cause embryo lethality that gives no offsprings. Gain-of-function strategy works efficiently to overcome these shortcomings, which enhances gene expression to generate mutation phenotypes (Weigel et al. 2000; Nakazawa et al. 2003).

Activation tagging was the first and most widely used gain-of-function mutation method which utilizes the enhancer element from the cauliflower mosaic virus (CaMV) 35S gene. T-DNA containing four folds 35S enhancers is transferred into the *Arabidopsis* genome and activates the nearby gene transcription (Weigel et al. 2000; Nakazawa et al. 2003). Activation tagging now has been successfully applied in rice (Jeong et al. 2002), tomato (Mathews et al. 2003), poplar (Fladung and Polak 2012) etc. A new strategy for activation tagging utilizes a recombinase reaction between two lines generated by the pEnLox/pCre vector system, which provides a new and easier way to analyze gain-of-function mutants (Pogorelko et al. 2008). Another strategy such as SARE (Sense/Antisense RNA Expression) overexpresses genes...
in sense or antisense manner directly at genome scale, which produces enhanced or suppressed mutants in the same mutant pools (Mou et al. 2002). The SARE system constructs an Arabidopsis cDNA library by inserting the cDNA fragments between the 35S promoter and NOS terminator. This expression library is then used to transform Arabidopsis through an Agrobacterium-mediated way. The initial application of this system has isolated a mutant overexpressing the sense cDNA fragment of a choline biosynthesis-related PEAMT gene (Mou et al. 2002). However, the combination of gain-of-function and loss-of-function mutants in the same pool would increase the difficulty of mutants screening.

All these methods require the homogenization of the mutation allele, and an effective transgenic technique to produce enough transgenic lines for saturating the genome (Peters et al. 2003). However, it is not easy to be achieved in the plants with a long life cycle such as the rubber trees (Hevea brasiliensis), which produce seeds at 4 or 5 years old (Priyadarshan and de Goncalves 2003). Besides, rubber trees are highly heterozygous and recalcitrant to transform, let alone to produce plenty of seeds at 4 or 5 years old (Priyadarshan and de Goncalves 2003). Besides, rubber trees are highly heterozygous and recalcitrant to transform, let alone to produce plenty of transgenic lines for screening mutations (Montoro et al. 2003; Jayashree et al. 2003; Blanc et al. 2006; Leclercq et al. 2010). So it is not feasible to utilize classical genetic approaches to identify genes in rubber tree. Here we report a new strategy to overexpress rubber tree genes in Arabidopsis, which has proved to be fast and effective for the identification of functional gene in H. brasiliensis.

Methods
Plant materials and growth conditions
Arabidopsis thaliana L. Heynh. ecotype Columbia (Col-0) was used in this study. Plant growth conditions were described elsewhere (Cheng et al. 2015).

Rubber tree (H. brasiliensis) clone 93-114 was used for RNA extraction (Cheng et al. 2008). This genotype was not EST sequenced before this work. The seedlings were grown in a green house with 12 h light/12 h dark photoperiod (120 μE m⁻² s⁻¹). When the seedlings were about 1 m high, with the second whirl leaves stabilized, the leaves and bark were harvested and frosted in liquid nitrogen for RNA extraction.

Construction of pXCS-LIB binary vector
The expression vector pXCS-LIB was derived from the pXCS-HASstrep (accession number AT812152) which was provided by Dr. Claus-Peter Witte (Witte et al. 2004). To generate pXCS-LIB, an adapter (AAGCTTGGCCATTACGGCCAAATAGGCCCGCCTCGGCCGAATTC, HindIII and EcoRI sites underlined) was ligated into the HindIII and EcoRI site in pXCS-HASstrep. The new constructed plasmid was sequenced using primer LibSeq (5′-TCCTTCGCAAGACCCCTTCCT-3′) to confirm right structure. The pXCS-LIB was digested by SfiI, and then dephosphorylated by calf intestinal alkaline phosphatase (Takara). The digested pXCS-LIB fragment was recovered and used for cDNA library construction.

Construction of a rubber tree cDNA library in pXCS-LIB
Rubber tree total RNA was extracted from mix sample of the leaf, bark and shoot tips by CTAB method (Cheng et al. 2015), and mRNA was isolated using a PolyATract mRNA Isolation System III (Promega). The cDNA was synthesized using the Clontech Creator SMART cDNA Library Construction Kit, and was then normalized with TRIMMER-DIRECT cDNA Normalization Kit (Ervogen). The normalized cDNA was then digested by SfiI and fractioned using CHROMA SPIN-400 Columns. The cDNA longer than 300 bp was recovered and ligated into the SfiI digested pXCS-LIB fragments. The ligation products were ethanol precipitated and dissolved in 5 μL double distilled water, and electroporated into 25 μL E.coli TOP10 competent cells using a Gibco BRL Cell Porator with the follow setting: capacitance 330 μF, voltage 350 V, impedance low ohms, charge rate fast, resistance 4 kΩ. After transformation, the cell was resuspended in 1 mL SOC medium and cultured at 37 °C for 45 min. One μL strain culture was diluted into 100 μL LB medium, and plated onto LB agar plate containing 50 mg/L carbenicillin. After overnight culture, the clones were counted and the library titer was calculated.

For cDNA insertion size determination, clones were randomly picked up and subjected to PCR analysis with primer with primer F-p (5′-TCCTTCGCAAGACCCCTTCCT-3′) and R-p (5′-TGAGGATGAGACCCACCCGC-3′). The products were resolved on agarose gel and the bands size was further calculated.

Transfer the binary plasmid library into Agrobacteria and generation of transgenic plants
The cDNA library was amplified on two hundreds 15 cm LB agar plates and all the clones were combined in 1000 mL liquid LB medium. The library plasmid was extracted from 100 mL of the amplified cDNA library using a Qiagen plasmid purification kit. This library plasmids were further introduced into Agrobacterium tumefaciens strain GV3101 (pMP90RK) by electroporation, and transformants were screened on YEB agar plates supplemented with 50 mg/L rifamycin, 20 mg/L gentamicin, 50 mg/L kanamycin, and 50 mg/mL carbenicillin (Koncz and Schell 1986). Transformants were allowed to grow on plates for 3 days and then were pooled and cultured for 2 h at 28 °C.

Arabidopsis Col-0 plants were transformed with these agrobacteria via flower dipping method (Clough and
Plants prepared for transformation were grown on the medium composed vermiculite and peat-moss (1:3) at 23 °C. Transformed plants were allowed to self-pollination and the seeds were harvested. Transgenic plants were selected by spraying 100 mg/L Basta (glufosinate ammonium) onto 5 d T1 plants for 3 times with 3 d intervals. The survived plants were cultured and the T2 seeds from each T1 plant were harvested individually.

**Mutant screening and isolation of the transferred gene**

T1 plants were examined for morphological aberrance. Individual T1 plants with specific phenotype were subjected to further analysis. For each mutant line, segregation of phenotype and Basta resistance was performed in the T2 and T3 generations. The lines with a 3:1 segregating ratio was regarded as single T-DNA insertion.

The overexpressed genes were isolated by amplifying the genomic DNA of the mutants using primers LIB5 (5′-ATGACGCACAATCCCACATAC-3′) and LIB3 (5′-TGTAGAGAGAGAGACTGTGGTGATTGGTGGTGGG-3′). The PCR products were cloned into pMD18-T (Takara) and sequenced.

**Confirmation that the transferred gene leads to the mutant phenotype**

To confirm the transferred gene induced the mutant phenotype, the PCR product of target gene was digested by SfiI and ligated into SfiI restrictive sites of pXCS-LIB vector. The constructed plasmid was introduced into Agrobacterium strain GV3101 (pMP90RK), and used to transform Col-0 wild type plants by flower dipping method. The T1 and T2 phenotype of each gene was examined.

**EST sequencing and COG annotation**

EST sequencing was conducted with ABI 3730 platform at BGI Company (Beijing, China). The raw sequencing data were first cleansed to get rid of vector sequence, low quality ESTs and chimeric sequences. Then the clean ESTs were assembled with CAP3 program using the parameters: identity, 0.95; minimal overlap, 50 bp (Huang and Madan 1999). The assembled unigenes were then used for BLASTx searches (E-value < 1e-5) and annotation against SWISSPROT, KEGG and COG databases.

**Availability of data and materials**

The datasets supporting the conclusions of this article are included within the article and its additional file. The EST sequences were submitted to GenBank as dataset.

**Results**

**Overview of the GAINA system**

The GAINA system overexpress rubber tree genes in Arabidopsis, and generate gain-of-function mutant pools for rubber tree genes. The gain-of-function mutant pools are then used for functional gene identification (Fig. 1). To obtain overexpression lines of rubber tree genes, a transgene-ready full length cDNA library was constructed. To fulfill this, we first constructed a binary vector designated as pXCS-LIB that is used for full length cDNA cloning in sense direction and is fully compatible with the Clontech Creator SMART cDNA Library Construction Kit (Fig. 2) (Zhu et al. 2001). The rubber tree full-length cDNAs were then cloned into the SfiI A and SfiI B sites, and resulted in the transgene-ready full length cDNA library. As the full length cDNAs were driven by a double 35S promoter and followed by a 35S poly A terminator, the cloned cDNAs were expressible in plants when transferred as part of the T-DNA fragment. The transformation ready cDNA library was then transformed into Arabidopsis in an Agrobacterium-mediated flowering dipping method. Independent transgenic lines were collected and each contained at least one overexpressed rubber tree gene. These lines constitute the gain-of-function Arabidopsis mutant pools of the rubber tree genome information facilitating functional genes identification in this non-model species.
Construction of a rubber tree cDNA library

The key point of GAINA system is to construct a high-quality transformation ready cDNA library with uniform abundance and broad representation. To achieve this goal, the RNA was extracted from mixed samples of leaf, bark and shoot tips, then the cDNA was further...
normalized. Besides, the cDNAs should contain intact CDS in the sense direction which ensures the cloned cDNAs are expressed and translated correctly. To meet these criteria, we constructed a normalized full-length cDNA library using the SMART™ and TRIMMER™ technology (Zhu et al. 2001; Bogdanov et al. 2010). The normalized full-length cDNA fragments were then cloned into the SfiI restrictive site in sense direction and transformed into TOP10 competent cells. An aliquot of the cDNA library is titrated which demonstrated this library contained $1.4 \times 10^6$ clones.

To evaluate the quality, diversity and insertion length of the constructed full length cDNA library, we randomly picked up thirty-six clones and analyzed with PCR amplification. As shown in Fig. 3, all the selected clones contained effective insertions. The cDNA insert length ranged from 0.5 to 2 kb with an average size of 1 kb, demonstrating this library was of high quality and rich diversity.

**EST sequence analysis**

The normalized full length cDNA library was subjected to EST sequencing from the 5′ terminal using Sanger sequencing technology. Totally 25020 EST sequences were obtained, of which 22585 were high-quality sequences. Using a CAP3 assembly program (Calikowski and Meier 2006), the ESTs obtained were assembled into 12114 unigenes (3233 contigs and 8881 singlets), with a redundancy rate of about 46%. The EST sequencing data demonstrated a high quality and wide representation of this library for rubber tree transcriptomes.

The ESTs sequences were also annotated using blastx against SWISSPROT, KEGG and COG databases. Figure 4 showed a profile of COGs annotation of all unigenes. Totally 3127 genes were annotated with COG database (e-value cut-off 1E-05), belonging to 757 COG ids. These COGs fell into 23 COGs classes, with posttranslational modification (16.9 %), translation (15.5 %) and general function prediction (14.5 %) as the major part. The COGs classes’ diversity of the annotated unigenes demonstrated the library represented genes with versatile functions.

**Multiple gain-of-function lines with visible phenotype aberrance**

The full length cDNA library was amplified on LB agar plates and the plasmids were extracted and transferred into *Agrobacterium* strain GV3101 (pMP90RK). The *Agrobacteria* were then used to transform *Arabidopsis* seedlings with flower-dip method. Each independent overexpression line was selected and regarded as one gain-of-function line of a particular gene. Totally about 2000 independent gain-of-function lines were obtained in an initial round of screening. Among these lines, more than one hundred plants showed obvious phenotype aberrance. As shown in Fig. 5, typical aberrant phenotypes included large-sized rosette leaves, small-sized rosette, twist leaves, short siliques etc. (Fig. 5b–g). The diverse aberrant phenotype of the transgenic lines indicated that the gain-of-function of rubber tree genes could result in *Arabidopsis* phenotype changes and thus be applicable to identify genes of particular interest.

**Isolation of the transferred gene**

The inserted rubber tree genes were then cloned from *Arabidopsis* gain-of-function lines using PCR method. Four lines (phenotype 5C to 5F) were characterized, each of which contained only one T-DNA insertion. Then the cloned fragments were subjected to sequencing, and were then annotated. The annotation results are listed in Table 1, in which two of them were genes with known functions, whereas the other two were annotated as unknown function. The sequences of the cloned genes...
were shown in Additional file 1: Data 1, Additional file 2: Data 2.

To test if overexpression of the rubber tree genes could reproduce the phenotypes found in the gain-of-function mutant pools, the phenotype 5D gene (metallothionein-like protein type 3) was re-overexpressed in Arabidopsis. The transgenic seedlings also displayed twisted leaves phenotype as found in gain-of-function mutant pools (Fig. 5h). These results demonstrated that the gain-of-function phenotype found in Fig. 5d did come from the overexpression of the transferred rubber tree metallothionein-like protein type 3 gene.

Discussion

Chemical and physical mutation is the most frequently used method to generate mutants. Though saturating a genome is relatively easy for chemical and physical mutation, it is difficult to map the mutation site, thus makes it very difficult to clone the target genes. Though some new methods for mapping the mutation emerged recent years, such as TILLING etc. (Till et al. 2003), it is still not easy to work in non-model plants. T-DNA insertion has been proved to be very convenient for the cloning of mutated target genes. However, the loss-of-function strategy makes this method to be limited in model plants. For the less-studied long life span trees, the T-DNA insertion mutation is not easy to be achieved. In non-model plants, overexpression in Arabidopsis is an alternative method to characterize the gene functions, which has been successfully applied in many plant species (Foucart et al. 2009; Kalamaki et al. 2009; Jiang et al. 2016). This gain-of-function strategy is very useful for the plants with low transformation efficiency.

In this study, we adopt a gain-of-function method (GAINA) to overexpress the rubber tree full-length cDNA library in Arabidopsis, and generate thousands of gain-of-function lines. These lines contain overexpressed rubber tree genes, and therefore can be used to identify genes with particular functions. The key of the GAINA system is to construct a full-length cDNA library in the sense direction, which ensures the successful expression after transferred into Arabidopsis. First we constructed a transformation-ready binary vector pXCS-lib which is compatible to full-length cDNA library construction. The full-length cDNA library was then cloned into pXCS-lib plasmid and utilized to transform Arabidopsis. To ensure the cDNA library mostly represent the rubber tree genes, a mixed RNA sample from several tissues was used for library construction. A normalization step was further used to gain uniform abundance for each gene in the library. These measures guaranteed the maximum likelihood to obtain gain-of-function mutation lines for each rubber tree genes. The EST sequencing of the full-length cDNA library also confirmed the wide representation of the constructed library.

Another key of the GAINA system is to generate enough Arabidopsis transgenic lines, which maximizes
the coverage of the rubber tree genes. In the initial experiment, two thousands independent transgenic lines were obtained, in which aberrant phenotypes were observed in many lines. This also demonstrated that the GAINA strategy was effective to identify functional genes in the rubber trees, especially for the genes involved in organ development, abiotic stress resistance. The next job is to obtain more GAINA transgenic lines, and to cover more rubber tree genes in this system.
In this study, four rubber tree genes are identified using GAINA system. The transferring of these genes caused visible phenotype aberrances in *Arabidopsis* (Fig. 5). Among these identified genes, only one (MT3) was previously well characterized. MT3 is low molecular weight, cycteine-rich proteins that bind metals such as Zn, Cu or Cd, and is proposed to participate in a variety of processes including metal ion homeostasis and tolerance (Benatti et al. 2014). MT3 is also suggested to protect cells against oxidative stress (Akashi et al. 2004). The overexpression of the rubber tree MT3 gene caused twisted leaves in GAINA system. This phenotype was reproduced in an independent *HbMT3* transgene experiment, which further confirmed the connection between phenotype and *HbMT3* overexpression, though the detail mechanism is yet to be unraveled.

Nowadays, there were numerous studies that transferred the non-model plants genes into *Arabidopsis* and characterized their functions (Foucart et al. 2009; Kalamaki et al. 2009; Polashock et al. 2010; Jiang et al. 2016; Wang et al. 2016; Liu et al. 2016). We also characterized rubber tree *HbCBF1* gene which showed conserved functions in regulating CBF pathway in *Arabidopsis* (Cheng et al. 2015). These studies utilized the reverse genetic approach to characterize gene functions. The GAINA system provides a forward genetic approach for batch identification of functional genes in non-model plants, which will greatly facilitate the genetic study in these species. However, a shortcoming of the GAINA system is that this method will not likely identify rubber tree specific genes, such as those involved in rubber biosynthesis, latex development. The study of these genes should rely on the molecular biology research progresses in *H. brasiliensis* (Kush et al. 1990; Duan et al. 2010; Chow et al. 2012). Even so, the GAINA system offers a powerful forward genetic tool for functional gene study and the identification of genes involved in particular pathways in non-model plant species.

**Conclusion**

In this study, we report a gain-of-function in *Arabidopsis* (GAINA) system that overexpresses a rubber tree full-length cDNA library in *Arabidopsis*, and generates thousands of transgenic lines. These gain-of-function lines prove to be beneficial for the functional identification of rubber tree genes. Therefore, the GAINA system offers a powerful forward genetic tool for functional gene study and the identification of genes involved in particular pathways in non-model plant species.

**Additional files**

**Additional file 1: Data 1.** The sequences of the cloned genes, phenotype SC, phenotype SD, phenotype SE, phenotype SF, from the plants of C, D, E, F in Figure 5.

**Additional file 2: Data 2.** Blastx data of the cloned genes, phenotype SC, phenotype SD, phenotype SE, phenotype SF, from the plants of C, D, E, F in Figure 5.

**Abbreviations**

GAINA: gain-of-function in *Arabidopsis*; MT3: metallothionein-like protein type 3.

**Authors’ contributions**

HC and HH designed the project; GJ, CHaibin and ZJ performed the experiments; HC wrote the manuscript and HH helped to draft the manuscript. All authors read and approved the final manuscript.

**Author details**

1 Key Laboratory of Ministry of Agriculture for Tropical Crops Physiology, Rubber Research Institute, Chinese Academy of Tropical Agricultural Science, Danzhou City, Hainan, People’s Republic of China. 2 Rubber Research Institute, Chinese Academy of Tropical Agricultural Science, Danzhou 571737, Hainan, People’s Republic of China.

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**Competing interests**

The authors declare that they have no competing interests.
Data archiving statement
The nucleotide sequences of the EST sequencing are submitting to NCBI repositories and the accession numbers will be supplied once available.

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