A 1232 bp upstream sequence of glutamine synthetase 1b from Eichhornia crassipes is a root-preferential promoter sequence

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Abstract

Background: Glutamine synthetase (GS) acts as a key enzyme in plant nitrogen (N) metabolism. It is important to understand the regulation of GS expression in plant. Promoters can initiate the transcription of its downstream gene. Eichhornia crassipes is a most prominent aquatic invasive plant, which has negative effects on environment and economic development. It also can be used in the bioremediation of pollutants present in water and the production of feeding and energy fuel. So identification and characterization of GS promoter in E. crassipes can help to elucidate its regulation mechanism of GS expression and further to control its N metabolism.

Results: A 1232 bp genomic fragment upstream of EcGS1b sequence from E. crassipes (EcGS1b-P) has been cloned, analyzed and functionally characterized. TSSP-TCM software and PlantCARE analysis showed a TATA-box core element, a CAAT-box, root specific expression element, light regulation elements including chs-CMA1a, Box I, and Sp1 and other cis-acting elements in the sequence. Three 5′-deletion fragments of EcGS1b upstream sequence with 400 bp, 600 bp and 900 bp length and the 1232 bp fragment were used to drive the expression of β-glucuronidase (GUS) in tobacco. The quantitative test revealed that GUS activity decreased with the decreasing of the promoter length, which indicated that there were no negative regulated elements in the EcGS1-P. The GUS expressions of EcGS1b-P in roots were significantly higher than those in leaves and stems, indicating EcGS1b-P to be a root-preferential promoter. Real-time Quantitative Reverse Transcription-Polymerase Chain Reaction (qRT-PCR) analysis of EcGS1b gene also showed higher expression in the roots of E.crassipes than in stems and leaves.

Conclusions: EcGS1b-P is a root-preferential promoter sequence. It can specifically drive the transcription of its downstream gene in root. This study will help to elucidate the regulatory mechanisms of EcGS1b tissue-specific expression and further study its other regulatory mechanisms in order to utilize E.crassipes in remediation of eutrophic water and control its overgrowth from the point of nutrient metabolism.

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Background

Eichhornia crassipes is a most prominent aquatic invasive plant [1], with negative effects on environment and economic development [2, 3]. Also, it is regarded as a valuable resource with several unique properties, and previous studies have reported that E. crassipes had high absorption efficiency of nitrogen (N), phosphorus (P) and heavy metals [4–6]. So it was used in the bioremediation of pollutants present in water [7, 8]. Furthermore, E. crassipes served as a good economic raw material for the production of feeding and energy fuel [9–11]. There was about 33% of crude protein accumulation in the leaves when grown in sewage wastewater, and it was also used in the production of fuels such as ethanol and methane along with some microorganisms [12, 13]. Thus, studying the biochemical metabolism of E. crassipes from the molecular level assists in further utilizing and controlling this weed.

Glutamine synthetase (GS) in plants acts as a key enzyme in N metabolism, converting inorganic N (NH₄⁺ or NO₃₋) that is absorbed from outside into organic N and further incorporating into other biomacromolecules by GS (Glutamine synthetase)-GOGAT (Glutamate Synthase) cycle [14]. GS could be divided into GS1, GS2 and GS3 according to its protein structure and gene sequence [15, 16], and higher plants have GS1 and GS2 forms in roots and leaves. GS2 is generally present in the leaf tissues, while GS1 in the roots and vascular tissues [11, 17]. Many studies have shown that GS1 in roots assimilates NH₄⁺ from soil into the plant and GS1 in leaves reassimilates NH₄⁺ generated during protein turnover in leaves, while GS2 assimilates NH₄⁺ derived from photosynthesis and nitrate reduction [18]. GS plays an important role in N assimilation which is essential to the plant growth and development. Plants have a refined regulation mechanism of GS expression, which is closely coordinated with the external condition and the development status of plants. Zhang et al. reported that the regulation of GS isozymes might promote flow strength and enhance N use efficiency (NUE) by a complex C-N metabolic mechanism [19]. In wheat, GS1-1 expression was upregulated in response to a reduction in N supply [20], whereas high NH₄⁺ supply specifically induced the expression of the GS1–3 isogene in barley and sorghum [21, 22]. GS expression was regulated by the external N application, but the extent of this regulation depended on the plant species, N source and plant tissue [18]. What is the regulation mechanism of GS expression? The study on the promoter of GS gene could reveal the temporal and spatial properties of GS expression, and further reveal the molecular mechanism of GS regulation at the transcriptional level, which could be better utilized to improve the absorption and utilization of N by plants.

Promoters are upstream sequences of the 5’ end of genes that regulates gene expression, contains RNA polymerase and transcriptional factor recognition and binding sites, and enables the initiation of transcription. Promoter sequences included core elements and regulatory elements, and the transcriptions of their downstream genes could be regulated by the conditions corresponding to these regulatory elements [23–25]. Heat shock elements of the soybean Gmhsp 17.3-B gene were involved in heat shock promoter activation during tobacco seed maturation [26]. The expression of isopentenyltransferase driven by the cold inducible AtCOR15a promoter could provide the sugarcane cultivated in tropical and subtropical region a greater tolerance to cold stress [27]. The root-specific promoter PsPR10 from Pinus strobus containing many abiotic regulatory elements could efficiently initiate the expression of downstream genes in root under the different hormonal or salt stress conditions [28]. So the study of the promoter and its regulation can help to elucidate the expression mechanism of its downstream gene. In this study, we isolated and characterized the promoter sequence of EcGS1b gene. For investigating the function of EcGS1b-P, transgenic tobaccos with GUS driven by EcGS1b-P were developed and it was found that EcGS1b-P could drive the gene expression preferentially in roots. To further validate this, the EcGS1b gene expression levels were measured by using a real-time qRT-PCR.

Results

Cloning and analysis of EcGS1b-P

To clone the regulatory regions of EcGS1b gene, the primer pairs were designed from the corresponding cDNA sequences. Using PCR gene walking on genomic DNA from E. crassipes, the upstream of EcGS1b gene was cloned thrice with the nested-PCR (Fig. 1) and then was sequenced. The EcGS1b gene upstream sequence (1232 bp) was obtained and named as EcGS1b-P. This sequence was submitted to NCBI database and assigned the accession number MT154418. Homology search using Blast programs revealed no similarities of known genes or promoters in the GenBank database. The transcriptional start site (TSS) was started from the 105th bp upstream to the ATG codon. The TSS distances from the TATA-box (−30 bp) and the CAAT-box (−96 bp) were consistent with those that were usually described for other plant promoters [29, 30]. Table 1 was the predicted result using TSSP-TCM software and PlantCARE, showing that some cis-acting elements involved in abiotic stress tolerance (MBS, HSE, LTR, circadian), endosperm and root specific expression elements (Skn-1_motif, ROOTMOTIFTAPOX1), light regulatory elements (chs-CMA1a, Box I and 5p1), and salicylic acid and gibberellin-responsive elements (TCA-
element, P-box) existed in this sequence (Fig. 2). So EcGS1b-P contained the core promoter, tissue specific expression elements, light regulation elements and other cis-acting elements, which might regulate the time and the space of GS1b expression.

**Histochemical analysis of GUS expressions driven by different length promoters of EcGS1b-P**

To investigate GUS expression driven by different lengths of EcGS1b-P, four different length EcGS1b-Ps (EcGS1b-P and its three 5’-deletion fragments, namely p1200, p900, p600 and p400) instead of 35 s promoter were fused with the reporter gene GUS in pBI121 vector (Fig. 3a) respectively. The fused vectors were verified by digesting the vector with HindIII and BamHI (Fig. 3b), respectively. After, pBI121s were transformed into tobaccos, and the transgenic tobaccos were confirmed by PCR. Histochemical analysis showed that all tissues of transgenic tobaccos with pBI121 vector which had 35 s promoter (positive control) appeared in conspicuous blue (Figs. 4a, 5a, 6a), which indicated that 35 s promoter could drive GUS expression well in all tissues of transgenic tobacco. All tissues of transgenic tobaccos with pBI101 vector which had no promoter to drive GUS expression (negative control, Figs. 4f, 5f, 6f) and the wild type tobacco (Figs. 4g, 5g, 6g) were

| Table 1 Functions of promoter elements |
|----------------------------------------|
| **Site names** | **Position** | **Strand** | **Function** |
| TATC-box | −1159 | + | cis-acting element involved in gibberellin reaction |
| MBS | −1043 | + | MYB binding site involved in drought induction |
| MBS | −352 | − | MYB binding site involved in drought induction |
| P-box | −535 | + | gibberellin response element |
| chs-CMA1a | −522 | + | part of a light response element |
| Box I | −277 | + | light response element |
| Box I | −159 | − | light response element |
| circadian | −399 | + | cis-acting regulatory element of circadian rhythm |
| TCA-element | −905/−301 | – | cis-acting element involved in salicylic acid response |
| Sp1 | −828 | – | light response element |
| Skn-1_motif | −617 | – | cis-acting regulatory element of endosperm expression |
| HSE | −503 | – | cis-acting element involved in heat stress response |
| LTR | −489 | – | cis-acting element involved in low-temperature response |
| Box-W1 | −656 | – | cis-acting element involved in fungus |
| ROOTMOTIFTAPOX1 | −1111/− 924/− 417 | + | root-specific expression element |
| ROOTMOTIFTAPOX1 | −925/− 710/− 418 | – | root-specific expression element |

Note: The A in ATG start codon was defined as + 1
white due to lack of GUS activity, which indicated neither transgenic tobacco with no 35S promoter nor non-transgenic tobacco showed GUS activity. All roots of transgenic tobaccos with EcGS1b-Ps showed blue (Fig. 4b, c, d, e), which indicated they had stronger GUS activity, especially in roots transformed with longer promoters (Fig. 4b, c). When the length of the 5′-upstream region of the EcGS1b gene was gradually declined, GUS expression in roots also showed decline. But in leaves, all transgenic tobaccos with EcGS1b-Ps (Fig. 5b, c, d, e) only exhibited a little blue color at the cut edge of the blades. This indicated that GUS expressions driven by different length EcGS1b-Ps in leaves were all very weak. In stems, all different length promoters of EcGS1b-P (Fig. 6b, c, d, e) showed white, which confirmed there were no GUS expressions in these stems.

### Quantitative analysis of different length promoters of EcGS1b-P by 4-methylumbelliferyl-β-D-glucuronide (MUG) assays

The impact of EcGS1b promoter length was tested on GUS activity in transgenic tobacco by MUG assay (Fig. 7a). GUS expression in roots was the highest among three different tissues no matter which promoter of these four different length promoters was used as the GUS driver. In roots, GUS expression levels declined with decreasing of promoter length. In contrast, there was no significant difference in different length promoters both in leaves and in stems ($P > 0.01$). The GUS expression level of transgenic plant with pBI121 vector (positive control) remained the highest, even higher than that of transgenic plant with EcGS1b-P. There was no significant difference in different tissues in the positive control.
In the negative control, GUS activities in both wild type and the transgenic plant with pBI101 vector were almost none in roots, leaves and stems. To determine whether GUS expression driven by EcGS1b gene promoter was influenced by light intensity, we examined GUS expressions in the roots of transgenic tobacco and wild type tobacco at different light intensities (7000 Lux, 3400 Lux, 1700 Lux, dark) for 3 days. As shown in Fig. 7b, no obvious variations in GUS expressions were observed at different light intensities ($P > 0.01$), and thus concluded that the GUS activity expressions were unaffected by the light intensity. In this experiment, we also found that GUS activity expressions of transgenic tobacco were related to the length of EcGS1b-P, confirming that the shorter the length of EcGS1b-P, the lower the GUS activity.

The expression of EcGS1b detected by qRT-PCR

The total RNA of *E. crassipes* was extracted and confirmed by using an electrophoresis, and then qRT-PCR was performed to investigate the expressions of EcGS1b in different tissues. The results showed that the relative expression of EcGS1b was much higher in roots than those in stems and leaves (Fig. 8). There was no significant difference of EcGS1b expression in stems and leaves ($P > 0.01$).

**Discussion**

Currently, the study on *E. crassipes* mainly focused on anaerobic fermentation, feed resources, fertilizer resources, water remediation and biogas production [31, 32]. There was only a little research regarding the physiological and biochemical characteristics of *E. crassipes* such as the determination of enzyme
activities related to N metabolism [33–36]. Moreover, there were few papers that reported on the mechanism of *E. crassipes* at the molecular level [37, 38]. In our previous paper [38], we have cloned three GS1 genes from *E. crassipes* and studied their expressions under different N conditions. In this study, we cloned GS1b promoter from *E. crassipes* and analyzed the expression of *EcGS1b* to investigate the regulation of *EcGS1b* expression.

GS is a key enzyme in the process of N metabolism, and the efficiency of this pathway is affected by its expression [14, 39]. The effects of GS1 gene on crop growth and yield were studied by genetic engineering techniques such as by transgenic and gene silencing techniques [40, 41]. The root dry weight, grain yield, N content and resistance to adversities such as cold, drought, salinity, and biotic stresses in the transgenic

![Fig. 4 GUS expression in transgenic tobacco root.](image)

![Fig. 5 GUS expression in transgenic tobacco leaf.](image)
wheat plants with GS gene were more than those in the wild type plants [42, 43]. The efficiency of photosynthesis could be maintained at normal level in the overexpression of GS plants under N stress conditions [44], which helps in reducing the use of N fertilizer, and making more environmental friendly and economical. The growth status of the transgenic poplar overexpressing GS1α gene of pine was better than that of the wild-type poplar [45]. But not all GS transgenic plant showed the positive effect. The effect of GS on transgenic plant was related with the coordination of N and carbon metabolism [46]. The GS activity might be downregulated via a chain of mechanisms, strongly affected by environmental conditions and metabolic imbalances [39]. Every GS has its own expression mechanism which can adapt to different N stress condition in different tissues in different plants. So studying the GS expression of *E. crassipes* and further exploring the regulation mechanism at the transcriptional level can help us to analyze GS function in order to elucidate the mechanism of efficient utilization of N in water by *E. crassipes*.

Root has many specifically expressed genes, which plays an important role in root structure and function. For instance, in Arabidopsis and Prunus, the root-specific gene *DRO1* influenced the root system architecture [47]. In Maize, 9-Lipoxygenase ZmLOX3 controlled development and maize resistance to root-knot nematodes as a root-specific suppressor [48]. A promoter can regulate the expression of its downstream gene at the transcriptional level. For example, in Arabidopsis, the promoter of *AtGln1;3* was recognized and bound to by the MYB transcription factor CCA1 to match the anticipated C supply [49, 50]. In rice, the OsMYB55 transcription factor bound to the promoter of *OsGSI;2* to initiate the transcription and mediated regulation of amino acid metabolism at high temperature [51]. Studying of root-specific promoters can facilitate analyses of gene functions in roots, further control plant development and growth [52]. The root-specific promoters also can be applied to initiate the transcriptions of root-specific genes instead of constitutive promoters such as 35 s promoter in the process of plant transgenic breeding [53]. Tissue-specific promoters enable adjustment of gene expression in a spatially controlled manner to avoid undesirable effects or excessive energetic costs to transgenic plants. So finding more tissue-specific promoters and exploring their regulation mechanisms are important for the expressions of tissue-specific genes in the breeding of transgenic plants. Much effort in mining tissue-specific promoter mainly focused on the model plants, crops and vegetables such as Arabidopsis, rice, maize, tomato and potato etc. Few studies were related with the aquatic higher plants although these aquatic higher plants are important in the remediation of eutrophication water in the ecological engineering.

In this study, *EcGS1b*-P from *E. crassipes* was isolated and analyzed with PlantCARE software. The results...
showed that the upstream of the promoter sequence contained many core elements such as TATA-box (−30 bp) and CAAT-box (−96 bp), and the distance from TATA-box or CAAT-box to TSS was consistent with that of the results described in the previous studies in other plant promoters [29]. Besides the core components TATA-box and CAAT-box, the EcGS1b-P also had root-specific function motif and cis-acting elements that were regulated by light, heat, hormones and temperature, which was also consistent with the previous studies [54–57]. It is worth to note that GS1b promoter contained root-specific expression element ROOT-MOTIFTAPOX, which indicated EcGS1b might be a root-preferential expression gene. Previous studies have reported many root-specific promoters or root-preferential promoters that contained ROOTMOTIFTAPOX1 functional elements, which could improve the gene expression in the roots [58–61].

Next, histochemical analysis and quantification of GUS activity in transgenic tobacco were used to verify the function of EcGS1b-P [62–64]. We found that GUS expression level was related to the length of the...
promoter fragment. The longer the promoter fragment was, the higher the GUS expression level was. This indicated that there were no negative regulated elements in the EcGS1b-P. This was consistent with the results of Niu et al and others. Niu's study also exhibited that blue color was declined with decreasing of ZmPEAMT promoter length in transgenic tobacco under no stress condition [65]. The 5′-truncated BvcPPOPs (F3, F4 and F5) also drove decreased GUS activity with the shortening of promoter length [66]. However, GUS activity was not positively correlated with the promoter length under the abiotic stress in Niu's study [65]. Hongli Zhang also has reported that the transcription activity of 467 bp (PZ7) core fragment of ZimPIS promoter remained the highest in transgenic tobacco among PZ1-PZ8 [64]. In these researches, the changes in the level of GUS activity were inconsistent with those in the promoter length. This contradictory conclusion might be due to the total effect of different function elements in the promoter. Some elements might have a positive effect, while others might have a negative effect [67, 68]. So, identification and characterization of core elements and other function elements of the promoter will assist us in understanding the molecular regulation mechanism of plant gene expression and further controlling its expression by molecular technique.

Previous research studies have showed that the tissue specific promoter could increase GUS activity in this tissue of transgenic plants. The GUS activity expression of the leaf-specific GapB promoter in transgenic tobacco plants in leaves was higher than that in stems and roots [69]. The GUS expression of the green-tissue promoter OrGSEp remained high in the leaf tissues of transgenic Arabidopsis at different growth stages [70]. The endosperm-specific LPAAT promoter has specially promoted the GUS expression in the endosperm of transgenic rice plant [71]. The pollens-preferential expression promoter OsUgp2 revealed a high GUS gene expression in the pollen of transgenic rice at the binucleate stage [72]. Our qualitative results of GUS staining results showed that the GUS activity expression driven by the EcGS1b-P remained the highest in the roots. There was no obvious GUS expression activity in the stems, and there was only a little in leaves transformed with longer promoters. Moreover, both quantitative results of GUS activity expression and the results of real-time RT-PCR experiments were consistent with that of the qualitative results of GUS staining. These results were also in line with the existence of ROOTMOTIFTAPOX1 elements, which was a root-specific motif in the EcGS1b-P that was analyzed above.

Generally, the GS gene promoter had multiple function motifs other than the root-specific motif, and so its regulatory mechanism might be complicated besides to the tissue specific, and this has been reported in previous studies. The GUS activity driven by 595RhVI11 and 468RhVI11 promoter was observed to be higher under light than in the dark, which might be due to the key light response elements GATA-box, I-box and GT-1box present in the promoter that responded to light in buds.

Fig. 8 Relative expression of EcGS1b Gene in different tissues detected by qRT-PCR. Note: Different letters in the group indicate significant differences in data within the group (P < 0.01)
The GUS activity of the green tissue-specific CsVDE promoter was increased after exposure to high light for 4 h, but decreased after 8 h illumination, which contained many light response elements such as Box-I, GAG-motif, G-box, AE-box, GA-motif, Sp1 and GT-1 motifs [74]. Moreover, the role of downstream signaling components such as MYC2 regulated the GUS expression of Z-box and G-box containing promoters at various wavelengths of light in different tissues [75, 76]. Although the EcGS1b promoter contained three light response elements BoxI, chs-CMA1a, and Sp1, the GUS activity expression level of transgenic plants with EcGS1b promoter was not induced by different light intensities in our study. The regulation of EcGS1b promoter might not be governed by light because it was a root-preferential promoter. The light response elements might be left over from genetic evolution.

**Conclusions**

In this study, a 1232 bp genomic fragment upstream of EcGS1b sequence from Eichhornia crassipes has been cloned, analyzed and functionally characterized. Sequence analysis showed that there were some core elements, root specific expression elements and other functional elements. Three 5′-deletion fragments of EcGS1b promoter (EcGS1b-P) with about 400 bp, 600 bp and 900 bp length and the 1232 bp fragment were respectively used to drive the expression of GUS in tobacco. Both quantitative test and histochemical analysis revealed GUS activity was decreased with decreasing of promoter length. The GUS expressions of EcGS1b-P in roots were significantly higher than those in leaves and stems, indicating EcGS1b-P to be a root-preferential promoter. Real-time qRT-PCR expression analysis of EcGS1b gene also showed higher gene expression in the roots of E. crassipes than in stems and leaves. In all, EcGS1b-P is a root-preferential promoter. This study will help to elucidate the regulatory mechanisms of EcGS1b tissue-specific expression and further study its other regulatory mechanisms in order to utilize E. crassipes in remediation of eutrophic water and control its overgrowth from the point of nutrient metabolism.

**Methods**

**Plant materials and growth conditions**

*E. crassipes* wild type with 2–3 new leaves was obtained from the lake near the west district living area of Guangdong University of Technology. The service and management office of this university permitted our sampling. This plant material was identified by Dr. MH Fu. A voucher specimen of this material had been deposited in South China Botanical Garden. *E. crassipes* was grown hydroponically in a 4 L container with the solution containing 7 mgL⁻¹ KH₂PO₄, 24.5 mgL⁻¹ MgSO₄·7H₂O, 25.5 mgL⁻¹ KNO₃ and 59 mgL⁻¹ Ca(NO₃)₂·4H₂O, and the solution was replaced every 3 days. After it was cultivated for 1–2 weeks and new white roots grew out, the fresh new roots could be used to extract DNA.

The tobacco (*Nicotiana benthamiana*) was germinated on Murashige and Skoog (MS) medium (pH 5.8) containing 0.8% agar at 25 °C with 16 h light/8 h dark cycles and used to do plant transformation experiment. The transgenic tobacco was grown under different lighting (dark, 7000 Lux, 3400 Lux, 1700 Lux) for 3 days and then used to perform the GUS activity experiment.

**Isolation of EcGS1b-P**

Genomic DNA was extracted from plantlets that were grown in the hydroponic system using GV-Plant Genomic DNA Extraction Kit (GENVIEW). The quality of the genomic DNA was assessed by 1% agarose gel electrophoresis. The design of PCR specific primers (GS1bSP1, GS1bSP2, GS1bSP3 sequences were presented in Table 2) was based on the the EcGS1b sequence (NCBI GeneBank accession number KJ881169). The PCR specific primer sequences were synthesized by Sangon Biotech. The 5′-upstream promoter region of EcGS1b was isolated using Genome Walking Kit (Takara). The reactions were performed in 50 μL containing 5 μL DNA, 0.1 μmolL⁻¹ of each primer, 8 μL dNTP (10 mmolL⁻¹ each), 5 μL 10x PCR Buffer, 0.5 μL LA Taq (5 U/μL) with the following conditions: 94 °C for 1 min, 98 °C for 1 min, 5 cycles of 94 °C 30 s, 64.8 °C for 1 min, 72 °C for 2 min, 15 cycles of 94 °C for 30 s, 25 °C for 3 min, 72 °C for 2 min; 94 °C for 30 s, 64.8 °C for 1 min, 72 °C for 2 min; 94 °C for 30 s, 64.8 °C for 1 min, 72 °C for 2 min; 94 °C for 30 s, 64.8 °C for 1 min, 72 °C for 2 min, enters into 94 °C for 30 s, 44 °C for 1 min, 72 °C for 2 min, and finally into 72 °C for 10 min. After that, 1 μL from the primary PCR products was used as the template for the second nested-PCR reaction and the remaining

**Table 2 A list of primers used to amplify different deletions of the EcGS1b gene promoter**

| Primer     | Sequence(5′-3′)                          |
|------------|-----------------------------------------|
| GS1bSP1    | GAGAGTGTCCTCGTGCTTTGCTTC                |
| GS1bSP2    | GCTTGAAGCCATATCATGATCCAC-3              |
| GS1bSP3    | TCCCCCTTGGTGTGTAGCACATGC-3              |
| F66        | 5′-CCCAAGCTTTCATATCCCCACCCACCGCAT-3′   |
| F340       | 5′-CCCAAGCTTCAGTGTCTGTCAGC-3′           |
| F635       | 5′-CCCAAGCTTATCATACTACATACATAC-3′       |
| F838       | 5′-CCCAAGCTTCATTGGTATGCCTGTAAGGATG-3′   |
| R1257      | 5′-TCAGGTTGATCCTGGGATAC-3′              |
| qPCR-F     | 5′-TCAGGTTGATCCTGGGATAC-3′              |
| qPCR-R     | 5′-TCATCAATGGCTGCTGAGGTGTAAGGATG-3′     |
| Eact-F     | 5′-CATTCATTGGCTGCTGAGGTGTAAGGATG-3′     |
| Eact-R     | 5′-GGATGAGCTGTAAGGATGTAAGGATG-3′       |
was stored at −20 °C. The second nested PCR conditions started at 15 cycles of 94 °C for 30 s, 61.6 °C for 1 min, 72 °C for 2 min; 94 °C for 30 s, 61.6 °C for 1 min, 72 °C for 2 min; 94 °C for 30 s, 44 °C for 1 min, 72 °C for 2 min and then enters into 72 °C for 10 min. And 1 μL from the second nested-PCR reaction product was used as the third nested-PCR reaction template with the following conditions: 15 cycles of 94 °C for 30 s, 64.6 °C for 1 min, 72 °C for 2 min; 94 °C for 30 s, 44 °C for 1 min, 72 °C for 2 min and then entered into 72 °C for 10 min. The product of the third nested-PCR reaction was purified using Agarose Gel DNA Purification Kit (TaKaRa) and then sequencing was performed.

Analysis of EcGS1b-P sequence
The EcGS1b-P was analyzed with the BLAST program of NCBI using EcGS1b-P sequence. Putative cis-acting regulatory elements and the transcription initiation site were predicted using TSSP-TCM software and PlantCARE (http://bioinformatics.psb.ugent.be/webtools/plantcare/html/), respectively.

Plasmid constructions and plant transformation
A pair of primers (R1257 and F66 presented in Table 2) containing the restriction sites HindIII and BamHI were employed to amplify the putative EcGS1b-P region upstream to ATG, followed by cloning the amplified fragment into the pMD19-T vector and then sequencing. After the pMD19-T vector with EcGS1b-P was digested by HindIII and BamHI, the EcGS1b-P was sub-cloned into pBI121. Similarly, 3 more 5′-deletion promoters of EcGS1b-P along with 5′UTR (p900, p600 and p400) were prepared using different primers (F340, F635, and F838 in Table 2). The different promoters sub-cloned into pBI121 were named as p1200::GUS, p900::GUS, p600::GUS, and p400::GUS, respectively. These promoter constructions were integrated into Agrobacterium tumefaciens EHA105 and then transformed into tobacco leaf discs. The pBI121 and pBI101 vectors were also integrated into EHA105, and then transformed into tobacco respectively as the positive control and the negative control. At least two independent transgenic lines for each construct were selected for histochemical assay and GUS activity measurement.

Histochemical assay of GUS activity
Different transgenic tobacco blades were grown in a chamber at 25 °C in dark for 2 days, and then were washed 5–6 times using sterile water. The assays of GUS expression were implemented according to the method described by Jefferson [77]. Different samples were placed in GUS staining solution at 37 °C for overnight. After dyeing, the samples were put into 75% ethanol solution for 48 h. Next, the decolorization of the leaves was observed with eyes and microscope.

GUS activity measurement
Different tissues of transgenic tobacco were milled in liquid N and then placed in GUS extraction solution (0.05 molL−1 Na2HPO4, 0.05 molL−1 NaH2PO4, 0.01 molL−1 EDTA, 1 mL 10% SDS, 100 μL Triton X-100, and 100 μL β-mercaptoethanol, to 100 mL H2O) at a ratio of 100 mg of sample to 1 mL of GUS extraction solution. The mixed samples were centrifuged at 12,000 g for 10 min, and then 50 μL supernatant was used for GUS activity. In addition, 20 μL supernatant was transferred to 1.5 mL centrifuge tube for measuring the protein concentration using the Bradford method [78]. The 250 μL reaction solution was added into GUS activity reaction containing 50 μL supernatant and 200 μL GUS reaction buffer (25 mg 4-MUG added to 25 mL GUS extraction solution) after heated at 37 °C (4-MUG buffer 2 mmolL−1 4-MUG). Then 200 μL of reaction mixture solution was added to 1.8 mL stop buffer containing 0.2 molL−1 NaCO3 and the fluorescence level was measured immediately. The remaining reaction mixture was incubated at 37 °C for 60 min and then 200 μL of reaction mixture was added to the stop buffer. The fluorescence level of 4-methylumbelliferone (4-MU), which was the breakdown product, was determined using a fluorescence spectrophotometer at an excitation/emission wavelength of 350 nm/455 nm specifically for 4-MU. The 4-MU concentration was then determined from the standard curve. GUS activity was expressed as 4-MU nmol per minute and per milligram protein. Same tissues in at least two different lines of the same construct were mixed to do the GUS activity assay, and each was carried out three replicates. The values were expressed as means ± standard deviation. The significance of the difference was tested using the Duncan method by SPSS 11.0 software (IBM, USA).

EcGS1b expression analysis by real-time qRT-PCR
Total RNA was extracted by CTAB using Plant RNA Kit (HUAYUEYANG BIOTECHNOLOGY), and then the mixed genomic DNA was removed. This was used to synthesize the first-strand cDNA in 20 μL of reaction mixture using PrimeScript™ RT reagent Kit with gDNA Eraser (Takara). Real-time PCR was performed using the SYBR Premix EX Taq™ II (Takara) with specific primers, and E.crassipes actin (Accession number: KC505366) was used as an internal reference gene for qRT-PCR to normalize the target gene expression (primers qPCR-F, qPCR-R, Eact-F, Eact-R given in Table 2). The real-time PCR reaction was performed on a Roche LightCycler96 PCR instrument with the following conditions: 94 °C for 5 min, 30 cycles of 94 °C for 30 s, 58 °C for 30 s, 72 °C
for 1 min, and finally enters into 72 °C for 10 min. The qRT-PCR was implemented three biological replicates, and each biological replicate was performed three technical replicates. The 2^ΔΔCT method was used for quantitative analysis. The values were expressed as means ± standard deviation. The significance of the difference was tested using the Duncan method by SPSS 11.0 software (IBM, USA).

Abbreviations
GS: Glutamine synthetase; E.crassipes: Eichhornia crassipes; GUS: β-glucuronidase; qRT-PCR: Quantitative Reverse Transcription-Polymerase Chain Reaction; GOGAT: Glutamate synthase; TSS: Transcriptional start site; MUG: 4-methylumbelliferyl-β-D-glucuronide; MS: Murashige and Skoog; 4-MU: 4-methylumbellifereone

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Authors' contributions
F-MH designed the research and had the funding support. Z-YS and L-XD performed the research, analyzed the data and wrote the draft. They had the same contribution to this paper. D-ZW did some experiments and analyzed some data. L-ZQ involved in revising the manuscript and analyzing some data. All Authors have read and approved the final manuscript.

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Availability of data and materials
The E𝑐GSIΔ-P sequence is deposited in GenBank of NCBI and assigned the accession number MT154418. This sequence can be accessible with the following link: https://www.ncbi.nlm.nih.gov/nuccore/MT154418. All other data generated or analyzed during this study are included in this published article.

Ethics approval and consent to participate
Not applicable.

Consent for publication
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Competing interests
The authors declare that they have no conflict of interest.

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References
1. Yan SH, Song W, Guo JY. Advances in management and utilization of invasive water hyacinth (Eichhornia crassipes) in aquatic ecosystems - a review. Crit Rev Biotechnol. 2017;37(2):218–28.
2. Abdul-Reem AM, Al-Abd AM, Shalaby E, Abul-Ela F, Naar-Allah AA, Mahmoud AM, El-Shemy HA. Eichhornia crassipes (Mart) Solms: from water parasite to potential medicinal remedy. Plant Signal Behav. 2011;6(6):834–6.
3. Chu JJ, Ding Y, Zhuang QJ. Invasion and control of water hyacinth (Eichhornia crassipes) in China. J Zhejiang Univ Sci B. 2006;7(8):623–6.
4. Akinbile CO, Yusoff MS. Assessing water hyacinth (Eichhornia Crassipes) and lettuce (Lactuca sativa L.) effectiveness in aquaculture wastewater treatment. Int J Phytoremediat. 2012;14(3):201–11.
5. Henares MNP, Camargo AFM. Treatment efficiency of effluent prawn culture by wetland with floating aquatic macrophytes arranged in series/ Eficiencia de wetland con macrofitas acuaticas flutuantes dispostas em serie Para o tratamento do efluente do cultivo de camarao. Braz J Biol. 2014;74(4):906–12.
6. Wu X, Wu H, Ye J. Purification effects of two eco-ditch systems on Chinese soft-shelled turtle greenhouse culture wastewater pollution. Environ Sci Pollut Res Int. 2014;21(8):5610–8.
7. Rezania S, Ponraj M, Baleghizadeh A, Mohammadi SE, Dinh MF, Talib SM, Sabbagh F, Sairain FM. Perspectives of phytoremediation using water hyacinth for removal of heavy metals, organic and inorganic pollutants in wastewater. J Environ Manag. 2015;156:125–33.
8. Spaúndo-Tavares LH, Floriêncio T, Scardelli-Truzzi B. Aquaculture biological waste as culture medium to cultivation of Anistrodesmus gracilis (Reinsch) Korshikov. Braz J Biol. 2018;78(5):79–87.
9. Das A, Ghosh P, Paul T, Ghosh U, Pati BR, Mondal KC. Production of bioethanol as useful biofuel through the bioconversion of water hyacinth (Eichhornia crassipes). Biotechnol Biofuels. 2015;8:110.
10. Lu W, Wang C, Yang Z. The preparation of high calorific fuel (HCF) from water hyacinth by deoxy-liquefaction. Bioresour Technol. 2009;100(24):6451–6.
11. de Vasconcelos GA, Veras RM, de Lima SJ, Cardoso DB, de Castro SP, de Morais NN, Souza AC. Effect of water hyacinth (Eichhornia crassipes) hay inclusion in the diets of sheep. Trop Anim Health Prod. 2016;48(3):539–44.
12. El-Shinnawi MM, El-Din MMA, El-Shimi SA, Badawi MA. Biogas production from crop residues and aquatic weeds. Resour Conserv Recy. 1989;3(1):33–45.
13. Mishima D, Kuniki M, Sei K, Soda S, Ike M, Fujita M. Ethanol production from candidate energy crops: water hyacinth (Eichhornia crassipes) and water lettuce (Pistia stratiotes L.). Bioresour Technol. 2008;99(7):2495–500.
14. Bernard SM, Habash DZ. The importance of cytosolic glutamine synthetase in nitrogen assimilation and new perspectives in glutamine synthetase in grasses. J Exp Bot. 2010;61(24):4529–39.
15. Swarbreck D, Defoin-Platel M, Hindle M, Saj J, Habash DZ. New perspectives on glutamine synthetase in grasses. J Exp Bot. 2011;62(4):1511–22.
16. Van RJ, Belthall H, Abrav V, Sewell BT. Proteolysis of the type II glutamine synthetase from Bacteroides fragilis suggests a role in cell wall synthesis. Acta Crystallogr F. 2011;67(3):358–63.
17. King SG, Walker EL, Conuzzi GM. Glutamine-synthetase genes of pea encode distinct polypeptides which are differentially expressed in leaves, roots and nodules. EMBO J. 1987;6(11):1–9.
18. Cren M, Hirel B. Glutamine synthetase in higher plants regulation of gene and protein expression from the organ to the cell. Plant Cell Physiol. 1999;40(12):1187–93.
19. Zhang ZY, Xiong SP, Wei Y, Meng XD, Wang XC, Ma XM. The role of glutamine synthetase isozymes in enhancing nitrogen use efficiency of N-efficient winter wheat. Sci Rep. 2017;7:12.
20. Caputo C, Criado MV, Roberts IN, Gelso MA, Barneix AJ. Regulation of glutamine synthetase 1 and amino acids transport in the phloem of young wheat plants. Plant Physiol Bioch. 2009;47(5):335–42.
21. El Omari R, Rueda-Lopez M, Avila C, Crespiollo R, Nihii M, Canovas FM, Ammonium tolerance and the regulation of two cytosolic glutamine synthetases in the roots of sorghum. Func Plant Biol. 2010;37(1):55–63.
22. Goodall AJ, Kumar P, Tobin AJ. Identification and expression analyses of cytosolic glutamine synthetase genes in barley (Hordeum vulgare L.). Plant Cell Physiol. 2013;54(4):492–505.
23. Benfey PN, Ren L, Chua NH. Tissue-specific expression from a GUS 35S enhancer subdomains in early stages of plant development. EMBO J. 1990;9(6):1677–84.
24. Shahmuradov IA, Gammerman AJ, Hancock JM, Bramley PM, Solovyev VV. PlantProm: a database of plant promoter sequences. Nucleic Acids Res. 2003;31(1):114–7.
25. Yamamoto YY, Ichida H, Matsui M, Obokata J, Sakurai T, Satou M, Seki M, Shimozaki K, Abe T. Identification of plant promoter constitutively by analysis of local distribution of short sequences. BMC Genomics. 2007;8(1):167.
26. Prändl R, Schöffl F. Heat shock elements are involved in heat shock promoter activation during tobacco seed maturation. Plant Mol Biol. 1996;31(1):157–62.
27. Belitangi NG, Guerzoni JTS, Moreira RMP, Vieira LGE. Improving low-temperature tolerance in sugarcane by expressing the ipt gene under a cold inducible promoter. Biotechnol Lett. 2012;34(12):1167–74.
28. Xu X, Guo S, Chen K, Song H, Liu J, Guo L, Qian Q, Wang H. A 796 bp PpR10 gene promoter fragment increased root-specific expression of the GUS reporter gene under the abiotic stresses and signal molecules in tobacco. Biotechnol Lett. 2010;32(10):1533–4.
29. Bezhani S, Sherameti I, Pfannschmidt T, Gilmuller R. A repressor with similarities to prokaryotic and eukaryotic DNA helicases controls the
assembly of the CAAT box binding complex at a photosynthesis gene promoter. J Biol Chem. 2001;276(26):23785–90.

30. Joshi CP. An inspection of the domain between putative TATA box and translation start site in 79 plant genes. Nucleic Acids Res. 1987;15(16):6463–9.

31. Qin H, Zhang Z, Liu M, Wang Y, Wen X, Yan S, Zhang Y, Liu H. Efficient assimilation of cyanobacterial nitrogen by water hyacinth. Bioresour Technol. 2017;241:1197–200.

32. Zhang YY, Liu HQ, Yan SH, Wen XZ, Qin HJ, Wang Z, Zhang ZY. Phosphorus removal from the hyper-eutrophic Lake Caohai (China) with large-scale water hyacinth cultivation. Environ Sci Pollut R. 2019;26(13):12975–84.

33. Andrade HM, Oliveira JA, Famese FS, Ribiero C, Silva AA, Campos FV, Neto JL. Arsenic toxicity: cell signalling and the attenuating effect of nitric oxide in Eichhornia crassipes. Biol Plant. 2016;60(1):173–80.

34. de Souza Reis IR, de Oliveira JA, Ventrecca MC, Otone WC, Marinato CS, de Matos LP. Involvement of glutathione metabolism in Eichhornia crassipes tolerance to arsenic. Plant Biol. 2020;22(2):346–50.

35. Malik A. Environmental challenge vs a Vis opportunity: the case of water hyacinth. Environ Int. 2007;33(1):122–38.

36. Moura Júnior EG, Pott A, Severi W, Zickel CS. Response of aquatic macrophytes biomass to limnological changes under water level fluctuation in tropical reservoirs. Braz J Biol. 2019;79(1):7.

37. Fu M, Jiang L, Li Y, Yan G, Jinping P. Identification of gene fragments related to nitrogen deficiency in Eichhornia crassipes (Pontederiaceae). Rev Biol Trop. 2014;62(4):1637–48.

38. Fu J, Jiang L, Yu C. Identification of G5IA, G5IB and G5IC genes from Eichhornia crassipes and their transcript analysis in response to different nitrogen sources. Pak J Bot. 2018;50(6):2197–204.

39. Thomsen HC, Ekselsson D, Møller IS, Schjoerring JK. Cytosolic glutamine synthetase: a target for improvement of crop nitrogen use efficiency? Trends Plant Sci. 2014;19(10):656–63.

40. Gao YJ, de Bang TC, Schjoerring JK. Cisgenic overexpression of cytosolic glutamine synthetase increases nitrogen utilization efficiency in barley and prevents grain protein decline under elevated CO2. Plant Biotechnol J. 2019;17(1):209–21.

41. Hu MY, Zhao XQ, Liu Q, Sun C, Liu H, Hong X, Zhang W, Zhang Y, Sun J, Zhang A, Yang W. The 160 bp insertion in the promoter of Rht-B1B plays a vital role in increasing wheat height. Front Plant Sci. 2016;7:307.

42. Srivastava AC, Blancaflor EB. The foployglutamate synthetase plastidial isoform is required for postembryonic root development in Arabidopsis. Plant Physiol. 2011;155(5):1237–51.

43. Chao D, Yuanchun M, Dan Z, Michael W, Zong-Ming C. Meta-analysis of the cis-regulation of the glnA promoter in anthocyanins-producing plants. Int J Mol Sci. 2015;16(11):27032–49.

44. Rueda-Lopez M, Crespillo R, Canovas FM, Avila C. Differential regulation of two glutamine synthetase genes by a single Dof transcription factor. Plant J. 2016;84(2):326–37.

45. Gallardo F, Fu J, Canton FR, Garcia-Gutierrez A, Canovas FM, Kirby EG. Expression of a conifer glutamine synthetase gene in transgenic poplar. Planta. 1999;210(1):19–26.

46. Wei YH, Shi AB, Jia XT, Zhang ZY, Ma XM, Gu MX, Meng XD, Wang XC. Nitrogen supply and leaf age affect the expression of TaGS1 or TaGS2 driven by a constitutive promoter in transgenic tobacco. Genes. 2018;9(8):18.

47. Koyama T, Ono T, Shimizu M, Jinbo T, Mizuno R, Tomita K, Mitsukawa N, Kawazu T, Kimura T, Ohmima K, et al. Promoter of Arabidopsis thaliana phosphate transporter gene drives root-specific expression of transgene in rice. J Biosci Bioeng. 2005;99(1):38–42.

48. Chao D, Yuanchan M, Dan Z, Michael W, Zong-Ming C. Meta-analysis of the effect of overexpression of dehydration-responsible element binding family genes on temperature stress tolerance and related responses. Front Plant Sci. 2018;9:15.

49. Li Z, Srivastava R, Tang J, Zheng Z, Howell SH. Cis-effects condition the induction of a major unfolded protein response factor, ZmZIP60, in response to heat stress in maize. Front Plant Sci. 2018;9:833.

50. Nitz J, Berkelfeld H, Puzio PS, Grundler FM, Pyklo I, a seedling and root specific gene and promoter from Arabidopsis thaliana. Plant Sci. 2001;161(2):337–46.

51. Nonogaki M, Sali K, Nambara E, Nonogaki H. Amplification of ABA biosynthesis and signaling through a positive feedback mechanism in seeds. Plant J. 2014;78(3):527–39.

52. Chen F, Schum J, Petsch M, Gershonien J, Bohlmann J, Pichersky E, Tholl D. Characterization of a root-specific Arabidopsis terpene synthase responsible for the formation of the volatile monoterpene 1,8-cineole. Plant Physiol. 2005;137(4):1556–66.

53. Chen L, Jiang B, Wu C, Sun S, Hou W, Han T. GmPRP2 promoter drives root-specific expression in transgenic Arabidopsis and soybean hairy roots. BMC Plant Biol. 2014;14(1):245.

54. Lou X, Li X, Li A, Pu M, Shoaib M, Liu D, Sun J, Zhang A, Yang W. The 160 bp insertion in the promoter of Rht-B1B plays a vital role in increasing wheat height. Front Plant Sci. 2016;7(307).

55. Srivastava AC, Blancaflor EB. The foployglutamate synthetase plastidial isoform is required for postembryonic root development in Arabidopsis. Plant Physiol. 2011;155(5):1237–51.

56. Shah S, Souare A, Deeba F, Sufiana T, Dokovic-Schulze S, Chen C, Naqui SMS. Transgenic analysis reveals 9′ abbreviated OsRGLP2 promoter(s) as responsive to abiotic stresses. Mol Biotechnol. 2017;59(11-12):459–68.

57. Siebertz B, Logemann J, Willmitzer L, Schell J. cis-analysis of the wound-inducible promoter wu1 in transgenic tobacco plants and histochemical localization of its expression. Plant Cell. 1989;1(10):961–8.

58. Zhang H, Hou J, Jiang P, Qi S, Xu C, He Q, Ding Z, Wang Z, Zhang K, Li K. Identification of a 467 bp promoter of maize phosphatidylchinositol synthase gene (ZmPls) which confers high-level gene expression and salinity or osmotic stress inducibility in transgenic tobacco. Front Plant Sci. 2016;7:42.

59. Niu GL, Gou W, Han XL, Qin C, Zhang LX, Abomohra AE, Ashraf M. Cloning and functional analysis of phosphoethanolamine methyltransferase promoter from maize (Zeas mays L.). Int J Mol Sci. 2018;19(11):191–203.

60. Yu Z, Han YN, Xiao XG. A PPO promoter from betalain-producing red Swiss chard, directs petiole- and root-preferential expression of foreign genes in anthocyanins-producing plants. Int J Mol Sci. 2015;16(11):27032–43.

61. Rueda-Lopez M, Crespo R, Canovas FM, Avila C. Differential regulation of two glutamine synthetase genes by a single DoFR transcription factor. Plant J. 2008;56(1):73–85.

62. Wang YL, Liu F, Wang W. Kinetics of transcription initiation directed by multiple cis-regulatory elements on the glnA2 promoter. Nucleic Acids Res. 2014;42(22):10360–8.

63. Kwon HB, Park SC, Peng HP, Goodman HM, Dewdney SH. Identification of a light-responsive region of the nuclear encoding the B subunit of chloroplast glycerol-3-phosphate dehydrogenase from Arabidopsis thaliana. Plant Physiol. 1994;105(3):357–67.

64. Xue M, Long Y, Zhao Z, Huang G, Huang K, Zhang T, Jiang Y, Yuan Q, Pei X. Isolation and characterization of a green-tissue promoter from common water hyacinth. Oryza rufipogon Griff., Int J Mol Sci. 2018;19(7):209–21.

65. Xu L, Ye R, Zheng Y, Wang Z, Zhou P, Lin Y, Li D. Isolation of the endornase-specific LPAAT gene promoter from coconut (Cocos nucifera L.) and its functional analysis in transgenic rice plants. Plant Cell Rep. 2010;29(9):1061–8.

66. Huang Z, Cao Z, He Y, Li Y, Liu X, Mu H. Functional analysis of a rice late pollen-abundant UDP-glucose pyrophosphorylase (OsSulg2) promoter. Mol Biol Rep. 2011;38(7):4291–302.
73. Rabot A, Portemer V, Peron T, Mortreau E, Leduc N, Hamama L, Couto-
Thevenot P, Atanassova R, Sakr S, Le Gourrierec J. Interplay of sugar, light
and gibberellins in expression of Rosa hybrida vacuolar invertase 1
regulation. Plant Cell Physiol. 2014;55(10):1734–48.
74. Li X, Sui X, Zhao W, Huang H, Chen Y, Zhang Z. Characterization of
cucumber violaxanthin de-epoxidase gene promoter in Arabidopsis. J Biosci
Bioeng. 2015;119(4):470–7.
75. Gangappa SN, Maurya JP, Yadav V, Chattopadhyay S. The regulation of the
Z- and G-box containing promoters by light signaling components, SPA1
and MYC2, in Arabidopsis. PLoS One. 2013;8(8):e62194.
76. Hoecker U, Tepperman JM, Quail PH. SPA1, a WD-repeat protein specific to
phytochrome a signal transduction. Science. 1999;284(5413):496–9.
77. Jefferton RA, Kavanagh TA, Bevan MW. GUS fusions: beta-glucuronidase as a
sensitive and versatile gene fusion marker in higher plants. EMBO J. 1987;
6(13):3901–7.
78. Bradford MM. A rapid and sensitive method for the quantitation of
microgram quantities of protein utilizing the principle of protein-dye
binding. Anal Biochem. 1976;72(1–2):248–54.

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