Heterologous expression of cyanobacterial PCS confers augmented arsenic and cadmium stress tolerance and higher artemisinin in Artemisia annua hairy roots

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Abstract
The present study provides the first report of heterologous expression of phytochelatin synthase from Anabaena PCC 7120 (anaPCS) into the hairy roots of Artemisia annua. Transformed hairy roots of A. annua expressing anaPCS gene showed better tolerance to heavy metals, viz., arsenic (As) and cadmium (Cd) owing to 143 and 191% more As- and Cd-accumulation, respectively, as compared to normal roots with a bioconcentration factor (BCF) of 9.7 and 21.1 for As and Cd, respectively. Under As and Cd stresses, transformed hairy roots possessed significantly higher amounts of phytochelatins and thiols probably due to the presence of both AaPCS (Artemisia annua PCS) and anaPCS. In addition, artemisinin synthesis was also induced in transformed hairy roots under heavy metals stresses. In-silico analysis revealed the presence of conserved motifs in both AaPCS and anaPCS sequences as well as structural modelling of PCS functional domain was conducted. Interaction of AaPCS and anaPCS proteins with CdCl2 and sodium arsenate gene ontology analysis gave insights to anaPCS functioning in transformed hairy roots of A. annua. The study provides transformed hairy roots of A. annua as an efficient tool for effective phytoremediation with added advantages of artemisinin extraction from hairy roots used for phytoremediation.

Keywords Phytochelatin synthase · Artemisia annua · Hairy root · Anabaena PCC 7120 · Arsenic · Cadmium

Introduction
Arsenic (As) and cadmium (Cd) are among the major soil pollutants due to their high toxicity and long persistence. Both As and Cd are non-essential elements and adversely affect plant growth and development even at very low concentration in the soil.

The entry of As in plants is concentration gradient dependent. Out of four oxidation states of As [i.e., As(0), As(-III), As(III) and As(V)], only arsenate [As(V)] and arsenite [As(III)] enter into the plant roots via different transporter proteins. Among arsenate and arsenite forms of As, arsenite is more toxic (Khalid et al. 2017). As(V) is a chemical analogue of phosphorus and hence uses Pi transporters, consisting of Pi transporter proteins (PHT), to enter into the plant (LeBlanc et al. 2013). Inside the cell, As (V) is converted into As (III) by the action of arsenate reductase (Bleecker et al. 2006). On the other hand, As(III) enters into the plant cells through Nodulin 26-like intrinsic proteins (NIPs) belonging to the class of plant aquaporins (Mukhopadhyay et al. 2014). Cd uptake takes place through several metal transporters such as ZIP (ZRT- and IRT-like proteins) family, YSL (Yellow Stripe-Like) transporters and NRAMP (natural resistance-associated macrophage protein) family of transporters. Cd is also known to enter plant symplast via passive transport or using Ca2+ channels present on guard cell plasma membrane (Ismael et al. 2019).

Once inside the plant, these heavy metals are immobilized, chelated and compartmentalized through metal-binding ligands which include organic acids, polypeptides or even small peptides. One such metal-binding polypeptide is phytochelatin (PC) which mitigate the harmful effects of heavy metal by chelation (Zhang et al. 2013). PCs are enzymatically synthesized from glutathione by...
phytochelatin synthase (PCS). Heavy metals significantly elevate the biosynthesis of PCs. Metal ions including both anions (for example arsenate) and cations (for example cadmium) tend to increase the level of PCs (Cobbett 2000). Cloning of PCS gene has been achieved in many plant species such as *Arabidopsis*, rice, and wheat, and overexpression of PCS has been linked with greater heavy metal tolerance (Vatamaniuk et al. 1999).

Plant’s potential to uptake heavy metals can be easily evaluated by calculating bioconcentration factor. Plants with a value of bioconcentration factor > 1 for any heavy metal has been suggested to be categorized as bio-accumulators and could be beneficial for phytoextraction (Usman et al. 2013).

Hairy roots are formed after infection with a Gram-negative bacterium *Agrobacterium rhizogenes*, which transfers its T-DNA region (transfer DNA) into the host cell, integrates with the plant genome and expresses. Cloning and expression of recombinant proteins can be achieved by using many systems such as bacterial, yeast expression system or mammalian cells. However, each of them poses some or other limitations such as risks of viral infections in bacterial systems, toxic molecules in the mammalian system or high production cost (Gutierrez-Valdes et al. 2020). For a couple of decade’s plants, systems are being used for several advantages including low cost. Hairy root cultures have emerged as a potential expression system with low maintenance cost and high efficiency. As hairy roots are derived from the wounded region of plants, they possess the genetic makeup of the plant part from which they are derived and hence do not behave like a true root. This property makes hairy root cultures as an excellent expression system. Hairy roots have found their potential application for phyto remediation of heavy metals (Agostini et al. 2013).

*Artemisia annua*, popularly known for its antimalarial drug artemisinin, has been found to act as a hyperaccumulator of arsenic. In our previous studies (Kumari et al. 2017; Rai et al. 2014), *A. annua* was found to survive well even at high As-concentration and have hyperaccumulating capacity. Many heavy metal stresses including both As (Kumari et al. 2017) and Cd (Li et al. 2012) have been reported to induce biosynthesis of artemisinin. Various reports have been published for hairy root induction in *A. annua* for enhanced secondary metabolite production (Patra and Srivastava 2014). *A. annua* hairy root potential for phyto remediation of heavy metals has not been investigated. In the present investigation, we have cloned a cyanobacterial PCS gene (from *Anabaena* PCC 7120) into hairy roots of *A. annua*. We hypothesize that transformed hairy roots overexpressing PCS gene can be used as an excellent tool for phyto remediation of arsenic and cadmium.

### Materials and methods

#### Plant material and hairy root induction

Uniform seeds of *Artemisia annua* were surface sterilized in sodium hypochlorite (10% w/v) solution for 15–20 min and washed three-to-four times in sterilized water. Seeds were then treated with ethanol (70% v/v) for 30 s and were then allowed to germinate on Murashige and Skoog (MS) medium containing sucrose (3% w/v) at pH 5.8. Germination initiated under the controlled condition of light and temperature (16/8 h light/dark regime and 25 ± 2 °C). The in-vitro raised plantlets were then used for induction of hairy roots by infecting the leaves/stem with *Agrobacterium rhizogenes* ATCC 15834 maintained in liquid YMB medium with O.D. at 600 nm ranged between 0.9 and 1.0 (Hooykass et al. 1977). The infected plantlets were then cultured on full-strength MS medium for 2 days and then transferred to half-strength MS medium containing cefotaxime (250 mg l⁻¹) under the same growth conditions. The plantlets were further sub-cultured by transferring them to new antibiotic supplemented medium for 2 weeks, and finally, the bacterial-free hairy roots were cultured on liquid MS medium. Hairy root cultures were incubated in dark inside the rotary shaker with constant rotation (100 rpm) and were sub-cultured twice a month into the fresh medium.

For experiments, three tissue samples were taken: normal roots (which were derived from plantlets), hairy roots (derived from leaves by *A. rhizogenes* infection) and transformed hairy roots (also derived from leaves by *A. rhizogenes* infection and expressing anaPCS gene).

#### Cyanobacterial strain and plasmids

*Anabaena* sp. PCC 7120 was photoautotrophically grown in BG-11 medium (without any source of combined nitrogen) prepared in Tris buffer with a photoperiod of 14:10 h under light (day light fluorescent tubes) intensity of 72 µmol m⁻² s⁻¹ PAR (photosynthetically active radiation) at 24 ± 2 °C (Rippka et al. 1979). The *E. coli* strains DH5α was used for cloning and BL21 (Novagen) for overexpression. The cells possessing recombinant plasmid were transferred on fresh Luria–Bertani (LB) medium supplemented with 100 µg ml⁻¹ ampicillin (Sambrook and Russell 2001). For cloning, pGEX-5X-2 was used as a vector (Chaurasia et al. 2008).
Cloning and confirmation of anaPCS gene and transformation in A. annua hairy roots

Total genomic DNA from Anabaena PCC7120 was isolated as per the standard CTAB method (Srivastava et al. 2007) and the region encoding phytochelin synthase (alr0975, denoted as anaPCS in the manuscript) was amplified using gene-specific primers having restriction sites for EcoRI and NotI (Chaurasia et al. 2008). The desired gene amplicon was then purified by the QIAquick gel extraction kit (Qiagen), and cloned in pGEX-5X-2 vector (Novagen) containing EcoRI and NotI restriction sites, and the resulting construct pGEX-5X-2-anaPCS was used to confirm anaPCS nucleotide sequence verification and expression analysis according to Chaurasia et al. (2008). For heterologous expression analysis, anaPCS gene was amplified using PF 5′–GCTCTAGAC GATAGTTAGAAACTCTTTA–3′, and PR 5′–GCAAGAG CTCCTAATCTTGTGTTTTACTTAC–3′, containing XbaI and SacI restriction sites and inserted into a pBII121 (Clontech, USA) under the control of promoter CMV35S and as binary vector introduced into the A. rhizogenes ATCC 15834 by electroporation for further transformation of A. annua hairy roots and expression studies. The confirmation of the resulting recombinant vector pBII121-anaPCS was done by performing double digestion of the plasmid with XbaI and SacI enzymes and its sequencing.

Plant stress treatments and measurement of arsenic and cadmium content

To assess the role of anaPCS in conferring heavy metal tolerance, A. annua hairy roots transformed with pBII121-anaPCS, non-transformed hairy roots (empty vector), and normal roots were subjected to arsenic (100 µM) and cadmium stress (50 µM) using sodium arsenate (Na2HAsO4·7H2O) and cadmium chloride (CdCl2), respectively, for 5 days. The treatments were divided into three sets, set 1 with no treatment (control), set 2 with 100 µM of arsenate (Na2HAsO4·7H2O) and set 3 with CdCl2. Each treatment set contains three groups of samples, untransformed hairy roots, transformed hairy roots and normal roots of the plant.

All the experiments were performed in triplicates. The accumulation of arsenic content in treated/non-treated hairy roots and normal roots was determined by the SEM–EDS method and was estimated by atomic absorption spectrometry (Perkin-Elmer A Analyst 600) as described by Kumari et al. (2018).

Estimation of phytochelin synthase, reduced glutathione and thiol

Phytochelin content was determined as described earlier (Rai et al. 2011). For glutathione estimation, 0.5 g tissue was homogenized in an extraction buffer containing sulphosalicylic acid (5% w/v). The absorbance of the reaction mixture containing potassium phosphate buffer (100 mM, pH 7.0), DTNB (5′-dithiobis-2-nitrobenzoic acid) and tissue extract was read at 412 nm (Anderson 1985). For total thiol estimation, 0.5 g tissue was homogenized in extraction buffer containing ascorbate buffer (20 mM) and EDTA (20 mM), and after centrifugation at 12,000×g for 15 min (4 ºC), 0.5 ml supernatant was mixed with reaction mixture that contained Tris–HCl buffer (200 mM) and DTNB (10 mM). Reaction mixture developed yellow color after incubation for 20 min at 30 ºC and absorbance was read at 412 nm (Nagalakshmi and Prasad 2001).

Measurement of antioxidative enzymes

Fresh tissues (500 mg) were homogenized with 5 ml Tris buffer (50 mM, pH = 7.0) that contains EDTA (2 mM), PMSF (1 mM), and PVP (1%). After centrifugation of homogenate at 12,000×g for 15 min, immediately the supernatant was used for the assay.

Antioxidative enzymes catalase, ascorbate peroxidase, glutathione reductase and peroxidases were measured as described earlier (Kumari et al. 2017; Pandey and Pandey-Rai 2014). All the spectrophotometric analysis was carried out using Hitachi U-2910 spectrophotometer.

Expression analysis of rolB and PCS gene and artemisinin biosynthetic pathway genes

Total RNA was isolated using Trizol reagent (Invitrogen) as per the manufacturer’s recommendation. One microgram of total RNA was used as template for cDNA synthesis with cDNA synthesis kit (Bio-Rad Laboratories, USA) following the manufacturer’s instructions.

RT-PCR analysis using primers listed in supplementary table 1 was performed by following the method described earlier (Pandey and Pandey-Rai 2014). All reactions were run in triplicate along with a negative control. Primer accuracy was tested using gDNA as positive control. PCR products were run on agarose gel and band intensities were measured using Quantity One software (Bio-Rad) installed with Gel Doc 2000 system (Bio-Rad).

PCR was performed in reaction mixture (50 µl final volume) containing 50 ng cDNA, 2.5 µl 10× PCR buffer with 15 mM MgCl2, 200 µM dNTPs, 10 pmol of each primer and 0.2 U Taq DNA polymerase (Bangalore Genei, India) in an iCycler (Bio-Rad, USA). The thermal amplification protocol
was as follows: 1 cycle at 95 °C for 5 min, and 30 cycles at 95 °C for 30 s, 59 °C (value varied for different primer sets) for 45 s and 72 °C for 45 s.

**Artemisinin estimation**

Artemisinin was extracted and prepared according to Zhao and Zeng (1985) and Pu et al. (2009) with minor modifications. Approximately 100 mg dry tissue was extracted in 20 ml petroleum ether by ultrasonication (35 min). The extract was evaporated and dissolved in 1 ml ethanol. After 1 min centrifugation at 10,000×g, 0.1 ml of the supernatant was treated with 0.4 ml of 0.2% (w/v) NaOH at 45 °C for 35 min and cooled to room temperature. The solution was acidified with 0.5 ml of 0.05 M acetic acid and filtered through a Millipore filter (0.45 µm) before HPLC analysis. HPLC analysis conditions: Waters 600E HPLC system equipped with 150 × 9.46 mm Hypersil BDS C8 column; injected samples (10 µl) were eluted with methanol/0.01 M Na2HPO4–NaH2PO4 buffer (pH 7) (2:3, v/v) at 1 ml min−1 and monitored at 260 nm. The concentration of artemisinin was measured using the standard curve.

**In-silico analysis for comparative phylogeny, structural and functional analysis of anaPCS and AaPCS gene.**

The sequence of phytochelatin synthase (PCS) gene in A. annua (AaPCS) and Anabaena PCC 7120 (anaPCS) was obtained using BLAST-P (http://blast.ncbi.nlm.nih.gov/Blast.cgi) by taking homology with the Arabidopsis phytochelatin synthase (AtPCS) gene (Locus: AAD41794.1) identified from TAIR (https://www.arabidopsis.org/). The sequences with maximum query coverage and percentage identity (> 90%) were selected for the phylogenetic tree construction and sequential characterization. The sequences were aligned using Bio-Edit tool (Hall 1999) and phylogenetic tree was constructed using MEGA 7 suite (Tamura et al. 2013). The result of BLAST-P search was further subjected to PDB database to evaluate the similarity indices of the sequences and confirmed by circos visualization tool (Krzywinski et al. 2009). All the identified probable PCS protein sequences were subjected to multiple sequence alignment analysis using CLC bio workbench and with the help of Clustal W (http://www.ebi.ac.uk/tools/msa/clustalw2/) for their sequential classification (Krzywinski et al. 2009). The functional domain regions occupied by PCS proteins were searched using InteProScan http://www.ebi.ac.uk/Tools/pfa/interproscan (Jones et al. 2014), NCBI CDD server http://www.ncbi.nlm.nih.gov/Structure/cdd/cdd.shtml (Marchler-Bauer et al. 2011) and ExPASy-Prosite scan http://prosite.expasy.org/scanprosite (de Castro et al. 2006). The occurrence of potential motifs in PCS protein in all the members was identified using MEME Suite 4.1.1.2 (Multiple EM for Motif Elicitation) (Bailey et al. 2006) by keeping the selection parameter to any number of repeats with motif width of 10 and 30 residues and by keeping a maximum number of motifs to 40.

**Structural modelling and model validation**

The 3D homology modelling of identified protein sequences for PCS protein in A. annua and Anabaena PCC 7120 (AaPCS and anaPCS) was done using template proteins available in protein database which were searched using BLAST-P programme of Protein Data Bank (www.rcsb.org/pdb/) (Berman et al. 2000). The result of BLAST-P analysis showed most significant and accurate templates for the modelling of functional domain structures of AaPCS and anaPCS proteins. Three most closely related proteins were selected for the homology modelling of AaPCS and anaPCS proteins using MODELLER module of Discovery Studio 3.0 (accelrys.com; Shahi et al. 2013). Five models were generated by the MODELLER using Crystal structure of anaPCS (template 1; PDB ID: 2BTW), Crystal structure of a putative C39-like peptidase from Bacillus anthracis (template 2; PDB ID: 3ERV) and Crystal structure of the N-terminal peptidase C39-like domain of the toxin secretion ATP-binding protein from Vibrio parahaemolyticus (template 3; PDB ID: 3B79). The model with lowest Discrete Optimized Protein Energy (DOPE) score was further refined for Cα traces by ModRefiner using two-step atomic level energy minimization module (Xu and Zhang 2011). The modelled proteins were then allowed to superimpose with each template using Superpose version 1.0 (Maiti et al. 2004) to access the topological details of the modelled proteins. The predicted models were further evaluated qualitatively in terms of their geometric analysis, stereochemical orientation and backbone conformation patterns of nonbonded atomic interactions. For the backbone conformations, the predicted models were subjected to Ramachandran plot analysis for evaluating the backbone dihedral φ (φ) and psi (Ψ) angles using PROCHECK module of PDBSum server http://www.ebi.ac.uk/pdbsum/ (Laskowski et al. 2005) which was further attested by RAMPAGE server (Lovell et al. 2003). The models were also evaluated by a single model method using ProSA (Wiederstein and Sippl 2007), Qmean (Bencert et al 2009), RESPROX (Resolution by Proxy), ERRAT (Colovos and Yeates 1993). For the quantitative evaluation, the models were subjected to VADAR (Volume, Area, Dihedral Angle Reporter) analysis (Willard et al. 2003). The protein models generated for the functional domain were further submitted to an online repository of PMDB (Castrignano et al. 2006) to obtain the accession identities.
Ligand modelling and protein docking

The molecular docking studies between ligands (Cadmium chloride; CdCl₂ and sodium arsenate; Na₂HAsO₄.7H₂O) and the AaPCS and anaPCS protein sequences were carried out using Hex 8.0 molecular docking server (Macindoe et al. 2010) to reveal the probable binding modes for ligand compounds with in the active site of PCS protein. The ligands, viz., CdCl₂ and sodium arsenate, were drawn in 2D structure using Chemdraw software (ChemDraw, CambridgeSoft Corporation, 875 Massachusetts Avenue, Cambridge, MA 02139, USA). The 3D structures of both AaPCS and anaPCS proteins were generated using MODELLER module of Discovery studio 3.0 and were further explored for molecular interaction studies with ligands using Hex 8.0 molecular docking server. The protein and ligand interaction studies were performed following correlation criterion, i.e., DARS + Shape + Electro, FFT Mode-3D fast lite and grid range of 0.6 with Receptor:Ligand:Twist:Distance range of 180:180:360:40. Several conformations of docked complexes were obtained; however, the docked complex having lowest binding energy values was analyzed by DS Studio 3.0 for identifying the key residues involved in the interaction with CdCl₂ and sodium arsenate.

Active site prediction

The prediction of transition metal-binding sites and identification of potential residues involved in the making of the active sites with AaPCS and anaPCS was done using metapocket server http://metapocket.eml.org (Huang 2009).

Statistical analysis

All the experiments were performed in three biological replications and subjected to analysis of variance (ANOVA). The mean differences were compared by performing Duncan’s multiple range test (DMRT) using SPSS software (SPSS Inc., Version 20.0) and values at P ≤ 0.05 were considered significant.

Results

Induction and establishment of normal roots, hairy roots and transformed hairy root

Plantlets of A. annua were successfully established under in-vitro conditions (supplementary Fig. 1) and produced visible roots after 15 days of inoculation onto the MS medium. Infection by A. rhizogenes carrying binary vector (pBI121-anaPCS) and co-culturing of infected A. annua leaves has resulted in transformed hairy root (containing anaPCS gene) induction after 3 week of inoculation at a relative transformation frequency of 71%. Normal hairy roots not carrying anaPCS gene were also obtained in 15 days by infecting A. annua leaves with A. rhizogenes and co-culturing leaf discs. The presence of rolB gene in normal hairy roots and transformed hairy roots (with anaPCS) confirmed their origin after A. rhizogenes infection. The rolB gene was absent in normal roots (Fig. 1a, b).

Transformed hairy roots expressing anaPCS show enhanced tolerance (accumulation) to heavy metal stresses

In A. annua, hairy roots accumulated very high amount of As and Cd as compared to normal roots under As- and Cd-stress, respectively (Fig. 2). However, hairy roots expressing anaPCS gene accumulated even higher amount of both heavy metals as compared to non-transformed hairy roots. Under As-stress, non-transformed hairy roots and transformed hairy roots accumulated 91.3% and 143.4% higher amount of As, respectively, as compared to normal roots of A. annua. Among hairy roots, 72.7% greater As accumulation was observed in transformed hairy roots expressing anaPCS gene. Similar trend was observed for Cd-accumulation, where non-transformed hairy roots and transformed hairy roots accumulated 36% and
191% more Cd as compared to normal roots. Further among hairy roots, transformed hairy roots showed 55.4% greater Cd-accumulation as compared to non-transformed hairy roots.

The bioconcentration factor (BCF) for As was found to be 3.71, 5.71 and 9.7 for normal roots, non-transformed hairy roots and transformed hairy roots, respectively (Fig. 3). Similarly, for Cd, bioconcentration factor was 9.4, 12.7 and 21.1 for normal roots, non-transformed hairy roots and transformed hairy roots, respectively. The results showed that bioconcentration of Cd is higher than As and hairy roots expressing *anaPCS* gene have greater BCF for both As and Cd as compared to non-transformed hairy roots.

Normal roots, hairy roots (with only *rol* gene) and transformed hairy roots (with *rol* and *anaPCS* gene) have morphological differences in terms of average weight and average weight increase and are listed in supplementary table 6. As visible in the Supplementary Fig. 10, branching pattern is also different among normal, hairy and transformed hairy roots. Transformed hairy roots (with *anaPCS* gene) had more branching as compared to normal and hairy roots (without *anaPCS* gene).

**Phytochelatins and phytochelatin synthase expression**

Heavy metal stresses enhanced the phytochelatin content in all the tissue samples; however, the concentration was significantly more in hairy roots rather than normal roots (Fig. 4a). As-stress boosted phytochelatin content up to 28.3%, 44.7% and 14.7% in normal roots, hairy roots and transformed hairy roots, respectively, as compared to their respective control. Cd-stress enhanced phytochelatin level up to 25%, 35.7% and 9.52% in normal, hairy roots and transformed hairy roots, respectively, in comparison to their respective control. Although the stress-induced percent increase of phytochelatin was comparatively less prominent in transformed hairy roots, the amount of phytochelatin was more. Under conditions of As- and Cd-stress, transformed hairy roots expressing *anaPCS* gene was shown to contain 35.3% and 37.7% more phytochelatin as compared to non-transformed hairy roots.

The expression of phytochelatin synthase was measured through RT-PCR. Expression of *anaPCS* was observed only in transformed hairy roots, and because normal roots and hairy roots were not carrying *anaPCS* gene copy, no expression was observed in them (Fig. 4b). Expression of *anaPCS* was reported to be induced up to 2.1- and 2.3-fold under As- and Cd-treatment, respectively, as compared to control. Expression of *AaPCS* (*Artemisia annua PCS*) was observed in all the tissue samples. *AaPCS* expression was significantly induced under As- and Cd-stress in hairy roots and transformed hairy roots; however, no significant elevation in *AaPCS* was observed in normal roots under heavy metal stress conditions.
Thiol and GSH level in normal roots, hairy roots and transformed hairy roots under As- and Cd-stress

In control samples (without heavy metal stress) of normal roots, hairy roots and transformed hairy roots, thiol content was found to 11.2, 12.2 and 21.1 µM g⁻¹ FW (Fig. 5a). Under condition of As-stress, thiol content was elevated up to 43.75%, 59.83% and 87.2% in normal roots, hairy roots and transformed hairy roots, respectively, as compared to their respective controls. However, under Cd-stress, the percentage increase was 39.2%, 38.5% and 52.6% for normal roots, hairy roots and transformed hairy roots, respectively, as compared to their respective control. Among hairy roots, thiol content was increased up to 102.5% under As-stress and 90.5% under Cd-stress in transformed hairy roots as compared to non-transformed hairy roots.

In contrast to thiol content, GSH level (Fig. 5b) was reduced under both heavy metal stresses in all tissue samples. Under As-stress 6.8%, 34.4% and 50.6% reduction in GSH level was observed in normal roots, hairy roots and transformed hairy roots, respectively, as compared to their respective control. Compared with the control, Cd-stress dropped GSH level up to 9.8%, 32.2% and 45.7% in normal roots, hairy roots and transformed hairy roots, respectively. As compared to non-transformed hairy roots, transformed hairy roots expressing anaPCS gene contained 30.5% and 26.2% lower levels of GSH under As- and Cd-stress, respectively.

Stress-induced antioxidative enzymes in normal roots, hairy roots and transformed hairy roots

Variable activities of four antioxidative enzymes were measured under heavy metal stress in all tissue samples (Fig. 6a-d). Under As-stress, catalase activity was reported to be induced up to 20%, 12.5% and 20% in normal roots, hairy roots and transformed hairy roots, respectively, as compared to control. Similarly, Cd-stress increased catalase activity up to 26.6%, 12.5% and 13.3% in normal roots, hairy roots and transformed hairy roots as compared to control. Although catalase activity under stresses was higher than control, their activity was similar among all three tissues. Transformed hairy roots did not show any significant change in catalase activity under heavy metal stresses in A. annua.

Under As-stress, APX activity was enhanced up to 21.9%, 67.6% and 34.7% in normal roots, hairy roots and transformed hairy roots, respectively, as compared to control. Under Cd-stress, APX activity was reported to be 17.4%, 72.3% and 5.37% in normal roots, hairy roots and transformed hairy roots, respectively, compared with the control. In comparison to non-transformed hairy roots, transformed hairy roots expressing anaPCS gene showed 31.9% greater APX activity under As-stress; however, Cd-stressed roots
transformed hairy root did not show any significant increment in APX activity as compared to non-transformed hairy roots.

Peroxidase activity was also enhanced up to 150%, 139.4% and 101.3% in As-stressed normal roots, hairy roots and transformed hairy roots, respectively, as compared to their respective control. Increment of 130.7%, 153.5% and 90.5% in peroxidase activity was reported in Cd-stressed normal roots, hairy roots and transformed hairy roots, respectively, as compared to control.

Stress-induced increment in GR activity was more pronounced in transformed hairy roots than non-transformed hairy roots. As-stress induced GR activity up to 5.5%, 18.6% and 151.7% in normal roots, hairy roots and transformed hairy roots, respectively, as compared to control. Similarly, Cd-stress enhanced GR activity up to 13.4%, 15.2% and 93.1% in normal roots, hairy roots and transformed hairy roots, respectively, as compared to control. Transformed hairy roots expressing *anaPCS* gene showed 108.5% and 64.7% greater GR activity under As- and Cd-stress, respectively.

Artemisinin content and expression of artemisinin biosynthetic pathway genes

Artemisinin level was found to be more in hairy roots than in normal roots in both normal and stressed conditions (Fig. 7a). Under As-stress, artemisinin level was enhanced up to 50%, 33.3% and 16.6% in normal roots, hairy roots and transformed hairy roots, respectively. On the other hand, Cd-stress boosted artemisinin level up to 45.41%, 25% and 22.2% in normal roots, hairy roots and transformed hairy roots, respectively, as compared to control. Figure 7b shows the expression pattern of four key artemisinin biosynthetic pathway genes, viz., *ADS, CYP71AV1, DBR2* and *ALDH1*, in all tissue samples. All the four biosynthetic pathway genes were found to
be significantly overexpressed in transformed hairy roots as compared to normal roots and hairy roots (non-transformed). Under As-stress, ADS expression was 7- and 6.4-fold more than normal roots and hairy roots, respectively. Similarly, in transformed hairy roots, Cd-stress enhanced ADS expression up to 2.8 and 3.2-folds as compared to normal roots and hairy roots, respectively. Similar over-expression patterns were observed for CYP71AV1, DBR2 and ALDH1, as well.

**Fig. 6** Antioxidative enzyme activities of **a** Catalase, **b** ascorbate peroxidase, **c** glutathione reductase, and **d** peroxidases, in normal roots, hairy roots and transformed hairy roots (expressing anaPCS gene) under arsenic and cadmium stress. Vertical bars denote mean ± SD and different alphabets show statistical significance among treatments (control/As-treatment/Cd-treatment) at *p* = < 0.05.

**Fig. 7** **a** Artemisinin content and **b** expression analysis of artemisinin biosynthetic enzymes (ADS, CYP71AV1, DBR2 and ALDH1) through RT-PCR, in normal roots, hairy roots and transformed hairy roots (expressing anaPCS gene) under arsenic and cadmium stress. Vertical bars denote mean ± SD and different alphabets show statistical significance among treatments (control/As-treatment/Cd-treatment) at *p* = < 0.05.
Database mining and phylogenomic analysis of AaPCS and anaPCS

To find *A. annua* and *Anabaena* PCC 7120 PCS homolog, BLAST-P search was performed using the *Arabidopsis* PCS gene (AAD41794.1) which led to the identification of six orthologs with high homology. The sequences (PWA96196.1) comprehending the highest query coverages (100%) and identity (73%) with the target sequence were selected for further studies. Comparative phylogenomic analysis of AaPCS genes performed by constructing a phylogenetic tree based on UPGMA (Fig. 8a) and maximum parsimony methods (Fig. 8b) revealed monophyletic origin with *S. tuberosum* (CAD68110.1) and with *O. sativa* (AAO13349.2), respectively. In contrasts, other members, viz., *B. juncea* (CAC37692.1) and *Anabaena* PCC 7120 (BAD10973.1), were observed to form a separate clade. The result of phylogenetic analysis among different plant species revealed homology of AaPCS with other homolog and ortholog members, established the evolutionary relationship between them and predicted that AaPCS is the closest homologue of StPCS.

Functional sites identification and domain analysis

The functional signature sequences underlying AaPCS and anaPCS were identified by PROSITE analysis which validated and confirmed that the query sequences of both AaPCS and anaPCS belongs to Papain-like cysteine peptidase superfamily. The position of functional signature sequences constituting phytochelatin catalytic domain from N-terminal end (NTD) was also retrieved. The results of InteProScan analysis further confirm the presence of phytochelatin catalytic domain from N-terminal domain in each AaPCS and anaPCS. In AaPCS, the N-terminal phytochelatin catalytic domain (NTD) prevailed from 7 to 234 amino acid residue where core signature sequences were occupied between Tyr7 and Arg234 (Supplementary Fig. 2), whereas in anaPCS, the N-terminal phytochelatin catalytic domain (NTD) existed between 25 to 237 amino acid residues consisting functional core residues in between Leu25 and Lys237 (Supplementary Fig. 3). The multiple sequence alignment (Supplementary Fig. 4) results depicted conserved residues within the N-terminal phytochelatin catalytic domain (NTD) of the protein along with some insertion and deletion of the amino acid residues at certain positions that may have led to the sequence divergence in other homologs and orthologs.

Fig. 8 Phylogenetic tree: a UPGMA method and b maximum parsimony method showing origin, evolutionary relationship and functional homology of phytochelatin synthase protein in *A. annua*, *S. tuberosum*, *A. thaliana*, *B. juncea*, *O. sativa* and *Anabaena* PCC 7120. The trees were generated using 1000 boot replication values based on maximum-likelihood methods.

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thus delineating their ancestral evolutionary origin. Furthermore, circos visualization tools confirmed the ancestral evolutionary similarity and differences across the members of phytochelatin synthase family (Fig. 9). At 50%, cut-off values, \( \text{AaPCS} \) displayed close relation with \( S. \text{tuberosum} \) and \( A. \text{thaliana} \), whereas \( \text{anaPCS} \) did not found in relation with any of the other homolog or ortholog members.

### Conserved motif analysis

To comprehend conservation and diversification of PCS proteins in \( A. \text{annua} \), putative functionally conserved network elements (motifs) of \( \text{AaPCS} \) were predicted using MEME software which predicted seven different motifs (Fig. 10). As anticipated, MEME motif analysis revealed that all the members in phylogenetic tree shared conjoint motif distribution pattern, signifying that PCS proteins within the same subclades might be involved in similar phytochelatin catalytic function. In our results, the motif scan analysis revealed significant motifs that comprised phytochelatin catalytic domain with in PCS proteins and statistical significance of the motifs was evaluated in terms of their \( E \)-value that speculate their adequate patterns and \( p \) value that conjectures how substantially each pattern correlate with the motif. The N-terminal catalytic domain (NTD) in \( \text{AaPCS} \) was represented by motif 3 (MAMASLYRLPSPPAIDF; \( p \) value 1.3e-23) and propensity of the amino acids constituting this motif was revealed based on size of the alphabets that correspond to amino acids. The motif distribution pattern revealed via motif scan analysis unveil commonalities in \( A. \text{annua} \), \( A. \text{thaliana} \), \( S. \text{tuberosum} \) and \( B. \text{juncea} \) (motif 3: MAMASLYRLPSPPAIDF; \( p \) value 1.7e-65). Furthermore, the motif results showed the presence of additional 1 motif in \( O. \text{sativa} \) (RRRLFRCAQAQ; \( p \) value 2.9e-11) and additional 2 motifs in \( \text{Anabaena PCC 7120} \) (QVARQGLTLD; \( p \) value 1.0e-12 and PNLIGFNENE; \( p \) value 1.1e-9) may have led to sequence divergence which could have been reason behind separate clustering of both \( \text{OsPCS} \) and \( \text{anaPCS} \) in the phylogenetic tree constructed in the present study.

### Structural modelling of PCS functional domain

In the present study, the N-terminal catalytic domain of both \( \text{AaPCS} \) and \( \text{anaPCS} \) was modelled using relevant templates chosen based on crystal resolution, sequence similarity and completeness. A total of five models each for \( \text{AaPCS} \) (Fig. 11a) and \( \text{anaPCS} \) (Fig. 11b) were generated by MODELLER module of DS Modeller (Supplementary Table 2). In the present study, the models retaining lowest RMSD value corresponding to \( C\alpha \) atoms within the template crystal structures and minimum electrostatic energy (calculated) were considered as good stable models which were selected for further interactions with \( \text{CdCl}_{2} \) and \( \text{Na}_{3}\text{AsO}_{4} \) (sodium arsenate) ligands modelled using ChemDraw tool. The predicted models were submitted to PMDB database and assigned PMDB IDs for our predicted structures of \( \text{AaPCS} \) (PM0083228) and \( \text{anaPCS} \) (PM0083229).

### Model evaluation and validation

The three-dimensional models both for \( \text{AaPCS} \) and \( \text{anaPCS} \) were generated by satisfying all spatial restraint using Modeller Discovery Studio Client 3.0. Five sets of models each for \( \text{AaPCS} \) and \( \text{anaPCS} \) were arranged based on their Discrete Optimized Protein Energy (DOPE) and molecular Probability Density Functions (PDFs). The models (both for \( A. \text{annua} \) and \( \text{Anabaena PCC 7120} \)) exhibiting minimum DOPE scores and optimum energy values were selected for further validation using various statistics. The model proteins were subjected to Ramachandran plot analysis using RAMPAGE and PDBSum servers to assess the behavior of amino acid residues. The RAMPAGE statistics shows the presence of 95.4% residues in most favored region with 3.2% in additionally allowed region and 1.4% in outlier region for \( \text{AaPCS} \), whereas for \( \text{anaPCS} \), the statistics revealed the occurrence of 98.2% residues in most favored region with 1.8% in additionally allowed region and 0.0% in outlier region. Furthermore, VADAR analysis revealed that both the modelled proteins, viz., \( \text{AaPCS} \) and \( \text{anaPCS} \) existed as helix type structure 68 and 80 (30 and 36%) with 57 and 59

**Fig. 9** Comparative analysis of similarities and differences in phytochelatin synthase (PCS) protein among \( A. \text{annua} \), \( S. \text{tuberosum} \), \( A. \text{thaliana} \), \( B. \text{juncea} \), \( O. \text{sativa} \) and \( \text{Anabaena PCC 7120} \). The Circos software was used to generate circular map based on similarity percentage matrices using Clustal W algorithm.
Fig. 10 The motif scan analysis showing distribution and presence/absence of common and uncommon motifs found in *A. annua*, *S. tuberosum*, *A. thaliana*, *B. juncea*, *O. sativa* and *Anabaena* PCC 7120 discovered through MEME and MAST analysis. **a** The block diagram showing the sequence of discovered motifs for AaPCS. The red arrows indicate the presence of uncommon motif in *O. sativa* and black arrows indicate the presence of uncommon motifs in *Anabaena* PCC 7120 which are absent in other members. **b** The sequential logo of the motif 1 showing consensus sequences present in all the representatives’ members.

Fig. 11 Predicted structures of functional domain of **a** *A. annua* PCS (AaPCS) and **b** *Anabaena* PCC 7120 PCS (anaPCS) generated and visualized through MODELLER module of Discovery Studio 3.0
(25 and 26%) and 96 and 80 (43 and 37%) coil type configuration having mean H-bond energy (−1.6 (SD = 1.1) against the expected values (-2.0 SD = 0.8). Furthermore, PROCHECK evaluation (Supplementary Fig. 6) revealed the occurrence of 87.6% (AaPCS) and 94.5% (anaPCS) residues in most favored region (A, B, L), with 8.8% (AaPCS) and 5.0% (anaPCS) in additionally allowed region (a, b, l, p). A protein model is generally considered as a good quality model if it has 90% of the residues in most favored region which corroborates the results of the present study. Additionally, ProSA results also validated the results of RAMPAGE and PROCHECK analysis, and confirmed that both the modelled proteins were precisely similar to template proteins. Moreover, RESPROX and QMEAN analysis confirmed the reliability of our predicted models by generating a composite score based on the template’s proteins (Supplementary Table 3).

**Protein–ligand interaction**

To further explore the core amino acid residues of AaPCS and anaPCS involved in the binding with the ligand, molecular docking studies were performed for the possible investigation of binding mode of ligand, viz., cadmium chloride (CdCl₂) to the given receptors. The ligand, viz., CdCl₂ was modelled using the ChemDraw tool and was docked with the modelled proteins through HexDock server. In the present study, most reliable protein models of AaPCS and anaPCS were docked and stability of the docked complex was measured in terms of binding energy. For AaPCS, the most stable docked complex has the binding energy of -134.58 kcal mol⁻¹, whereas in case of anaPCS, the most stable complex was docked with the binding energy of -132.45 kcal mol⁻¹. The probable amino acid residues which are potentially involved in the interaction with the ligand are depicted in Fig. 12a. The docking studies for AaPCS revealed that the key residues, viz., Met²⁶, Met³⁸, Glu³⁹, Phe⁴², Val⁶⁵ and Leu⁶⁶ actively participated in the interaction with CdCl₂ (Fig. 12b). In contrast, the residues involved in the interaction of anaPCS with CdCl₂ were Thr¹⁰⁸, Leu¹⁰⁹, Leu¹¹², Ile¹⁵², Val¹⁵³, Asn¹⁵⁴, Ile¹⁶⁸ and Val¹⁷⁹ (Fig. 12d). Key amino acid residues involved during sodium arsenate interaction with AaPCS were Phe⁸⁴, Met⁸⁸, Asp⁹⁹, Lue¹¹¹ (Fig. 12c). However, sodium arsenate and anaPCS interaction involved Thr¹⁰⁸, Leu¹⁰⁹, Leu¹¹², Ile¹⁵², Val¹⁵³, Asn¹⁵⁴, Ile¹⁶⁸, Gly²¹⁷, Val²¹⁹ (Fig. 12e).

The result of molecular docking study revealed different amino acid residues were involved in the interaction of AaPCS with the ligand as those observed for anaPCS, thus clearly portraying that members of PCS family specifically

![Fig. 12 a](image1) Comparative analysis of docked complex with experimentally resolved X-ray diffraction structures of functional domain of AaPCS and anaPCS with cadmium chloride (CdCl₂) and sodium arsenate. b, e Structure of docked complexes in A. annua as visualized in Discovery Studio 3.0. d, e Structure of docked complex in Anabaena PCC 7120 as visualized in Discovery Studio 3.0.
binds to ligand with varying amino acid residues that discloses a common interacting pattern.

**Protein–protein interaction**

The functional protein–protein interactive network of both AaPCS and anaPCS was predicted using STRING server at medium confidence level, and the results were further validated by comparing protein–protein interaction results by accessing PTIR database. At medium confidence level, AaPCS found to have strong interaction (Fig. 13a) with chloroplastic glutamatecysteine ligase (PGSC0003DMT400031827; score value 0.973 which regulates salicylic acid and phytoalexin biosynthesis, thereby regulating tolerance response through GSH-dependent developmental pathway), chloroplastic glutathione synthetase (PGSC0003DMT400073627; score value 0.955, which is exclusively involved in the glutathione biosynthesis and also regulates metal binding with in the plants) and lastly with gamma-glutamyl transpeptidases (PGSC0003DMT400022803; score value 0.955, which is mainly involved in the catabolism of glutathione, regulation of post-translational processing and cellular localization). On the other hand, anaPCS actively interacted (Fig. 13b) with ABC transporter ATP-binding protein (anaPCS; score value 0.999, which is involved in nutrients uptake including metals by utilizing energy from ATP binding), ABC transporter permease (anaPCS; score value 0.996, regulates metal transport) and binding protein of ABC transporter component (anaPCS; score value 0.995, which is also involved in the import and export of various substrates including metal). From the results of the present study, it is evident that all the functional interactive partners of both AaPCS and anaPCS proteins belong to transmembrane protein families exhibiting metal transport activity thus signifying the coordinated role of PCS proteins in the transportation of heavy metals. The complete list of the interacting partners from both AaPCS and anaPCS have been shown in supporting information (supplementary table 4 and 5).

**Superimposition results**

Several studies have reported that protein structures are conserved to a greater extent compared to sequences. Keeping these facts in minds, the predicted structures of both AaPCS and anaPCS were superimposed over each other to predict their topological properties (Supplementary Fig. 7). The superimposition was performed using BLOSUM 62 matrix with gap penalty of 10 and extended penalty of 0.5. The superimposition results indicated that a total of 143 amino acid residues are substituted in between AaPCS and anaPCS with covering overall sequence similarity of 46.1% which was also further confirmed by the RMSD values of alpha carbon (2.06 Å) and backbone atoms (2.07 Å). In

![Fig. 13](image) Functional interactive network of a AaPCS and b anaPCS with other protein family members as found on STRING server where the colored nodes describe query proteins from first shell interactors and white nodes form second shell interactors. The large node size represents characterized proteins and smaller nodes for uncharacterized proteins.
contrast, when AaPCS was allowed to be superimposed with the template 2BTW, the calculated RMSD values were 1.03 Å for alpha carbon and 1.09 Å for backbone atoms. Similarly, superimposition of anaPCS over template 2BTW showed RMSD values of 1.41 Å both for alpha carbon and backbone atoms. The superimposition results confirmed the conservation of PCS functional domain structures as well as sequence across the divergent PCS members. However, structural alignment of functional domain of both AaPCS and anaPCS revealed some identical and non-identical (light blue) substitutions that extricated AaPCS domain from anaPCS. The amino acid substitutions were observed at position such as Ser5, Val10, Pro12, Glu34, Asn47 and Gly75 in AaPCS when compared to anaPCS, where Gln5, Thr12, Asp241 and Phe242 were present.

**Gene ontology enrichment analysis**

The available protein sequences, viz., AaPCS and anaPCS, were subjected to CATH-Gene3D analysis to predict their subcellular localization and their functional annotation using gene ontology terms. Gene ontological (GO) terms serve as structural and functional descriptors characterizing genes and their products on the basis of their cellular, biological and molecular processes. The ReviGO analysis measured the biological function (Supplementary Fig. 8) of AaPCS protein revealed that 44% of the AaPCS protein is involved in the phytochelatin biosynthetic pathway (GO:0046938) and is potentially involved in the detoxification of metals such as arsenic, cadmium and copper (GO:0010273, GO:0046686, GO:0046685). The molecular function measured in terms of gene ontology revealed that the predicted proteins (56%) are mostly engaged in glutathione gamma-glutamyl cysteinyl transferase activity (GO:0016756) and respond well under metal stress condition (GO:0005507, GO:0046870). On the other hand, CELLO2GO analysis (Supplementary Fig. 9) revealed that 56.8% of the proteins are localized into plasma membrane (score value 2.839) involved in phytochelatin biosynthetic process (40.6%) and metal ion binding (35.7%). For anaPCS, 17.8, 11.1, 4.4, 8.9 and 2.2% of proteins are involved in phytochelatin biosynthetic process, detoxification of copper, response to arsenic, detoxification of cadmium ion and arsenic transport, respectively.

**Discussion**

In the present study, we aimed to enhance the heavy metal tolerance of *A. annua* hairy roots by expression of *Anabaena* PCC7120 phytochelatin synthase (*anaPCS*) gene. In their natural habitat, cyanobacteria including *Anabaena* are constantly exposed to adverse environmental condition such as high temperature, salinity, varied pH range and heavy metals (Rai et al. 2014) which make them an ideal system for extracting stress-tolerant genes. Cloning of *anaPCS* was achieved in *E. coli* (Chaurasia et al. 2008) and *anaPCS* over-expression in *Anabaena* had shown multiple stress tolerance in the cyanobacteria (Chaurasia et al. 2017). The present study shows first-ever report of expression of cyanobacterial gene in *A. annua*.

We began with three different tissue samples, i.e., normal roots (derived from plantlets), hairy roots and transformed hairy roots (expressing *anaPCS* gene). Presence of rolB, presence of rolB and *anaPCS* and absence of both rolB and *anaPCS* confirmed the type of tissues as hairy roots, transformed hairy roots and normal roots, respectively (Figs. 1 and 4B).

Transformed hairy roots show significantly high amount of PCS expression owing to presence of two copies of PCS gene: one its own and another from *Anabaena* PCC 7120. As shown in Fig. 4, both As- and Cd-stress significantly induced the expression of *anaPCS* and AaPCS in transformed hairy roots, and as a result, As- and Cd-accumulation capacity was drastically enhanced in transformed hairy roots which accumulated 143.4 and 191% higher As and Cd than control. The value of bioconcentration factor (BCF) which gives an idea about capacity to accumulate heavy metals was significantly higher for transformed hairy roots. The BCF values of up to 21 and 9.7 for Cd and As, respectively, clearly explain that Cd is accumulated more than As; however, BCF values for both the heavy metals were significantly higher for transformed hairy roots as compared to non-transformed hairy roots or normal roots. In support of the present study, heterologous expression of *NnPCS1* from lotus (with proved phytoremediation potential) conferred Cd tolerance in *A. thaliana* (Liu et al. 2012). Similarly, PCS gene from many other plants such as garlic and Indian mustard has been expressed in *A. thaliana* (Guo et al. 2008; Gasic and Korban 2007) and enhanced Cd and As tolerance significantly. In addition, heterologous expression of PCS gene from an aquatic macrophyte has provided better As and Cd tolerance in Tobacco (Shukla et al. 2012).

Both As and Cd are non-redox active metals which, however, cause significant oxidative damage to cell. Plant cells possess excellent defense system comprising of non-enzymatic and enzymatic antioxidant to mitigate their adverse effects.

Differential responses of four antioxidative enzymes were observed under As- and Cd-stress in all tissue samples. In transformed hairy roots, all antioxidative enzymes except catalase were drastically induced under heavy metal stresses. At different locations in the cell, *H₂O₂* is converted to water and oxygen through APX, GPX and catalase. Only a slight increase in catalase activity after As and Cd exposure in all tissue samples (normal roots, hairy roots and transformed hairy roots) may probably be due to its high sensitivity
towards superoxide radicals (Cakmak 2000). Catalase sensitivity under heavy metal stresses have also been reported in several plant species (Cao et al 2004; Gupta et al 2009) including A. annua (Rai et al 2011). Lesser catalase activity in heavy metal stressed transformed hairy roots was compensated by APX which possessed 31.9 and 34% greater activity compared to normal roots and hairy roots, respectively. Transformed hairy roots possess better defense against heavy metal stress attributed to its capability to maintain high level of APX, GR and POD in untransformed and stressed conditions. High activity of GR has probably contributed towards better As and Cd tolerance in transformed hairy roots of A. annua, because GR plays a significant role in heavy metal stresses by maintaining high GSH:GSSG ratio in the cell via recycling GSSG to GSH. As- and Cd- stress induced antioxidative enzyme activities including that for APX, GR and POD in several plants (Kumar et al 2014; Rai et al 2011; Zhang et al 2007).

In addition to antioxidative enzymes, low-molecular-weight thiols also play critical role in maintaining the overall oxidative balance in the cell. Their maximum induction in transformed hairy roots (with anaPCS gene) under As- and Cd- stresses (Fig. 5a) is indicative of better stress tolerance. GSH is one of the low-molecular-weight thiols with at least two crucial functions: synthesis of PCs and maintaining the ascorbate–glutathione cycle, both as part of antioxidative defense (Cobbett 2000). In the present study, transformed hairy roots expressing anaPCS showed approximately four-fold and twofold higher PCs level as compared to normal roots and non-transformed roots. As PCs are known to form metal-complexes which are sequestered into the vacuoles, boosted PCs level in transformed hairy roots suggests their potential to efficiently detoxify As and Cd. This can also be correlated with higher accumulation of As and Cd in transformed hairy roots than other samples and high GR activity which generates enough GSH to be used for PCs synthesis. Overexpression of AaPCS and anaPCS in transformed hairy roots supports the higher PCs level. However, in the present study, GSH levels were dropped under As- and Cd-stress. As product of PCS gene is being synthesized in high amount, lower GSH level in transformed hairy roots was probably because of the overconsumption for PCs synthesis and not due to its inhibition. The result finds support with Samane et al. (2007), Ben Ammar et al. (2008) and Meng et al. (2019) who have found lower GSH but higher PCs level in heavy metal stressed Arabidopsis thaliana, tomato, and lettuce, respectively.

In-silico analysis gave several confirmatory pieces of evidence for the proper functioning of a cyanobacterial anaPCS in a higher plant A. annua. The presence of conserved catalytic domain (Fig. 9) as well as functional domain structures (Supplementary Fig. 7) among anaPCS and AaPCS may have led cyanobacterial anaPCS to function properly in A. annua hairy roots. Protein–protein interaction analysis confirmed anaPCS involvement in heavy metal transport as their functional interacting partners belong to transmembrane protein families having metal transport activity. This further get confirmation through gene ontology analysis where proportions of anaPCS proteins have been shown to be involved in biological processes like phytochelatin biosynthesis, arsenic transport and detoxification of cadmium ions.

In addition to better tolerance towards As and Cd, transformed hairy roots also showed the significantly greater synthesis of its popular antimalarial compound artemisinin (Fig. 7). Artemisinin biosynthesis have been shown to be improved under both arsenic (Kumari et al. 2018) and cadmium stress (Li et al. 2012). Owing to its immense therapeutic potential as antimalarial, anticarcinogenic, antimicrobial and many others, its global demand is always high and needs to be supplied enough to meet high demand (Pandey and Pandey-Rai 2016). Recently artemisinin has been suggested to be repurposed for the treatment of COVID-19 and other related respiratory disorders which will further increase its demand (Cheong et al. 2020). Higher artemisinin biosynthesis with enhanced heavy metal tolerance in hairy roots of A. annua expressing anaPCS gene can serve a dual purpose.

Conclusion

Heterologous expression of a cyanobacterial phytochelatin synthase (anaPCS) in hairy roots of A. annua was successfully achieved which led to enhanced heavy metal (arsenic and cadmium) tolerance. Under conditions of heavy metal stresses, greater accumulation of arsenic and cadmium, higher biosynthesis of phytochelatins, and activation of enzymatic and non-enzymatic antioxidants have made transformed hairy roots (expressing anaPCS) more tolerant to arsenic and cadmium stresses. Due to conserved catalytic domain and functional domain among anaPCS and AaPCS, the former could have performed in a different system (higher plant). In-silico analysis revealed involvement of anaPCS protein in heavy metal transport, phytochelatin biosynthesis and detoxification of cadmium and arsenic. Enhanced artemisinin biosynthesis in transformed hairy roots further gave scope of their probable use in phytoremediation and extraction of artemisinin from used hairy roots.

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**Declarations**

**Conflict of interest** Authors declare no conflict of interest.

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