Metagenomic insights into plant growth promoting genes inherent in bacterial endophytes of *Emilia sonchifolia* (Linn.) DC

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Abstract

Studies on the genome of endophytes reveal the metabolic potential of endophytic microbiome including both culturable and unculturable fractions. The metagenome analysis through the Illumina HiSeq platform gives access to the genetic data encrypted for the molecular machinery, which takes part in plant growth promotion activity of the endophyte in various aspects including production of plant growth hormones and enhancing nutrient availability for the host plant. The present work was undertaken to identify the genes involved in plant growth promotion activities from the endophytes of *Emilia sonchifolia* (Linn.) DC. through metagenome analysis. Metagenomic studies include the analysis of functional annotations which aid in the detection of biocatalysts taking part in the metabolic pathway of host plants. The annotations of expressed genes in different databases like NCBI Nr, KEGG, eggNOG and CAZy resulted in enlisting the vast array of information on the genetic diversity of the endophytic microbiome. The metagenome analysis of endophytic bacteria from the medicinal plant *E. sonchifolia* unveiled characteristic functional genes involved in plant growth promotion such as nitrogen metabolism (nif) and siderophore production (enterobactin category), ipdC and tnaA (IAA producing), ACC deaminase coding genes (regulation of elevated ethylene levels in host tissues), Mo-Nitrogenase, nitrous-oxide reductase (nosZ), nitrate reductase (narG, napA), nitrite reductase (nirD) (nutrient assimilation and absorption) enterobactin siderophore synthetase components F and D and acid phosphatase genes. This clearly explains the effective plant-microbe relationship and the role of bacterial endophytic microbes in regulating the growth of host plants.

Keywords

Functional annotation, Gene prediction, Metagenome analysis, Nutrient uptake, Siderophores

Introduction

Endophytes are non-pathogenic microbiome occupying internally in plant tissues and occupy in many plants. Many researchers screened the endophytic microbes associated with medicinal plants for analysing the active and passive role of these microorganisms in the synthesis of host-derived bioactive compounds (1, 2). Its multilevel interactions like plant-microbe, microbe-microbe, and microbe-environment enhanced the complexity of endophytic microbiome analysis (3). The identification of closely related endophytic microbes and their functional genes becomes difficult because of their genome plasticity (4). Recent developments in molecular biology and metagenome analy-
sis further widened our understanding of microbial genetic diversity associated with such plants. The metagenome analysis of microbial communities explains the community composition, perspectives of plant-microbe interactive development, physiological and biosynthetic potential of the association (5). Metagenome data mining opens new avenues in DNA/gene-level information followed by the identification and expression pattern of genes. The genomic analyses provide information on many of the unanswered problems in endophytism like the reason for the coexistence of different microbes, the extent of plant-microbe interaction and the plant-microbe symbiotic co-evolution (6). The construction of metagenomic libraries followed by phylogenetic analysis can explicate the diversity of microbial communities associated with different plants.

High throughput sequencing techniques like Illumina shifted the end barriers further in the genomic studies and it provided a new reliable platform for researchers to screen the hidden world of microbes living inside the microhabitat in plant tissues. Illumina HiSeq technology generates a large quantity of genomic data from which relatively high-quality data on protein-coding genes are generated. This sequenced data on comparison with the known database can identify the functional roles of those identified genes. This genomic information thus created will lead to the identification of metabolic capacities of microbes and to tackle many of the hidden areas of plant-microbe interactions.

In order to keep a stable symbiotic relationship, the endophytic bacteria enhance the production of some growth regulators for the host. In this way, they activate the host metabolic machinery by the production of some compounds that sustain their endophytic mode of life (7). Endophytes produce plant growth regulators like Indole Acetic Acid (8), ACC deaminase (9), and enhance the availability of nutrients like phosphorous (10), nitrogen (11) and iron (12). Indirect ways of plant growth promotion occur through the ability of endophytes to prevent the growth of pathogenic microorganisms (13, 14) and they help the plant tolerate different stress conditions (15, 16). Endophytic association resulted in the enhancement of expression of selected genes of the host (17) and the alterations in metabolism were beneficial to both partners. Hence, these associations are considered as symbiotic rather than pathogenic.

Most endophytic bacteria reported were from the group Proteobacteria which are soil bacteria (18, 19). Microbial ecology and diversity studies on Illumina MiSeq/HiSeq platform reduces the errors and increases the likelihood of finding rare and beneficial endophytes with higher phylogenetic resolution at a comparatively lower cost (20, 21, 22). Because of very effective therapeutic applications in traditional medicines, E. sonchifolia was screened thoroughly to find out the phytochemical components (23, 24). Therefore, high throughput screening will reveal the role of endophytes in the biosynthetic potential of the host. During the biodiversity analysis, two different phyla identified from E. sonchifolia were Proteobacteria and Firmicutes (25). The present investigation on endophytic genes and their functional role unveiled the role of the endophyte in the plant growth promotion activity of this beneficial medicinal plant.

Materials and Methods

DNA extraction and library construction
DNA was extracted from surface-sterilized Emilia sonchifolia (Linn) DC. It was quantified (Qubit®4.0 fluorometer, Invitrogen, Carlsbad, CA, USA) and fragmented randomly by sonication (Covaris 220). The adaptors were indexed which helped for the easy identification of reads in Illumina platform. Amplification of the fragments was done by PCR for 8 cycles using P5 (5’ ATGTACCCGCAACCAGATCTCAC 3’) and P7 (5’ CAAGCAGAGACAGCATACTGAG 3’) primers. These primers are universal primers and both primers had specific sequences which can anneal with flow cell to perform bridge PCR and the P7 primer carried a six base index which helped for multiplexing. The VAHTSTM DNA clean beads were used for cleaning the PCR amplified products and they were quantified by Qubit®4.0 fluorometer (Invitrogen, Carlsbad, CA, USA). These next-generation sequencing library preparations were done by following the manufacturer’s protocol (VAHTS Universal DNA library preparation kit for Illumina).

Illumina HiSeq sequencing
The indexed libraries prepared were loaded on the Illumina HiSeq instrument and the different indices were multiplexed and sequenced according to the manufacturer’s instructions (Illumina, San Diego, CA, USA). Sequencing was carried out in 2 x 150 paired-end (PE) configuration; image analysis and base calling were conducted by the HiSeq control software (HCS), OLB, and GA pipeline-1.6 (Illumina) on the HiSeq instrument. Bcl2fastq analysed the original Image data (V2.17.14) for base calling and quality analysis and saved in fastq format.

Processing and assembly of data
The Phred quality score was calculated based on ASCII standards, and the quality score (Q20) less than 20 were discarded. The pass filtered data saved in fast q format files one for read 1 and the other for read 2 were selected for creating paired-end data. GC content was also calculated to reduce the AT-GC sequencing bias. Next-generation data quality software Cutadapt (v.1.9.1) was used to trim adaptors, eliminate low-quality reads and N-rich reads. We also removed the primers and reads with lengths less than 75bp in this filtering process. The chance for the contamination of reads with host sequence was removed with the help of BWA (v0.7.12) software which filtered out host sequences based on the host genome.

Metagenome assembly and gene prediction
The clean data was used for the whole genome de novo assemblies and that was processed using MEGAHIT (v 1.1.3), with different K-mer (39, 59, 79, 119). After assembling, the scaffold with the biggest N50 was selected for further gene prediction analysis. The reads were analysed
for coding genes to specify gene predictions of metagenomic or unknown microorganisms using Prodigal (v 3.02). Sequence clustering was done with CD-HIT (v4.5.6) which reduced the redundancy of the predicted gene sequences and clustering of unique gene sequences was processed at 95% identity and 90% coverage level by default. Pre-processed reads were then aligned to a non-redundant set of genes with the help of Soap Aligner (v2.21) generated gene abundance or read coverage of the genes. The number aligned and normalised reads were calculated based on gene length which was used to measure the gene abundance.

**Gene functional annotation**

Gene functional annotations were predicted by aligning the predicted gene with different databases like Nr database (non-redundant protein database), KEGG pathway database (Kyoto Encyclopedia of Genes and Genomes database), eggNOG (evolutionary genealogy of genes: Non Supervised Orthologous Groups, Version 4.0) and CAZy database (Carbohydrate Active enZYmes Database, Diamond Version 0.8.15.77 and BLAST Version 2.2.31+) were used for the database search and alignment to predict the gene functional annotations. Gene annotation resulted from each database was used to categorise relative abundance of different functional categories..

**Results**

The original data analysed using Bcl2fast q (v2.17.1.14) software, and it checked the base quality of the first 25 bases in a read and determined the conversion of data to FASTA format. The pass filtered data without error and PHRED score higher than 20 (Q20) were kept. The quality score of a base and the reads mean quality distribution were calculated.

**Metagenome Assembly and Gene Prediction**

We assembled clean quality optimized data to generate a scaffold, and it generated the detailed assembly results. After the assembly of metagenome data, 92250 reads were generated with an average length of 1432.89bp. The Prodigal (v3.02) software analysed the genome data, especially for gene prediction from metagenomic data or the data of unknown organisms. The sequence clustering generated unique sequences and the unigene statistics were analysed. The average length of the annotated read was 672, and it created 82bp and 161694 sequence reads. Soap Aligner detected the coding regions of the assembled scaffolds (Version 2.2.1). The count of each unigene was marked and calculated the unigene abundance. The unigene sequence ID K141-73893-1 gave a maximum count of 69. The sequence data was submitted in NCBI Biosample database SRA with accession numbers SAMN11617377 and SAMN11161726.

**Gene functional annotation**

The protein sequences of the predicted gene were compared with the protein database to get the gene functional annotations. The protein sequence of the predicted gene and the reference gene from the database showed significant similarity while searching the NCBI Nr (non-redundant) database. The sequence alignment length was set as over 60% similarity between predicted gene and a threshold error rate of 1e-5. This database search revealed 148289 annotated sequences which showed sequence similarity. In the Nr (non-redundant) database annotation pattern repre-

| Functional role of gene | NR_description | Identity (%) | Evalue |
|-------------------------|----------------|--------------|--------|
| IAA Production Tryptophanase | g[76323301301301301301 ref|96.8 4.20E-09
|                         | g[44836684030130130130 ref|93.3 3.80E-252
| IAA Production Acetoin | g[9467055130130130130 ref|95.6 1.20E-83
|                         | g[10226653630130130130 ref|99.6 5.40E-139
|                         | g[5448050830130130130 ref|93.9 2.00E-153
|                         | g[5580888280130130130 ref|99.2 1.30E-137

Table 1. Nr (non-redundant) database annotation pattern representing the protein involved in plant growth promotion and the species in which that protein belongs.

In the present data analysis, the functional potential of the endophytic microbiome was analysed using the KEGG database. From this, we identified 250 pathways and divided them into six major categories viz. cellular processes, environmental information processing, genetic information processing, human disease, metabolism and the organisinal system. Each category was further subdivided and analysed. The summary of annotated genes and major functional annotations of genes according to KEGG database was shown in Table 2.

The functional annotation based on Orthologous Groups was done against the eggNOG (v4.5) database. In the gene annotations, 25 different functional categories like energy production and conservation, general function, cell motility, function unknown etc were included. CAZy is the database used to analyze the carbohydrate-active enzymes. These include genes of six major functional categories like glycoside hydrolases (GHS), glycosyl transferases (GTs), polysaccharide lyases (PLs), carbohydrate esterasers (CES), auxiliary activities (AAS), and carbohydrate-binding modules (CBMs). The microbial carbohydrate metabolism of endophytes was clearly understood from CAZy annotations and some of the major genes of carbohydrate metabolism were recognized in the annotation. The plant growth promotion activity of endophytic bacteria...
was resolved through metagenome analysis because it disclosed many proteins coding genes and transport systems that enhance the growth of the host plant. Plant growth enhancement occurs either through the production of plant growth hormones or the endophytes augment the nutrient uptake and utilization. The endophytic metagenome carries genes like *ipdC*, *tnaA*, *ytrE*, *trpC*, *F* and these gene products were taking part in the production or degradation of phytohormones like indole-3-acetic acid and ethylene. Genes involved in the pathway of zeatin biosynthesis indicates the role of endophytes in the production of plant growth regulators. Activation of plant defence generates the plant hormone salicylic acid. The presence of salicylate hydroxylase annotation was found in the metagenomic data. The annotated genes were identified in the phenyl propanoid biosynthesis, terpenoid backbone biosynthesis, and inositol phosphate metabolism, which lead to the production of plant growth regulators. The annotated genes like *speA*, *B* and *E* give compounds like arginine decarboxylase etc. which will increase the fitness of the plant cell and it promotes the growth of both endophyte and host (Table 2).

| Functional role of the gene | Gene_Name | Definition | EC | Pathway |
|----------------------------|-----------|------------|----|---------|
| IAA synthesis | *tnaA* | tryptophanase | EC4.1.99.1 | ko00380 Tryptophan metabolism |
|                    | *ipdC* | indolepyruvate decarboxylase | EC4.1.1.74 | |
|                    | *trpC* | indole-3-glycerol phosphate synthase/ phosphoribosyl anthranilate isomerase | EC4.1.1.48, 5.3.1.24 | ko00400 Phenylalanine, tyrosine and tryptophan biosynthesis |
|                    | *nitrilase* | | | EC3.5.5.1 |
| Ethylene | *E3.5.99.7* | 1-aminocyclopropane-1-carboxylate deaminase | EC3.5.99.7 | ko00270 Cysteine and methionine metabolism |
| Acetoin | *ytrE,F* | acetoin utilization transport system ATP-binding protein | EC1.2.5.1 | ko00620 Pyruvate metabolism; |
| | *acu C,A,B* | acetoin utilization protein Acu C/B/A | -- | ko02010 ABC transporters; |
| Salicylic acid | *E1.14.13.1* | salicylate hydroxylase | EC1.14.13.1 | ko00626 Naphthalene degradation; |
| Zeatin | *miaA*, TRIT1 | tRNAdimethylallyltransferase | EC2.5.1.75 | ko00908 Zeatin biosynthesis |
| PGP | *speA* | arginine decarboxylase | EC4.1.1.19 | |
| | *speB* | agmatinase | EC3.5.3.11 | ko00330 Arginine and proline metabolism |
| | *speE, SRM* | spermidine synthase | EC2.5.1.16 | |
| | *yjbB* | Phosphate: Na+ symporter | | -- |

Fig. 1. Phosphate phosphinate metabolic pathway indicating annotations of genes in endophytic metagenome which take part in phosphate solubilization and uptake

Table 2. Annotated genes and their major functional annotations involved in plant growth promotion based on KEGG database.
Table 3. Annotated genes and their major functional annotations involved in phosphorus solubilization according to KEGG database.

| Gene Name | Gene functional product | EC | Pathway |
|-----------|------------------------|----|---------|
| pqqA, pqqB, pqqC, gcd, pxpA | pyrroloquinoline–quinone biosynthesis protein A, B, D, E, quinoprotein glucose dehydrogenase, exopolyporphosphate / guanosine-5'-triphasphate, 3'-diphosphate pyrophosphatase, inorganic pyrophosphatase | EC:1.3.3.11, EC:1.1.5.2, EC:3.6.1.11 3.6.1.40 | ko00030 Pentose phosphate pathway, ko00230 Purine metabolism, ko00190 Oxidative phosphorylation |
| pphB | two-component system, OmpR family, phosphate regulation response regulator PhoB | -- | ko02020 Two-component system |
| pheB | two-component system, OmpR family, phosphate regulon sensor histidine kinase PhoR | EC:2.7.13.3 | -- |
| pstS | phosphate transport system substrate-binding protein | -- | ko02010 ABC transporters; ko02020 Two-component system |
| pstB | phosphate transport system ATP-binding protein | EC:3.6.3.27 | -- |
| pstA/C | phosphate transport system permease protein | -- | ko02010 ABC transporters |
| phoH, phoL | phosphate starvation-inducible protein PhoH and related proteins | -- | -- |
| ybjB | Phosphate :Na+ symporter | -- | -- |
| phoR | low-affinity inorganic phosphate transporter | -- | -- |
| phoA, phoB | alkaline phosphatase | EC:3.1.3.1 | ko02020 Two-component system; ko00562 Inositol phosphate metabolism |
| E3.1.3.8 | 3-phytase | EC:3.1.3.8 | -- |
| appA | 4-phytase / acid phosphatase | EC:3.1.3.26 3.1.3.2 | -- |

Table 4. Annotated genes and their functional role in nitrogen metabolism based on KEGG database.

| Gene Name | Gene functional product | EC | Pathway |
|-----------|------------------------|----|---------|
| nifH | nitrogenase iron protein NifH | EC:1.18.6.1 | ko00910 Nitrogen metabolism |
| nifD | nitrogenase molybdenum-iron protein alpha chain | -- | ko02020 Two-component system; |
| nifK | nitrogenase molybdenum-iron protein beta chain | -- | -- |
| nifA | Nif-specific regulon protein | -- | -- |
| nifN | nitrogenase molybdenum-iron protein NifN | -- | -- |
| nifE | nitrogenase molybdenum-iron protein NifE | -- | -- |
| nifX, nifY, T, Z, B | nitrogen fixation protein NifX, NifY, T, Z, B | -- | -- |
| nifW | nitrogenase-stabilizing/protective protein | -- | -- |
| iscU, nifU | nitrogen fixation protein NifU and related proteins | -- | -- |
| flaA, nifF, isiB | flavodoxin I | -- | -- |
| nosZ | nitrous oxide reductase | EC:1.7.2.4 | ko00910 Nitrogen metabolism; ko00910 Nitrogen metabolism; ko02020 Two-component system; |
| narG, narZ, nxnA | nitrate reductase / nitrite oxidoreductase, alpha subunit, nitrate reductase gamma subunit, nitrate reductase (NADH) large subunit | EC:1.7.1.99.4, EC:1.7.1.79.4, EC:1.7.1.15 | ko00910 Nitrogen metabolism; ko02020 Two-component system; |
| nirB | nitrite reductase (NADH) large subunit | -- | ko00910 Nitrogen metabolism; |
| nirD | nitrite reductase (NADH) small subunit | EC:1.7.1.15 | ko00910 Nitrogen metabolism; |
| nirC | nitrite transporter NixC | -- | ko00910 Nitrogen metabolism |
| ncd2, npe | nitronatochromeoxigenase | EC:1.13.12.16 | ko00910 Nitrogen metabolism |
| napA | periplasmic nitrate reductase NapA | EC:1.7.99.4 | ko00910 Nitrogen metabolism |
| arc | carbamate kinase | EC:2.7.2.2 | ko00910 Nitrogen metabolism |
| gldG | glutamyl synthase (NADPH/NADH) small chain | EC:1.4.1.13 1.4.1.14 | ko00910 Nitrogen metabolism; 01230 Biosynthesis of amino acids; |
| cynT, can | carbonic anhydrase | EC:2.4.1.11 | ko00910 Nitrogen metabolism; |
| NRT, narK, nfxB, nasA | MFS transporter, NNP family, nitrate/nitrite transporter | -- | ko00910 Nitrogen metabolism; |
| glnA, ntrB | two-component system, NtrC family, nitrogen regulation sensor histidine kinase GlnL | EC:2.7.13.3 | ko02020 Two-component system; |
the iron uptake capacity of the endophytes. Many genes involved in the biosynthesis of siderophore group of non-ribosomal peptides (Fig. 3) were also observed in this study.

**Discussion**

The plant growth promotion activities of endophytic microbes were under research for application. We can incorporate it into the agricultural sector for the growth enhancement of economically valuable crops. Plant growth-promotion activities were mainly studied in endophytes from cultivable crops like rice, wheat, sugarcane etc. Plant growth-promoting activity and increase in productivity and biomass of rice plants were analysed after inoculation with *Azospirillum* sp. B510 (26, 27). Endophytes increase the growth of their host by various mechanisms. They can take part in the production of different classes of plant growth hormones (28, 29, 30) or they enhance the plant growth by

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**Table 5.** Annotated genes and their functional role in iron acquisition based on KEGG database.

| Gene Name | Gene functional product | EC | Pathway |
|-----------|-------------------------|----|---------|
| entF,D    | Enterobactin synthetase component F,D | EC:6.3.2.14 | ko01053 Biosynthesis of siderophore group non-ribosomal peptides, |
| dbhF      | Non-ribosomal peptide synthetaseDhbF | -- | -- |
| entA      | 2,3-dihydro-2,3-dihydroxybenzoate dehydrogenase | EC:1.3.1.28 | -- |
| entE, dbhE | 2,3-dihydroxybenzoate-AMP ligase | EC:6.3.2.14 2.7.7.58 | -- |
| entC      | isochorismate synthase | EC:5.4.4.2 | -- |
| fes       | enterochelin esterase and related enzymes | -- | -- |
| ent8, dbh8, vibB, mxcF | Bifunctional isochorismate/aryl carrier protein | EC:3.3.2.1 6.3.2.14 | -- |
| entS      | MFS transporter, ENTS family, enterobactin (siderophore) exporter | -- | entS |
| flu       | Catecholate siderophore receptor | -- | -- |
| ABC.FEV.A | iron complex transport system ATP-binding protein | EC:3.6.3.34 | ko02010 ABC transporters; |
| FTR, FTH1, efeU | high-affinity iron transporter | -- | -- |
| TC.FEV.OM | iron complex outer membrane receptor protein | -- | -- |
| isca, ISca1 | iron-sulfur cluster assembly protein | -- | -- |
| iscR      | Rnf2 family transcriptional regulator, iron-sulfur cluster assembly transcription factor | -- | -- |
| feoC,B    | ferrous iron transport protein C,B | -- | -- |
| fepA,     | ferric enterobactin receptor | -- | ko02010 Two-component system; |
| fhF       | ferric iron reductase protein FhF | -- | -- |
| irr       | Fur family transcriptional regulator, iron response regulator | -- | -- |
| sitA      | manganese/iron transport system substrate-binding protein | -- | -- |
| sitB      | manganese/iron transport system ATP-binding protein | -- | -- |
| sitCD     | manganese/iron transport system permease protein | -- | ko02010 ABC transporters; |
| efeO      | iron uptake system component FeoO | -- | -- |
increasing the availability of nutrients or will enhance its uptake (31).

The best-known phytohormone produced by endophytes is IAA which is synthesised from tryptophan through indole pyruvate (32). The presence of two notable enzymes, indole pyruvate decarboxylase (ipdC) and tryptophanase (tnaA) confirmed the IAA production capacity of the endophytic microbiome. Along with this, enzyme salicylate hydrolase showed the chance for conversion of tryptophan to IAA and salicylic acid (SA). Nitrilase is an enzyme (EC 3.5.5.1) that was reported to be involved in the biosynthesis of IAA from indole-3-acetonitrile, which is a tryptophan-independent pathway (33). The different enzymes and pathways for IAA biosynthesis explained the role of IAA as signalling molecules along with plant growth promotion (34). Acetoin and 2, 3-butanediol were reported to have plant growth promotion activity in Arabidopsis and their biosynthesis was recognized from rhizobacteria (35). A small fraction of acetoin production occurs by the activity of poxB gene during the pyruvate metabolism (36). The endophytic metagenome study carries the gene for pyruvate dehydrogenase (poxB), acetoin utilization proteins C, B, A (acuc, B, A) and ATP binding protein (ytrE, F) involved in the acetoin utilization transport system. Many endophytes produce cytokinins (37). In this study, enzymes involved in zeatin biosynthetic pathway like tRNA dimethylallyl transferase (miaA) were detected.

Most endophytic bacterial studies described the enzyme 1-aminocyclopropane-1-carboxylate (ACC) deaminase (9, 38, 39) which is produced during stress conditions to manage the elevated levels of ethylene. It cleaves the precursor of ACC to α-ketobutyrate and ammonium (40) and reduces the conversion of ACC to ethylene which affects the root growth. The presence of the enzyme ACC deaminase was also identified in this study. Studies using endophytic plant growth-promoting bacterium Burkholderia phytofirmans PsJN showed that the mutation in ACC deaminase gene inactivates the root elongation ability of the endophytic bacterium (41).

Spermidine is a plant growth promoting substance reported from the rhizobacterium Bacillus subtilis OkB105 (42). Three different genes speE (spermidine synthase), speB (Argmatinase), speA (arginine decarboxylase) which were involved in the synthesis of spermidine, spermine and putrescine respectively were also detected in the present study. Terpenoids play an important role in plants in the assembly of reaction centres in photosynthesis, stress tolerance (43) and in defence mechanisms (44). Terpenoids include a vast family of compounds or natural products. Different pathways like mevalonate and isoprenoid were identified that synthesise terpenoids. Different microbial organisms were screened for the large-scale production of terpenoids. Microbes like Yeast, E. coli (45) and Streptomyces (46, 47) were used and proved as a promising method for the large-scale production of terpenoids. In this study, the metagenome analysis revealed that endophytic genome carries genes that take part in the mevalonate pathway of terpenoid biosynthesis and so they were interacting with the host plant for the growth promotion of the host.

Nitrogen and phosphorus are important nutrients for plant growth. Because of the unavailability of phosphorus in its insoluble form, it acts as a limiting factor for plant growth (48, 49). The endophytic bacteria can solubilise poorly soluble inorganic phosphate and can enhance plant...
growth. The plant growth promotion activity of phosphate solubilising bacteria like *Pseudomonas* (50) and *Enterobacter* (51) were reported. Endophytic bacteria could solubilise both organic and inorganic phosphate and the genome of different plant growth-promoting endophytic bacteria contains genes for specific enzymes and regulatory systems taking part in it (52). Phytases are enzymes involved in the solubilisation of organic phosphate and they were thermally stable. The storage form of phosphate in plants was phytate (myoinositol 1, 2, 3, 4, 5, 6-hexakisphosphate). Phytases are enzymes that can remove the phosphate group from the phytate. Along with that, it prevents the chelate-forming capacity of phytase with mineral nutrients and thus making other mineral nutrients also available (10, 53). appA gene identified and isolated from *E. coli* which codes for acid phosphatase (54). The solubilisation of inorganic or mineral phosphate in bacteria was found to be associated with the synthesis of organic acids and the direct oxidation of glucose to gluconic acid (GA) (55, 56). The enzyme taking part in this oxidation reaction is glucose dehydrogenase and pyrroloquinoline quinine (PQQ) as a cofactor (ppqABCD). Metagenome analysis of endophytic bacteria from this medicinal plant shows endophytes with great potential to solubilise both organic and inorganic phosphate. The genome contains *appA* genes for 4-phytase/acid phosphatase, 3-phytase which take part in the solubilisation of organic phosphate. The complete operon *ppqABCD*E for the co-factor pyrroloquinoline quinine (PQQ) and the gene *gcd* for glucose dehydrogenase were present and it highlights the solubilisation potential of inorganic phosphate. Presence of exopolyphosphatases (*ppx-gppA*) and inorganic phosphatase (*ppa*) shows its potential to solubilise the phosphate to make them available to plants. Along with all these phosphate solubilisation capacities the metagenome contains genes for phosphate starvation inducible proteins (*PhoH/L*), phosphate transport system substrate / ATP binding protein (*pstS/B*) and low affinity inorganic phosphate transporter (*pit*).

Many researchers conducted the studies on the nitrogen fixation ability of endophytic bacteria. In the present study, the dominant family of endophytic bacteria was *Enterobacteriaceae*. Many members from family *enterobacteriaceae* were already reported as nitrogen fixers. *Enterobactor* sp. And *Klebsiella* sp. from sugarcane showed *nifH* gene for nitrogen fixation (57). The processes of nitrogen fixation in most of the nitrogen-fixing microbes occur with the help of *Mo*-Nitrogenase. This enzyme is composed of two metallo-proteins *NifDK* and *NifH* (58). The genes *nifD*, *nifK*, *nifH*, and the protein coding genes regulating and assembling *Mo*-Nitrogenase like *nifU*, *nifA*, *nifN*, *nifE*, *nifX*, *nifQ*, *nifT*, *nifS* were present in this study. Along with genes for nitrogen fixation, there were enzymes for dissimilatory nitrate reduction to ammonia. The additional mechanism for nitrogen assimilation occurs in two steps viz. denitrification and reduction (59). We also found the different enzymes involved in the metagenome analysis like nitrous-oxide reductase (*nosZ*), nitrate reductase (*narG, napA*), nitrite reductase (*nirD*) etc. All these enzymes were also reported from an endophytic *Enterobactor* sp. SA187 isolated from *Indigofera argentea* (60). Glutamine is one of the amino acid metabolites then takes a major role in ammonium assimilation. Cellular nitrogen status regulates the amount of glutamine metabolism in a bacterial cell. This showed the role of glutamine in the assimilation of nitrogen and as a signalling molecule in nitrogen metabolism (61).

Complete genome analysis of plant growth-promoting endophytic bacterium *Enterobactor* sp.638 isolated from Poplar contains different iron uptake systems (62). The microorganism developed different solutions for iron acquisition like the production of chelators (siderophores). Siderophore act as a transport vehicle for iron (63). The presence of microbial siderophore act as a direct supply of iron for plants and a sufficient amount of iron-related to the immune system of the plant helps to prevent some diseases. The biocontrol efficiency of *Pseudomonas fluorescens* against fusarium wilt in tomato was analysed, and it was also based on siderophore production. The *P. fluorescens* was found effective in siderophore production and prevention of fusarium wilt (64). The presence of genes for enterobactin siderophore synthetase components F and D (*entF, D*), other proteins and enzymes involved in its production (*entE, C, B*) and enterobactin exporter (*entS*) and discloses the iron acquisition potential of the endophytic bacteria from *E. sonchifolia*.

**Conclusion**

The metagenome analysis of endophytic bacterial genome revealed high genetic diversity and diverse functional genes. The endophytic microbiome incorporates different genes which were potentially involved in the plant growth promotion activities like the production of plant growth regulators, enhancement of nitrogen availability by its assimilation through various pathways, solubilisation of phosphate and increase in the iron acquisition by the production of siderophore. Different genes with a functional role as participation in nitrogen metabolism (*nif* and siderophore production (enterobactin category) were noticed in the annotations. IAA production capacity of the endophytic microbiome was indicated by enzyme coding gene annotations like *ipDC* and *tnaA*. Presence of enzyme ACC deaminase coding gene in the metagenomic data shows the endophytic role in regulating elevated ethylene levels in host tissues. A great majority of genes having functional role in nutrient assimilation and absorption like genes for assembling *Mo*-Nitrogenase, nitrous-oxide reductase (*nosZ*), nitrate reductase (*narG, napA*), nitrite reductase (*nirD*), enterobactin siderophore synthetase components F and D and acid phosphatase explains the effective plant-microbe interactive relationship and the role of bacterial endophytic microbes in regulating the growth of host plant.

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Authors’ contributions

AM conceived, designed and coordinated the experiments and corrected the manuscript. SKU carried out the experiment, analysed the data and written the manuscript. Both authors read and approved the final manuscript.

Conflict of interests

Conflict of interest: The authors have no conflict of interest to declare.

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