Comparative *de novo* transcriptome analysis of flower and root of *Oliveria decumbens* Vent. to identify putative genes in terpenes biosynthesis pathway

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The *Oliveria decumbens* Vent. is a wild, rare, annual medicinal plant and endemic plant of Iran that has metabolites (mostly terpenes) which make it a precious plant in Persian Traditional Medicine and also a potential chemotherapeutic agent. The lack of genetic resources has slowed the discovery of genes involved in the terpenes biosynthesis pathway. It is a wild relative of *Daucus carota*. In this research, we performed the transcriptomic differences between two samples, flower and root of *Oliveria decumbens*, and also analyze the expression value of the genes involved in terpenoid biosynthesis by RNA-seq and its essential oil’s phytochemicals analyzed by GC/MS. In total, 136,031,188 reads from two samples of flower and root have been produced. The result shows that the MEP pathway is mostly active in the flower and the MVA in the root. Three genes of *GPP*, *FPPS*, and *GGPP* that are the precursors in the synthesis of mono, di, and triterpenes are upregulated in root and 23 key genes were identified that are involved in the biosynthesis of terpenes. Three genes had the highest upregulation in the root including, and on the other hand, another three genes had the expression only in the flower. Meanwhile, 191 and 185 upregulated genes in the flower and root of the plant, respectively, were selected for the gene ontology analysis and reconstruction of co-expression networks. The current research is the first of its kind on *Oliveria decumbens* transcriptome and discussed 67 genes that have been deposited into the NCBI database. Collectively, the information obtained in this study unveils the new insights into characterizing the genetic blueprint of *Oliveria decumbens* Vent. which paved the way for medical/plant biotechnology and the pharmaceutical industry in the future.

**KEYWORDS**

co-expression network, computational genetics, illumine, medicinal plant, terpenoid
Introduction

The *Oliveria decumbens* Vent. is an aromatic annual herb from the Apiaceae family and endemic to Iran. The flower of the plant is prescribed in Persian Traditional Medicine (PTM) in the form of dry powder/infusion for dyspepsia, pyrexia, and diarrhea also as an anti-inflammatory and antibacterial agent (Alizadeh Behbahani et al., 2018; Eftekari et al., 2021). The height of the plant varies from 20 to 45 cm. Mostly collected from the >2500 m heights from the skirts of central Zagros mountains in Fars, Chaharmahal & Bakhtiari, Kohgilouyeh & Boyerahmad, Boushehr, and Khuzestan provinces. The top chemical compounds have been reported in the plant including thymol, carvacrol, gamma-terpinene, and para-cymene (Khoshbakht et al., 2020). Each one of them has significant chemotherapeutic agent potential and also antibacterial properties (Ashraf, 2020; Ashraf et al., 2022). In this plant, they are available in high percentages which makes it a great candidate as a plant-based anti-cancer agent. In a very latest study, Khodavirdipour and his team elucidated the anti-cancer and anti-metastatic effect of this rare medicinal plant on HT-29 colorectal cell line (Khodavirdipour et al., 2021).

The core structure of terpenes consists of IPP (isopentenyl pyrophosphate) and its isomer, DMAPP (dimethylallyl pyrophosphate), where both of them synthesize via the cytosolic pathway, MVA (mevalonate pathway), and non-mevalonate plastid pathway, MEP (methylerythritol 4-phosphate). MVA pathway starts with 3 units of Acetyl-CoA and finally ends up in the production of FPP (Farnesyl pyrophosphate) (Bergman et al., 2019). The FPP has been formed by a condensation reaction of 2 units of IPP and a single unit of DMAPP and is an immediate precursor to all terpenes and terpenoids and sterols. Pyruvate and glyceraldehyde 3-phosphate are precursors in plastid-based MEP pathways. GPP (geranyl pyrophosphate) is a precursor to monoterpene, diterpene, and tetraterpenoids (carotenoids) and is produced by condensation of one unit DMAPP and one unit IPP. In later stages, the linear carbon backbone of GPP and FPP is converted into the main backbone of terpenes by TPS (terpene synthase) enzymes. Saponins are a terpenoid subclass and the biggest class of plant metabolites. They are active ingredients in more than 100 families of sea and soil endophytic fungi. The complex structure of Saponins is portrayed in physical, chemical, biological, and medicinal variations and resulted in Saponin’s importance in the health industry, cosmetics, and pharmaceutical sectors (Netala et al., 2014). Despite research conducted in gene profiling on genes involved in terpenes biosynthesis, still many mechanisms behind the secondary metabolite synthesis. Identification of key genes involving pathways is unknown and needed in-depth analysis (Sun et al., 2010; Hwang et al., 2015). There are numerous studies conducted to identify the genes involved in terpenes synthesis by the NGS method. In one of the studies using the NGS aiming to identify the involved genes in Saponins synthesis in *P. grandiflorum*, they produced more than 58 million reads, and by employing the de novo assembly genes, expression profiles in triterpenes synthesis have been determined (Ma et al., 2016). In another research on Siraitia grosvenorii to find the genes in the biosynthesis of triterpenes, more the 48 million reads have been analysed from the 2 samples from different time frames. The first one is 50 days and the second one 70 days after the flowering by the Illumina solexa method. The results suggest that there are seven gene families of Cytochromes P450 (CYP450) and five gene families of Uridine diphosphate glucose (UDPG) in magrosides biosynthesis (Tang et al., 2011). 18α-glycyrrhetinic acid and 18β-glycyrrhetinic acids are the two core and key structural medicinal compounds accumulated in *Glycyrrhiza* glabra. In RNA-seq research, the effects of three different level X-ray emissions were evaluated on the triterpenes biosynthesis pathway. Finally, 1376 genes, under influence of treatment, had the highest upregulation, and the genes involved in this pathway vigorously affected the synthesis above triterpenes (Gao et al., 2020). The comparative transcriptomic analysis of *Momordica charantia* was performed by RNA-seq. The findings were showing that despite fruits being the main accumulation sites of Cucurbitacins, but the highest upregulation and expression of genes happens in the leaves of the plants. This phenomenon displays that the biosynthesis of the Cucurbitacins occurs in the leaves and is transferred and amasses in the fruits (Takase et al., 2019). In another project, root, stem, and the leaves of *Entada phaseoloides* analyzed by NGS RNA-seq methods and yield more than 53 million reads have been produced. The results suggest that the highest upregulation of genes involved in triterpenes synthesis was observed in the stem of the plant (Liao et al., 2020). Other studies were also performed on mungbean, soybean, and Indian barnyard millet (Li et al., 2017; Jayakodi et al., 2019; Sudha et al., 2022). There is no study so far conducted to identify the key genes responsible for terpenes biosynthesis and, by and large, the reason behind the medicinal properties of the *Oliveria decumbens* Vent. by the latest sequencing techniques, and current research is the first of its kind in research on de novo assembly study on the rare endemic plant of Iran. This study designed and performed to identify the genes involved in the terpenoid biosynthesis pathway and build an expression profile of those genes in both root and flower of the *Oliveria decumbens* aiming to compare the intensity of metabolite synthesis in root and flower and also to find the exact site of production and accumulation of those secondary metabolites in both organs. Besides that, a complete metabolomic analysis of root and flower of the plant was performed for the first time to identify medicinally valued compound.

Materials and methods

Plant material

The study was carried out in compliance with all relevant guidelines and regulations. The plant samples include the root material.
and flower of the wild plant *Oliveria decumbens* Vent. collected from the >2000m heights of the Zagros mountains of Kazeroun city-Fars province at the end of May 2020, identified by Dr. A.R. Khosravi, Division of Plant Biology, Department of Biology, Faculty of Sciences, Shiraz University-Fars province (herbarium #56024).

**Essential oil isolation and GC-MS analysis**

The shadow-dried powder of the two samples (120 gm of root and 150 gm of flower) was exposed to hydro-distillation for 4 h by a Clevenger device. The obtained oil was again dried with anhydrous sodium sulfate and then later kept at 4°C.

GC-MS was carried out by Agilent GC (7890A)/MS (5975) with a capillary column, He gas carrier with a flow rate of 1.5 ml/ min, flame ionization detector, and a split ratio of 1:25. The temperature was set at 50°C for the column for 1 min and then for 265°C for the heating purpose at the rate of 2.5°C/min, then stored at 265°C for 20min. detector temp at 300°C, injector temp at 265°C flow of Helium at 35 ml/min with the airflow rate of 400 ml/min. The MS set off at 70 eV ionization energy. Retention times have been recorded and calculated by utilizing the n-alkanes retention times which have been injected in the same condition after the oil (www.agilent.com). The final found chemical compounds were reported by the “Agilent’s ChemStation Integrator” algorithm (www.agilent.com).

**RNA isolation, sequencing, and trimming**

The total RNA extraction of two samples (flower and root) was performed on healthy and suitable samples by the Japelagi method (Japelagi et al., 2011). The flower parts used for this study were the petal and last 10–12 cm of the root of the plant. Quality and quantity of the extracted RNA were analyzed and suitable samples have been sent to Macrogen (Seoul-South Korea) for RNA-seq by Illumina HiSeq 2000 platform with paired-end and read length of 101 bp at a sequencing depth of 10G. All the primary analyses include trimming, *de novo* assembly, mapping, nucleotide BLAST, and statistical analysis done by CLC Genomics Workbench 20.0.4 (http://www.qiagenbioinformatics.com). The software has been accessed for analyzing and statistical purposes from September to early November 2021.

**De novo transcriptome assembly, mapping, and annotation**

After the *de novo* assembly of both the root and flower of the plant, they separately assembled on the genome for the mapping. The statistical analysis was done by the Z test method ((Kal et al., 1999). Unigenes BLAST was performed based on the NCBI nucleotide database and finally, genes were selected by highest e-value and identity percentage, for gene ontology and network analyses and performed Convert Gene ID (homology search) based on the *Arabidopsis thaliana* as a model organism. To convert gene identifications (GI) of identified genes to homolog GIs in *Arabidopsis thaliana*, the ensemble plant database and web-based Biomart software were used ((Kinsella et al., 2011; Hussain et al., 2021)).

**Gene ontology and co-expression network analysis**

Cytoscape 3.7.2 software (Shannon et al., 2003) and BinGo plug-in (Maere et al., 2005) were used for gene ontology purposes. Gene co-expression network for selected genes from both root and flower of *Oliveria decumbens* Vent. was performed and re-constructed by STRING web service (Szklarczyk et al., 2017) Cytoscape v.3.7.2. Network topology was evaluated and analyzed by the NetworkAnalyzer 2.7 plug-in of Cytoscape software (Assenov et al., 2008).

**Validation of RNA-seq results**

To validate, based on the functional annotation importance, five genes related to plants are involved in terpenoid synthesis in *Oliveria decumbens* Vent. has been selected for validation through quantitative real-time PCR (qRT-PCR), and based on assembled sequence, the suitable primer for real-time-PCR has been designed by Primer3 plus web tools (Table 1) (Untergasser et al., 2007). The Actin7 gene has been chosen as the reference gene. Consequently, the DNA-free RNA was used for first-strand cDNA synthesis by transcriptor First Strand cDNA Synthesis Kit (Cinnagen, Tehran, Iran) following the manufacturer’s instructions. PCR amplification was performed using 2× SYBR Green PCR Master Mix (Roche Applied Science, Penzberg, Germany) and Light-Cycler® 480 (TaKaRa, Kusatsu, Shiga, Japan) following the standard protocol Relative expression of desired genes was obtained based on their Ct and analyzed by Pfaff formula and calculated by 2^−ΔΔCT (Livak and Schmittgen, 2001).

**Results and discussion**

**Metabolite components analysis by GC/MS in both flower and the root of *Oliveria decumbens***

GC/MS data which is analyzed by “Agilent’s Chemstation Integrator” showed that the compounds found in the flower
include (in order of %, higher to lower): gamma-terpinene, p-cymene, limonene, thymol, beta-pinene, carvacrol, (23S)-ethylcholest-5-en-3beta-ol, clionasterol, beta-myrcene, myristicin, alpha-pinene, alpha-thujene, alpha-terpinene, 4,5-dimethyl-1,2-phenylenediamine, and 3-carene; while the chemical composition of *Oliveria decumbens* Vent. root deciphered for the very first time and shed light for future in detail *in vitro* and *in vivo* research. The top 15 compounds of the root include linolyl acetate, linalool, alpha-pinene, 1,8-cineol, alpha-terpinolene, camphor, p-cymene, camphene, p-cymene, alpha-terpinene, gamma-terpinene, nonadecane, carvacrol, beta-Myrcene, and isoborneol (Tables 2, 3).

Totally, 136,031,188 reads were produced, and after trimming and removing the adaptor sequences, 135,009,232 high-quality reads were used for *de novo* assembly (Table 4). After the *de novo* assembly, both
sequenced samples of root and flower were separately mapped on the assembled genome. Out of 74,140,058 reads, 63,038,304 reads for the flower and out of 61,264,048 reads of root, 50,603,051 reads were successfully mapped (Table 5). The mapping result showed that out of 27,118 unigenes, 23,219 unigenes have shown a significant differential expression at \( p < 0.01 \). To a great extent, 10,685 unigenes of root showed at least double upregulation of gene compared to the flower of...
Oliveria decumbens vent. and vice versa. 3,449 unigenes of flower displayed at least double upregulation in comparison to the root. 1,060 unigenes were specifically expressed in a single sample only (either root or flower). Based on the mapping results, upregulated unigenes were selected by thresholds such as \( p < 0.01 \), fold change >10, and < -10, and also a minimum of 5000 expression copies per unigenes in both flower and root samples. 191 and 189 unigenes from the flower and the root, respectively, selected for the BLAST (Supplementary Material S1). The selected unigenes primarily BLAST with all organisms. The BLAST findings determined that Oliveria decumbens Vent. has the most similarity to the Daucus carota with 78% similarity. In the next step, the GENE ID of identified genes is converted into GENE ID of homolog genes in Arabidopsis thaliana as a model organism (Supplementary Material S2).

### Gene ontology

The result of upregulated genes in both flower and root of the Oliveria decumbens Vent. showed that 115 cellular components, 772 biological pathways, and 98 molecular functions at the significant level of 5% had at least one representative in the upregulated genes sector. As shown in Figure 1, the flower of the Oliveria decumbens Vent. is three times more active than root tissue. It has more active
organelles and displays distinct and diverse biological pathways and molecular functions. Undoubtedly, one of the reasons shall be the diverse organelles of the flower with distinguished functions and sundry cellular tissues in comparison to the root structure (Supplementary Material S3).
The biosynthesis of isoprenoids is a vital process in all living organisms. Isoprenoids also have economic importance in agriculture and industries. Prenyl-PP (prenyl diphosphate) are precursors of all isoprenoids. Unlike the biosynthesis of isoprenoids in other living organisms, prenyl diphosphate is synthesized by two distinct pathways: MEP and MVA. The 2-C-methyl-D-erythritol 4-phosphate (MEP) pathway in plastids and the cytoplasmic pathway of the mevalonate (MVA) (Figure 2). However, all living organisms use only one of the above pathways, except plants (Yang et al., 2012). Plants optimize the regulation of isoprenoid biosynthesis according to ATP availability and fixed carbon and via the MEP pathway in the plastid or the MVA pathway of cytoplasm. Furthermore, survival pressure in different environments shall ease the various metabolites of specialized complex isoprenoid-derived biosynthesis through the use of both MEP and MVA pathways (Vranová et al., 2013). Understanding the expression of genes involved in both the above pathways in different tissues shall provide a clear perspective of the process. As it has shown in Table 4, genes involved in the MEP pathway show the highest upregulations in the flower tissue, and in the contrast, most genes involved in the MVA pathway are observed in the root tissue. With regards to that, flower tissue mostly consists of plastid and chloroplast compared to the root tissue; the high level of MEP activity in the flower was not unexpected. The gene ontology results also confirm the greater activity of the genes involved in the MEP pathway in the plastid and chloroplast of the flower than in the root.

TABLE 4 Brief reports on quality and quantity of sequencing and de novo assembly.

| Parameter                      | Length |
|--------------------------------|--------|
| Unigene measurements (excluding scaffolded regions) |        |
| N75                            | 750    |
| N50                            | 1,259  |
| N25                            | 2,094  |
| Minimum                        | 53     |
| Maximum                        | 20,380 |
| Average                        | 978    |
| Count                          | 32,734 |
| Total                          | 32,011,608 |
| Unigene measurements (including scaffolded regions) |        |
| N75                            | 836    |
| N50                            | 1,368  |
| N25                            | 2,235  |
| Minimum                        | 500    |
| Maximum                        | 20,380 |
| Average                        | 1,181  |
| Count                          | 27,118 |
| Total                          | 32,036,864 |

Analyzing the expression of genes involved in terpenes biosynthesis pathway in *Oliveria decumbens* Vent.

The biosynthesis of isoprenoids is a vital process in all living organisms. Isoprenoids also have economic importance in agriculture and industries. Prenyl-PP (prenyl diphosphate) are precursors of all isoprenoids. Unlike the biosynthesis of isoprenoids in other living organisms, prenyl diphosphate is synthesized by two distinct pathways: MEP and MVA. The 2-C-methyl-D-erythritol 4-phosphate (MEP) pathway in plastids and the cytoplasmic pathway of the mevalonate (MVA) (Figure 2). However, all living organisms use only one of the above pathways, except plants (Yang et al., 2012). Plants optimize the regulation of isoprenoid biosynthesis according to ATP availability and fixed carbon and via the MEP pathway in the plastid or the MVA pathway of cytoplasm. Furthermore, survival pressure in different environments shall ease the various metabolites of specialized complex isoprenoid-derived biosynthesis through the use of both MEP and MVA pathways (Vranová et al., 2013). Understanding the expression of genes involved in both the above pathways in different tissues shall provide a clear perspective of the process. As it has shown in Table 4, genes involved in the MEP pathway show the highest upregulations in the flower tissue, and in the contrast, most genes involved in the MVA pathway are observed in the root tissue. With regards to that, flower tissue mostly consists of plastid and chloroplast compared to the root tissue; the high level of MEP activity in the flower was not unexpected. The gene ontology results also confirm the greater activity of the genes involved in the MEP pathway in the plastid and chloroplast of the flower than in the root.
in the root (Pütter et al., 2017). Acetoacetyl-CoA is converted to HMG-CoA (β-hydroxy-β-methylglutaryl-CoA) by HMG synthase (HMGs EC 2.3.3.10). The expression of the HMGs gene in the root is higher than in flowers, like the AACT gene (Figure 2). HMG-CoA in two reductive steps converted to MVA (mevalonic acid) by 3-hydroxy-3-methylglutaryl-CoA reductase (HMG; EC 1.1.1.34). In the *Oliveria decumbens* Vent. too, expression of this gene in the root tissue was higher compared to the flower tissue. Entire plant proteins of HMG bind to the ER (endoplasmic reticulum) with their catalytic site facing the cytosol (Darabi et al., 2012). In two back-to-back reactions catalyzed by phosphor-MVA kinase (PMK; EC2.7.4.2; At1g31910) and MVA kinase (MK; EC 2.7.1.36; At5g27450), MVA is later phosphorylated to MVA 5-diphosphate. PMKs and MKs are encoded by single genes in most plants. PMKs are mainly active in peroxisomes while cytosol is the chief activity site of MKs (Zheng et al., 2020). The expression of both genes was almost equal in the flower and the root of *Oliveria decumbens* Vent. MKV (mevalonate kinase) is considered a potential regulatory enzyme in the isoprenoid biosynthesis pathway (Lluch et al., 2000). Evaluation and analysis by the Northern blot technique confirmed that the mRNA of this gene prefers to accumulate in inflorescences and the root of the plant (Lluch et al., 2000). In the following, the diphosphatevalonate decarboxylase enzyme (MVD EC 4.1.1.33) catalyzes the decarboxylation of mevalonate 5-diphosphate. The expression comparison of this gene in the flower and the root of *Oliveria decumbens* Vent. shows significant upregulation in the root compared to the flower. As is shown in Table 5, out of eight genes involved in the MVA biosynthesis pathway, seven genes have the highest upregulation in the root tissue, and only the MVK gene shows slight upregulation in the flower of the plant (around 10%). The highest upregulation is going to the HMG gene by 4568 reads. Also, this gene with 1906 reads had the highest level of expression out of all genes in the MVA biosynthesis pathway in the flower tissue. The four end genes of the MVA pathway including PMK, MVD, PMD, and IPK, in both flower and root tissues, had the lowest expressions, compared to the genes of the beginning of the pathway. Generally, the total expression of MVA pathway genes in the root was recorded as 11,689 reads and 6457 reads for the flower, which shows that expression difference is almost double in the root compared to the flower (Table 6).

Analyzing the genes involved in the MEP pathway in the root and the flower of *Oliveria decumbens* Vent.

In the beginning, the MEP pathway is known as the Rohmer or non-mevalonate pathway (Banerjee and Sharkey, 2014). The MEP pathway initiated with the synthesis of 1-deoxy-β-xylulose 5-phosphate from n-glyceraldehyde 3-phosphate and pyruvate catalyzed by 1-deoxy-β-xylulose 5-phosphate synthase (DXS; EC 2.2.1.7). In most the species of the plant, DXS is encoded by several paralogs of the gene. The entire enzymes of the MEP pathway (except DXS) in *A. thaliana* and nearly all other plants are encoded by single-copy genes (Vranová et al., 2013). Expression of DXS gene in the flower of *Oliveria decumbens* Vent. was around 13 times its expression in the root. DXP reductoisomerase (DXR; EC 1.1.1.267) is the second enzyme in the MEP pathway which uses NADPH for reduction. The expression of the DXR gene in the flower of the *Oliveria decumbens* Vent. found to be double its expression in the root. 2-C-Methyl-D-erythritol 4-phosphate converted to CDP-ME by methyerythritol phosphate cytidyllitransferase (MCT; EC 2.7.7.60). The expression value of this gene was almost equal in both flower and the root of the *Oliveria decumbens* Vent. The position of the hydroxyl group in CDP-ME’s C2 position is phosphorylated by 4-(cytidine 5’-diphospho)-2-C-methyl-D-erythritol kinase (CMK; EC 2.7.1.148; at2g26930). The expression of the CMK gene in the flower was double of the root. 2-C-Methyl-d-erythritol-2,4-cyclopropylphosphate (MeCPP) enzyme (MDS EC 4.6.1.12) catalyzed 4-CDP-2-C-methyl-D-erythritol 2-phosphate (CDP-ME2P) to 2-C-Methyl-d-erythritol-2,4-cyclopropylphosphate (MePP). The MDS gene shows double expression in the flower compared to the root of the plant. Later, MeCPP converted to HMBP by 4-hydroxy-3-methylbut-2-enyl-diphosphate synthase (ferredoxin) (HDS EC 1.17.7.1), and finally, HMBP converted to a mixture of IPP and DMAPP by 4-hydroxy-3-methylbut-2-enyl diphosphate reductase (HCR; EC 1.17.1.2). The expression of both HDS and HDR genes in the flower of *Oliveria decumbens* is more than in the root. The synthesized IPP and DMAPP in the MVA pathway are used in isoprenoid synthesis in the cytosol and the mitochondria while the IPP and DMAPP-which are synthesized by the MEP pathway-used in isoprenoid synthesized in the plastids (Berthelot et al., 2012). Conversion of DMAPP from IPP and the balance between DMAPP and the IPP are controlled by IPP-isomerase (IPPI; EC 5.3.3.2) in a reversible reaction (Berthelot et al., 2012). The RNA-seq results of the *Oliveria decumbens* Vent. showed the double difference in expression of IPP gene in the lower compared to the root. DMAPP is further used for the synthesis of shorter chains prenyl-PPs such as GPP (geranyl diphosphate), FPP, and GGPP by adding the IPP. GPP, FPP, and GGPP are the main synthesis core of all isoprenoids’ final products. GPP synthase enzyme (GPPS EC 2.5.1.1) produces the GPP as the monoterpenoid precursor. FPP synthesizes by the farnesyl pyrophosphate synthase (FPPS EC 2.5.1.10) as the main precursor for triterpenoids, sterols, sesquiterpenoids, brassinosteroids, oligoprenols, polyrenols, and dolichols for prenylation of proteins. FPPS shall bind IPP to both GPP and DMAPP as substrates. GGPP pool in plants is used for the synthesis of gibberellins, chlorophylls, diterpenoids, plastoquinone, phylloquinone, carotenoids, tocopherols, oligoprenols, abscisic acid, polyphenols, and dolichols for...
protein geranylgeranylation. GGPP is synthesized by the GGPP synthase (GGPPS; EC2.5.1.29), which is performed as a homodimer and utilizes all three allylic prenyl-PPs (GPP, FPP, and DMAPP) as substrate. The analysis of expression profile of involved genes in the MEP pathway of the Oliveria decumbens Vent. shows that the average expression of key genes in the root were 10,464 reads vs. flower's 27,329 reads. It is absolutely clear now that unlike the root tissue, MEP is the main pathway for the synthesis of terpenes in the flower tissue of Oliveria decumbens Vent. Of the eight genes in this pathway, seven genes had higher expression in the flower tissue, except the MCT gene which is with a very little margin expressed higher in the root. The total expression of MEP and MVA genes were 22,153 reads in the root vs. 33,777 reads of the flower. It can be concluded that generally more terpenes are synthesized in flower of the Oliveria decumbens Vent. than the root tissue. The three genes of GPP, GGPP, and FPPS which at the end of the MEP and MVA pathways catalyze the production of precursors for the synthesis of mono, di, and triterpenoids, respectively, had higher expression levels than flowers at the root. The mean expression of these three genes was 856 reads in the root vs. 604 reads of the flower (Table 6).

Synthesis of monoterpenoids, diterpenoids, and triterpenoids in Oliveria decumbens Vent.

The largest class of secondary metabolites of plants belong to terpenoids (terpenes). More than 36,000 distinct structures of this class have been reported so far (Dong et al., 2016). The terpene’s structure is extremely changeable and displays hundreds of different carbon skeletons. Nevertheless, this extensive structural diversity has a regular biosynthesis feature: entire terpenes are procured from the basic assembly process of 5-carbon atom isoprene units. The terpene compounds categories consist of one (C5 hemiterpenes), two isoprene units (C10 monoterpenes), three isoprene units (C15 sesquiterpenes), four isoprene units (C20 diterpenes), six isoprene units (C30 triterpenes), eight isoprene units (C40 tetraterpenes) or even more units namely called polyterpenes (>C40) (Dong et al., 2016). Limonene and its active serum-oxygenated metabolite derivatives, dihydroperillic acid and perlic acid, have been shown antiproliferative properties through their chemopreventive actions during the promotion stage of liver and breast carcinogenesis (Cox-Georgian et al., 2019). This is because of suppression of tumor cell proliferation, increase in tumor cell death rate, and/or induction of tumor cell differentiation (Cox-Georgian et al., 2019). As it is shown in Figure 3, a wide range of monoterpenes are formed by numerous enzymes in the GDP substrate in plants. Till now, a great number of genes encoding the catalyzing enzymes in monoterpenes biosynthesis in plants have been cloned; for example, linalool synthase (Jia et al., 1999), (-)-limonene synthase (Bohlmann et al., 1997), (+)-limonene synthase (Maruyama et al., 2002), myrcene synthase (Fischbach et al., 2001), (-)-camphene synthase, (-)-β-phellandrene synthase, (-)-limonene/(−)-α-pinene synthase, and terpinolene synthase (Bohlmann et al., 1999). It seems that enzymes leading to the production of primary monoterpenes skeletons appear to be active in plastids because all of the genes in this pathway that have been cloned to date have plastid target signals located in parenchymal cell chloroplasts and secretory cell leucoplasts (Aharoni et al., 2005). Comparison of the expression of the main genes involved in monoterpenoid synthesis in the root and the flower of the Oliveria decumbens Vent. shows a higher expression of all the genes except one gene in the flower tissue compared to the root tissue. The geranyl diphosphate
phosphohydrolase gene is present in free monoterpenes alcohols biosynthesis in the cytosolic pathway that promotes the fragrance of the aromatic plants. This process lacks the activity of terpene synthase but shows the diphosphohydrolase function with farnesyl diphosphate and geranyl diphosphate as substrates (Magnard et al., 2015). Table 7 shows the expression of several genes for the synthesis of mono, di, and triterpenoids. Of the 23 genes identified in Oliveria decumbens Vent. transcriptome, 19 genes showed higher expression in the flower tissue. In contrast, three genes in the root sample showed the higher expression, namely geranyl diphosphate phosphohydrolase, geranylinalool synthase, and ent-kaurene synthase. The expression value of the Vetispiradiene synthase gene was equal in both root and flower tissues of the Oliveria decumbens. Most of the genes showed only one homolog in the transcriptome of Oliveria decumbens. But the Germacrene D synthase gene had seven homologs, which could be due to the high similarity of the sequence of other genes in the same family, such as Germacrene-A synthase (EC: 4.2.3.23), Germacrene-B synthase (EC: 4.2.3.71), Germacrene-C synthase (EC: 4.2.3.60), Germacrene A oxidase (EC: 1.14.14.95) and germacradanol/ geosmin synthase (EC: 4.2.3.22). As there was no sequence in the NCBI database for any of these genes in the Apiaceae, it was not possible to differentiate them by sequencing. The total expression of the main genes in the synthesis of mono, di, and triterpenoids in the flower tissue was 25,511 reads vs. just 3753 reads for the root tissue of the Oliveria decumbens. These findings clearly show that the main site for the synthesis of these compounds is the flower with a huge gap compared to the root. Meanwhile, three genes of (R)-limonene synthase, (S)-limonene synthase, and beta-bisabolene synthase are expressed only in the flower sample and had zero expression in the root tissue. Monoterpenes are important aromatic molecules that are widely distributed in nature (more than 400 structures) that shall be extracted by the perfume industry from the leaves, flowers, and fruits of many plants (Chen et al., 2003). Volatile compounds as a mixture extracted from diverse metabolite classes of plants such as terpenes, fatty acid derivatives, sulfur or nitrogen-containing compounds, and phenylpropanoids have been shown to attract different insects (even some mammals) which are pollen vectors (Dudareva and Pichersky, 2000). In a study to analyse the volatile metabolites secreted from the vegetative parts and flowers of Arabidopsis by headspace sampling method. The result showed that a mixture of volatile terpenes is emitted particularly from inflorescences and distinguished the majority of plentiful monoterpenes including β-mycene, linalool, and limonene involved in the biosynthesis. It is discovered that most of the sesquiterpenes such as (+)-thujopsene, (-)-(E)-β-caryophyllene, (E)-β-farnesene, α-humulene, (–)-cuparene, and (+)-β-chamigrene were solely emitted from the flowers (Chen et al., 2003). Also, a comparison of limonene synthase gene expression showed that this gene is expressed exclusively in the flower of Oliveria decumbens Vent. Diterpenoids are defined as molecules deriving from the condensation of four isoprenyl units and containing 20 carbon atoms. Like terpenoids, they are ubiquitous throughout entire plants and many of them biosynthesized from geranylgeranyl diphosphate that consequently creates phytanes (acyclic), halimanes, clerodanes and labdanes (bicyclic), abietanes, rosanes, podocarpanes, pimaranes, cassanes, vouacapanes (tricyclic), kaurens, stemodanes, atisanes, trachylobanes, ingenanes and tiglanes (macrocyclic diterpenes) based on their cyclization (Malla et al., 2019). While biosynthesis of diterpenes has been substantially studied in fungi and plants as well, bacteria recently identified for their unique production of diterpenoids and are probably to keep an underexplored reservoir of new DTSs (Smanski et al., 2012). Diterpenoids belong to the abietane and pimarane series. Along with aromatized nor-abietane diterpenoids, they have been determined for showing anti-tubercular activity (Ulubelen et al., 1997). Triterpenoid saponins are synthesized through the isoprenoid pathway via 2,3-oxidosqualene cyclization to offer dammarane triterpenoid or oleananthe skeletons. The backbone of triterpenoid later undergoes numerous modifications such as glycosylation, substitution, and oxidation, detected by infrared with glycosyltransferases, cytochrome P450-dependent monooxygenases, and other enzymes (Haralampidis et al.,

**TABLE 5 Brief reports on mapping of flower and the root of the Oliveria decumbens Vent. with the de novo assembly transcriptome.**

| Mapping       | Flower | Root |
|---------------|--------|------|
|               | Number of fragments | Percentages of total | Number of fragments | Percentages of total |
| Counted fragments | 59,642,553 | 80.74 | 48,020,831 | 78.55 |
| unique fragments | 58,572,566 | 79.29 | 46,814,066 | 76.57 |
| non-specifically | 1,069,987 | 1.45 | 1,206,765 | 1.97 |
| Uncounted fragments | 14,230,033 | 19.26 | 13,115,815 | 21.45 |
| Total fragments | 73,872,586 | 100.00 | 61,136,646 | 100.00 |
Simple triterpenes are parts of specialized membranes and surface waxes and shall feasibly function as signaling molecules, where saponins (complex glycosylated triterpenes) furnish protection against pests and pathogens. Conjugated triterpenes or even simple ones have a wide range of uses in the health, biotechnology, and food sectors (Thimmappa et al., 2014). With regards to that the *Oliveria decumbens* plant has not been sequenced so far and the present study is the first transcriptome sequencing of this kind, finding the sequence of the main genes for the terpenoid synthesis requires doing the BLAST with genes at NCBI database on transcriptome of *Oliveria decumbens* plant. Alignment of already deposited genes for taxonomically remote families had no results. Therefore, inevitably the terpene synthetase genes sequence BLAST against Apiaceae family.

**Table 7** shows the expression of some terpene synthetase genes that can identify the relevant genes on the transcriptome of *Oliveria decumbens* Vent. by alignment. As can be seen, the expression of these genes in the flower of the *Oliveria decumbens* is much higher than in the root of the plant. In Persian traditional medicine, more emphasis has been placed on the medicinal effects of the *Oliveria decumbens* flower (Eftekhari et al., 2019; Khodavirdipour et al., 2021). Undoubtedly, however, a significant portion of terpenoids is transported between plant tissues in various ways after synthesis. The terpenoid transportation shall happen in at least three ways: transport in host cells, transport across the cell membrane, and intracellular transport (Fang and Xiao, 2021). Terpenoids are usually extracted from different parts of plants such as roots, leaves, or fruits. It must be considered that many of them were accumulated in the cuticular wax of different tissues of plants (Szakiel et al., 2012; Pensec et al., 2016; Pina et al., 2016). The cuticular wax consists of an epicuticular layer on the surface and an intracuticular layer implanted in the cutin matrix. The triterpenoid contents in the intracuticular layer are sometimes higher than in the epicuticular layer. A realizable elucidation for the triterpenoid’s uneven distribution in the cuticular wax tissue is the separation of the liquid-liquid phase since the long-chain compounds such as cyclic triterpenoids exhibit different conformation and molecular polarities (Buschhaus and Jetter, 2002). Simple triterpenes are parts of specialized membranes and surface waxes and shall feasibly function as signaling molecules, where saponins (complex glycosylated triterpenes) furnish protection against pests and pathogens. Conjugated triterpenes or even simple ones have a wide range of uses in the health, biotechnology, and food sectors (Thimmappa et al., 2014). With regards to that the *Oliveria decumbens* plant has not been sequenced so far and the present study is the first transcriptome sequencing of this kind, finding the sequence of the main genes for the terpenoid synthesis requires doing the BLAST with genes at NCBI database on transcriptome of *Oliveria decumbens* plant. Alignment of already deposited genes for taxonomically remote families had no results. Therefore, inevitably the terpene synthetase genes sequence BLAST against Apiaceae family.

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Comparison of the expression network of upregulated genes in the roots and flowers of *Oliveria decumbens*

The bigger size of the node is representative of higher betweenness centrality and its the exact term in bioinformatics analysis the Closeness centrality and blueish color showing the lower closeness centrality in the network. Also, the thicker and more reddish connective line, represents the higher Edge Betweenness; the thinner and tending to light blue color showing the lower edge centrality. Figure 4 shows the expression network of upregulated genes in the flowers of the *Oliveria decumbens*. Topology analysis of the network showed that GAPA-2, PRK, and PETC genes each with 35 connections, and FBA1 and GAPA genes each with 34 connections had the highest number of connections with other genes in the network. The two elements of closeness centrality and betweenness centrality are very useful in analyzing network topology. The centrality of a node within a complex network is called the closeness centrality, which is calculated based on the shortest communication lines between the nodes. In contrast, the term closeness centrality is the shortest distance from one node to other nodes (Shiri et al., 2018; 2020). In other words, the closer the amount of these two components are to each other, the more important that gene is in the network. Based on the closeness centrality component, 23 genes recorded the highest value, which is the same as score one, some of which are: PSBP-3, LACS2, and THI1. Also, the highest betweenness centrality values for CRU2, AT1G03890, CYP704A2, LACS2, and CYP86A7 genes with the following values 0.166667, 0.166667, 0.083333, 0.047619, and 0.047619 recorded, respectively (Supplementary Material S4). For genes expressed in the flowers, one main network and four sub-networks were observed. GAPA-2 genes are involved in the Calvin-Benson cycle (photosynthetic reductive pentose phosphate pathway). This gene by NADPH catalyzes the reduction of 1,3-diphosphoglycerate (Li et al., 2019). The studies show that this gene has a key role in responding to abiotic stress in wheat, and this stress resistance process shall be complemented by the H2O2-mediated ABA signaling pathway (Li et al., 2019). PRK regulates the germination of pollen and growth of pollen tube polar (Chang et al., 2013). The product of the PETC gene is a vital and key protein for phototrophy with the help of light dissipation of light energy and luminal acidification (increase of PH gradient) resistance to photo-oxidative damage. This product belongs to the cytochrome b6-f complex—which facilitates the electron transfer between photosystem I (PSI) and photosystem II (PSII)—and is responsible for state transfer and circular electron flow around the PSI (Munekage et al., 2001; Jahns et al., 2002; Maivard et al., 2003). Fructose-1,6-biphosphate aldolase (FBA) in chloroplast catalyzes the development of both sedoheptulose 1,7-bisphosphate and fructose 1,6-bisphosphate within the Calvin-Benson cycle (Carrera et al., 2021). While the expression of the FBA1 gene is most important in the aerial part of the plant, FBA3, its homolog, is expressed in the root (Carrera et al., 2021). PSBO1 and PSBP1 genes play a role in the photosystem II regulatory processes (Yi et al., 2005). One of the essential enzymes for the photosynthesis is RCA gene product named Rubisco activase. This enzyme removes the inhibitory sugar phosphates from the Rubisco’s active site, a necessary process for carbon fixation and activation of Rubisco (Nagarajan and Gill, 2018). THI1 gene has a key role in thiamine precursor thiazole biosynthesis. Catalyzes the conversion glycin and NAD to ADT (adenosine diphosphate 5-(2-hydroxyethyl)-4-methyl thiazole-2-carboxylic acid, an adenylation thiazole intermediate (Li et al., 2016). Analyzing the role of the other genes in the upregulated gene expression in the flower demonstrated that the most of genes are involved in the photosynthesis process, responses to biological and non-biological stresses, and also specific genes for the development of the flower tissue and synthesis of the fatty acids. Figure 5 shows the upregulated gene expression network in the root. 44 genes construct the four-gene network of the root. Analyzing the network topology elucidated that CCoAOMT1, IRX3, PAL4, TUA6, and IRX1 had the highest number of connections, respectively, with 9, 8, 7, 7, and 7 with other genes in the network. Based on the closeness centrality factor, TUA6, HSP81-1, TUA4, PATL1, and Hsp81.4 genes, respectively, expressed with 0.888889, 0.8, 0.8, 0.8 and 0.727273 values. Also in terms of the betweenness centrality component, PHI-1, PATL1, CCOAOMT1, and AT3G59080 genes had the highest values with 0.75, 0.666667, 0.50614, 0.5, and 0.46286 respectively (Supplementary Material S5). CCoAOMT1 (caffeoyl coenzyme A-dependent O-methyltransferase 1) is known to be one of the main OMTs (methyltransferases) in the lignin monomers biosynthesis, sinapoylalcohol. The CCoAOMT1 is exclusively and strongly expressed in the roots and stems of vascular tissues (Do et al., 2007; Kai et al., 2008; Fellenberg et al., 2012). IRX1 and 3 genes are involved in cellulose biosynthesis.
In higher plants, cellulose is synthesized by a multi-enzyme complex comprising multiple catalytic subunits and other proteins at the plasma membrane. In IRX3, synthesis of cellulose is deficient in the thickened secondary cell walls of interfascicular tissues and xylem in the stem (Ha et al., 2002). The product of the PAL4 gene is a crucial and essential enzyme in plant metabolism catalyzing the preliminary reaction in the biosynthesis of a wide variety of natural products based on the phenylpropane skeleton from L-phenylalanine (Cochrane et al., 2004).

TUA4 and TUA6 are key genes involved in tubulin synthesis in plants. Tubulin is the microtubule’s fundamental constituent, and it is bound to two GTP moles, one at a non-exchangeable site on the alpha chain and one at an exchangeable site on the beta chain. The mutant studies of these genes in Arabidopsis demonstrated that despite the detection of no obvious abnormalities in aerial parts of plants, root growth was seriously affected. The root tip cells contained abnormal microtubule structure, atypical proliferation, and the impracticality of normal cell division. These cellular defects may cause significant radial expansion of the root tip and also prevent elongation of the root (Bao et al., 2001; Matsumoto et al., 2010). Hsp81.4 and HSP81-2 are genes of molecular chaperones that boost structural maintenance, maturation, and specific target protein’s proper regulation that is involved in signal transduction and cell cycle control. It undergoes a functional cycle that is linked to its ATPase activity. The said cycle is likely to induce client proteins conformational changes, by that means promoting their activation. This functional cycle interconnect interactively with different co-chaperones that regulate its recognition of substrate, chaperone function, and ATPase cycle (Schenck et al., 2013).

Hsp81.4 and HSP81-2 are key genes involved in the membrane transfer events associated with cell plate proliferation or maturation, hinting at the involvement of phosphoinositides in cell plate biogenesis (Peterman et al., 2004). In general, it can be concluded that the genes of expression network of *Oliveria decumbens* root are mostly

| Gene name | Protein name | Enzyme code | Number of unigenes | Expression value in flower | Expression value in root |
|-----------|--------------|-------------|--------------------|---------------------------|-------------------------|
| NUDX1     | Geranyl diphosphate phosphohydrolase | 3.6.1.68 | 3 | 256 | 548 |
| GES       | Geraniol synthase | 3.1.7.11 | 1 | 502 | 132 |
| TPS14     | (S)-linalool synthase | 4.2.3.25 | 2 | 164 | 22 |
| MYRS      | Myrcene/ocimene synthase | 4.2.3.15 | 1 | 12,206 | 31 |
| LIS       | (R)-linalool synthase | 4.2.3.26 | 1 | 127 | 116 |
| sLIMS     | (S)-limonene synthase | 4.2.3.16 | 1 | 344 | 0 |
| rLIMS     | (R)-limonene synthase | 4.2.3.20 | 1 | 605 | 0 |
| PFS       | (-)-Alpha-pinene/(-)-camphene synthase | 4.2.3.119 | 1 | 375 | 16 |
| QH6       | (-)-Beta-pinene synthase | 4.2.3.120 | 1 | 3609 | 39 |
| FES       | (-)-Endo-fenchol synthase | 4.2.3.10 | 1 | 256 | 122 |
| SABS      | Sabinene-hydrate synthase | 4.2.3.11 | 1 | 3006 | 78 |
| TPS26     | (-)-Alpha-terpine synthase | 4.2.3.111 | 1 | 137 | 4 |
| CAMS      | (+)-Camphene synthase | 4.2.3.116 | 1 | 138 | 23 |
| TPS3      | (-)-Camphene synthase | 4.2.3.117 | 1 | 668 | 363 |
| TPS1      | (+)-Bornyl diphosphate synthase | 5.5.1.13 | 3 | 319 | 157 |
| CPS1      | Ent-copalyl diphosphate synthase | 4.2.3.19 | 1 | 75 | 125 |
| KS1       | Ent-lauren synthase | 4.2.3.144 | 1 | 340 | 389 |
| GLS       | Germacrindiol synthase | 4.2.3.75 | 7 | 348 | 304 |
| GERDS     | Germacrene D synthase | 4.2.3.21 | 1 | 93 | 93 |
| PVS1      | Vetispiradiene synthase | 4.2.3.55 | 1 | 285 | 0 |
| TPS1      | Beta-bisabolene synthase | 5.4.99.8 | 1 | 507 | 450 |
| CAS1      | Cycloartenol synthase | 4.2.1.125 | 1 | 963 | 698 |
## TABLE 8 List of sequences identified in *Oliveria decumbens* in this research so far and deposited to the NCBI database.

| Accession number | Gene name |
|------------------|-----------|
| MK392541         | Gamma-terpinene synthase mRNA, partial cds |
| MK392542         | Terpene synthase mRNA, partial cds |
| MK392543         | 1-Deoxy-α-D-xylulose 5-phosphate reductoisomerase mRNA, partial cds |
| MK392544         | Cytochrome P450 mRNA, partial cds |
| MK392545         | (R)-limonene synthase mRNA, partial cds |
| MN398148         | E3 ubiquitin-protein ligase UPL 2-like mRNA, complete cds |
| MN398149         | Midasin mRNA, complete cds |
| MN398150         | Putative magnesium transporter NIPAB mRNA, complete cds |
| MN398152         | Protein decapping 5 mRNA, complete cds |
| MN398153         | RRP12-like protein mRNA, complete cds |
| MT002142         | Protein cathepsin B mRNA, complete cds |
| MT002143         | Protein indole-3-acetic acid-amido synthetase GH3.6 mRNA, partial cds |
| MT002144         | Protein DNA-directed RNA polymerase II subunit 1 mRNA, partial cds |
| MT002145         | Hypothetical protein mRNA, complete cds |
| MN954564         | Hypothetical protein mRNA, complete cds |
| MN954565         | Protein guanylate kinase 2 (GK-2) mRNA, partial cds |
| MN937350         | NADH-plastoquinone oxidoreductase subunit 5 protein (nadB) mRNA, complete cds |
| MN937351         | Curvature thylakoid 1A protein (CURT1A) mRNA, complete cds |
| MN937352         | PIN-LIKE 3-like protein (PILS3) mRNA, partial cds |
| MN900853         | Strictosidine synthase-like 10-like protein mRNA, complete cds |
| MN900854         | ATP synthase CF1 alpha subunit protein mRNA, complete cds |
| MN900855         | photosystem II 47 kDa protein mRNA, complete cds |
| MN984141         | Two-component response regulator-like APRR7 protein mRNA, complete cds |
| MN984142         | CRIB domain-containing protein RIC6-like protein mRNA, partial cds |
| MN984143         | Exportin 1A protein mRNA, complete cds |
| MN984144         | maturase protein mRNA, partial cds |
| MN892226         | Cytochrome P450 7A6-like protein mRNA, partial cds |
| MN891826         | WRKY transcription factor 7 protein mRNA, complete cds |
| MN882159         | Non-specific lipid-transfer protein mRNA, complete cds |
| MN882160         | Hypothetical protein mRNA, complete cds |
| MN882161         | Hypothetical protein mRNA, complete cds |
| MN864743         | Non-functional NADPH-dependent codeinone reductase 2-like protein mRNA, complete cds |
| MN861110         | ATP-dependent helicase BRM-like protein mRNA, complete cds |
| MN861111         | Heat shock cognate 70 kDa protein 2 mRNA, complete cds |
| MN848166         | Chloroplast cytochrome P450 monoxygenase 97A3 mRNA, complete cds, nuclear gene for chloroplast product |
| MN818813         | BTB/POZ domain-containing protein mRNA, complete cds |
| MW435622         | Acetyl-CoA acetyltransferase cytosolic 1 (AAT1) mRNA, partial cds |
| MW435623         | Dammarenediol II synthase protein (DDS) mRNA, partial cds |
| MW447134         | Cycloartenol synthase (CAS1) mRNA, partial cds |
| MW447135         | Dammarenediol II synthase protein transcript variant X2 (DDS) mRNA, partial cds |
| MW447136         | Dammarenediol II synthase protein transcript variant X3 (DDS) mRNA, partial cds |
| MW447137         | Farnesyl pyrophosphate synthase 1 (FPS1) mRNA, partial cds |
| MW447138         | 3-Hydroxy-3-methylglutaryl coenzyme A reductase (HMGR) mRNA, partial cds |
| MW447139         | Hydroxymethylglutaryl-CoA synthase (HMGCS1) mRNA, partial cds |
| MW447140         | Diphosphomevalonate decarboxylase (MVD2) mRNA, complete cds |
| MW447141         | Mevalonate kinase (MVK) mRNA, partial cds |
| MW447142         | Phosphomevalonate kinase (PMVK) mRNA, partial cds |

(Continued on following page)
**TABLE 8 (Continued) List of sequences identified in Oliveria decumbens in this research so far and deposited to the NCBI database.**

| Accession number | Gene name |
|------------------|-----------|
| MW447143         | Squalene synthase (FDFT1) mRNA, partial cds |
| MW447144         | Squalene monoxygenase transcript variant x1 (SQLE) mRNA, partial cds |
| MW447145         | Squalene monoxygenase transcript variant x2 (SQLE) mRNA, partial cds |
| MW447146         | Squalene monoxygenase transcript variant x3 (SQLE) mRNA, partial cds |
| MT920432         | ATP-dependent zinc metalloprotease FTSH 7 mRNA, partial cds |
| MW620662         | *Oliveria decumbens* 1-deoxy-D-xylulose-5-phosphate synthase (DXS) mRNA, partial cds |
| MW620663         | *Oliveria decumbens* enolase 1 (ENO1) mRNA, partial cds |
| MW620664         | *Oliveria decumbens* fructose-bisphosphate aldolase 1 (FBA1) mRNA, partial cds |
| MW620665         | *Oliveria decumbens* alcohol dehydrogenase 1 (ADH1) mRNA, complete cds |
| MW620666         | *Oliveria decumbens* solaneryl diphosphate synthase 1 (SPS1) mRNA, complete cds |
| MW620667         | *Oliveria decumbens* 4-hydroxy-3-methylbut-2-en-1-yl diphosphate synthase (ISPG) mRNA, complete cds |
| MW620668         | *Oliveria decumbens* 3-hydroxy-3-methylglutarate-coenzyme A reductase (HMGR) mRNA, complete cds |
| MW620669         | *Oliveria decumbens* hydroxymethylglutaryl-CoA synthase (HMGCS1) mRNA, partial cds |
| MW620670         | *Oliveria decumbens* isopentenyl-diphosphate Delta-isomerase 1 (IPPI) mRNA, complete cds |
| MW620671         | *Oliveria decumbens* 4-hydroxy-3-methylbut-2-enyl diphosphate reductase (ispH) mRNA, partial cds |
| MW620672         | *Oliveria decumbens* phosphoglycerate kinase 3 (PGK3) mRNA, complete cds |
| MW620673         | *Oliveria decumbens* pyruvate kinase (At4g26390) mRNA, complete cds |
| MW620674         | *Oliveria decumbens* phosphomevalonate kinase (PMVK) mRNA, partial cds |
| MW620675         | *Oliveria decumbens* gamma-terpinene synthase mRNA, partial cds |

**FIGURE 4**

The co-expression network of upregulated genes in the flower of *Oliveria decumbens* Vent. The bigger size of the node is representative of higher betweenness centrality and the darker the red color is representative the higher the Closeness centrality and blueish color showing the lower closeness centrality in the network. Also, the thicker and more reddish connective line, represents the higher Edge Betweenness, the thinner and tending to light blue color showing the lower edge centrality.
active in the processes of root growth and development, responses to environmental stresses, and biosynthesis of compounds such as L-phenylalanine.

Validation

The qRT-PCR method was used to validate the expression of seven genes involved in the terpenoid synthesis in the *Oliveria decumbens*. As it is shown in Figure 6, the expression of all seven genes is highly consistent with the expression of the other genes and their expression from the RNA-Seq results. It is important to consider that the slight difference in obtained expression values from the qRT-PCR method with the RNA-Seq results is negligible due to different parameters such as the primers efficiency, quality of the PCR kit, and the accuracy of the thermocycler.

Conclusion

In this study, by producing more than 136 million reads of both samples of root and flower of the medicinal plant *Oliveria decumbens* Vent., transcriptomic differences between the tissue samples were investigated. The phytochemical analysis also confirmed the presence of the above gene products such as medicinally valuable compounds. The results showed that the MEP pathway is the most active in the flowers and the MVA pathway is the most active pathway in the root. These findings confirm that the synthesis of these compounds in the flowers of *Oliveria decumbens* has a great advantage over the root. The gene ontology analysis results showed that in general, the flower of the *Oliveria decumbens* is almost three times more active than the root tissue and has more active organs and more diverse biological pathways and molecular functions. Genes of the root are often active in the processes of root growth and development, response to environmental stresses, and the biosynthesis of compounds such as L-phenylalanine; while the genes of flower sample demonstrated that most of them are involved in the photosynthesis process, response to biological and abiotic stresses, specific genes for the growth, and development of flower organelles and synthesis of fatty acids.
Data availability statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

Author contributions

AK is performed conceptualization, data curation, formal analysis, project administration, software, and writing the manuscript; RS has supervised the entire project and validate the findings; MH and MAH reviewed the final manuscript. All authors approved the final version of the manuscript.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2022.916183/full#supplementary-material
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