



**Full Length Article**

## Identification of SSR Markers and Putative Genes Associated with Chlorogenic Acid in *Vaccinium uliginosum* through Transcriptome Analysis

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### Abstract

Bog bilberry (*Vaccinium uliginosum*) is a perennial woody deciduous member of the *Vaccinium* genus, subfamily Vacciniaceae, family Ericaceae. *V. uliginosum* has been domesticated as a promising fruit crop for significant healthful functions from the last few decades. In the present study, more than 2G clean data was obtained from the leaf transcriptome of *V. uliginosum*, which was sequenced by using second generation high-throughput technology, and assembled into 43,507 unigenes with an average length of 588 bp. Of these, 30,689 (70.538%) unigenes were annotated by at least one of Nr, Swiss-Port, KOG or KEGG databases, and 30,641, 21,518, 19,273 and 13,334 unigenes had a significant hit ( $p < 1E-5$ ) in these databases, respectively. In addition, 2,339 unigenes were identified as candidate resistance genes, which belong to 19 families. A total of 9,680 SSR loci from 7,780 unigenes were identified with AG/CT repeat motifs accounted for 72.19%. But most importantly, eight candidate unigenes that encoded five key enzymes and responsible for the bio-synthesis of chlorogenic acid were identified, which is an important bioactive component with high medicinal function. The molecular markers developed in this study would be helpful for further studies about the molecular genetics, molecular ecology and marker assisted breeding of *V. uliginosum*. The putative resistance genes and chlorogenic acid identified in this study would help to deepen the understanding on molecular mechanism of ecological adaptation and medicinal functions of *V. uliginosum*. © 2019 Friends Science Publishers

**Key words:** Blueberry; Bioinformatics; Functional annotation; Transcriptome; Unigenes

### Introduction

*Vaccinium uliginosum*, commonly called bog bilberry or blueberry, is a member of family Ericaceae. It is a deciduous shrub (Kähkönen *et al.*, 2001). Its typical habitat is open biotopes like wet heaths and bogs in circumboreal regions and it tends to grow on acidic, poorly drained and wet soils (Jacquemart, 1992). Compared to its two relatives, bilberry and blueberries, *V. uliginosum* is less utilized in commercial berry products, but recently great interest developed worldwide for their high anthocyanin content as well as flavonols, especially myricetin and quercetin (Latti *et al.*, 2010; Li *et al.*, 2011). The fruit of *V. uliginosum* has been used as edible berries, and added into juices, jams, pies, jelly, and wine. *V. uliginosum* has been used as folk medicinal materials for centuries in some circumboreal countries (Stefko *et al.*, 2014; Wang *et al.*, 2014).

Anthocyanin in *V. uliginosum* have many health benefits, such as anti-memory decline effects (Krikorian *et al.*, 2010), anti-light damage to the retina (Yin *et al.*, 2012) antioxidant activity (Kähkönen *et al.*, 2001) and anticancer, anti-vascular disease effects (Vendrame *et al.*, 2014). Li *et al.* (2011) reported these berries as a good source of food colorants. Leaves of *V. uliginosum* contain essential oils, hyperoside (also known as quercetin-3-O-galactoside) (Wang *et al.*, 2008; Chen *et al.*, 2015), and chlorogenic acid (also known as 3-O-caffeoylquinic acid, Chlorogenate, Caffeoyl quinic acid, trans-5-O-Caffeoyl-D-quinic acid), and these three bioactive compounds are very beneficial to the human health (Umezu, 2012; Zhang *et al.*, 2014). Among them, the biosynthetic pathways of chlorogenic acid have been revealed, and three metabolic pathways that could generate chlorogenic acid have been found (Campa *et al.*, 2003).

*V. uliginosum* has been cultivated for about two decades, and the studies on *V. uliginosum* have increased during that time, including ecology (Auffret *et al.*, 2010), population genetics (Albert *et al.*, 2005), botany (Rossum *et al.*, 2012; Stephens *et al.*, 2012), breeding (Lyrene and Olmstead, 2012), molecular biology (Primetta *et al.*, 2015), physiology (Boesgaard *et al.*, 2012), phyto-chemistry (Kellogg *et al.*, 2010; Su *et al.*, 2016), and pharmacology or pharmaceuticals (Han *et al.*, 2012; Kim and Choung, 2014; Park *et al.*, 2016; Yoon *et al.*, 2016). However, there is no genomic information on *V. uliginosum*, which could delay the study of molecular biology or genetics of *V. uliginosum*.

Transcriptome sequencing has been widely used for various crop plants to detect molecular markers at large scale. Molecular markers are essential tools of modern agriculture to conduct studies on various aspects of plants (Varshney *et al.*, 2005; Zheng *et al.*, 2017; Nadeem *et al.*, 2018). This is the first report on the leaf transcriptome of *V. uliginosum* by next generation high-throughput sequencing technology. Our analysis result would not only provide a lot of information about functional genes to deepen the understanding about the medicinal functions of *V. uliginosum*, the SSR marker we identified from transcriptome data would also offer more molecular tools for further studies on the molecular genetics, molecular ecology and marker assisted breeding of *V. uliginosum*.

## Materials and Methods

### Plant Material and Total RNA Extraction

A fully grown plant of *V. uliginosum* (two years old with about 0.8 m high), collected from Greater Khingan Range and planted at the research center of blueberry engineering technology, Majiang county, Guizhou province, China. Fresh leaves of *V. uliginosum* were used for total RNA isolation from by classic TRIzol method. RNA was quantified by using Bio-analyzer 2100 (Agilent Technologies, Santa Clara C.A.), and quality was checked on agarose gel electrophoresis.

### Construction of cDNA Library and Sequencing

To construct the cDNA sequencing library, the following protocol was used. Sera-mag Magnetic Oligo (dT) Beads (Illumina) were used to purify the poly (A) RNA from 20 mg total RNA. Purified mRNA was treated with fragmentation buffer. Random hexamer primers were used for cDNA synthesis. cDNA was synthesized by using QIA rapid PCR kit (QIAGEN). The cDNA was amplified and sent for sequencing to Illumina sequencing platform (Illumina HiSeq 2000, Guangzhou, China). Average length of sequencing read was 2\*125 bp. Whole set of raw sequencing reads was deposited into the Short Read Archive (SRA) database of the National Center for Biotechnology Information (NCBI) [accession number SRP157131].

### Data Filtering and *De Novo* Assembly

Low-quality reads (reads with Q-value  $\leq 20$  containing more than 50% bases) were discarded and adaptors were also trimmed. De-novo assembly of remaining high-quality sequences into unigenes was done by using the short reads assembling program Trinity (https://github.com/trinityrnaseq) with the default parameters.

### Functional Annotation

Assembled unigenes were searched for functional annotation against the non-redundant protein database (Nr, <http://www.ncbi.nlm.nih.gov/>) by using BLASTx and the Swiss-Port database (<http://www.expasy.ch/sprot>), with a cut off E-value of  $10^{-5}$ . Gene ontology (GO) enrichment analysis by Blast2GO (Conesa *et al.*, 2005) used to classify the unigenes into functional categories and WEGO program (Ye *et al.*, 2006) was used to generate GO trees. Then the unigenes were queried against the STRING database v. 9.05 (<http://string-db.org>) for forecasting about clusters of orthologous groups (COGs) and possible functions using BLASTx with a cut off E-value of  $10^{-10}$ . Finally, complex biological functions and the metabolic pathways associated with the unigenes were searched against Kyoto Encyclopedia of Genes and Genomes database (KEGG, <http://www.genome.jp/kegg>) (Kanehisa and Goto, 2000)

### Detection of SSR Markers and Primer Design

SSR markers were identify by using MISA program (<http://pgrc.ipk-gatersleben.de/misa/>) among all assembled genes with the following parameters: motifs ranging from the minimum number of repeat units (6 for di-, 5 for tri-, and 4 for tetra-, penta- and hexa-nucleotides), and the maximum interruption length between two SSRs of 100 bases. Primer 3 ([http:// primer3.ut.ee/](http://primer3.ut.ee/)) were used to design SSR primers. GC contents ranged from 40% to 60%, and the expected PCR product sizes ranging from 100 to 280 bp.

### Detection of Candidate Genes Encoding Enzymes Involved in Biosynthesis of Chlorogenic Acid

Physiological pathways for chlorogenic acid was accessed at KEGG database which was specifically annotated to *V. uliginosum* transcriptome. Biosynthetic route map concluded by Campa *et al.* (2003) and Lepelley *et al.* (2007) were followed for the detection of homologous unigenes. Corresponding unigenes were confirmed by blast search at NCBI database.

## Results

### Sequencing and *De Novo* Assembly

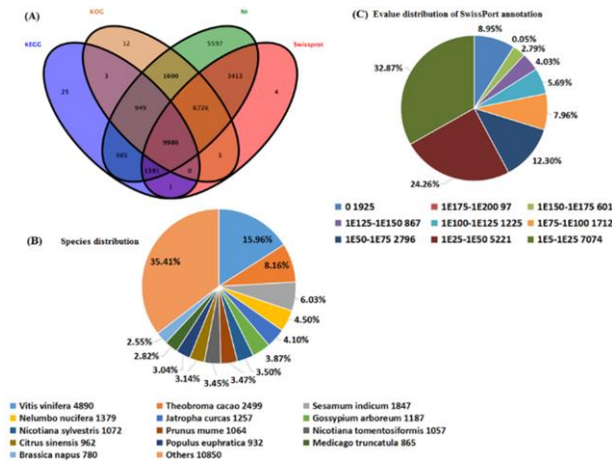
Illumina HiSeq produced 20,356,360 raw reads, from

**Table 1:** Summary of RNA-sequencing of *V. uliginosum*

	Total reads	Total reads nt	Max/min length	Mean length	Q20 (percentage)	Q30 (percentage)	GC (percentage)
Raw data	20356908	2544613500	125/125	125	2440185074 (95.90%)	2349684086 (92.34%)	1188117625 (46.69%)
Clean data	20214388	2508184666	125/50	123	2428839064 (96.84%)	2341476440 (93.35%)	1170793118 (46.68%)

**Table 2:** Assembled results of the leaf transcriptome of *V. uliginosum*

Total number (>200 bp)	Total number (>500 bp)	Total number (>1kp)	Total number (>2kp)	Total number (3kp)	GC ratio	N50 number	Total size	Mean length
43507	16406	6657	1153	206	44.2273	9198	25615894	588



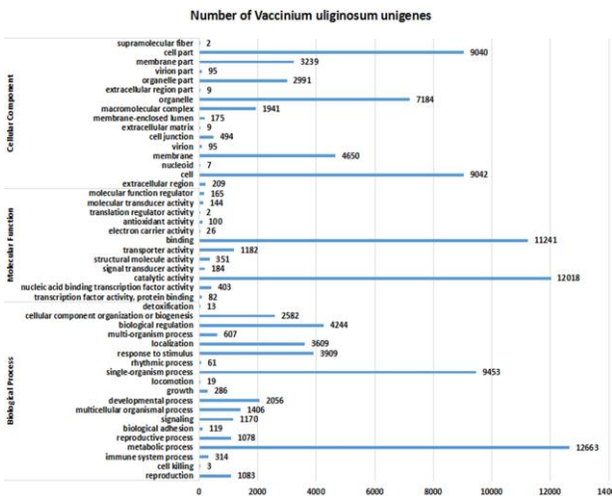
**Fig. 1 (A):** Venn diagram of BLAST hits for unigenes against KOG, Nr, Swiss-Port and KEGG database (E-value $\leq$ 1.0e-05). Numbers in the circles indicate the number of unigenes annotated by single or multiple databases, **(B)** Species distribution of the top BLAST hits for the assembled unigenes (E-value of 1.0e-05). **(C)** E-value distribution of BLAST hits Swiss-Port dataset for each unique sequence (E-value  $\leq$ 1.0e-05)

adaptors, accounted for 0.70% of total raw data (Table 1). Average length of 123 bp was found in clean reads (total nucleotides: 2,508,184,666 bp), 96.84% bases reached to the Q20 quality standard and 93.35% bases reached to the Q30 quality standard. The GC content of total clean data was 46.68% (Table 1).

High quality sequences were assembled into 43507 unigenes with a length more than 200bp. Among them, 27,101 (62.29%) were 200–500 bp in length, 16,406 (37.71%) unigenes were more than 500 bp, 6,657 (15.30%) unigenes were more than 1000 bp while, 1153 (2.65%) unigenes were more than 2000 bp and only 206 (0.47%) unigenes were more than 3000 bp long. The size of N50 was 807 bp and its number was 9189. The total nucleotides of whole assembled unigenes were 25,615,894, and the average length of unigenes was 588 bp (Table 2).

### Functional Annotation

Among 43,507 assembled unigenes, there were 30,641 (70.43%) unigenes which were showed significant similarity to the known protein in Nr data base, while and 21,518 (49.46%) unigenes showed significant similarity in Swiss-Port datasets. Furthermore, 19,273 (44.30%) and 13,334 (30.65%) unigenes could be annotated by the KOG and KEGG databases for *V. uliginosum*, respectively (Fig. 1A). A total of 30,689 unigenes accounted for 70.54% of the total unigenes that were annotated by at least one of the above datasets and the remaining 12,818 accounted for 29.46% were not annotated to above datasets (Fig. 1A). The number of known proteins showed significant similarities with *V. uliginosum* unigenes, and the 10 top-hit species with unigenes hit were *Vitis vinifera* (4890, 15.96%), *Theobroma cacao* (2499, 8.16%), *Sesamum indicum* (1847, 6.03%), *Nelumbo nucifera* (1379, 4.50%), *Jatropha curcas* (1257, 4.10%), *Gossypium arboreum* (1187, 3.87%), *Nicotiana sylvestris* (1072, 3.50%), *Prunus mume* (1064, 3.47%), *N. tomentosiformis* (1057, 3.45%) and *Citrus sinensis* (962, 3.14%) (Fig. 1B). 42.87% annotated sequences with high identity with their best hits (smaller than 1e-50) were found in the Swiss-Port database when compared in the E-value distribution, whereas 24.26% ranged from 1e-25 to 1e-50 and another 32.87% ranged from 1e-25 to 1e-5 (Fig. 1C).



**Fig. 2:** GO functional classification of *V. uliginosum* Unigenes

which 20,214,388 high quality sequences were obtained after filtration of low quality 141,972 reads with

GO classification assigned 109,755 GO terms to 21,782 unigenes with BLAST hits to annotated proteins in the Nr database (Fig. 2). These GO terms were classified into three main GO categories and 47 sub-categories by WEGO. 19 sub-categories belong to biological process, 12 sub-categories belong to molecular function and 16 sub-categories belong to cellular component, indicating a diverse range of functional genes in the *V. uliginosum*. Among sub-categories belonging to biological process, the unigenes were more frequent in metabolic process and single-organism process and the numbers of corresponding GO terms were 12,663 and 9,453. Among the sub-categories belonging to molecular function, the unigenes were most frequent in the catalytic activity and binding categories, and the number of corresponding GO terms reached to 12,018 and 11,241, respectively. Cell part and cell were more frequent sub-categories in the cellular component category, and the numbers of GO terms were 9,042 and 9,040, respectively (Fig. 2).

Subsequently, 19,273 unigenes were aligned to the appropriate COG clusters and 31,461 functional annotations were obtained and grouped into 25 functional categories, while some unigenes exhibited multiple COG functional annotations. Among these categories, the number of unigenes involving general functional category were the most frequent, followed by posttranslational modification, protein turnover, chaperones and signal transduction mechanisms (Fig. 3).

Finally, to represent active biological pathways in *V. uliginosum*, 30,641 annotated unigenes were aligned to the reference plant pathways in the KEGG database. Among them, 13,334 unigenes were assigned to 129 KEGG pathways, illustrating the broad overview of the groups of genes located in transcriptome. The top five pathways with most unigenes were “Carbon metabolism” (427 members), “RNA transport” (364 members), “Spliceosome” (361 member), “Plant-pathogen interaction” (339 members) and “Biosynthesis of amino acids” (572 members). All KEGG annotation information of unigenes is presented in Table 3.

Resistance gene (R-gene) is the key molecular mechanism that plants adapt to cope with the environment. In the current study, we identified 2,339 unigenes as candidate R-genes belonging to 19 gene families. Among them, 731 unigenes belong to the RLP family, which accounted for 21.56% of total candidate R-gene, followed by N, NL and TNL families, which accounted for 15.86, 13.25 and 9.53% respectively, and the rest 15 families accounted for 30.10% of total candidate R-gene unigenes (Table 4).

### EST-SSR Kinds and Number

SSRs are one of the most popular molecular markers, which widely used in the life science studies such as molecular

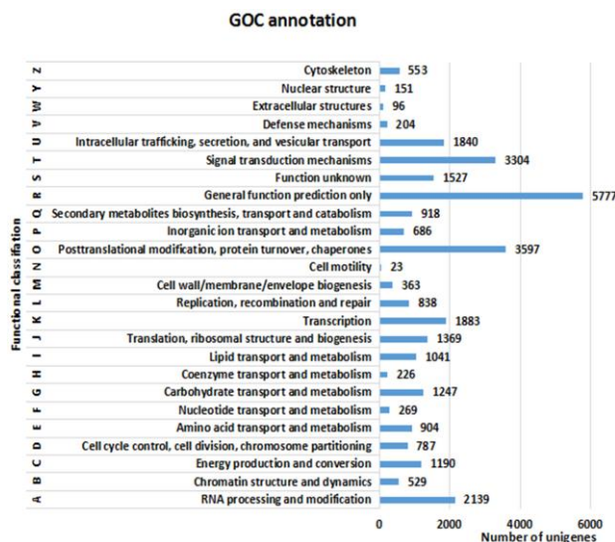


Fig. 3: KOG functional classification of *V. uliginosum* Unigenes

genetics, molecular ecology, association study and marker assisted breeding. From the *V. uliginosum* leaf transcriptome, we identified 9680 SSR sequence segments from 7780 unigenes. Of these SSR, two, three, four, five and six base repeat types of SSRs were 7423, 1907, 147, 76 and 127 that accounted for 76.68%, 19.70%, 1.52%, 0.08% and 1.31%, respectively. The highest repeat number of two base type was 6, three base type was 5 and four, five and six base types were four (Table 5). Among all detected SSRs, the AG/CT motif kind exhibited the highest frequency, which accounted for 72.19%, followed by AAG/CTT, AC/GT, and ACC/GGT motif kind that accounted for 6.42, 3.36 and 3.33%, respectively (Table 6).

### The Candidate Gene Involving in Chlorogenic Acid Biosynthesis

KEGG database (<http://www.kegg.jp/kegg/kegg1.html>) was explored by query ‘chlorogenic acid’ under the retrieve item ‘DBGET’, and three ‘KEGG compound’ encoded as C00852, C10468 and C17417 related to the chlorogenic acid were found. C00852 was found to be involved in three pathways including ‘Phenylpropanoid biosynthesis’ (map00940), ‘Flavonoid biosynthesis’ (map00941), ‘Stilbenoid, diarylheptanoid and gingerol biosynthesis’ (map00945) that associated with the biosynthesis of chlorogenic acid. Homologous unigenes were detected after annotation which encode five key enzymes for bio-synthesis of chlorogenic acid, including Unigene0000447, Unigene0000448 and Unigene0000449 encoded phenylalanine ammonia-lyase (PAL), Unigene0008683 encoded 4-coumarate-CoA ligase (4CL), Unigene0014113 and Unigene0014114 encoded trans-cinnamate 4-monooxygenase (CYP73A), Unigene0015838 encoded shikimate O-hydroxycinnamoyltransferase (HCT) and Unigene0035395 encoded coumaroylquininate

**Table 3:** KEGG functional classification of *Vaccinium uliginosum* Unigenes

KEGG A class	KEGG B class	Pathway	Count (7036)
Cellular Processes	Transport and catabolism	Endocytosis (270), Peroxisome (132), Phagosome (126), Regulation of autophagy (74)	
Environmental Information Processing	Membrane transport	ABC transporters (84)	
Environmental Information Processing	Signal transduction	Plant hormone signal transduction (265), Phosphatidylinositol signaling system (102)	
Genetic Information Processing	Folding, sorting and degradation	Protein processing in endoplasmic reticulum (346), Proteasome (81), Protein export (78), RNA degradation (232), SNARE interactions in vesicular transport (45), Sulfur relay system (21), Ubiquitin mediated proteolysis (259)	346
	Replication and repair	Base excision repair (59), DNA replication (64), Homologous recombination (71), Mismatch repair (70), Non-homologous end-joining (15), Nucleotide excision repair (123).	
	Transcription	Basal transcription factors (84), RNA polymerase (89), Spliceosome (361), Aminoacyl-tRNA biosynthesis (108), mRNA surveillance pathway (266), Ribosome (293), Ribosome biogenesis in eukaryotes (221), RNA transport (364)	
Metabolism	Amino acid metabolism	Alanine, aspartate and glutamate metabolism (82), Arginine and proline metabolism (75), Arginine biosynthesis (41), Cysteine and methionine metabolism (125), Glycine, serine and threonine metabolism (124), Histidine metabolism (25), Lysine biosynthesis (22), Lysine degradation (62), Phenylalanine metabolism (68), Phenylalanine, tyrosine and tryptophan biosynthesis (63), Tryptophan metabolism (36), Tyrosine metabolism (58), Valine, leucine and isoleucine biosynthesis (28), Valine, leucine and isoleucine degradation (79)	
	Biosynthesis of other secondary metabolites	Anthocyanin biosynthesis (26), Betalain biosynthesis (1), Caffeine metabolism (6), Flavone and flavonol biosynthesis (9), Glucosinolate biosynthesis (2), Flavonoid biosynthesis (39), Isoquinoline alkaloid biosynthesis (40), Isoflavonoid biosynthesis (6), Monobactam biosynthesis (18), Phenylpropanoid biosynthesis (135), Stilbenoid, diarylheptanoid and gingerol biosynthesis (24), Tropane, piperidine and pyridine alkaloid biosynthesis (35)	
	Carbohydrate metabolism	Amino sugar and nucleotide sugar metabolism (188), Ascorbate and aldarate metabolism (66), Butanoate metabolism (27), Citrate cycle (TCA cycle) (92), C5-Branched dibasic acid metabolism (13), Starch and sucrose metabolism (257), Galactose metabolism (110), Fructose and mannose metabolism (89), Glycolysis / Gluconeogenesis (193), Glyoxylate and dicarboxylate metabolism (102), Inositol phosphate metabolism (112), Pentose and glucuronate interconversions (57), Pentose phosphate pathway (74), Propanoate metabolism (66), Pyruvate metabolism (144)	
	Energy metabolism	Carbon fixation in photosynthetic organisms (125), Nitrogen metabolism (36), Oxidative phosphorylation (207), Photosynthesis (76), Photosynthesis - antenna proteins (22), Sulfur metabolism (52)	
	Global and Overview	Biosynthesis of amino acids (334), Carbon metabolism (427), Degradation of aromatic compounds (10), Fatty acid metabolism (122), 2-Oxocarboxylic acid metabolism (68)	
	Glycan biosynthesis and metabolism	Glycosylphosphatidylinositol(GPI)-anchor biosynthesis (38), Glycosaminoglycan degradation (35), Glycosphingolipid biosynthesis - ganglio series (16), Glycosphingolipid biosynthesis - globo series (17), N-Glycan biosynthesis (86), Other types of O-glycan biosynthesis (9), Other glycan degradation (51)	
	Lipid metabolism	alpha-Linolenic acid metabolism (63), Arachidonic acid metabolism (26), Biosynthesis of unsaturated fatty acids (48), Cutin, suberine and wax biosynthesis (13), Ether lipid metabolism (45), Fatty acid biosynthesis (70), Fatty acid elongation (41), Fatty acid degradation (71), Glycerolipid metabolism (100), Glycerophospholipid metabolism (135), Linoleic acid metabolism (28), Sphingolipid metabolism (59), Steroid biosynthesis (39), Synthesis and degradation of ketone bodies (8)	
	Metabolism of cofactors and vitamins	Biotin metabolism (26), Folate biosynthesis (26), Lipoic acid metabolism (6), Nicotinate and nicotinamide metabolism (42), One carbon pool by folate (35), Pantothenate and CoA biosynthesis (33), Porphyrin and chlorophyll metabolism (72), Thiamine metabolism (28), Riboflavin metabolism (16), Ubiquinone and other terpenoid-quinone biosynthesis (66), Vitamin B6 metabolism (26)	
	Metabolism of other amino acids	beta-Alanine metabolism (75), Glutathione metabolism (102), Cyanoamino acid metabolism (54), Selenocompound metabolism (25), Taurine and hypotaurine metabolism (16),	
	Metabolism of terpenoids and polyketides	Brassinosteroid biosynthesis (25), Carotenoid biosynthesis (39), Diterpenoid biosynthesis (26), Limonene and pinene degradation (13), Monoterpenoid biosynthesis (18), Sesquiterpenoid and triterpenoid biosynthesis (21), Terpenoid backbone biosynthesis (84), Zeatin biosynthesis (21)	
	Nucleotide metabolism	Purine metabolism (259), Pyrimidine metabolism (192)	
Organismal Systems	Environmental adaptation	Circadian rhythm - plant (77), Plant-pathogen interaction (339)	

(coumaroylshikimate) 3'-monoxygenase (CYP98A, C3'H), and codes of these enzymes in KEGG dataset were 4.3.1.24, 6.2.1.12, 1.14.13.11, 2.3.1.133 and 1.14.13.36, respectively (Table 7)

## Discussion

*Vaccinium* L. (Ericaceae) comprising of approximately 450 species is a morphologically diverse genus of terrestrial or

epiphytic shrubs and lianas. Primarily these species occur in the cooler areas of northern hemisphere. *Vaccinium* species have been highly concerned in the recent years because the genus contains three kinds of new fruit with excellent health care values *viz.*, blueberry, cranberry and lingonberry. Among three kinds of domesticated *Vaccinium* fruits, the blueberry genomics research has been widely conducted, and these studies have greatly enhanced the blueberry

**Table 4:** Candidate R-gene families and corresponding number of unigenes in *V. uliginosum*

Class	Count	Class	Count	Class	Count	Class	Count	Class	Count
RLP	731	N	371	NL	310	TNL	223	CNL	189
RLK	188	RLK-GNK2	104	CN	63	Other	43	T	41
Mlo-like	26	Pto-like	24	L	10	RPW8-NL	6	RLP-Malectin	5
RLK-Malectina	2	RLP-Malectina	1	RLK-Pto-like	1	RLK-Malectin	1	Total	2339

**Table 5:** SSR types identified from the *V. uliginosum* leaf transcriptome

Number of repeat unit	Di-	Tri-	Tetra-	Penta-	Hexa-	Total
4	0	0	100	60	79	239
5	0	1044	34	13	29	1120
6	1238	467	12	2	4	1723
7	1033	215	1	0	8	1257
8	1030	43	0	1	7	1081
9	993	28	0	0	0	1021
10	799	34	0	0	0	833
11	361	9	0	0	0	370
12	69	10	0	0	0	79
13	17	20	0	0	0	37
14	104	11	0	0	0	115
>=15	1779	26	0	0	0	1805
Total	7423	1907	147	76	127	9680

**Table 6:** Motif kinds of SSR and its count in *V. uliginosum* leaf transcriptome

Motif kinds	Count	Frequency	Motif kinds	Count	Frequency
AC/GT	325	0.0335743801652893	ACT/AGT	18	0.0018595041322314
AG/CT	6988	0.721900826446281	AGC/CTG	203	0.0209710743801653
AT/AT	98	0.0101239669421488	AGG/CCT	339	0.0350206611570248
AAC/GTT	56	0.00578512396694215	ATC/ATG	133	0.0137396694214876
AAG/CTT	621	0.0641528925619835	CCG/CGG	104	0.0107438016528926
AAT/ATT	40	0.00413223140495868	AAAG/CTTT	28	0.00289256198347107
ACC/GGT	322	0.0332644628099174	AAAT/ATTT	20	0.00206611570247934
ACG/CGT	71	0.00733471074380165	others	314	0.0324380165289256

**Table 7:** The key enzymes responsible for the bio-synthesis of chlorogenic acid and corresponding unigenes identified in *V. uliginosum*

Code in KEGG dataset	Name	Description	ORTHOLOG	Corresponding unigenes code
4.3.1.24	PAL	phenylalanine ammonia-lyase	K10775	Unigene0000447 Unigene0000448 Unigene0000449
6.2.1.12	4CL	4-coumarate--CoA ligase	K01904	Unigene0008683
1.14.13.11	CYP73A	trans-cinnamate 4-monooxygenase	K00487	Unigene0014113 Unigene0014114
2.3.1.133	HCT	shikimate O-hydroxycinnamoyltransferase	K13065	Unigene0015838
1.14.13.36	CYP98A, C3'H	Coumaroylquininate (coumaroylshikimate) monooxygenase	3'- K09754	Unigene0035395

growth, physiological active components metabolism, molecular breeding of blueberry, biological engineering technology and understanding the molecular mechanism of cold tolerance (Rowland *et al.*, 2012). Genome draft of high bush blueberry has also been published as integrated genome browser (Gupta *et al.*, 2015), and genome database of American cranberry (*V. macrocarpon*) also available (Polashock *et al.*, 2014). However, there is no report about the genome and transcriptome analysis of *V. uliginosum*, and this is the first corresponding report until now.

Because its unique geography distribution, *V. uliginosum* is often used as a representative plant to study the ecology, physiology and evolution of polar/Arctic plants (Alsos *et al.*, 2003; Albert *et al.*,

2005; Boesgaard *et al.*, 2012; Faubert *et al.*, 2012; Preece and Phoenix, 2014). During the last few decades, molecular markers, revealing polymorphism at the DNA level, have played an important role in studies of plant biotechnology and their genetics and ecology (Kumar *et al.*, 2009; Nadeem *et al.*, 2018). In the recent years, some SSR markers have been developed using pyrosequencing method for *V. uliginosum* (Mayer *et al.*, 2014); however, the number was limited. In this study, we exploited huge numbers of EST-SSRs markers from whole transcriptome data. The molecular markers play indispensable roles in many research areas such as molecular breeding, conservation genetics, molecular ecology and association studies (Li *et al.*, 2016; Nadeem *et al.*, 2018; Wang *et al.*, 2019).

Therefore, the results of this study would greatly help for above mentioned studies about *V. uliginosum*.

R-genes evolved from a phylogenetically ancient form of immunity, which is common in animals and plants (Chisholm *et al.*, 2006). It is a major protection way to defend themselves against the attack of pathogens based on the mechanism of gene-for-gene interactions between plants and their infectious agents (Flor, 1971). The R-gene identification had attracted the attention of scientists for its potential value for improving crop adaptability (Harris *et al.*, 2013; Dong *et al.*, 2019). In this study, we detected 2339 unigenes related to 19 families as the candidate of resistance gene, and this would greatly deepen our understanding about the molecular mechanism of resistance in *V. uliginosum*. Moreover, corresponding unigenes are potential molecular tool for the resistance breeding and genetic engineering of *V. uliginosum* and its relative species.

Not only the berries of *V. uliginosum* are rich in antioxidants, especially anthocyanins and other polyphenolic compounds. They are medicinally important for cardiovascular diseases, obesity, aging diseases, urinary tract infections and periodontal disease, diabetes and in reduction of the risk of cancers or assist in treatment of cancers (Su, 2012). Its leaves are also rich in bioactive ingredients such as chlorogenic acid (3-O-caffeoylquinic acid) (Chen *et al.*, 2015). Chlorogenic acid (CGA) is the ester of caffeic acid and quinic acid, functioning as an intermediate in lignin biosynthesis (Boerjan *et al.*, 2003). Three pathways of CGA bio-synthesis have been exploited, from the primary substrate 'Phenylalanine' to primary production 'Caffeoyl quinic acid' CGA, and there are few key enzymes including PAL, C4H, 4CL, C3'H, CQT, COMT and CCoACMT (Campa *et al.*, 2003). From leave transcriptome data of *V. uliginosum*, we have identified homologous genes which encoded key enzymes such as PAL, C4H, 4CL, C3'H and CYP73A that responsible for the biosynthesis of CA. Moreover, we also have detected some unigenes that were involved into other biosynthesis pathways of antioxidation components of other secondary metabolites, such as Anthocyanins and flavonoids (Table 3). These results are important to deepen our knowledge about the molecular mechanism of healthful functions hidden in *V. uliginosum* leaves.

## Conclusion

In this study, about 2G transcriptome data of *V. uliginosum* was obtained and analyzed for unigenes assemblage and functional annotation. To date this is the first functional information reported on *V. uliginosum*. The transcriptome data assembled into 43,507 unigenes with an average length of 588 bp of these, 30,689 (70.538%) unigenes were annotated by Nr, Swiss-Port, KOG or KEGG databases, and 30,641, 21,518, 19,273 and 13,334 unigenes had a significant hit ( $p < 1E-5$ ) in above four databases, respectively. 2339 unigenes were identified as candidate

resistance genes, which belong to 19 families. A total of 7780 unigenes was identified as 9680 SSR loci, and AG/CT repeat motifs was the most common SSR. Overall, eight candidate unigenes encoded five key enzymes that were responsible for the bio-synthesis metabolic pathway of chlorogenic acid, which is one of the important bioactive components with high medicinal function. This transcriptome analysis is a significant contribution for molecular marker development, functional gene discovery and studies of differential gene expressions in *V. uliginosum*.

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