Identification of *Viscum album* L. miRNAs and prediction of their medicinal values

Wenyan Xie¹*, Jacob Adolf², Matthias F. Melzig¹

¹ Institut für Pharmazie, Freie Universität Berlin, Berlin, Germany, ² Technische Hochschule Wildau, Wildau, Germany

*wenyan.xie@fu-berlin.de

Abstract

MicroRNAs (miRNAs) are a class of approximately 22 nucleotides single-stranded non-coding RNA molecules that play crucial roles in gene expression. It has been reported that the plant miRNAs might enter mammalian bloodstream and have a functional role in human metabolism, indicating that miRNAs might be one of the hidden bioactive ingredients in medicinal plants. *Viscum album* L. (Loranthaceae, European mistletoe) has been widely used for the treatment of cancer and cardiovascular diseases, but its functional compounds have not been well characterized. We considered that miRNAs might be involved in the pharmacological activities of *V. album*. High-throughput Illumina sequencing was performed to identify the novel and conserved miRNAs of *V. album*. The putative human targets were predicted. In total, 699 conserved miRNAs and 1373 novel miRNAs have been identified from *V. album*. Based on the combined use of TargetScan, miRanda, PITA, and RNAhybrid methods, the intersection of 30697 potential human genes have been predicted as putative targets of 29 novel miRNAs, while 14559 putative targets were highly enriched in 33 KEGG pathways. Interestingly, these highly enriched KEGG pathways were associated with some human diseases, especially cancer, cardiovascular diseases and neurological disorders, which might explain the clinical use as well as folk medicine use of mistletoe. However, further experimental validation is necessary to confirm these human targets of mistletoe miRNAs. Additionally, target genes involved in bioactive components synthesis in *V. album* were predicted as well. A total of 68 miRNAs were predicted to be involved in terpenoid biosynthesis, while two miRNAs including val-miR152 and miR9738 were predicted to target viscotoxins and lectins, respectively, which increased the knowledge regarding miRNA-based regulation of terpenoid biosynthesis, lectin and viscotoxin expressions in *V. album*.

Introduction

MicroRNAs (miRNAs) are a class of single-stranded non-coding RNA molecules of approximately 22 nucleotides that play crucial roles in gene expression [1]. They generally bind to complimentary sequences in the 3’ untranslated region (UTR) of specific protein-coding genes, inducing mRNA cleavage or translational repression [2]. MiRNAs are highly pleiotropic
and a single miRNA can recognize hundreds of mRNA transcripts, allowing them to regulate a diverse range of biological pathways [3,4]. In 2012, Zhang et al. reported miRNAs derived from plant-based dietary can function as active signalling molecules to regulate mammalian genes [5]. A recent study demonstrated that a medicinal plant-derived miRNA, MIR2911, can be acquired by mice via GI tract and target influenza A virus and protect the mice against influenza virus infections [6]. These findings provide thrilling clues that miRNAs might act as bioactive constituent mediating the cross-kingdom regulation [7].

Viscum album L. (Loranthaceae) commonly known as mistletoe or European mistletoe, is a hemi-parasitic evergreen shrub that grows on a number of host trees including apple, oak, poplar and other trees [8]. Mistletoe has been used medicinally in Europe for centuries. In ancient Greece, Hippocrates (460–377 BC) used the mistletoe to treat disorders of the spleen and complaints associated with menstruation. Around 150 AD, the Roman naturalist Celsus prescribed mistletoe to treat abnormal growths (including possible swellings and cancer). During the middle ages, mistletoe was considered as a golden herb for the treatment of epilepsy. In the 16th century, mistletoe was applied for many conditions including epilepsy, diseases of kidneys and spleen, ulcers, bone fractures and labour-pain. According to the homeopathic Materia Medica, mistletoe was applied for “weakness of the heart” and oedema in the 18th century. However, in the late 19th century when the modern medicine rose, these medicinal applications of mistletoe did not gain considerable attention. Until 1907, Gaultier scientifically proved the anti-hypertensive effect of mistletoe extract. In the 1920s, mistletoe was recommended as a possible treatment for cancer. Thereafter, the medicinal use of mistletoe has awakened [9].

Nowadays, V. album extracts are most frequently used in adjuvant cancer therapy in German-speaking countries [10]. Preparations from V. album extracts for this purpose are commercially available in Europe, such as Iscador®, Eurixor®, Helixor® and Abnoba viscum®. Three components of mistletoe, namely viscotoxins, lectins and terpenoids, which showed significant immune-system-stimulating activity and cell-killing activity, were suggested to be responsible for its anti-cancer effect [11–13]. In folk medicine, V. album has been mainly practiced for the treatment of cardiovascular diseases such as hypertension and diabetes [14,15], but its clinical efficacy has not been established [16]. Studies have shown that V. album extracts possess potent cardioprotective, hypoglycemic, anti-hypertensive and vasodilator effects both in vitro and in vivo [17–22], nitric oxide pathway, calcium signaling pathway and cholinergic pathway might be involved [18,20,23]. Although various secondary metabolites such as flavonoids, saponins, tannins, alkaloids, phenylpropanoids are present in V. album [15,20], the bioactive constituents that might be responsible for its cardiovascular protective effects remain to be elucidated [19,24].

In this study, we considered that miRNAs might be involved in the pharmacological activities of V. album. The conserved and novel miRNAs from V. album have been identified using Illumina platform technologies. The putative human targets have been predicted using bioinformatics tools, and their potential roles in human biological pathways and diseases have been elucidated. The results indicated that mistletoe miRNAs might possess beneficial effects against some human diseases such as cancer, cardiovascular diseases and neurological disorders, which might explain the medicinal use of mistletoe in ancient time, and provide scientific support for folk medicinal use and clinical use of mistletoe in modern medicine.

Furthermore, to promote understanding of miRNA-based regulation of bioactive ingredients in V. album, the genes involved in terpenoids, lectins and viscotoxins biosynthesis in V. album have been characterized, and their corresponding regulatory miRNAs have been predicted, which might facilitate bioengineering research in the production of mistletoe pharmacologically active components.
Materials and methods

Ethics statement

The mistletoe plants were collected from Niefern-Öschelbrunn (Baden-Wuerttemberg) Germany with permission of Birken AG. We confirm that the experiments in this study did not involve any endangered or protected species.

Plant materials

One-year-old leaves and stems were randomly collected from different individual *V. album* grown on *Malus domestica* L. trees in Niefern-Öschelbrunn (Baden-Wuerttemberg) Germany in November 2015. The mistletoe plants were snapped in liquid nitrogen and stored at -80˚C until use.

RNA isolation, library construction and high-throughput sequencing

Total RNA was extracted from plant leaves and stems using RNA isolation reagent (amsbio, USA) according to the manufacturer’s protocol. The RNA samples with high purity (OD260/280 between 1.8 and 2.2) and high integrity (RNA integrity number of 6.5 or higher) were used to construct the sRNA library. The mRNA and small RNA library preparations and sequencing were performed by BGI (Beijing Genomics Institute, Shenzhen, China). For mRNA library construction, mRNA in the sample was enriched and fragmented. The RNA fragments were served as templates for cDNA synthesis. The cDNA fragments were ligated with sequencing adapters, and amplified by PCR to construct the cDNA library for paired-end sequencing. Small RNAs (18 to 30 nt) were gel purified and ligated to the 3’ and 5’ adaptor. The ligated products were used for cDNA synthesis, followed by acrylamide gel purification and PCR amplification to generate small RNA library. The Agilent 2100 Bioanalyzer (Agilent, USA) was used for quantification and qualification of the sample library. Finally, the library was sequenced using Illumina HiSeq 4000 sequencing platform (Illumina Inc., San Diego, CA, USA).

Sequence data analysis

The raw reads obtained from Illumina sequencing were processed by trimming low-quality reads, reads with 5’ adapter contaminants, reads without 3’ adapters, reads without an insert fragment, reads containing poly A, and reads shorter than 18 nt. Other RNAs (rRNA, tRNA, snRNA and snoRNA) were removed by blasting against the GenBank database (http://blast.ncbi.nlm.nih.gov) and the Rfram database (http://rfam.xfam.org/). The remaining clean reads were used to detect conserved and novel miRNAs.

The reads obtained by RNA-seq sequencing were filtered by adaptor sequences, duplication sequences, and low quantity reads. De novo transcriptome assembly was performed by Trinity [25]. The Trinity program first assembles reads of a certain length that overlap to form longer fragments without gaps called contigs. These contigs were further processed for sequence clusters using the sequence clustering software TGICL [26] to obtain unigenes that could no longer be extended on either end. The sequence dataset generated in this study is available at the sequence read archive (SRA) of National Center for Biotechnology Information (NCBI) under the accession numbers of SUB2752327 and SUB2754679.

Identification of the conserved and putative novel miRNAs

The clean data were used in a BLAST search against known plant miRNAs in the miRBase 21.0, and matched sequences were considered as conserved *V. album* miRNAs. The small RNAs that were unaligned to any databases were defined as unannotated sequences.
miRNAs of *Viscum album* L.

The novel miRNAs were identified by mapping unannotated sequences to the *V. album* transcriptome using Mireap software (http://sourceforge.net/projects/mireap/). The parameters setting for the identification of novel miRNA were: 1) the sequences used to predict novel miRNAs were from the unannotated sequences that were matched to the transcriptome of *V. album*; 2) The sequences and their structures satisfy the criteria of forming hairpin miRNAs, and that the mature miRNAs were present in one arm of the hairpin precursors; 3) Hairpin precursors did not contain large internal loops or bulges; 4) Secondary structures of hairpins had free energy of hybridization $\leq -18$kcal/mol; and 5) The number of mature miRNAs with predicted hairpins were $\geq 5$ in the alignment result [27]. The novel miRNAs were named using a “miR” prefix to denote miRNAs, a three-letter prefix to denote the species (e.g. “val” representing *V. album*) and a unique sequential number [1].

**Plant targets prediction for both conserved and novel miRNAs**

The *V. album* conserved and novel miRNA candidates were searched against the *V. album* transcriptome database using psRobot (http://omicslab.genetics.ac.cn/psRobot/) and TargetFinder (http://targetfinder.org/) with default parameters to identify potential miRNA target genes. The target candidates were searched against protein database Nr using BLASTX with E-values less than $e^{-5}$ to predict their possible functions. To classify the function distribution of these potential targets, Gene Ontology (GO) annotation and functional classification were conducted using Blast2GO and WEGO [28,29].

**Human target gene prediction for the novel miRNAs**

The novel miRNAs are unique to *V. album* and differ from those found in other plant species, and might be responsible for the unique medicinal value of *V. album*. These novel miRNAs were therefore used for human targets prediction. In addition, to minimize false positives, the novel miRNAs were further filtered with following conditions: (1) the maximal free energy allowed for the miRNA precursor was -30kcal/mol; (2) the length of precursors were no more than 200nt; (3) the reads for mature miRNAs were at least 20. The human mRNA sequence were download from the UCSC genome browser (http://hgdownload.soe.ucsc.edu/goldenPath/hg19/bigZips/refMrna.fa.gz). Four commonly used animal target prediction algorithms including TargetScan, miRanda, PITA, and RNAhybrid were employed to predict putative human genes, and only those identified by all four softwares were selected for further study [30–33]. The target genes were mapped to the KEGG database to identify significantly enriched metabolic pathways or signal transduction pathways in target genes compared with the whole genome background. A corrected p value $<0.05$ was set as the threshold.

**Real-time quantitative PCR**

Five conserved and five novel miRNAs were randomly selected and validated by stem-loop RT-PCR as previously described by Chen et al. [34]. The stem-loop primers for reverse transcription and primers for PCR were listed in S1 Table. First-strand cDNA synthesis was performed using TaqMan MicroRNA Reverse Transcription Kit (Thermo Scientific). The reaction was carried out at 16˚C for 30min, at 42˚C for 10min, followed by heat-inactivation at 85˚C for 5min. Quantitative real-time PCR was conducted using the PowerUp™ SYBR® Green Master Mix (Thermo Scientific) and PikoReal Real-Time PCR System (Thermo Scientific). The reactions were carried out under the following amplification conditions: activation at 50˚C for 2min, 95˚C for 2min, followed by 40 cycles of denaturation at 95˚C for 15s, annealing at 55˚C for 15s, and extension then at 72˚C for 30s. All reactions were performed in three independent
biological samples with three technical repeats. The melting curve was generated to test the specificity of PCR products and avoid the false-positive peaks. No template control and no reverse transcription control were included in all reactions.

**Results**

**Analysis of small RNA**

In total, 7,635,733 raw reads were initially obtained. After data processing, 7,355,262 clean reads (96.33% of all raw reads) were kept for subsequent analysis. As shown in Fig 1, the clean reads exhibited an uneven length distribution, with the majority (~85%) ranging from 19 to 25nt in length. The most abundant was the small RNAs of 24nt, followed by those of 22, 21 and 20nt. In addition, 342,263 (7.75%) unique small RNAs were mapped to the transcriptome data of *V. album*. After annotating and removing the non-coding RNAs, including rRNAs, tRNA, snRNAs and snoRNA, 33,369 reads remained for the identification of conserved miRNAs, and 4,306,925 unannotated reads were used for the prediction of novel miRNAs (Table 1).

![Length distribution of small RNAs from *V. album*](https://doi.org/10.1371/journal.pone.0187776.g001)

Table 1. Distribution of small RNAs among different categories of *V. album*.

| Category   | Unique small RNAs | Percent (%) | Total small RNAs | Percent (%) |
|------------|-------------------|-------------|------------------|-------------|
| total reads| 441,5441          | 100         | 7,355,2622       | 100         |
| matched reads* | 342,263      | 7.75        | 4,472,7990       | 60.81       |
| miRNA      | 33,369           | 0.76        | 424,5509         | 5.77        |
| rRNA       | 58,531           | 1.33        | 686,0606         | 9.33        |
| snRNA      | 2,030            | 0.05        | 92,161           | 0.13        |
| snoRNA     | 1,534            | 0.03        | 38,977           | 0.05        |
| tRNA       | 13,052           | 0.30        | 1,141,541        | 1.55        |
| unannotated| 4,306,925        | 97.54       | 6,117,3828       | 83.17       |

*The reads that matched to the *V. album* transcriptome.*

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Identification of conserved miRNAs in *V. album*

Evolutionarily conserved miRNAs are present in diverse plant species and play essential roles in plant development and adaptation to adverse environments. The conserved nature of plant miRNAs provides the possibility of finding homolog sequences of miRNAs in different plant species. In this study, 699 conserved miRNAs were identified in *V. album* with a total read number of 5511469, of which 44% were detected with more than 100 reads. The most abundant miRNAs were miR166a-3p with 687103 reads, and miR166 with 673128 reads, followed by miR9778 (614703 reads), miR4993 (539038 reads) and miR159a (486726 reads) (S2 Table). Some miRNAs were lowly expressed with abundance of less than ten reads such as miR8004, miR5799 and miR6296 (S2 Table).

Identification of putative novel miRNAs in *V. album*

To identify novel miRNA candidates in *V. album*, the unannotated small RNA sequences were matched against the assembled unigene sequences of *V. album*. A total of 1373 miRNAs with reads varied from 5 to 11875 were identified as novel miRNA candidates (S3 Table). The length of novel miRNAs ranged from 20 to 23nt, and the precursors ranged from 50 to 372bp in length, with an average of 178bp. The average minimum free energy (MFE) value obtained for these pre-miRNAs was -58.7kcal/mol, which is comparable with the MEF values of precursors for trifoliate orange (*Citrus trifoliata* L. Raf.) (-52.41kcal/mol) [35], Arabidopsis thaliana L. (-57kcal/mol) [36], and Ginkgo biloba var. epiphylla Mak (-46.0kcal/mol) [37]. The first nucleotide bias of these candidate miRNAs was common 5’ terminal uridine (U) nucleotide, which is a typical feature of miRNAs [36,38]. The most abundant novel miRNA candidate was val-miR218 with 11875 reads in *V. album*, followed by val-miR11 and val-miR1338. Only 4.9% of novel miRNAs were counted more than 20 reads. Although the expression levels of novel miRNA candidates were much lower than the conserved miRNAs, the species-specific functions they played should not be ignored.

Experimental validation of conserved and novel miRNAs in *V. album*

Stem-loop RT-qPCR was employed to validate the gene expression data from Illumina sequencing. As illustrated in Fig 2, miR166a-3p was the most abundant miRNA among tested miRNAs, followed by miR159a, miR6135c, val-miR218, miR4414-3p, miR831-5p, val-miR1017, val-miR832, val-miR633 and val-miR1087, respectively. The results from sequencing showed that miR166a-3p, miR159a, miR6135c, val-miR218, miR4414-3p, miR831-5p, val-miR1017, val-miR832, val-miR633 and val-miR1087 with reads of 687103, 673128, 614703, 539038, 486726, 42040, 11875, 1099, 852, 736, 562 and 285, respectively (S2 and S3 Tables). The expression trend of tested miRNAs was consistent with the Illumina sequencing results, indicating that the gene expression data of miRNAs by sequencing technique was reliable.

Bioinformatics prediction of *V. album* targets for miRNAs

Based on the *V. album* transcriptome, a total of 16188 and 17078 target genes were identified for 593 conserved miRNAs and 1373 novel miRNAs, respectively (S4 Table). To evaluate the putative functions, the targets were mapped to Nr database. Many of putative targets were annotated as transcription factors that play important roles in plant growth and development, such as TATA-binding protein (TBP)-associated factor 4 as a potential target of miR5246, miR838-3p, val-miR314 and val-miR1299; transcription initiation factor TFIID as a target of miR838-3p; basic leucine zipper (bZIP) transcription factors predicted to be targeted by miR5380c, val-miR1128, val-miR 885, val-miR273 and val-miR331; MADS-box protein might
be targeted by miR8130-5p, miR396e-3p, miR8168, miR477g, miR5293, miR5371-5p and val-miR287.

Besides, some targets encoded proteins involved in stress responses, for example, heat shock protein 70 as target of miR6425a-3p; WRKY transcription factor as target of miR5380c, miR5298b and val-miR799; SNF7 family protein as target of miR477g, miR838-3p and val-miR953; E3 ubiquitin-protein ligase COP1 as target of val-miR132. Other predicted targets encode proteins associated with pollen tube development (val-miR284 for mitochondrial Rho GTPase, val-miR111 for gamma-aminobutyrate transaminase), secondary metabolites synthesis (miR172e-3p and val-miR632 for phenylalanine ammonia-lyase, miR5491, val-miR954 and val-miR477 for omega-hydroxypalmitate O-feruloyl transferase) and immune response (miR8136, val-miR360, val-miR260 and val-miR500 for silencing defective 1 family protein), indicating *V. album* miRNAs may be involved in a broad range of physiological and pathological functions.

GO analysis assigned these putative targets into three main categories in terms of biology processes, cellular components, and molecular functions (S5 Table). Based on biology processes, these genes were classified into 23 categories, and the most three over-represented GO terms were “cellular process”, “single-organism process” and “metabolic process”. Categories based on cellular component revealed that these genes were related to 17 cellular parts, of which they are mostly related to “cell”, “cell part” and “organelle”. Based on molecular function, the genes were classified into 14 categories, of which they are mostly involved in “binding” and “catalytic activity” and “transporter activity”.

**Biosynthesis of bioactive components in *V. album* and their putative regulatory miRNAs**

Terpenoids are one of the main components of mistletoe. The lipophilic extract of *V. album* that contained oleanolic acid, betulinic acid, ursolic acid and beta-amyrin acetate showed
potent anti-tumor effects [8, 39]. To date, the pathways involved in terpenoids biosynthesis as well as the miRNAs that might regulate these pathways in *V. album* are still unclear. Terpenoid precursors can be biosynthesized through the mevalonate (MVA) pathway and/or the methylerythritol phosphate (MEP) pathway in different organisms [40,41]. In current study, most enzymes involved in mevalonate pathway including acetyl-CoA C-acetyltransferase (AACT), 3-hydroxy-3-methylglutaryl-CoA (HMG-CoA) synthase, HMG-CoA reductase, mevalonate kinase, phosphomevalonate kinase and diphosphomevalonate decarboxylase have been identified at transcriptome level (S6 Table). Meanwhile, the genes encoding all the enzymes of the methylerythritol phosphate (MEP) pathway were present in *V. album*, including deoxyxylulose-5-phosphate (DOXP) synthase, DOXP reductase, 4-diphosphocytidyl methylerythritol (CDP ME) synthase, CDP ME kinase, methylerythritol cyclodiphosphate (MEcPP) synthase, hydroxymethylbutenyl 4-diphosphate (HMBPP) synthase, HMBPP reductase. These findings indicated that both MVA pathway and MEP pathway were involved in terpenoids backbone biosynthesis in *V. album*.

The enzymes of MVA pathway including AACT, HMG-CoA synthase, HMG-CoA reductase and diphosphomevalonate decarboxylase were predicted to be targeted by miR5042-3p, val-miR720; miR477g; miR6196, miR395o-3p, val-miR187; and miR5246, respectively. The DOXP reductase in upstream processes of MEP pathways was predicted to be target of miR8673. All of these aforementioned enzymes are involved in the biosynthesis of isopentenyl diphosphate/dimethylallyl diphosphate, the precursors of the all the downstream end terpenoids. Additionally, some enzymes associated with the biosynthesis of sesquiterpenoid, triterpenoid, monoterpenoid and diterpenoid might be targeted by miRNAs. For example, miR3932b-5p; miR6451, miR9748; miR7820, miR8714, miR9748; miR5258, miR2106 might be able to target beta-amyrin synthase, farnesol dehydrogenase, (+)-neomenthol dehydrogenase and ent-kaurene oxidase, respectively, which are responsible for catalyzing the generation of common triterpene beta-Amyrin, sesquiterpenoid farnesol, monoterpenoid neomenthol and diterpene ent-kaurene, respectively.

Mistletoe lectins (MLs) are complex molecules comprising both protein and carbohydrates that are capable of binding to cells and inducing biochemical changes in cells. Three commonly known toxic lectins MLI, MLII, and MLIII are type II ribosome inactivating proteins (RIPs), and have been reported to stimulate immune system and induce apoptosis in tumor cells [11,42,43]. They share a common primary structure homology but differ in molecular mass and carbohydrate specificity [44]. It was speculated that ML I-III might be encoded by the same gene and process differently during post-translational modification [45]. The sequences CL9238.Contig1 and CL9238.Contig2 are highly homolog to lectin I precursor (99.11%) and lectin precursor (92.31%), respectively (Table 2), and might be responsible for the encoding of ML I-III in *V. album*. A structurally unrelated chitin-binding mistletoe lectin that consists of 49 amino acids has been characterized recently [46]. It is less toxic than ML I-III [46]. This lectin shows complete identity (100%) with the protein encoded by Unigene23246, indicating Unigene23246 might be the origin of chitin-binding mistletoe lectin. Besides, two other lectins including Mannose/glucose-specific lectin and Curculin-like (mannose-binding) lectin were also identified in mistletoe, which might possess pharmacological effects [47].

Viscotoxins are small proteins belonging to plant thionins, exhibiting cell-killing activity and possible immune-stimulating activity [11,13,15]. To date, seven different isoforms of viscotoxins have been characterized (A1, A1, A3, B, B2, C1 and 1PS), and they differ mainly in their sequence of amino acids. The viscotoxin composition of *V. album* depends on its host tree [8]. Consistent with a previous study [48], our results confirmed that viscotoxins A3 and
viscotoxin B, which is probably encoded by CL146.Contig1 and CL10031.Contig1, were present in *V. album* growing on *Malus*.

As for the potential regulatory miRNAs, only miR9748 and val-miR152 were predicted to target Curculin-like (mannose-binding) lectin family protein and viscotoxin, respectively.

**Bioinformatics prediction of human gene targets for *V. album* novel miRNAs**

Stringent filters were further applied for the novel miRNA candidates that might target human genes. As listed in Table 3, novel miRNA candidates with more than 20 reads, precursors with minimum free energy of less than -30kcal/mol, and precursors with length of no more than 200nt were considered as the most genuine novel miRNA candidates and were later used for human target prediction.

As listed in Table 4, the putative human targets vary by different programs. TargetScan, miRanda, PITA, and RNAhybrid predicted thousands of potential human genes for every novel miRNA, while PITA predicted potential targets for 29 miRNAs, which was probably caused by different algorithms and parameters used in different miRNA target prediction programs. It has been suggested that not a single program was consistently superior than the others among current miRNA target prediction programs [49]. Based on the combination of the methods, the intersection of 30697 potential genes with 59266 miRNA-target pairs were used in subsequent bioinformatics analysis (S7 Table).

Molecular components often interact with each other in a complex reaction network to perform certain biological functions. Pathway enrichment analysis identifies significantly enriched metabolic pathways and signal transduction pathways in proposed targets comparing with the whole genome background, which helps further elucidating genes biological functions. The predicted targets were mapped to the KEGG database and categorized into 305 pathways of which 33 signaling pathways were significantly enriched (Table 5). Notably, these highly enriched KEGG pathways are associated with some human diseases, especially cancer, cardiovascular diseases and neurological disorders.

Five significant enriched signaling pathways are highly related to cancer including transcriptional misregulation in cancer (ko05202), vascular endothelial growth factor (VEGF) signaling pathway (ko04370), pathways in cancer (ko05200), pancreatic cancer (ko05212) and non-small cell lung cancer (ko05223). Tumorigenesis is a multistep process involving a series of genetic alterations [50]. Transcription factors play instrumental functions in driving these pathways.

**Table 2. Genes involved in lectin and viscotoxin expressions and their putative regulatory miRNAs.**

| Gene name                                      | Accession no. | Putative genes | Identity   | E value   | miRNAs  |
|------------------------------------------------|---------------|----------------|------------|-----------|---------|
| lectin precursor                               | AAR25545.1    | CL9238.Contig1  | 99.11%     | 0         | NA      |
| lectin precursor                               | AAR25551.1    | CL9238.Contig2  | 92.31%     | 3.73E-34  | NA      |
| Chitin-binding lectin                          | P81859.1      | Unigene23246    | 100%       | 1.78E-26  | NA      |
| Mannose/glucose-specific lectin family protein | XP_006372325.1| Unigene7638     | 62.77%     | 2.02E-27  | NA      |
| Curculin-like (mannose-binding) lectin family protein | XP_007021734.1| Unigene24650    | 54.77%     | 5.98E-137 | miR9748 |
| thionin precursor                              | AAB29761.1    | CL146.Contig4   | 96.49%     | 1.12E-60  | NA      |
| Viscotoxin-A3                                  | AAB29759.1    | CL146.Contig1   | 90.99%     | 2.01E-53  | val-miR152 |
| Viscotoxin-B                                   | P08943.2      | CL10031.Contig1 | 85.15%     | 2.21E-47  | NA      |

NA represents not available.

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Table 3. Potential novel miRNAs from *V. album* used for human target prediction.

| miRNA   | Reads | Sequences                     | ML(nt) | PL(nt) | MFE (kcal/mol) |
|---------|-------|-------------------------------|--------|--------|----------------|
| val-miR218 | 11875 | GAUGAUCGCCAGUCGCCAGGAGGA      | 21     | 119    | -63.1          |
| val-miR11  | 1436  | CACUGUAACACUUUUUGAAGAAGG     | 22     | 85     | -30.2          |
| val-miR1338 | 1081  | CGCAAGAGCUUAAUAGAUGAU         | 21     | 143    | -43.03         |
| val-miR856 | 1021  | UAAGUGUGCGUUGUIAUCAGAUA      | 22     | 105    | -56.41         |
| val-miR718 | 882   | UUUUGUCUUGUGAAGAUGUCUUUUGU   | 21     | 161    | -79.2          |
| val-miR1017 | 852   | UCCCAACUCGGACUCUGAGGUC        | 21     | 189    | -48.4          |
| val-miR832 | 736   | UAAUGUCCAGUCUCUGACUACC       | 21     | 136    | -49.9          |
| val-miR457 | 585   | UAGCCGGUUCUUCUCAACGCGC       | 21     | 182    | -74.77         |
| val-miR633 | 562   | UCAAUGAUCUGGUUGUGGCGCUU      | 21     | 105    | -53.3          |
| val-miR1370 | 436   | UAUCAGUACUGUIGUGAGGAG        | 20     | 98     | -48.9          |
| val-miR588 | 435   | CGAUCGUAAUAUCAAGAAGUUU       | 21     | 126    | -50.4          |
| val-miR539 | 366   | UAAGUCUGCAGACAGUGGUGCA       | 21     | 131    | -44.7          |
| val-miR1087 | 285   | GGGGAUAUGCAACUUGGGACCC       | 21     | 179    | -97            |
| val-miR944 | 278   | UUUUCUUGGUUGUGUUGUGGU        | 21     | 156    | -82.66         |
| val-miR1048 | 267   | CGUGGGAACCUCGGCAGUGG         | 20     | 64     | -51            |
| val-miR262 | 227   | UUAAUUCUGAAGUUGUCUCG        | 21     | 131    | -63.1          |
| val-miR765 | 201   | CAGAUGUGAGAAGAAGGCAC         | 21     | 146    | -70.8          |
| val-miR1052 | 181   | CAAGACAUUACUGUGCCUC          | 21     | 117    | -36.13         |
| val-miR421 | 103   | UUUCAGUAAUGUUGUUGGAC         | 21     | 119    | -30.4          |
| val-miR885 | 103   | UUGAUGAUCACUAGUGUGGCUC       | 21     | 104    | -43.3          |
| val-miR64  | 101   | AGGGUGUGGAGUACUGGGGGA        | 21     | 85     | -43.7          |
| val-miR790 | 93    | UAGCCAGAAGCUAGACUUGGCUC     | 21     | 111    | -54.6          |
| val-miR648 | 84    | UUGUGAAUAGAUCUCCACAGU        | 22     | 81     | -31.6          |
| val-miR333 | 80    | AUAUGUGUGGAGUAAUGGCA         | 21     | 85     | -30            |
| val-miR503 | 76    | UCAUCAAARAGUUGUGUCGCA        | 21     | 149    | -80            |
| val-miR552 | 48    | CAUUGGACUGAUCUGAGAC         | 21     | 150    | -54            |
| val-miR1306 | 40    | CAAUGGAGGCGCGACUGCUGGCUG    | 21     | 108    | -68.61         |
| val-miR269 | 36    | GACUACGAGUACGAGACCGGGG       | 22     | 142    | -51.1          |
| val-miR1328 | 33    | CGAAGAUGUAGGCAAGGCGAC        | 22     | 127    | -39.8          |
| val-miR1127 | 32    | CCCACACUCUAAGAUUGGGUG        | 20     | 170    | -67.8          |
| val-miR1110 | 28    | CCGCGAGGCGACAGCGGCCCGG       | 22     | 91     | -46.8          |
| val-miR855 | 27    | GAACUGAGUGCGACUGUAGCCUC      | 22     | 129    | -36.3          |
| val-miR1086 | 26    | UGGCCCGGAGAUGCUUGGACGCGC    | 21     | 141    | -57.9          |
| val-miR6    | 26    | AUAGUGAUGAUGCAAAUUGGAC       | 21     | 148    | -46.41         |
| val-miR954 | 26    | UCCUGAUGCGCGAGGGCGACGCAC    | 21     | 130    | -56.6          |
| val-miR198 | 24    | UGACGAGUUGGACAAAGAAACU       | 21     | 148    | -69.39         |
| val-miR560 | 24    | UUCUCAGAUCUCUGCAGUAGC       | 21     | 180    | -56.4          |
| val-miR82  | 24    | UGAUGUCCCGCACUUCUGGCCCUC    | 21     | 178    | -58.5          |
| val-miR1342 | 23    | ACCUGUAGCGCGUAGGGCCAGCA     | 21     | 104    | -33.39         |
| val-miR92  | 22    | AAAGAUCAGAUAUACAGGAGCUCU    | 22     | 59     | -36.5          |
| val-miR550 | 21    | UCUUUUGGUAUUAUGGAGGAC        | 21     | 117    | -59.3          |
| val-miR615 | 21    | UUUCUGUCGCGACUCUGGAGA       | 21     | 125    | -42.5          |
| val-miR163 | 20    | GAUCGAGGUAUAGUAACUACU       | 20     | 197    | -84.8          |
| val-miR834 | 20    | CGGCACUGCGUCUUCGUGCC        | 20     | 102    | -34.6          |

ML: mature miRNA length; PL, precursor length; MFE, minimum free energy.

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Table 4. Statistics of miRNA target predictions.

| miRNA target prediction program | miRNA number | Target genes | miRNA-target pairs |
|---------------------------------|--------------|--------------|-------------------|
| Targetscan                       | 44           | 46555        | 581567            |
| miRanda                          | 44           | 36284        | 95353             |
| PITA                             | 29           | 48154        | 1009051           |
| RNAhybrid                        | 44           | 48354        | 1524020           |
| Intersection                     | 29           | 30697        | 59266             |

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Table 5. Highly enriched KEGG pathways for putative human targets.

| Pathway terms                        | Number of target genes | Rich factor | Qvalue     | Pathway ID |
|--------------------------------------|------------------------|-------------|------------|------------|
| MAPK signaling pathway               | 793                    | 0.795       | 7.71E-10   | ko04010    |
| Transcriptional misregulation in cancer | 1866            | 0.755       | 1.89E-08   | ko05202    |
| Calcium signaling pathway            | 495                    | 0.810       | 2.41E-08   | ko04020    |
| VEGF signaling pathway               | 271                    | 0.850       | 2.43E-08   | ko04370    |
| ECM-receptor interaction             | 365                    | 0.824       | 6.91E-08   | ko04512    |
| Purine metabolism                    | 1684                   | 0.750       | 8.92E-07   | ko00230    |
| Glutamatergic synapse                | 312                    | 0.823       | 9.30E-07   | ko04724    |
| Focal adhesion                       | 692                    | 0.773       | 1.37E-05   | ko04510    |
| Neurotrophin signaling pathway       | 421                    | 0.794       | 1.52E-05   | ko04722    |
| Pyrimidine metabolism                | 1488                   | 0.746       | 4.01E-05   | ko00240    |
| Measles                              | 309                    | 0.794       | 4.00E-04   | ko05162    |
| Morphine addiction                   | 219                    | 0.808       | 7.92E-04   | ko05032    |
| GnRH signaling pathway               | 273                    | 0.796       | 7.92E-04   | ko04912    |
| Phosphatidylinositol signaling system| 229                    | 0.801       | 1.49E-03   | ko04070    |
| Arginine and proline metabolism      | 404                    | 0.774       | 1.57E-03   | ko00330    |
| Type II diabetes mellitus            | 188                    | 0.810       | 1.57E-03   | ko04930    |
| Axon guidance                        | 440                    | 0.769       | 1.91E-03   | ko04360    |
| GABAergic synapse                    | 214                    | 0.799       | 2.69E-03   | ko04727    |
| Pathways in cancer                   | 874                    | 0.746       | 3.37E-03   | ko05200    |
| Herpes simplex infection             | 407                    | 0.768       | 3.71E-03   | ko05168    |
| Retrograde endocannabinoid signaling | 230                    | 0.785       | 9.05E-03   | ko04723    |
| Glycosaminoglycan biosynthesis—chondroitin sulfate | 40                  | 0.909       | 1.20E-02   | ko00532    |
| Alanine, aspartate and glutamate metabolism | 93                  | 0.830       | 1.47E-02   | ko00250    |
| Influenza A                          | 448                    | 0.755       | 1.69E-02   | ko05164    |
| Amphetamine addiction                | 246                    | 0.776       | 1.69E-02   | ko05031    |
| Leishmaniasis                        | 174                    | 0.791       | 1.70E-02   | ko05140    |
| Fc epsilon RI signaling pathway      | 211                    | 0.781       | 1.76E-02   | ko04664    |
| B cell receptor signaling pathway    | 252                    | 0.771       | 2.64E-02   | ko04662    |
| Pancreatic cancer                    | 195                    | 0.780       | 2.83E-02   | ko05212    |
| beta-Alanine metabolism              | 226                    | 0.771       | 3.70E-02   | ko00410    |
| Nicotine addiction                   | 86                     | 0.819       | 3.73E-02   | ko05033    |
| Toxoplasmosis                        | 267                    | 0.763       | 4.63E-02   | ko05145    |
| Non-small cell lung cancer           | 147                    | 0.786       | 4.63E-02   | ko05223    |

Rich factor represents the ratio of the number of predicted genes and the number of all genes in the pathway.

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gene expressions, whereas misregulation of transcription factors can cause the acquisition of tumor-related properties [51]. For example, runt related transcription factor 1 (RUNX1) and lysine methyltransferase 2A (MLL1) are essential for chromosomal translocations in acute myeloid leukemia [52,53], which were predicted as targets of val-miR1086, val-miR765, val-miR615; and val-miR834 val-miR765, val-miR550, val-miR1127, val-miR954, val-miR1086, val-miR421, respectively. It is well known that the tumor protein p53 is a major tumor suppressor, and the mutation of p53 can provoke tumor imitation [54]. The miRNAs such as val-miR1086 and val-miR1127 were predicted to regulate the expression level of p53. By targeting these genes associated with transcriptional misregulation in cancer, these novel miRNA candidates may prevent cancer initiation and progression. Angiogenesis induction is one of the major hallmarks of cancer. It is generally accepted that VEGF is a major driver of the angiogenic process in physiological and pathological processes. VEGF and its receptors are often found overexpressed in tumors [55,56]. Suppression of the essential molecules in VEGF signaling pathway, such as phospholipase C gamma 1 (PLCG1) possibly targeted by val-miR615 and val-miR1086, tyrosine-protein kinase Src possibly targeted by val-miR834 and val-miR1086, may block the angiogenic activity of tumor tissue, resulting in tumor vascular regression and anti-tumor effects.

Diabetes mellitus is one of the most prevalent metabolic disorders. It is characterized by hyperglycemia, and long-term hyperglycemia may lead to systemic complications, such as macrovascular diseases, coronary artery disease, peripheral arterial disease and stroke [57]. Insulin receptor (INSR) dysregulation is a well-established defect in type II diabetes mellitus (ko04930). Mitogen-activated protein kinase 1(ERK) and inhibitor of nuclear factor kappa B kinase (IKK) are serine kinases that can directly inactivate insulin receptor substrate (IRS) through serine phosphorylation, and impair insulin sensitivity [58,59]. The putative inhibition of ERK and IKK by val-miR615 and val-miR834 might restore the impaired INSR singling. Calcium is a critical mediator of excitation–contraction coupling in cardiac cells, and cellular calcium signaling dysfunction is central to the pathophysiology of a wide range of cardiac diseases [60]. Based on computational analysis, a range of key molecules in calcium signaling pathway (ko04020) were targeted by mistletoe miRNAs, for example, calcium voltage-gated channels (CaV1, CaV2 and CaV3) were mutual putative targets of val-miR1086 and val-miR765, calcineurin (CaN) was predicted to be target by val-miR834 and val-miR1086. Through regulation of these potential therapeutic targets [61,62], mistletoe miRNAs might be responsible for its cardiovascular protective effect.

It is noteworthy that the putative human genes targeted by mistletoe miRNAs were also involved in several pathways associated with the nervous system, such as neurotrophin signaling pathway (ko04722), morphine addiction (ko05032), glutamatergic synapse (ko04724), GABAergic synapse (ko04727), axon guidance (ko04360), amphetamine addiction (ko05031) and nicotine addiction (ko05033). Mistletoe had been beneficial for the treatment of epilepsy, depression, sleep disorders and labour-pain in middle ages [63], however, since deficiency of scientific evidence, mistletoe preparations are not applied for neurological diseases in modern medicine. Based on our bioinformatics prediction, mistletoe novel miRNAs might target critical neurotransmitter receptors and neurotransmitter transporters, such as gamma-aminobutyric acid type B receptor (GABA_B) as a putative target of val-miR1086; glutamate metabotropic receptors (mGluRs) as putative targets of val-miR1342, val-miR954, val-miR550, val-miR560, val-miR765, val-miR1086, val-miR550, val-miR1328; dopamine transporter (DAT) as a putative target of val-miR765 and val-miR1110. It is possible that mistletoe miRNAs could influence neurotransmission by affecting transport of neurotransmitters including GABA, glutamic acid and dopamine, which might explain the traditional use of mistletoe to treat epilepsy, insomnia and other neurological disorders. One critical concern regarding treatment of
neurological diseases is the blood-brain barrier that represents a problem to any therapy involving systemic delivery of oligonucleotides [64]. Recent publications indicated that exosomes could transfer across blood-brain barrier, serving as an efficient vehicle to deliver miRNAs to the recipient neurocytes [65–67]. Therefore, with the assistance of plant- or animal-derived molecules or nanoparticles such as exosomes, mistletoe miRNAs might reach nervous system and exert its function.

Interestingly, it is recorded that mistletoe has been used by North American Indians to treat measles and dog bites [68], indicating the antiviral potential of mistletoe. Indeed, the predicted targets were involved in several virus and parasitic infections including measles (ko05162), herpes simplex infection (ko05168), influenza A (ko05164), leishmaniasis (ko05140) and toxoplasmosis (ko05145). Specifically, Toll-like receptors that are sentinel receptors of the host innate immune system to detect the presence of microbial infection [69], were predicted to be targeted by val-miR765 and val-miR1328. Furthermore, val-miR1086 and val-miR954 might target TNF receptor (TNFR) and interferon gamma receptor (IFNGR), and thus prevent overexuberant inflammatory response. Although mistletoe is not used for pathogenic diseases nowadays, the antiviral potential of mistletoe might be recognized.

Discussion

Endogenous microRNAs (miRNAs) are a class of single-stranded non-coding RNA molecules of approximately 22 nucleotides that play crucial roles in gene expression. In mammals, an estimated 60% of all protein-coding genes may contain miRNA binding sites [3,4]. MiRNA dysregulation is frequently associated with human diseases such as cancer, cardiovascular diseases, central nervous system diseases and metabolic disorders [70,71]. To date, miRNA-based novel therapeutics have been developed for the treatment of human diseases, and several preclinical studies on therapeutic miRNA replacement have been initiated [72,73], indicating miRNA-based therapeutics are coming of age.

Herbal medicine is globally accepted as a valid alternative system of therapy. Though ancient medical treatises have documented a large number of medicinal plants, their bioactive constituents and corresponding interactions with human have not been comprehensively characterized. New plant bioactive molecules are being discovered. In recent years, regulation of human genes by plant miRNAs has attracted great attention. Rice miRNAs were suggested to enter mammalian bloodstream and have a functional role in human metabolism [5]. The MIR2911 from honeysuckle were found to target influenza viruses and protect mice from influenza [6]. Plant derived miR159 significantly suppressed breast cancer cell proliferation by targeting transcription factor 7 (TCF7) [74]. Oral application of a cocktail that consisted of plant-based tumour suppressor miRNAs was able to reduce tumour burden in mice [75]. These studies indicate that miRNAs derived from plants may function as bioactive constituents to regulate human health.

*V. album* is a European medicinal plant surrounded by legends and myths. It has been used in folk medicine for the treatment of cancer, cardiovascular disease, and other symptoms. In modern medicine, *V. album* has been mainly used as an anti-tumor therapy, which is attributed to the anti-cancer and immune stimulating activities of its bioactive components including viscotoxins, lectins and terpenoids. However, the active ingredients that might be responsible for its cardiovascular protective effects as well as other beneficial applications remain to be clarified.

Here we propose that miRNAs in *V. album* might serve as an independent category of active ingredients and provide beneficial effects for human consumers. Since *V. album* genome
information is limited, we conducted RNA-seq and sRNA-seq to identify and characterize the conserved and possibly novel miRNAs from *V. album*. Bioinformatics tools have been applied to understand their possible functions in plant biological processes and potential roles in human gene regulation.

By using high-throughput sequencing technology, a total of 699 conserved miRNAs and 1373 novel miRNAs with a length of 21–24 nt have been identified from *V. album*. The reliability of the sequencing data was confirmed by qRT-PCR. In *V. album*, these miRNAs were involved in various biological processes including plant growth, development, signal transduction and stress responses. Transcription factors are involved in important plant developmental processes. The MADS-box transcription factors are crucial for floral development [76], and might be controlled by several miRNAs such as miR8130-5p, miR396e-3p and miR8168. WRKY transcription factors are involved in various plant processes, especially in coping with diverse biotic and abiotic stresses [77], and were predicted to be targeted by miR5380c, miR5298b and val-miR799. One miRNA could target several transcription factors, such as miR838-3p was predicted to target TBP-associated factor and TFIID; miR5380c might target bZIP transcription factors and WRKY transcription factors, indicating their multiple roles in plant processes. Besides, the annotated targets involved in various metabolic processes, stimulus response, catalytic activity, and other biological processes were also predicted, suggesting that miRNAs play essential roles in plant growth and development.

Mistletoe lectins and viscotoxins are pharmaceutical proteins present in *V. album*, and they have been considered to be mainly responsible for the anti-tumor activity of *V. album*. Previous studies have isolated and identified lectins and viscotoxins from mistletoe at protein level [44,46,78,79]. In this study, the expressions of these bioactive components were confirmed at transcriptome level. ML I-III might be translated from CL9238.Contig1 and/or CL9238.Contig2, while Unigene23246 probably encodes chitin-binding mistletoe lectin. However, the miRNAs that might target CL9238.Contig1, CL9238.Contig2 and Unigene23246 were not identified in our study. It has been reported that the amount of MLs in the leaves of *V. album* showed maximum in December [80]. It is possible that at this time, their corresponding regulatory miRNAs are too low to be detected. Mannose/glucose-specific lectin and curculin-like (mannose-binding) lectin, which differ from known mistletoe lectins (ML I-III and chitin-binding lectin), have been newly identified from mistletoe. However, the expression and bioactivities of these two newly identified mistletoe lectins need to be further validated.

For viscotoxins, CL146.Contig1, which is highly homologous to Viscotoxin-A3 (90.99%), was predicted to be target of val-miR152. However, no miRNAs were identified to target CL10031.Contig1 and CL146.Contig4, which were annotated as Viscotoxin-B and thionin precursor, respectively. Except through post-transcriptional regulation by miRNAs, the expressions of viscotoxins might as well be controlled by transcriptional regulation.

The pharmacological properties of *V. album* also attributed to the presence of triterpene acids, especially oleanolic acid, betulinic acid and ursolic acid, which have been reported to enhance the toxicity of mistletoe lectins in tumor cells [39,42]. The biosynthesis of terpenoids in *V. album* has not yet been elucidated. Our study identified majority of genes encoding enzymes that involved in both MVA pathway and MEP pathway, indicating terpenoids biosynthesis in *V. album* was via both pathways. The compounds isopentenyl pyrophosphate (IPP) and dimethylallyl pyrophosphate (DMAPP), produced in upstream pathway, are the common precursors for all the downstream end terpenoids. The miRNAs such as miR5042-3p, miR477g and miR6196 that predicted to target the upstream enzymes, might be involved in the regulation of IPP and DMAPP levels. Some putative targets of miRNAs were downstream enzymes
in mono-, sesqui, di-, and triterpenoid biosynthetic pathways, such as beta-amyrin synthase, farnesol dehydrogenase, neomenthol dehydrogenase and ent-kaurene oxidase might be targeted by miR3932b-5p; miR6451, miR9748; miR7820, miR8714, miR9748; miR5258, miR2106, respectively.

Bioinformatics predictions have been employed to identify the potential human targets of plant miRNAs. Seven miRNAs from medicinal plant *Moringa oleifera* L., have been predicted to involve in cell cycle, apoptosis and metabolic regulation in humans [81]. Approximately 50 human target genes associated with energy metabolism, lipoprotein metabolism, and other biological process have identified as the target genes of a rice miRNA (MIR168a) [5]. In this study, bioinformatics tools were applied to identify the putative human target genes of 44 *V. album* specific miRNAs. A huge number of 30697 putative genes were predicted and then mapped to the KEGG database to find their roles in human metabolism and human diseases. A total of 14995 putative targets were highly enriched in 33 KEGG pathways. Among them, five pathways were highly related with cancer including transcriptional misregulation in cancer (ko05202), VEGF signaling pathway (ko04370), pathways in cancer (ko05200), pancreatic cancer (ko05212), non-small cell lung cancer (ko05223), while Type II diabetes mellitus (ko04930) and calcium signaling pathway (ko04020) were associated with cardiovascular and metabolism diseases. By targeting essential molecules involved in these pathways, *V. album* specific miRNAs might possess pharmaceutical effects against cancer, cardiovascular and metabolism diseases, which might provide scientific support for the folk and clinic use of mistletoe.

The use of mistletoe for the treatment of neurological disorders and infections has been recorded in ancient times. However, since there is no scientific evidence explaining its effects, mistletoe is not used for these purposes in modern medicine. Interestingly, bioinformatics predictions showed that some of the putative targets relate to several neurological pathways, including neurotrophin signaling pathway (ko04722), morphine addiction (ko05032), glutamateergic synapse (ko04724), GABAergic synapse (ko04727), axon guidance (ko04360), amphetamine addiction (ko05031) and nicotine addiction (ko0533). Some infections related pathways such as measles (ko05162), herpes simplex infection (ko05168), influenza A (ko05164), leishmaniasis (ko05140) and toxoplasmosis (ko05145) were also highly enriched. These findings might provide an explanation for the traditional medicine use of mistletoe in middle ages, and inspire the modern medicine use of mistletoe.

Experimental validation of predicted plant miRNA-human mRNA interaction is necessary in upcoming investigations. However, a series of questions remain to be answered. Would herbal miRNAs be stable during herbal preparation and human digestion process? Would herbal miRNAs be selectively absorbed by the human gastrointestinal tract? How would plant miRNAs be recognized by human cells? How would plant miRNAs be loaded into mammalian RNA Induced Silencing Complex (RISC), in which the miRNAs exert their function together with Argonaute proteins? Our study implied that medicinal plant specific miRNAs might contribute to their corresponding pharmaceutical effects, and our next step would be to focus on the detection of herbal miRNAs in various herbal preparations, evaluation of the capability of herbal miRNAs to transfer intestinal barriers, and investigation of their intracellular fate in human cells.

In summary, this study comprehensively identified the miRNAs from medicinal plant *V. album*, and characterized the genes and their potential regulatory miRNAs for the synthesis of bioactive components such as viscotoxins, lectins and terpenoids, helping to develop a deeper understanding of biosynthesis of active ingredients in mistletoe. Computational predictions indicated the anti-tumor potential, cardiovascular protective and neurological protective
effects of *V. album* specific miRNAs, and initiated further investigation to elucidate the regulatory function of plant miRNAs in human health and diseases.

**Supporting information**

S1 Table. Primers for qRT-PCT.
(XLSX)

S2 Table. Conserved miRNAs identified in *V. album*.
(XLSX)

S3 Table. Novel miRNA candidates from *V. album*.
(XLSX)

S4 Table. Predicted *V. album* targets for conserved and novel miRNAs.
(XLSX)

S5 Table. GO analysis of target genes for miRNAs in *V. album*.
(XLSX)

S6 Table. Target genes for miRNAs involved in terpenoids biosynthesis.
(XLSX)

S7 Table. Putative human targets of *V. album* novel miRNAs.
(XLSX)

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**Author Contributions**

Conceptualization: Wenyan Xie, Matthias F. Melzig.

Data curation: Wenyan Xie, Jacob Adolf.

Funding acquisition: Matthias F. Melzig.

Investigation: Wenyan Xie.

Methodology: Wenyan Xie, Jacob Adolf.

Project administration: Wenyan Xie.

Supervision: Wenyan Xie, Matthias F. Melzig.

Writing – original draft: Wenyan Xie.

Writing – review & editing: Jacob Adolf, Matthias F. Melzig.

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