De Novo Transcriptome Analysis of Warburgia ugandensis to Identify Genes Involved in Terpenoids and Unsaturated Fatty Acids Biosynthesis

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Abstract

The bark of Warburgia ugandensis (Canellaceae family) has been used as a medicinal source for a long history in many African countries. The presence of diverse terpenoids and abundant polyunsaturated fatty acids (PUFAs) in this organ contributes to its broad range of pharmacological properties. Despite its medicinal and economic importance, the knowledge on the biosynthesis of terpenoid and unsaturated fatty acid in W. ugandensis bark remains largely unknown. Therefore, it is necessary to construct a genomic and/or transcriptomic database for the functional genomics study on W. ugandensis. The chemical profiles of terpenoids and fatty acids between the bark and leaves of W. ugandensis were compared by gas chromatography-mass spectrometry (GC-MS) analysis. Meanwhile, the transcriptome database derived from both tissues was created using Illumina sequencing technology. In total, about 17.1 G clean nucleotides were obtained, and de novo assembled into 72,591 unigenes, of which about 38.06% can be aligned to the NCBI non-redundant protein database. Many candidate genes in the biosynthetic pathways of terpenoids and unsaturated fatty acids were identified, including 14 unigenes for terpene synthases. Furthermore, 2,324 unigenes were discovered to be differentially expressed between both tissues; the functions of those differentially expressed genes (DEGs) were predicted by gene ontology enrichment and metabolic pathway enrichment analyses. In addition, the expression of 12 DEGs with putative roles in terpenoid and unsaturated fatty acid metabolic pathways was confirmed by qRT-PCRs, which was consistent with the data of the RNA-sequencing. In conclusion, we constructed a comprehensive transcriptome dataset derived from the bark and leaf of W. ugandensis, which forms the basis for functional genomics studies on this plant species. Particularly, the comparative analysis of the transcriptome data between the bark and leaf will provide critical clues to reveal the regulatory mechanisms underlying the biosynthesis of terpenoids and PUFAs in W. ugandensis.
Introduction

Warburgia ugandensis Sprague, which belongs to a member of Canellaceae family, is a small evergreen tree distributed in eastern and southern Africa. Warburgia bark has been commonly used as traditional medicines for the treatment of gastro-intestinal disorders, cold, cough and sore throat, fever, malaria, respiratory and odontological problems in African countries [1]. It has been reported that the extract of Warburgia bark or leaf exhibits a variety of pharmacological effects, such as anti-bacterial, anti-fungal, anti-mycobacterial, antioxidant, and anti-inflammatory activities [2–6]. In addition, W. ugandensis is also used for fodder, insecticide, and toothbrush et al [1].

Several phytochemical studies reported the presence of diverse terpenoids in the bark and leaf of W. ugandensis, including waburganal, muzigadial, ugandenial A, and ugandensidial (cinnamodial) et al. [7–12]. The abundant terpenoids, especially those drimane- and colorotane-type sesquiterpenoids, were suggested to be relevant to its various medicinal effects [2, 3, 7, 12, 13]. Terpenoid is the largest class of plant natural products and plays crucial roles in plant growth development and ecological applications [14]. The biosynthesis of plant terpenoids can be divided into three stages as follows [15, 16]. In the first stage, the universal precursors, isopentenyl diphosphate (IPP) and dimethylallyl diphosphate (DMAPP), are formed via the 2-C-methyl-D-erythritol 4-phosphate (MEP) pathway and/or the mevalonate (MVA) pathway; second, geranyl diphosphate synthase (GPPS) and/or farnesyl diphosphate synthase (FPPS) convert the universal isoprenes to the terpenes precursors, geranyl diphosphate (GPP) and farnesyl diphosphate (FPP), respectively. In the last stage, the unique backbone of the terpenoids are generated under the catalysis of different kinds of terpenoid synthases (TPSs), including monoterpen and/or sesquiterpene synthases. Molecular characterization of TPSs has been intensively studied in many plant species, such as Arabidopsis thaliana, Malus domestica, Oryza sativa, and Vitis vinifera [17–21]. However, compared with the proceeding of chemical characterization of terpenoids, the study on the molecular characterization of TPSs, the terpenoid-forming enzymes, is far away backward. For example, in W. ugandensis, although several terpenoids have been structurally characterized, there is no TPS characterized in this plant at molecular level so far.

Polyunsaturated fatty acids (PUFAs) are a class of fatty acids that contain two or more double bonds in their backbones. Some PUFAs such as linoleic acid (18:2) and linolenic acid (18:3) cannot be synthesized in mammals or humans, and human only consumes them through the diet. PUFAs, especially those omega-3 (Ω-3) types, are important for maintaining human body’s normal metabolism and provide numerous health benefits, including anticancer properties, prevention and treatment of coronary heart disease, hypertension and type 2 diabetes [22, 23]. It has been reported that the crude extract of W. ugandensis shows anti-mycobacterial activity and its stem bark can be used to treat tuberculosis, and these effects are probably due to the presence of linoleic acid (18:2) and drimane sesquiterpenoids [7]. In plant, fatty acids are generally synthesized from acetyl-CoA in a three-step process to form palmitic acid (16:0) or stearic acid (18:0). Then, stearic acid (18:0) is desaturated to oleic acid (18:1), linoleic acid (18:2), or linolenic acid (18:3) by specific fatty acid desaturases (FADs), including FAD2 and FAD3, which are key enzymes to control the biosynthesis of unsaturated fatty acids [24]. Finally, linoleic acid (18:2) or linolenic acid (18:3) can be further converted into a variety of complex PUFAs under the catalysis of a series of desaturases and elongases [25, 26].

In recent years, due to the high-throughput, accuracy, and reproducibility, mRNA sequencing (RNA-Seq) technology has emerged as a powerful tool to profile the genome-wide transcriptional pattern in different tissues and/or at different developmental stages [27], and discover novel genes in specific biological processes, especially in those non-model organisms.
without genomic sequences [28–30]. To date, the genomic information of *W. ugandensis* is not available yet, and only 20 nucleotide sequences (or ESTs), including five partial cDNAs encoding for putative sesquiterpene synthases, have been deposited in the NCBI GenBank database. Considering the medicinal and economic importance of *W. ugandensis*, it deserves to lunch genomic studies on this plant species, especially to investigate the biosynthesis of its medicinal components, e.g. terpenoids and PUFAs. To start this project, we reported here the construction of *W. ugandensis* transcriptome database derived from its leaf and bark, which are the tissue sources for its medical use. As a result, a total of 17.1 G clean nucleotides were obtained and *de novo* assembled into 72,591 unigenes thereafter. This database will provide an important resource in identifying genes encoding enzymes in terpenoids and PUFAs biosynthetic pathways in *W. ugandensis*.

**Results**

**Chemical analysis of terpenoids and fatty acids in the leaf and bark of *W. ugandensis***

Since terpenoids and PUFAs are thought to be potential bioactive compounds of *W. ugandensis*, those compounds were analyzed and compared between its bark and leaf. As shown in Fig 1A, five sesquiterpenoids, including copaene, caryophyllene, farnesene, humulene, and cubebene, were detected by gas chromatography-mass spectrometry (GC-MS). Except that farnesene is exclusively synthesized in the leaf, the other four sesquiterpenoids are presented in both tissues with their concentrations being relatively higher in the leaf. Eight monoterpenoids were detected by this GC-MS analysis. The content of myrcene and ocimene is significantly higher in the leaf than that in the bark, while more of other monoterpenoids such as α-pinene and camphene is detected in the bark, especially limonene is only found in the bark but not in the leaf (Fig 1B).

PUFAs consist of the major proportion of fatty acids in *W. ugandensis* bark and leaf, accounting for 41.34% and 67.76% of their total fatty acids, respectively (Fig 1C). In the bark, the most abundant of PUFA is linoleic acid (18:2), which accounts for 28.59% of the total fatty acids; while linolenic acid (18:3) is the highest amount of PUFA in the leaf, reaching up to 52.77% of the total fatty acids. Palmitic acid (16:0) accounts for roughly equal proportions (about 19%) in the both tissues. The remaining types of fatty acids, such as stearic acid (18:0), oleic acid (18:1), and erucic acid (22:1), are detected in a higher concentration in the bark relative to the leaf.

**Transcriptome sequencing and *de novo* assembly**

Two cDNA libraries from the bark and leaf of *W. ugandensis* were constructed by the Illumina sequencing technology, which produced 88,241,216 and 85,560,090 raw reads in the bark and leaf, respectively (Table 1). After removing the low-quality reads and adapter sequences, 87,093,692 clean reads for the bark and 84,389,152 clean reads for the leaf were obtained; and then a total of 17.1 G data were further *de novo* assembled into 72,591 unigenes with an average size of 1,051 bp and an N50 of 1,467 bp. The length distribution of the *W. ugandensis* unigenes showed that approximately 31% of the unigenes was presented in a length of more than 1 Kb (S1 Fig). These results showed that the throughout and sequencing quality were high enough for further bioinformatics analyses.

**Functional annotation and classification of unigenes**

For functional annotation of the *W. ugandensis* transcriptome, the unigenes were searched against diverse databases using BLASTX program. Only 38.06% of unigenes (27,631) were
annotated in the NCBI non-redundant (Nr) protein database at a cut-off E-value of $10^{-5}$ (Table 2). The relative low number of annotated unigenes was probably due to no publicly available genome or the limitations of EST information for species in the Canellaceae family.
Out of these hits, 16,366 unigenes (59.23%) had an E-value of less than $10^{-45}$, indicating they had very strong similarities to known proteins.

The KOG database refers to Cluster of Orthologous Groups of gene products. Every protein in the KOG is assumed to be evolved from an ancestor protein, and thus they are either orthologs or paralogs. The KOG annotation showed that 16,636 (22.92%) unigenes were classified into 25 KOG categories. The cluster for "General function prediction only" represented the largest group, followed by "posttranslational modification, protein turnover, chaperones", and "signal transduction mechanisms" (Fig 2). Additionally, 1097 unigenes (6.59% of total KOG hits) were categorized into "secondary metabolites biosynthesis, transport and catabolism".

To further functionally categorize the *W. ugandensis* transcriptome, a total of 21,702 (29.90%) of the Nr-annotated unigenes were assigned into different GO terms according to their associated biological processes, cellular components and molecular function. In the "biological process" category, the term "metabolic process" and "cellular processes" were predominant; and in the "cellular component" group, "cell parts" and "cell" were found to be the largest classes; while under the "molecular function" class, the majority of unigenes were assigned to be involved in "catalytic activities" and "binding" (Fig 3).

To better understand the function of the assemble unigenes in biological pathways; they were searched against the KEGG pathway database. It was found that only 6,489 (8.94%) unigenes were assigned to 327 KEGG pathways (See S1 Table). The highest represented pathway was "carbon metabolism" (ko01200, 550 unigenes), followed by "ribosome" (ko03010, 510 unigenes) and "biosynthesis of amino acids" (ko01230, 349 unigenes). Since sesquiterpenoids and PUFAs were thought to be medicinal components of *W. ugandensis*, we focused our efforts to survey the unigenes whose functions related to terpenoid and fatty acid metabolism (Table 3). As a result, 53 unigenes were identified for "terpenoid backbone biosynthesis" (ko00900), 20 unigenes for "sesquiterpenoid and triterpenoid biosynthesis" (ko00909), 13 unigenes for "monoterpenoid biosynthesis" (ko00902), 94 unigenes for "fatty acid biosynthesis" (ko00061), and 100 unigenes for "biosynthesis of unsaturated fatty acids" (ko01040).

### Table 1. Statistic of sequencing and de novo assembling of transcriptome in *W. ugandensis*.

| Item         | Sample | Number (n) | Sequence (bp) | Valid ratio (%) | GC (%) | Q30 (%) | N50 (bp) | Average length (bp) |
|--------------|--------|------------|---------------|-----------------|--------|---------|----------|---------------------|
| raw read     | Leaf   | 88241216   | 8824121600    | —               | —      | —       | —        | —                   |
|              | Bark   | 85560090   | 8556009000    | —               | —      | —       | —        | —                   |
| clean read   | Leaf   | 87093692   | 8693658441    | 98.52           | 45.00  | 88.89   | —        | —                   |
|              | Bark   | 84389152   | 8422552230    | 98.44           | 45.50  | 89.38   | —        | —                   |
| Total        | Unigenes | 72591     | 76284840      | —               | —      | —       | 1467     | 1050.89             |

doi:10.1371/journal.pone.0135724.t001

### Table 2. Statistics of annotations for assembled unigenes in *W. ugandensis*.

| Database | Account | Percentage (%) |
|----------|---------|----------------|
| Nr       | 27631   | 38.06          |
| SWISS-PROT | 20145   | 27.75          |
| KOG      | 16636   | 22.92          |
| GO       | 21702   | 29.90          |
| KEGG     | 6489    | 8.94           |
| Total    | 27995   | 38.6           |

*The percent of annotated unigenes in the total 72591 assembled unigene.*

doi:10.1371/journal.pone.0135724.t002
Putative structural genes involved in terpenoid backbone biosynthesis pathway

Although mono- and sesqui-terpenoids were detected in *W. ugandensis*, the structural genes involved in these biosynthetic pathways have been largely unknown so far. Analysis of the transcriptome data allowed us to identify 17 unigenes coding for four enzymes in the mevalonate pathway and 20 unigenes for five enzymes in the MEP pathway, which synthesizes the common precursor isopentenyl diphosphate (IPP) for downstream terpenoid pathways (Fig 4). Like those in other organisms, the structural genes for upstream terpenoid pathways are presented as multi-members in *W. ugandensis*. For example, six unigenes were predicted to code for hydroxymethylglutaryl-CoA reductase (HMGR), which is the first committed enzyme in the mevalonate pathway to form mevalonate (S2 Table); six unigenes were for 1-deoxy-D-xylulose-5-phosphate synthase (DXS) that catalyzes the conversion of pyruvate and D-glyceraldehyde-3-phosphate into 1-deoxy-D-xylulose-5-phosphate (DOXP) in the first step in the MEP pathway; two unigenes were for isopentenyl-diphosphate delta-isomerase (IDI), which catalyzes the isomerization of isopentenyl diphosphate to dimethylallyl diphosphate; seven unigenes were annotated to encode prenyltransferases in terpenoid pathways, of which three unigenes were for GPPSs and four unigenes for FPPSs.

Identification of putative TPSs in the *W. ugandensis* transcriptome

TPSs are the primary enzymes responsible for catalyzing the conversion of prenyl diphosphates (GPP or FPP) into monoterpenes (C10), sesquiterpenes (C15), or diterpenes (C20). The multiple diversities of terpene carbon skeletons can be largely attributed to a great number of different TPSs [15]. The PFAM motif PF01397 (N-terminal TPS domain) [31] was used to search against the assembled *W. ugandensis* unigenes. As a result, a total of 16 unigenes were found to encode putative TPS, of which 9 unigenes were predicted to contain full-length coding sequences (S3 Table). To infer their possible functions and better describe the evolutionary relationships, phylogenetic analysis of the putative TPSs protein sequences and their homologs in other plants was performed, which indicated that they were divided into three main clades.
In the first group, seven TPSs (WuSps1-7) are likely to be involved in sesquiterpene biosynthesis, which are clustered with two available W. ugandensis TPSs sequences (WargTPS-c, ACJ46047.1; WargTPS-g, ACJ46048.1) and a β-cubebene synthase from Magnolia grandiflora.

(Fig 5). In the first group, seven TPSs (WuSps1-7) are likely to be involved in sesquiterpene biosynthesis, which are clustered with two available W. ugandensis TPSs sequences (WargTPS-c, ACJ46047.1; WargTPS-g, ACJ46048.1) and a β-cubebene synthase from Magnolia grandiflora.
(ACC66281.1) in the same subclass. Seven unigenes (WuMts1-7) were found to be probably involved in monoterpene biosynthesis, and scattered into different subgroups. For example, four WuMts (WuMts1-4) are clustered with a *Litsea cubeba* thujene synthase (AEJ91555.1), while WuMts5 and WuMts7 forms a separate subgroup with several linalool- or nerolidol synthase homologs from other plants. The remaining two TPSs (WuDts1 and WuDts2) with full-length sequence were predicted to participate in diterpene biosynthesis, exhibiting closer relationships with several ent-copalyl diphosphate synthases and ent-kaurene synthases.

**Putative genes related to unsaturated fatty acids metabolism**

The high pharmaceutical effects of *W. ugandensis* were reported to be associated to its ability to biosynthesize considerable amounts of PUFAs including linoleic acid (18:2) [1, 7, 8]. To gain a better understanding of the mechanisms underlying PUFAs biosynthesis, we summarized the unigenes related to unsaturated fatty acids metabolism pathway in the database. For identification of genes in unsaturated fatty acid metabolic pathway, the assemble unigenes were searched against the KEGG pathway database. As a result, 100 genes were identified for “biosynthesis of unsaturated fatty acids” pathway (ko01040), most of which appeared to encode enzymes responsible for fatty acid desaturation and chain elongation (Table 3, S4 Table). For example, we identified nine unigenes encoding putative 3-oxoacyl-[acyl-carrier-protein] reductase (EC 1.1.1.100), which is a key enzyme participated in PUFAs biosynthesis, catalyzing the conversion of 3-hydroxyacyl-[acyl-carrier-protein] into 3-oxoacyl-[acyl-carrier-protein] with NAD⁺ or NADP⁺ as acceptor. In addition, 21 genes were found to encode putative fatty acid desaturases, of which 13 unigenes were annotated as stearoyl-CoA desaturase, the enzyme catalyzing the removal of two hydrogen atoms of stearic acid (18:0) to form oleic acid (18:1), seven unigenes were predicted to code for delta-12 fatty acid desaturase (or ω-6 fatty acid desaturase), a key enzyme in converting oleic acid (18:1) into linoleic acid (18:2), and one unigene was predicted to encode ω-3 fatty acid desaturase, the enzyme catalyzing linoleic acid (18:2) to form linolenic acid (18:3).

**Differentially expressed unigenes between *W. ugandensis* bark and leaf**

To identify the differentially expressed genes (DEGs) between the bark and leaf, the FPKM (fragments per kb per million reads) method was used to calculate the expression levels of the unigenes. The results showed that 12,704 unigenes were specifically expressed in the bark while 2,420 unigenes were specific for the leaf (S2 Fig). A total of 2,324 unigenes were expressed in both tissues but their expressions were dramatically different between the leaf and bark (with
fold changes larger than two and \( p \leq 0.05 \). Among these DEGs, 818 unigenes were up-regulated, while 1,506 were down-regulated in the \( W. \) ugandensis bark as compared to the leaf (S2 Fig and S5 Table). GO enrichment analysis of the DEGs indicated that a total of 302 GO terms were significantly enriched as compared to the genomic background (S6 Table). Additionally, 15 metabolic pathways were significantly over-represented by the KEGG pathway enrichment analysis (S7 Table). Among them, photosynthesis (ko00195) process was the most enriched pathway, and the remaining pathways were mainly related to secondary metabolites biosynthesis including monoterpenoid biosynthesis (ko00902), phenylpropanoid biosynthesis (ko00940), and novobiocin biosynthesis (ko00401), and fatty acid metabolism (cutin, suberine and wax biosynthesis, ko00073; fatty acid elongation, ko00062).
Fig 5. Phylogenetic tree of terpene synthases. Phylogenetic analysis of 14 putative *W. ugandensis* WuTPS protein sequences with their homologs from other plants indicates that they are clustered into three
main clades including: monoterpenoid synthase (WuMts), sesquiterpenoid synthase (WuSps), and diterpenoid synthase (WuDts). WarbTPS-c (ACJ46047.1, putative sesquiterpene synthase, *W. ugandensis*); WarbTPS-g (ACJ46048.1, putative sesquiterpene synthase, *W. ugandensis*); WuSp5, CL29511Contig1; WuSp4, CL3178Contig1; WuSp3, CL160Contig1; WuSp5, comp68897_c0_seq1; WuSp6, CL4969Contig1; WuSp7, CL30258Contig1; WuMts1, CL27268Contig1; WuMts2, CL27398Contig1; WuMts3, CL30385Contig1; WuMts4, CL276Contig2; WuMts5, CL1Contig9269; WuMts6, CL29966Contig1; WuMts7, CL14869Contig1; WuDts1, CL126Contig1; WuDts2, CL28648Contig1; Citsi _Germacrene_ D (XP_006494713.1, (-)-germacrene D synthase-like isoform X2, *Citrus sinensis*); Popeu _Valencene_ (XP_011015484.1, valencene synthase-like, *Populus euphratica*); Nelnu _Germacrene_ D (XP_010258444.1, (-)-germacrene D synthase-like, *Nelumbo nucifera*); Eletr _Coapane_ (ADK94034.1, alpha-coapane synthase, *Eleutherococcus trifoliatius*); Gosar _Germacrene_ D (KHGO4103.1, (-)-germacrene D synthase, *Gossypium arboreum*); Vitvi _Germacrene_ A (ADR66821.1, Germacrene A synthase, *Vitis vinifera*); Citja _Elemene_ (BAP74389.1, delta-elemene synthase, *Citrus jambhiri*); Theca _Cadinene_ (EOY12648.1, Delta-cadinene synthase isozyme A, *Theobroma cacao*); Ricco _Cubebene_ (EFJ551654.1, delta-cadinene synthase isozyme A, *Ricinus communis*); Aquisi _Guaiane_ (AIT75875.1, putative delta-guaiane synthase, *Aquilaria sinensis*); Vitvi _Linalool_ (ADR74212.1, (3S)-linalool synthase, *Vitis vinifera*); Maggr _Cubebene_ (ACC66281.1, beta-cubebene synthase, *Magnolia grandiflora*); Cinos _Linalool_ (AFK09265.1, S-(+)-linalool synthase, *Cinnamomum osmophloeum*); Nelnu _Nerolidol_ (XP_010248179.1, (3S,6E)-nerolidol synthase 1-like, *Nelumbo nucifera*); Vitvi _Nerolidol_ (XP_010646919.1, (3S,6E)-nerolidol synthase 1, *Vitis vinifera*); Vitvi _Linalool_ (ADR74212.1, (3S)-linalool synthase, *Vitis vinifera*); Actpo _Linalool_ (ADD81295.1, linalool synthase, *Actinidia polygama*); Nelnu _Ent-copalyl_ (XP_010277558.1, ent-copalydiphosphate synthase, *Nelumbo nucifera*); Theca _Ent-copalyl_ (XP_007050589.1, Copalyl diphosphate synthase, *Theobroma cacao*); Moro _Ent-copalyl_ (XP_001009049.1, En-t-copalydiphosphate synthase, *Morus notabilis*); Gosar _Ent-copalyl_ (KHGO1750.1, Ent-copalydiphosphate synthase, *Gossypium arboreum*); Nelnu _Ent-kaurene_ (XP_010260722.1, ent-kaurene synthase, *Nelumbo nucifera*); Phoda _Ent-kaurene_ (XP_008809130.1, ent-kaurene synthase, *Phoenix dactylifera*); Ricco _Ent-kaurene_ (XP_002533694.1, Ent-kaurene synthase B, chloroplast precursor, *Ricinus communis*); Popeu _Ent-kaurene_ (XP_011014299.1, ent-kaurene synthase, *Populus euphratica*); Nicta _Epi-Aristolochene_ (3M02.A, 5-Epi-Aristolochene Synthase, *Nicotiana tabacum*); Soltu _Vetispiradiene_ (Q9XJ32.1, Vetispiradiene synthase 1, *Solanum tuberosum*); Litgu _Ocimene_ (AEJ91554.1, trans-ocimene synthase, *Litsea cubeba*); Litgu _Thujene_ (AEJ91555.1, alpha-thujene synthase, *Litsea cubeba*); Citli _Limonene_ (AAM53946.1, (+)-limonene synthase 2, *Citrus limon*); Vitvi _Ocimene/Mycyne_ (ADR74206.1, (E)-beta-ocimene/myrcene synthase, *Vitis vinifera*); Quel _Pinene_ (CAKSS168.1, pinene synthase, *Quercus ilex*).

doi:10.1371/journal.pone.0135724.g005

Quantitative real-time reverse transcription PCR (qRT-PCR) analysis of DEGs

To confirm the difference in expression of unigenes detected by RNA-seq, 12 DEGs that were predicted to be coding for key enzymes involved in terpenoid and unsaturated fatty acid metabolic pathways were selected for qRT-PCR validation (S8 Table). As a result, six unigenes exhibited higher expression levels in the bark than those in the leaf, including FabF, ACOT8, ACAD, DXS, HMGGR and WuMts1; while the remaining six unigenes, such as FAD7 (a ω-3 fatty acid desaturase) and FAD6 (a ω-6 fatty acid desaturase) showed lower expression levels in the bark than those in the leaf (Fig 6). The data of qRT-PCRs mostly match the results by RNA-seq, which suggests the reliability of the transcriptome database (S8 Table).

Discussion

Pharmacological studies on *W. ugandensis* have indicated that sesquiterpenoids and PUFAs are active compounds for its medical use [2]. The bark of the plant is mostly used for its medical treatments while its leaf is also utilized in some cases [1]. These literatures drove us to compare the chemical profiles of terpenoids and PUFAs between the bark and leaf. Five sesquiterpenoids (copaene, carophyllene, farnesene, humulene, and cubebene) and two monoterpenoids (myrcene and ocimene) were detected on the condition of our GC-MS analysis. Unexpectedly, their contents are significantly higher in the leaf relative to the bark (Fig 1). On the other hand,
the total proportion of PUFAs in the leaf also showed a little higher level than that in the bark. Due to over-exploitation of the bark of the plant for its medical use, the population of the Warburgia species has considerably declined throughout their natural habitats [32]. The phytochemical study of W. ugandensis here hinted that its leaf might be also an important resource for the active compounds and could be used as a good replacement for bark, as the higher yield of the leaf holds great potential for its pharmacological applications [33].

In recent years, RNA-seq has been applied to several medicinal plants without available genomic information, such as Eugenia uniflora L. and Litsea cubeba [28, 30]. Before we lunch this project, little genomic information and very few ESTs sequences of W.ugandensis are available in public databases. Here, de novo transcriptome analysis of W. ugandensis bark and leaf was performed using the Illumina sequencing platform. This work will provide an important public genomic resource and facilitate further functional genomic studies on this plant.

Terpenoid biosynthetic pathway has been well studied, and many genes in this pathway have been identified in other plants [17, 28, 34, 35]. In this work, a total of 37 unigenes were found to be participated in terpenoid backbone biosynthesis, which showed high similarity to those homologs from other plants. For downstream terpenoid pathways, 14 putative TPSs were found in the W. ugandensis transcriptome, the number of the identified TPSs seemed to be reasonable in medicinal plants. For example, 14 TPSs in L. cubeba and 17 TPSs in T.laciniata were identified by use of de novo RNA sequencing platform [20, 30]. Phylogenetic tree analysis of the identified TPSs showed that most of them clustered with WuSps (WuSps1-6 represent sesquiterpene synthases) and WuMts (WuMts1-6 represent monoterpane synthases), which correlates well with the fact that sesquiterpenes and monoterpenes are the major terpenoids in W. ugandensis. For the pathway of PUFAs, it has been well dissected in some oil plants by RNA-seq strategy [36–38]. Fatty acid desaturation and long chain elongation are the key processes in PUFAs formation. In this study, 21 unigenes encoding putative fatty acid desaturases and seven unigenes for long chain elongases are present in the W.ugandensis database (S4 Table). Similarly, 21 unigenes coding for fatty acid desaturases were previously identified in Sacha Inchi by RNA-seq [37].

Fig 6. qRT-PCR analyses. qRT-PCR was performed to validate the 12 differentially expressed unigenes related to terpenoid biosynthesis and/or unsaturated fatty acid metabolisms. The gene names, serial numbers and primer sequences used for qRT-PCR analysis are shown in S8 Table. doi:10.1371/journal.pone.0135724.g006
Comparative transcriptome analysis showed that most of DEGs were enriched in the photosynthesis, followed by several secondary metabolism pathways, including phenylalanine metabolism and phenylpropanoid biosynthesis that provide the precursors for lignin formation. This result was consistent with the fact that photosynthesis reactions often take place in leaf, while lignin mostly accumulates in bark. In the datasets, it was found that seven out of eight WuSpSs (except for WuSpS4) had higher expression levels in the leaf than in the bark according to their FPKM values (S3 Table), which correlates well with our phytochemical results, in which higher concentrations of sesquiterpenoids were detected in the leaf than in the bark (Fig 1A). For example, both WuSpS2 and WuSpS7, exhibiting highly homologous to a beta-cube-bene synthase from M. grandiflora, were found to be expressed at significantly higher levels in the leaf than in the bark, which might result in a higher level of cube-bene accumulated in its leaf. Furthermore, the chemical profiles indicated that linoleic acid (18:2) and linolenic acid (18:3) are the major constituents of PUFAs in W. ugandensis, but their proportions are very distinct between the leaf and bark, which is probably due to the difference in the expression of fatty acid desaturases in PUFAs biosynthesis. Among the 21 putative fatty acid desaturases in the database, the expression levels of FAD6 and FAD7 were significantly higher in the leaf than in the bark. FAD6 is the enzyme to desaturate oleic acid (18:1) to form linoleic acid (18:2) while FAD7 is the enzyme to convert linoleic acid (18:2) to linolenic acid (18:3). Thus, higher transcript levels of FAD6 and FAD7 in the leaf with relatively lower levels in the bark led to a relatively lower concentration of oleic acid (18:1) and linoleic acid (18:2), and a higher level of linolenic acid (18:3) accumulated in the leaf than in the bark.

Conclusions

W. ugandensis has been used as a traditional medicine source in some African countries for a long history. Here we report the construction of a comprehensive transcriptome dataset (17.1G) derived from two different tissues (bark and leaf) by using Illumina sequencing technology. The database would accelerate functional genomic research on this plant. In particular, many candidate genes for structural enzymes in terpenoids and PUFAs biosynthesis were identified in this study, which would help people to understand the biosynthesis of terpenoids and PUFAs in W. ugandensis.

Methods

RNA extraction, cDNA library construction and sequencing

Warburgia ugandensis Sprague materials (bark and mature leaves) were collected from plants grown in the greenhouse of Wuhan Botanical Garden, Chinese Academy of Sciences, and immediately frozen in liquid nitrogen and stored at −80°C until use. Total RNA was extracted using the EASYspin plant RNA extraction kit (Aidlab Biotechnologies, Co., Ltd, China). RNA integrity and quantity were evaluated by the electrophoresis on 1% agarose gel and the detection on Agilent 2100 Bioanalyzer (S4 Fig). Equal amounts (about 5 μg) of RNA from three independent biological repeats for each sample were mixed together and then send to Shanghai Oebiotech Corporation (Shanghai, China) for the sequencing by Illumina HiSeq2000 platform. The pair-end method was applied in this work, and the read length was 100 bp for each read. The quality of raw data was evaluated using the FastQC software (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/), and low quality reads were discarded by use of NGS QC Toolkit (v2.3.3) software (http://59.163.192.90:8080/ngsqctoolkit/). First, removing all low quality reads with phred score less than 20 and read length shorter than 35 bp. Second, trimming bases having phred quality score less than 20 at 3’ end of the read. Third, trimming the sequences containing ambiguous ‘N’ bases, and reads shorter than 35 bp would be
discarded. Then, the clean reads were assembled into contigs or unigenes using Trinity software (version: trinityrnaseq_r20131110) with the specific parameter set to “—seqTypefq—min_contig_length 200—JM 400G” and all other parameters set to default [39]. The raw transcriptome data was available at the sequence read archive (SRA) of NCBI with the accession number SRX970743.

Functional annotation

The function of the unigenes was annotated by alignment of the unigenes with the NCBI non-redundant (Nr), SwissProt, and KOG (http://www.ncbi.nlm.nih.gov/COG) databases using Blastx with a threshold \( E \) value of \( 10^{-5} \). The proteins with the highest hits to the unigenes were used to assign functional annotations thereto. Gene ontology (GO) classification was performed by the BLAST2GO program [40]. The unigenes were mapped to the Kyoto Encyclopedia of Genes and Genomes (KEGG) database (http://www.genome.jp/kegg/) to annotate their potential metabolic pathways.

Identification of terpene synthases in the *W. ugandensis* transcriptome

The terpene synthase N-terminal motif (pfam01397) was downloaded from the Pfam database (http://pfam.sanger.ac.uk/), and used as a bait to search against the *W. ugandensis* transcriptome. TPS domain sequences containing the pfam01397 motif were obtained using BLASTX with an \( E \)-value threshold of \( 10^{-5} \), and the full length coding unigenes of TPS were predicted by ORF finder on line (http://www.ncbi.nlm.nih.gov/gorf/gorf.html). Several homologous TPS sequences in other plants [28, 30] including two *W. ugandensis* sesquiterpene synthase sequences (ACJ46047.1, ACJ46048.1), were downloaded from the Nr protein database and used as references to perform a phylogenetic analysis with our identified TPS. The phylogenetic tree was constructed by MEGA 5.0 using neighbor-joining method with 1000 bootstrap trials, and other specific parameters were set by default.

Differential expression analysis of unigenes

The FPKM (fragments per kb per million reads) method was used to calculate the expression abundance of the unigenes [41]. The fold-changes of each gene expression between leaf and stem bark were evaluated by their ratio of the FPKMs. The DEGs were determined by the R package DESeq (http://bioconductor.org/packages/release/bioc/html/DESeq.html), which provides methods to test the differential expression by use of the negative binomial distribution and a shrinkage estimator for the distribution’s variance, with setting the threshold of FDR (false discovery rate) \( \leq 0.05 \) and \( \log2\text{Ratio} \geq 1 \) [42]. Using a hypergeometric test (with \( p \) value \( < 0.05 \)), the significantly enriched GO terms in DEGs were analyzed by comparing gene numbers for each certain GO term with the genome background of *W. ugandensis*. DEGs were also used in pathway enrichment analysis by mapping all the DEGs to KEGG database, the number of genes in each pathway was calculated and compared to the whole genome background by the threshold of \( p \) value \( < 0.05 \).

GC-MS analysis

About 0.3 g fresh weight of *W. ugandensis* stem barks or leaves were collected, and immediately powdered in liquid nitrogen frozen. The plant tissues were extracted in 500 \( \mu \)l hexane by ultrasonification for 30 min. The crude extracts were centrifuged at 12,000 g for 10 min, then 2 \( \mu \)l of the supernatant was analyzed on an Agilent 7890A GC machine (Agilent Technologies, Waldbronn, USA) and an Agilent 5975C mass selective detector. Separation was performed on an
Agilent HP-5MS column (0.25 mm × 30 m × 0.25 μm), and the carrier gas was helium with a flow rate of 1.2 ml/min. The initial GC oven temperature was set at 50°C for 2 min, followed by a 5°C /min ramp to 120°C and a 30°C /min ramp to 240°C, and held at 240°C for 5 min. Compounds were determined by mass spectral comparison with the chemical database installed in the GC machine.

qRT-PCR analysis

After treating with DNase I (Thermo, USA) to remove residual genomic DNA, 2 μg of total RNA was converted to the first-strand cDNA by use of SuperScript III reverse-transcriptase (Invitrogen, USA). The cDNA products were then diluted in 100-folds with deionized water prior to PCRs. The qRT-PCR reaction was performed on an ABI 7500 Real-Time PCR Detection System using a FastStart Universal SYBR Green Master Mix (Roche). The thermal cycling conditions were as follows: 95°C for 10 min for one cycle; at 95°C for 15 s, then at 60°C for 1 min for 40 cycles. Three independent biological replicates were performed for each sample. The relative mRNA levels were normalized with respect to inner control genes (actin) and expressed relative to the corresponding values of leaf (control), which were given an arbitrary value of 1. Primer pairs were designed using Primer 5 software, and the primer sequences were available on-line (S8 Table).

Supporting Information

S1 Fig. Length distribution of unigenes in the *W. ugandensis* transcriptome.
(TIF)

S2 Fig. Summary of differentially expressed genes between the bark and leaf. A: the number of the unigenes that are exclusively expressed in the bark or leaf, and the unigenes that are expressed in both tissues; B: the number of the differentially expressed genes (DEGs).
(TIF)

S3 Fig. Putative structural genes involved in unsaturated fatty acid biosynthesis pathway. The values in the bracket indicate the number of unigenes in the corresponding gene families. KAS, ketoacyl-ACP synthase; KAR, ketoacyl-ACP reductase; HAD, hydroxyacyl-ACP dehydrase; EAR, enoyl-ACP reductase; SAD, stearoyl-ACP desaturase; FAD6, oleate desaturase; FAD7, linoleate desaturase; ACOT, acyl-ACP thioesterase.
(TIF)

S4 Fig. The RNA integrity and quantity were evaluated by the electrophoresis on 1% agarose gel (A), and the detection of leaf (B) and bark (C) RNA by using Agilent 2100 Bioanalyzer.
(TIF)

S1 Table. KEGG pathway annotation of unigenes.
(XLSX)

S2 Table. Putative structural genes involved in terpenoid backbone biosynthesis pathway.
(XLSX)

S3 Table. The unigenes coding for putative terpene synthases in terpenoid biosynthesis.
(XLSX)

S4 Table. Unigenes annotated to be involved in unsaturated fatty acids metabolism.
(XLSX)
S5 Table. List of DEGs between *W. ugandensis* stem bark and leaf. (XLSX)

S6 Table. GO enrichment analysis of DGEs. (XLSX)

S7 Table. KEGG pathway enrichment analysis of DGEs. (XLSX)

S8 Table. qRT-PCR primers used in this study. (XLSX)

Acknowledgments
We thank Dr. Qingfeng Wang for identification *Warburgia ugandensis* plants in this study.

Author Contributions
Conceived and designed the experiments: YZ. Performed the experiments: XW CZ. Analyzed the data: XW. Contributed reagents/materials/analysis tools: XY DM. Wrote the paper: YZ XW.

References
1. Maroyi A. The genus *Warburgia*: A review of its traditional uses and pharmacology. Pharm Biol. 2014; 52:378–391. doi:10.3109/13880209.2013.837935 PMID: 24188508
2. Drage S, Mitter B, Trols C, Muchugi A, Jamnadass RH, Sessitsch A, et al. Antimicrobial drimane sesquiterpenes and their effect on endophyte communities in the medical tree *Warburgia ugandensis*. Front Microbiol. 2014; 5:13.
3. Kuglerova M, Tesarova H, Grade JT, Halamova K, Wanyana-Maganyi O, Van Damme P, et al. Antimicrobial and antioxidative effects of Ugandan medicinal barks. Afr J Biotechnol. 2011; 10:3628–3632.
4. Ngure PK, Tonui WK, Ingonga J, Mutai C, Kigondu E, Ng’ang’a Z, et al. *In vitro* antileishmanial activity of extracts of *Warburgia ugandensis* (Canellaceae), a Kenyan medicinal plant. J Med Plants Res. 2009; 3:61–66.
5. Wube AA, Bucar F, Gibbons SZ, Asres K, Rattray L, Croft SL. Antiprotozoal activity of sesquiterpenes from *Warburgia ugandensis* towards *Trypanosoma brucei rhodesiens* and *Plasmodium falciparum in vitro*. Planta Med. 2008; 74:994–994.
6. Kubo I. Antifungal sesquiterpene dialdehydes from the *Warburgia* plants and their synergists. In: Atta ur R, editors. Studies in natural products chemistry. Part D: Elsevier; 1995. pp. 233–249.
7. Wube AA, Bucar F, Gibbons S, Asres K. Sesquiterpenes from *Warburgia ugandensis* and their antimycobacterial activity. Phytochemistry. 2005; 66:2309–2315. PMID: 16153670
8. Kiyo D, Gray AI, Waterman PG. A comparative study of the stem-bark drimane sesquiterpenes and leaf volatile oils of *Warburgia ugandensis* and *W. Stuhlmannii*. Phytochemistry. 1990; 29:3535–3538.
9. Xu M, Litaudon M, Krief S, Martin MT, Kasenene J, Kiremire B, et al. A new drimane-type sesquiterpenoid from *Warburgia ugandensis*. Molecules. 2009; 14:3844–3850. doi:10.3390/molecules14103844 PMID: 19924033
10. Rajab MS, Ndegewa JM. 11 alpha-hydroxy muzigadiolide, a novel drimane sesquiterpene from the stem bark of *Warburgia ugandensis*. B Chem Soc Ethiopia. 2000; 14:45–49.
11. Brooks CJW, Draffan GH. Sesquiterpenoids of *Warburgia* species—I: *Warburgin* and *warburgiadione*. Tetrahedron. 1969; 25:2865–2885.
12. Opiyo SA, Manguro LOA, Okinda-Owuor P, Ateka EM, Lemmen P. 7 alpha-Acetylugandensolide and antimicrobial properties of *Warburgia ugandensis* extracts and isolates against sweet potato pathogens. Phytochem Lett. 2011; 4:161–165.
13. Van Wyk BE. The potential of South African plants in the development of new medicinal products. S Afr J Bot. 2011; 77:812–829.
Sando T, Takaoka C, Mukai Y, Yamashita A, Hattori M, Ogasawara N, et al. Cloning and characterization of monoterpene, sesquiterpene and diterpene-codon code status and future opportunities. Plant Biotechnol J. 2013; 11:169–196. doi: 10.1111/pbi.12022 PMID: 23171352

15. Tholl D. Terpene synthases and the regulation, diversity and biological roles of terpene metabolism. Curr Opin Plant Biol. 2006; 9:297–304. PMID: 16600670

16. Degenhardt J, Kollner TG, Gershenzon J. Monoterpene and sesquiterpene synthases and the origin of terpene skeletal diversity in plants. Phytochemistry. 2009; 70:1621–1637. doi: 10.1016/j.phytochem.2009.07.030 PMID: 19793600

17. Chen H, Li G, Kollner TG, Jia QD, Gershenzon J, Chen F. Positive Darwinian selection is a driving force for the diversification of terpenoid biosynthesis in the genus Oryza. BMC Plant Biol. 2014; 14:239. doi: 10.1186/s12870-014-0239-x PMID: 25224158

18. Cheng AX, Xiang CY, Li JX, Yang CQ, Hu WL, Wang LJ, et al. The rice (E)-beta-caryophyllene synthase (OsTPS3) accounts for the major inducible volatile sesquiterpenes. Phytochemistry. 2007; 68:1632–1641. PMID: 17524436

19. Aubourg S, Lechamay A, Bohlmann J. Genomic analysis of the terpenoid synthase (AtTPS) gene family of Arabidopsis thaliana. Mol Genet Genomics. 2002; 267:730–745. PMID: 12207221

20. Drew DP, Dueholm B, Weitzel C, Zhang Y, Sensen CW, Simonsen HT. Transcriptome analysis of Thapsia laciniata Rouy provides insights into terpenoid biosynthesis and diversity in Apiaceae. Int J Mol Sci. 2013; 14:9080–9098. doi: 10.3390/ijms14059080 PMID: 23897865

21. Martin DM, Aubourg S, Schouwey MB, Daviet L, Schalk M, Toub O, et al. Functional Annotation, Genome Organization and Phylogeny of the Grapevine (Vitis vinifera) Terpene Synthase Gene Family Based on Genome Assembly, FLcDNA Cloning, and Enzyme Assays. BMC Plant Biol. 2010; 10:226. doi: 10.1186/1471-2229-10-226 PMID: 20964856

22. Murray M. Omega-3 polyunsaturated fatty acids and their metabolites as inhibitors of mammalian tumorigenesis. Phytochem Rev. 2014; 13:139–156.

23. Warude D, Joshi K, Harsulkar A. Polyunsaturated fatty acids: Biotechnology. Crit Rev Biotechnol. 2006; 26:83–93. PMID: 16809099

24. Ohriroge JB, Jaworski JG. Regulation of fatty acid synthesis. Ann Rev Plant Physiol Mol Biol. 1997; 48:109–136.

25. Truksa M, Vrinten P, Oiu X. Metabolic engineering of plants for polyunsaturated fatty acid production. Mol Breeding. 2009; 23:1–11.

26. Ruiz-Lopez N, Syanova O, Napier JA, Haslam RP. Metabolic engineering of the omega-3 long chain polyunsaturated fatty acid biosynthetic pathway into transgenic plants. J Exp Bot. 2012; 63:2397–2410. doi: 10.1093/jxb/err454 PMID: 22911311

27. Xiao M, Zhang Y, Chen X, Lee EJ, Barber CJ, Chakrabarty R, et al. Transcriptome analysis based on next-generation sequencing of non-model plants producing specialized metabolites of biotechnological interest. J biotechnol. 2013; 166:122–134. doi: 10.1016/j.jbiotec.2013.04.004 PMID: 23602801

28. Guzman F, Kulcheski FR, Turchetto-Zolet AC, Margis R. De novo assembly of Eugenia uniflora L. transcriptome and identification of genes from the terpenoid biosynthesis pathway. Plant Sci. 2014; 229:238–246. doi: 10.1016/j.plantsci.2014.10.003 PMID: 25443850

29. Yang L, Ding G, Lin H, Cheng H, Kong Y, Wei Y, et al. Transcriptome analysis of medicinal plant Salvia miltiorrhiza and identification of genes related to terpenoid biosynthesis. PLoS One. 2013; 8:10.1371

30. Han XJ, Wang YD, Chen YC, Lin LY, Wu QK. Transcriptome sequencing and expression analysis of terpenoid biosynthesis genes in Litsa cubeba. PLoS One. 2013; 8:e76890. doi: 10.1371/journal.pone.0076890 PMID: 24130803

31. Finn RD, Tate J, Mistry J, Coggill PC, Sammut SJ, Hotz HR, et al. The Pfam protein families database. Nucleic Acids Res. 2008; 36:D281–288. PMID: 18939703

32. Botha J, Witkowski ETF, Shackleton CM. The impact of commercial harvesting on Warburgia salutaris (‘pepper-bark tree’) in Mpumalanga, South Africa. Biodivers Conserv. 2004; 13:1675–1688.

33. Drewes SE, Crouch NR, Mashimbye MJ, de Leeuw BM, Horn MM. A phytochemical basis for the potential use of Warburgia salutaris (pepper-bark tree) leaves in the place of bark. S Afr J Sci. 2001; 97:383–386.

34. Ma Y, Yuan L, Wu B, Li X, Chen S, Lu S. Genome-wide identification and characterization of novel genes involved in terpenoid biosynthesis in Salvia miltiorrhiza. J Exp Bot. 2012; 63:2809–2823. doi: 10.1093/jxb/err466 PMID: 22291132

35. Sando T, Takaoka C, Mukai Y, Yamashita A, Hattori M, Ogasawara N, et al. Cloning and characterization of mevalonate pathway genes in a natural rubber producing plant, Hevea brasiliensis. Biosci Biotechnol Biochem. 2008; 72:2049–2060. PMID: 18685207
36. Dussert S, Guerin C, Andersson M, Joet T, Tranbarger TJ, Pizot M, et al. Comparative transcriptome analysis of three oil palm fruit and seed tissues that differ in oil content and fatty acid composition. Plant Physiol. 2013; 162:1337–1358. doi:10.1104/pp.113.220525 PMID: 23735505

37. Wang XJ, Xu RH, Wang RL, Liu AZ. Transcriptome analysis of Sacha Inchi (Plukenetia volubilis L.) seeds at two developmental stages. BMC Genomics. 2012; 13:716. doi:10.1186/1471-2164-13-716 PMID: 23256450

38. Troncoso-Ponce MA, Kilaru A, Cao X, Durrett TP, Fan JL, Jensen JK, et al. Comparative deep transcriptional profiling of four developing oilseeds. Plant J. 2011; 68:1014–1027. doi:10.1111/j.1365-313X.2011.04751.x PMID: 21851431

39. Grabherr MG, Haas BJ, Yassour M, Levin JZ, Thompson DA, Amit I, et al. Full-length transcriptome assembly from RNA-Seq data without a reference genome. Nat Biotechnol. 2011; 29:599–600.

40. Gotz S, Garcia-Gomez JM, Terol J, Williams TD, Nagaraj SH, Nueda MJ, et al. High-throughput functional annotation and data mining with the Blast2GO suite. Nucleic Acids Res. 2008; 36:3420–3435. doi:10.1093/nar/gkn176 PMID: 18445632

41. Mortazavi A, Williams BA, Mccue K, Schaeffer L, Wold B. Mapping and quantifying mammalian transcriptomes by RNA-Seq. Nat methods. 2008; 5:621–628. doi:10.1038/nmeth.1226 PMID: 18516045

42. Reiner A, Yekutieli D, Benjamini Y. Identifying differentially expressed genes using false discovery rate controlling procedures. Bioinformatics. 2003; 19:368–375. PMID: 12584122