Chromosome-Level Genome Assembly Provides New Insights into Genome Evolution and Tuberous Root Formation of Potentilla anserina

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Abstract: Potentilla anserina is a perennial stoloniferous plant with edible tuberous roots in Rosaceae, served as important food and medicine sources for Tibetans in the Qinghai-Tibetan Plateau (QTP), China, over thousands of years. However, a lack of genome information hindered the genetic study. Here, we presented a chromosome-level genome assembly using single-molecule long-reading sequencing, and the Hi-C technique. The assembled genome was 454.28 Mb, containing 14 chromosomes, with contig N50 of 2.14 Mb. A total of 46,495 protein-coding genes, 169.74 Mb repeat regions, and 31.76 Kb non-coding RNA were predicted. P. anserina diverged from Potentilla micrantha ~28.52 million years ago (Mya). Furthermore, P. anserina underwent a recent tetraploidization ~6.4 Mya. The species-specific genes were enriched in Starch and sucrose metabolism and Galactose metabolism pathways. We identified the sub-genome structures of P. anserina, with A sub-genome was larger than B sub-genome and closer to P. micrantha phylogenetically. Despite lacking significant genome-wide expression dominance, the A sub-genome had higher homoeologous gene expression in shoot apical meristem, flower and tuberous root. The resistance genes was contracted in genome-wide expression dominance, the A sub-genome had higher homoeologous gene expression in shoot apical meristem, flower and tuberous root. The resistance genes was contracted in A sub-genome, which probably drives the tuber formation. The genomics and transcriptomics data generated in this study advance our understanding of the genomic landscape of P. anserina, and will accelerate genetic studies and breeding programs.

Keywords: Potentilla anserina; allopolyploids; genome assembly; gene family; sub-genome expression dominance; starch metabolism

1. Introduction

P. anserina (Poan) is a stoloniferous herb in Rosaceae which is widely distributed in the western regions of China, particularly in the Qinghai-Tibetan Plateau (QTP) [1]. It can produce different levels of stolon with roots and leaves at nodes during the growing season (Supplementary Figure S1). By virtue of fast clonal growth of the stolon and robust root system, Poan had become one of the dominant species in the alpine meadow. The aerial parts of Poan are considered as excellent forage for wild animals and livestock. The roots of Poan usually swell and form edible tubers in QTP [2]. It has been served as a tonic food and folk medicine for ages [3,4]. The tuberous root is rich in starch [4], amino acids [3], tannins [5], flavonoids [6], and triterpenes [7], and is a crucial plant-based...
food source for Tibetan people. Quite a few pharmacological studies have proved the medicinal functions of the extracts and constituents from Poan, including hepatoprotective [1], immunomodulatory [8], antitussive [4], and expectorant [4] effects. These research results confirmed that Poan possesses essential medicinal application value and has huge industrialization potential.

The Potentilla genus belongs to the Potentilleae tribe, Rosaceae family, including Anserina and other five major clades, which have been confirmed by both morphological and molecular data [9–11]. Up to now, over 300 species and numerous naturally occurring hybrids were recognized, showing an intricate relationship within Potentilla genus [11]. However, only one species in this genus was sequenced and assembled [12], with a contig-level genome assembly, which is far from enough to allow the phylogenetic study of such a diverse and important genus.

The lack of genomic information undoubtedly limited the molecular genetic study of Poan. Here, we reported a chromosome-level genome assembly of Poan (2n = 4x = 28), combining PacBio long reads, Illumina paired-end (PE) reads and high-resolution Hi-C data. Based on the reference genome, comparative genomics and phylogenetic analysis among Poan and its relatives was carried out. Moreover, transcriptomic analysis among different tissues were utilized for gene annotation and investigation of expression pattern of key genes involved in starch metabolism. All the data generated in this study will serve as a valuable resource for unraveling the molecular mechanism behind special traits of Poan and elucidating the evolution of Potentilla genus.

2. Materials and Methods

2.1. Plant Materials and High-Throughput Sequencing

Poan was collected in Guolog Tibetan Autonomous Prefecture, Qinghai province, China. The specimen used for sequencing were cultured in the greenhouse of the key laboratory of molecular breeding in the Northwest Institute of Plateau Biology, Chinese Academy of Sciences (CAS). High-quality genomic DNA was extracted from fresh young leaves using the Qiagen DNeasy Plant Mini Kit (Qiagen, Germantown, MD, USA). The quantity and quality of extracted DNA were examined with NanoDrop 2000 spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA) and electrophoresis on 1% agarose gel, respectively.

Above all, PE reads with an insert size of 350 bp were generated using the NEB Next Ultra DNA Library Prep Kit according to the manufacturer’s instructions and further sequenced on Illumina HiSeq X Ten platform. For the long-reads library, 20 Kb SMRTbell libraries were constructed according to the BluePippin Size Selection System protocol as described by Pacific Biosciences (Pacific Biosciences, CA, USA) and then sequenced on PacBio sequel II platform with 5 SMRT cells. For Hi-C sequencing, one Hi-C library was constructed under the following procedure: chromatin was fixed with fresh formaldehyde to obtain the crosslink of DNA and protein. The crosslink was digested with Mbo I restriction endonuclease and 5’ overhangs were filled in with biotinylated nucleotides. After fixation, the adjacent fragments were ligated. Then, crosslinks were unlocked, and the DNA was extracted and sheared to a mean fragment size of 350 bp. Fragments with the biotin tag were captured using streptavidin beads and sequenced on the Illumina HiSeq X Ten platform.

In order to ensure the accuracy of gene annotation, the normal roots, tuberous roots, leaf, stem, shoot apical meristem (SAM), and flower of the Poan plant were collected for RNA-seq library construction. Then, all tissues mentioned above from another Poan plant were harvested for Iso-seq library construction. The RNA-seq libraries were generated using the NEB Next Ultra RNA Library Prep Kit for Illumina and sequenced on Illumina HiSeq 4000 to produce PE reads. The Iso-seq libraries were constructed and sequenced as the long-reads sequencing procedure mentioned above. To investigate the karyotype of Poan, we selected the stem apex tissue of Poan. The stem apex was treated with 0.1% colchicine for 4 h, then fixed for 20 to 24 h with Carnoy Fixative (acetic acid: absolute
alcohol = 1:3). After that, the specimen was dissociated using 1 mol/L hydrochloric acid in water bath at 60 °C for 10 min. Finally, the specimen was dyed with nucleic acid dye (DAPI) for 10 min. The karyotype of Poan was scanned by a OLYMPUS BX63 automated fluorescence microscope under 100 × field of view.

2.2. Estimation of Genome Size and Genome Assembly

The qualified Illumina PE reads were used to estimate the genome size and heterozygosity through k-mer analysis. We adopted kmer_freq (https://github.com/fanagislab/kmerfreq (accessed on 10 May 2021)) and GCE (https://github.com/fanagislab/GCE (accessed on 10 May 2021)) for 17-mer analysis. The genome size was calculated according to the equation: genome size = k-mer_number/peak_depth. The ploidy of Poan was confirmed using Smudgeplot [13].

De novo genome assembly was performed using a FALCON (v0.2.2) assembler [14]. Firstly, the qualified PacBio subreads were self-corrected through a pre-assembled sequence. Then, overlaps were detected and filtrated for graph construction. Finally, contigs were generated using the graph. The initial contigs were aligned and polished with pbalign and Arrow programs as implemented in SMRT Link (v8) software (https://www.pacb.com/support/software-downloads/ (accessed on 8 April 2021)) with PacBio subreads. The refined contigs were further polished for two rounds using Pilon (v1.24) [15] (–fix snps, indels) based on Illumina PE reads, generating the draft genome assembly of Poan.

The qualified Hi-C reads were mapped to the draft genome assembly. We employed Hi-C pro (v3) [16] to obtain the valid pairs after duplication removal, sorting, and quality assessment. Subsequently, the valid Hi-C reads together with the draft genome assembly were processed following the 3D-DNA (20150322) pipeline [17] (-r 2 -i 2000). Finally, we obtained an explicit cross-linked pattern of sequences, which exhibited chromosomal arrangement. The cross-linked maps were visualized and manually checked using Juicebox (v1.11.08) [18]. To evaluate the assembled genome quality and continuity, we used three different methods. First, short PE reads were mapped back to the assembly genome using BWA [19] with the default parameters. Then, BUSCO (v5) [20] assessment was carried out (eudicots_odb10, --offline). Finally, LTR Assembly Index (LAI) [21] scores were calculated by LTR_Retriever (v2.8) [22] with the default parameters.

2.3. Non-Coding RNA and Repeat Identification

To identify the non-coding RNAs in the genome, tRNAscan-SE (v2.0.8) [23] and BLAST (v2.9.0+) [24] were used to detect the tNRA and rRNA, respectively. Meanwhile, the miRNA and snRNA were obtained using Infernal (v1.1.4) [25] based on the Rfam database [26]. The transposable element (TE) of genome were annotated in two ways including an ab initio method and homology-based search. RepeatModeler (v1.0.11) [27] and LTR_FINDER (v1.07) [28] with the default parameters were used for ab initio TE identification. For homology-based search, RepeatMasker (v4.1.0) (http://www.repeatmasker.org (accessed on 1 June 2021)) with default parameter was used to find the TE against the Repbase [29]. Furthermore, full-length LTR-RTs were collected for insertion time analysis using in-house Perl scripts.

2.4. Gene Prediction and Functional Annotation

Protein coding genes of the Poan genome were predicted combining three approaches, including ab initio prediction, homology-based alignment and transcriptome-based prediction. The gene prediction were performed on the basis of repeat-masked genome assembly. For ab initio prediction, we adopted AUGUSTUS (v3.3.3) [30] (–species = poan) and SNAP [31] (default parameter) to train and predict gene structure. For homology-based prediction, protein sequences from Arabidopsis thaliana, Oryza sativa, Malus domestica, and Fragaria vesca were downloaded from Phytozome database [32] and aligned to Poan genome sequences using tblastn [24] (expect (E) value cutoff: 1 × 10^{-5}). Blast hits were filtered and integrated by in-house Perl scripts. Next, the sequences in Poan genome corresponding
to targeted protein were extended upstream and downstream by 2000 bp to represent a protein-coding region. Finally, GeneWise (v2.4-1) [33] with -both -gff -quiet -silent -sum parameters was used to identify the exon and intron boundary. For transcriptome-based predictions, we build a comprehensive transcriptome database using genome-guided and de novo methods according to PASA pipeline (v2.4.1) with default parameter (https://github.com/PASApipeline/PASApipeline (accessed on 8 August 2021)). The Illumina PE reads were assembled under genome-guided mode using Trinity (v2.10.0) [34] and the PacBio long-reads were polished and clustered into full length transcripts through IsoSeq (v3) pipeline with default parameter (https://github.com/PacificBiosciences/IsoSeq/ (accessed on 8 August 2021)). At last, all evidences from three methods were integrated into a complete gene set without redundancy by EVidenceModeler (v1.1.1) with the default parameters [35].

Functional annotation of the predicted genes was actualized with BLASTP against various protein databases, including Pfam [36], Swiss-Prot (https://www.ebi.ac.uk/uniprot/ (accessed on 15 August 2021)), KEGG (https://www.genome.jp/kegg/ (accessed on 15 August 2021)), TrEMBL (https://www.ebi.ac.uk/uniprot/ (accessed on 15 August 2021)), and eggNOG [37].

2.5. Comparative Genomics and Phylogenetic Analysis

The nucleotide and amino acid sequences of seven Rosaceae species including: Fragaria vesca (Frve), Malus domestica (Mado), Potentilla micrantha (Pomi), Prunus persica (Prpe), Pyrus communis (Pyco), Rosa chinensis (Roch), and Rubus occidentalis (Ruoc) were downloaded from the GDR database [38], while the sequence of Arabidopsis thaliana (Arth) was downloaded from the Phytozome database [32]. We used OrthoFinder (v2.5.4) [39] with default parameters to identify orthologous genes and gene family membership among nine species with default parameters. The species-specific gene families and expanded families were annotated and enriched using the topGO package (https://bioconductor.org/packages/release/bioc/html/topGO.html (accessed on 6 September 2021)) and KOBAS (v3) [40] based on the GO and KEGG databases.

For phylogenetic analysis, 286 single-copy genes were selected and aligned using MUSCLE (v3.8.31) with default parameters [41]. Subsequently, the aligned sequences were concatenated to generate a super-gene for each species using an in-house script. At last, we extracted the nucleotides at position 2 (phase 1) of each codon to construct phylogenetic tree using RAxML (v8.2.12) [42]. After maximum likelihood (ML) tree searching, one optimized ML tree were sequentially constructed with -m PROTGAMMAJTTF and bootstrap value of 1000, with Arth as the outgroup. Furthermore, in order to investigate the speciation and divergence time of these species, we used the MCMCTree program with the default control file in PAML (v4.10.0) [43] to construct the time tree based on the ML tree. The calibrated timescales were obtained from TimeTree website [44]: Pyco–Mado (6–43 Mya), Arth–Mado (98–117 Mya). The gene synteny and gene duplication analysis was performed using the JCVI pipeline (https://github.com/tanghaibao/jcvi (accessed on 20 September 2021)).

2.6. Whole Genome Duplication (WGD) Analysis

To analyze the WGD event among Poan, Pomi and Frve, we characterized homologous genes using BLASTP (E-value cutoff: $1 \times 10^{-5}$). Then by running MCScanX [45] with default parameters, we identified collinear blocks that included at least thirty collinear gene pairs. Finally, we calculated the Ks values of paralogous and orthologous gene pairs using PAML (v4.10.0) with default parameters, and depicted the distribution plot using R software. The WGD time was calculated according to the formula $\text{Time} = \frac{\text{Ks}}{2r}$ ($r = 7 \times 10^{-9}$ mutations per site per year).

2.7. Sub-Genome Analysis and Expression Bias of Homeologs

To distinguish the genome structure of Poan, three different methods were combined. First, we investigated gene synteny within the Poan genome by JCVI, finding the existence
of sub-genome structure. Then, we compared gene synteny between Pomi and Poan, confirming the order of sub-genomes. Finally, the homoeologous gene pairs (1:1) between sub-genome of Poan were obtained using Orthofinder. We further constructed a phylogenetic tree for each homoeologous gene pairs. The sub-genome with closer genetic distance to Pomi was denoted as the A sub-genome, and the remainder of the sub-genome was denoted as the B sub-genome. These gene pairs within the sub-genome were defined as homoeologous genes which were derived from latest WGD.

The homoeologous gene pairs were further used for expression analysis. We calculated the TPM value of homoeologs in tuberous root, normal root, stem, SAM, leaf, and flower. Log2(A/B) > 0 indicated the biased expression of A sub-genome homoeologs, and Log2(A/B) < 0 indicated the biased expression of B sub-genome homoeologs.

2.8. Identification of Resistance Genes and Starch Biosynthesis Related Genes

The conserved domains were collected from the Pfam database [36]. R genes were identified using HMMER (v3.3.1) [46] (E-value cutoff: $1 \times 10^{-10}$). The protein sequences of key genes involved in starch biosynthesis were collected from the NCBI gene bank (https://www.ncbi.nlm.nih.gov/ (accessed on 12 October 2021)). All the target sequences were aligned with Poan, Pomi, Frve, Roch, Ruoc gene set using BLASTP (E-value cutoff: $1 \times 10^{-10}$). After elimination of redundancy (identity less than 35% and coverage less than 60%), we confirmed the copy number of key genes in each species. For transcriptome analysis, low-quality reads and adaptors were filtered and trimmed using Trimmomatic (v0.39) [47]. Qualified reads were aligned to the genome assembly by HISAT2 (v2.1.0) [48]. Gene expression was measured as TPM using the featureCounts program (v2.0.1) [49] and R scripts. The expression level of key genes was displayed through heatmap package (clustering_method = ‘ward.D’, scale = “row”).

3. Results
3.1. Genome Sequencing and Assembly

Young fresh leaves from the Poan plant were collected for de novo genome sequencing. To generate a high-quality genome assembly, we adopted hybrid methods including deep PacBio long-read sequencing, Illumina short-read sequencing, and Hi-C chromosome conformation capture. A total of 49.87 Gb qualified PacBio long-reads (~108×), with reads N50 of 20.26 Kb, 44.45 Gb clean PE reads (~96×), and 47.04 Gb clean Hi-C data (~102×) were obtained (Supplementary Table S1).

We first investigated genome size using Illumina paired-end (PE) reads. The k-mer (k = 17) depth and frequency results showed that the Poan genome was 459.1 Mb with a heterozygosity rate of 0.7% (Supplementary Table S2). In addition, the sharp peak (82) in the k-mer frequency curve was expected from the homozygous region of genome, and the smooth peak (168) indicated the allotetraploidy of Poan [50,51] (Figure 1). The ploidy was further analyzed using the reported method for reference-free profiling of polyploid genomes [13], suggesting that the possibility of AABB was 0.48 (Figure 1), which confirmed the allotetraploidy of Poan. In addition, karyotype analysis showed that the sequenced Poan genome contained 28 chromosomes (Supplementary Figure S2). Considering the basic chromosome number in Rosaceae species was seven; therefore, Poan was an allotetraploid plant (2n = 4x = 28).

For genome assembling, all 49.93 Gb long-reads were self-corrected and assembled to a 487.44 Mb draft graph containing 1126 contigs, with a contig N50 value of 2.13 Mb. Eventually, the refined contigs were reordered and anchored into 14 chromosomes using Hi-C data (Supplementary Figure S3). The final assembly was 454.28 Mb, covering 98.95% of predicted genome size.
3.2. Assessment of Genome Quality

In order to assess the quality of genome assembly, we adopted different methods. The Illumina PE reads were mapped back to the genome with a 99.49% mapping ratio, 94.39% of which was properly paired (Supplementary Table S3). We also counted coverage depth of each base and found that, of the mapped base pair, 98.86% had more than five depths. Then, BUSCO evaluation showed that 2288 (98.3%) completed BUSCOs and 7 (0.3%) fragmented BUSCOs were detected in the Poan genome (Supplementary Table S4), respectively, indicating a high completeness. Furthermore, LTR are the predominant type of transposable element (TE), which is poorly assembled in draft genomes. Ou et al. [21] proposed a metric called LTR Assembly Index (LAI) to evaluate assembly continuity. We therefore calculated the LAI scores of Poan 14 chromosomes. The average LAI score of Poan genome was 17.81, belonging to reference genome level. The highest median chromosome LAI score was 20.36 (chromosome 8), and the lowest median was 16.22 (chromosome 4) (Figure 2a). All these results implied the high integrity and continuity of the assembled Poan genome.
3.3. Genome Annotation

To obtain a high-quality gene set, we combined three approaches, including transcriptomic evidence, homology-based alignment and ab initio prediction. As a result, a total of 46,495 protein-coding genes were predicted. The average gene length, exon length, exon number per gene were 1963 bp, 230 bp and 5, respectively, which is similar to Pomi (Supplementary Table S5). The gene density was high in the distal region (Figure 2b). To obtain a better sense of functionality, 42,797 of the genes were successfully assigned to at least one public functional database. In detail, 92.05%, 82.05%, 46.53%, 79.36%, 39.32% of the protein-coding genes were assigned to Nr, Swiss-Prot, KEGG, TrEMBL, GO, respectively (Supplementary Figure S4). In addition, based on the Rfam database [26], 58.85 Kb, 83.08 Kb, 98.15 Kb, and 77.52 Kb of miRNA, tRNA, rRNA, and snRNA were identified in the Poan genome, respectively (Table 1).

Table 1. Statistics of Non-coding RNAs and TE sequences.

| Type          | Number | Total Length (bp) | Proportion in the Genome (%) |
|---------------|--------|-------------------|------------------------------|
| miRNA         | 479    | 58,847            | 0.013                        |
| tRNA          | 1116   | 83,079            | 0.018                        |
| rRNA          | 148    | 98,151            | 0.022                        |
| snRNA         | 655    | 77,523            | 0.017                        |
| Total         | 2398   | 317,600           | 0.07                         |
| Retroelements | 166,482| 118,458,221       | 26.08                        |
| LINEs         | 39,955 | 12,791,517        | 2.82                         |
| SINEs         | 2373   | 269,946           | 0.06                         |
| LTR elements  | 124,154| 105,396,758       | 23.20                        |
| Copia         | 56,850 | 53,227,095        | 11.72                        |
| Gypsy         | 31,449 | 38,942,113        | 8.57                         |
| DNA transposons| 167,514| 4,317,7431        | 9.50                         |
| Others        | 27,886 | 6,454,876         | 1.42                         |
| Total         | 368,838| 169,737,457       | 37.26                        |

TE has been considered to be the contributor of genetic variations in the plant genome [52]. Therefore, we used the ab initio method and homology-based method with Repbase [29] to annotate the repeats. Taken together, a total of 169.74 Mb (~37.26%) TEs were identified (Figure 2b). The most abundant TE was LTR-RTs accounting for 23.2% (105.4 Mb) in total. Among LTR-RTs, Gypsy and Copia were two biggest TE sub-families, reaching 11.72% and 8.57% of the Poan genome, respectively. The DNA transposons, long interspersed nuclear elements, and short interspersed nuclear elements covered 9.50%, 2.82% and 0.06% of the genome, respectively (Table 1). The burst of LTR amplification usually affects the genome size and structure [53,54]. Therefore, we investigated the LTR insertion time of Poan, and two relatives (Pomi, and Frve). Interestingly, the LTR burst in Poan occurred ~0.18 Mya, which is slightly later than Frve (~0.24 Mya) (Supplementary Figure S5). However, Pomi, another sequenced species in Potentilla genus, owned a totally different pattern of LTR-RT burst that happened ~1.36 Mya (Supplementary Figure S5), which was significantly earlier than Poan and Frve.

3.4. Comparative Genomics and Evolutionary Analysis

Rosaceae is an economically important family with high diversity. The protein sequences of Poan and seven Rosaceae species, including Frve, Mado, Pomi, Prpe, Pyco, Roch, and Ruoc, were collected for comparative genomics analysis, with Arabidopsis thaliana (Arth) as the outgroup. Taken together, we found 29,221 gene families, of which 10,997 were shared by the Rosaceae species and 676 were specific to Poan (Figure 3a). GO enrichment analysis showed that the Poan-specific gene families were enriched in “response to light intensity”, “response to UV-B”, “lipid metabolic process” and other related biological processes (Supplementary Table S6). KEGG enrichment showed that the Poan-specific
gene families were mainly enriched in “Biosynthesis of secondary metabolites”, “Galactose metabolism”, and “Starch and sucrose metabolism” pathways (Supplementary Table S7). We also identified 88 gene families which were specific to the Potentilla genus. In addition, a total of 201 and 185 gene families were shared by woody plants and herbaceous plants, respectively (Supplementary Table S8).

To investigate the evolution of the Poan genome, we selected 286 single copy genes to construct the phylogenetic tree. It is obvious that Pomi was the most closely relative of Poan. The Fragaria genus was close to the Potentilla genus. Poan, Pomi, Frve, Roch, and Ruoc were clustered into a clade that represented the herbaceous plants among the Rosaceae family, while Mado, Prpe, and Pyco represented the woody plants (Figure 3b). The herbaceous plants diverged from woody plants about ∼90.83 Mya (Figure 3b). The Potentilla genus separated from Fragaria genus about ∼40.68 Mya, followed by the divergence between Poan and Pomi (∼28.52 Mya). Based on the tree and multiple alignment sequences, we identified the expansion and contraction of gene families, which are crucial for environmental adaption of the plant. A total of 11,701 significantly expanded gene families (p < 0.05) and 3259 contracted gene families (p < 0.05) were found in the Poan genome. The expanded gene families were significantly enriched in 153 GO terms (biological process) including “response to hormone”, “hormone-mediated signaling pathway”, “reproductive shoot system development” and others (Supplementary Table S9).

**Figure 3.** Evolution of the Poan genome. (a) Orthologous gene families among Rosaceae plants. (b) Phylogenetic relationship of Poan and eight other species. Herbaceous plants are shown with a purple background and woody plants are shown with a yellow background. (c) Gene duplication events among nine different species. (d) Ks distribution between Poan, Pomi and Frve.

Gene duplication plays a key role in novel functions and important traits of the plant genome [55,56]. We therefore investigated the gene duplication event among these species. Interestingly, the results showed that Poan possessed the most (858) WGD/segmental duplication involving 34,572 genes and the fewest (1279) singletons (Figure 3c).

The homologous gene pairs in Poan, Pomi, and Frve were collected for whole genome duplication analysis. The synonymous substitutions per site (Ks) distance of these gene pairs revealed that Poan, Pomi and Frve shared a common ancient WGD (Ks ∼1.05–1.2), while Poan underwent an additional WGD (Ks ∼0.08), leading to the tetraploidization. The
shared ancient WGD occurred at ∼70–80 Mya, while the tetraploidization of Poan occurred at ∼6.4 Mya, which was later than the divergence between Poan and Pomi, as well as Poan and Frve (Figure 3d).

3.5. Sub-Genome Structure and Expression Bias Analysis

Given that the Poan is an allotetraploid plant, it is necessary to dissect the sub-genome structure of the Poan. We adopted inter- and intra-species gene synteny and evolutionary analyses. First, we identified 16,748 gene pairs within the Poan genome, showing a high collinearity among 14 Poan chromosomes (Figure 4a). The 14 chromosomes were clearly divided into two groups, indicating the existence of the sub-genome structure (Figure 4a). Furthermore, there were 26,032 gene pairs between Poan and Pomi, 51% and 13% of which showed a 1:2 or 1:3 synteny pattern (Pomi:Poan) (Supplementary Figure S6). The top 10 contigs of Pomi also showed a high collinearity with two sub-genomes of Poan (Supplementary Figure S7).

![Figure 4. Sub-genome structure and expression dominance analysis. (a) Intra-genome gene synteny within Poan 14 chromosomes. (b) The phylogenetic tree of single copy genes between Pomi and Poan sub-genomes. (c) Sub-genome constitution of Poan. (d) Ka/Ks values of two sub-genomes. (e) Expression values for co-expression of homoeologous gene pairs.](image-url)

In order to distinguish the two sub-genomes, 6345 single copy gene pairs between Pomi and Poan sub-genomes were collected for constructing phylogenetic trees. A group of seven chromosomes (1,3,5,7,9,11,13) who had a closer genetic relationship with Pomi genes were denoted as A1-7, and the remaining group was B1-7 (Figure 4b). The statistic of two sub-genomes showed that the A sub-genome was larger in size, gene number, and TE s, whereas it was similar in gap number and GC content (Figure 4c).

Allotetraploidy can coordinate two different genomes through genetic modifications to exhibit stronger adaptability and vigor [57]. To investigate the expression pattern and evolution of homoeologous genes, sequence alignment and RNA-seq analyses was carried out. Above all, a total of 7701 homoeologous gene pairs were identified. The non-synonymous substitution rate (Ka) and Ks of homoeologous gene pairs showed that...
the homoeologous genes in the A sub-genome had a significantly lower Ka/Ks ratio (mean Ka/Ks = 0.34) than the B sub-genome (mean Ka/Ks = 0.38) (Figure 4d), indicating that the A sub-genome might evolved differently to the B sub-genome. Then, we calculated the expression level of homoeologous genes in various tissues containing tuberous root, normal root, stem, SAM, leaf, and flower. In total, 43,112 (92.7%) genes were expressed (TPM > 1 in at least one tissue). The overall expression of homoeologous genes in the A sub-genome was slightly higher than in the B sub-genome (Supplementary Figure S8). Of the 7701 homoeologous gene pairs, 6490 (84.27%) pairs had expression differences greater than two-fold change in at least one tissue, including 3321 and 3169 homoeologs with higher expression in the A and B sub-genomes, respectively. Furthermore, 1134 and 1075 homoeologs of the A and B sub-genomes had higher expression values in all the six tissues, respectively. In addition, 97 and 117 homoeologs exclusively expressed in the A and B sub-genome, respectively. When the homologous gene pairs were both expressed, the median expression level of A homoeologs was similar to B homoeologs in normal root and stem, and was higher in SAM, flower, and tuberous root, but was lower in leaf (Figure 4e).

3.6. Resistance Gene Number Variation

QTP was characterized by dry climate, low temperature and strong ultraviolet, which was unfavorable for the plant pathogens and insect pests. Resistance genes containing nucleotide binding site (NBS) domains provide resistance to pathogens [58]. We therefore characterized the Resistance (R) genes in Poan and Frve. A total of 528 R genes were identified, including 1 TIR-NBS, 18 CC-NBS, 29 coiled-coil-NBS-LRR (CNL), 80 NBS-LRR genes, 66 NBS, 334 LRR-RLK genes. Based on the same parameter, the R gene number was 1321 in Frve and Pomi, which was over two times that of Poan and three times that of Pomi. There were 268 copies of TIR-NBS-LRR in the Frve genome, while it could not be detected in Poan. For NBS-LRR, there were 213, 80 and 0 copies in Frve, Poan and Pomi, respectively. In addition, the copy number of TIR-CC-NBS-LRR in Pomi (109) is far more than Frve (1) and Poan (0).

3.7. Genes Involved in Starch Biosynthesis

One of the most important trait of Poan is the tuberous root. Compared to the normal root, the tuberous root of Poan contains more starch. Therefore, we focused on key enzymes related to starch metabolism pathway. In root tissues, starch synthesis is derived from imported sucrose [59]. The sucrose was transformed into starch under a series of reactions [60,61]. The key enzymes involved in starch biosynthesis were expanded in Poan genome, including glucose-1-phosphate adenylyltransferase (AGP), the first key regulatory and rate-limiting enzyme in the starch biosynthesis pathway, starch synthetase (SS), granule-bound starch synthase1 (GBSS1), and disproportionating enzyme1 (DPE1). The gene numbers of AGP, SS, GBSS1, DPE1 were 11, 8, 4, and 2, respectively, which is significantly more than those of Pomi, Frve, Roch, Ruoc (Figure 5a,b). Of the remaining starch biosynthesis-related genes, Poan still had the advantage of gene copy numbers (Figure 5b). Interestingly, the number of sucrose transporters (SUT) was increased to 13 in Poan (Figure 5b), four of which were tandem duplicated genes.
Figure 5. Analysis of key genes involved in starch metabolism. (a) A diagram depicting the main reactions involved starch metabolism. (b) Key gene copy numbers among Poan, Pomi, Frve, Roch, and Ruoc. (c) Heatmap of the expression level of genes related in starch metabolism in different tissues.

We further investigated the expression pattern of these key genes in different tissues. The results showed that seven AGP copies were highly expressed in tuberous root (Figure 5c), while 1 and 3 AGP copies were active in SAM and leaf, respectively (Figure 5c), indicating the gene subfunctionalization after allotetraploidization. Two DPE1 genes had higher expression levels exclusively in tuberous root. GBSSI was highly expressed in both tuberous root and leaf (Figure 5c). Of other key genes, isoamylase (ISA), and the starch branching enzyme (SBE) were simultaneously exhibited up-regulated expression in tuberous root, while SS was expressed in leaf, stem and tuberous root (Figure 5c). Sucrose synthase (SUS) and SUT were quite active in different tissues (Supplementary Table S11).

4. Discussion

Poan is an important plant in QTP with immense economical and ecological value. In this study, we assembled a chromosome-level genome assembly and performed a series of investigation for uncovering the special features of Poan using genomic and transcriptomic information. The BUSCO assessment and LAI score results consistently indicated that the genome assembly of Poan showed a high contiguity and completeness, even in the repeat region. Our genome assembly is much better than Pomi assembly, another sequenced Potentilla species, whose genome assembly only reached contig level (contig N50 = 33.5 Kb) [12]. The Pomi genome has 2,674 contigs with 0.07% heterozygosity, which is quite homozygous compared to Poan genome [12]. In addition, the Poan genome size (454 Mb) was larger than Pomi (327 Mb) and had more genes due to the allotetraploidization [12]. The high-quality Poan genome assembly should be attributed to our massive sequencing data including PacBio long-reads, short PE reads and Hi-C data, and constantly revised assembling methods. The chromosome-level Poan genome should be the phylogenetic backbone of Potentilla genus and valuable resources for investigation of chromosome fusion and fission in Rosaceae family.

Based on the high-quality genome assembly, we investigated the evolution of Poan genome. The divergence between Poan and Pomi happened at ∼28.52 Mya. The WGD analysis suggested that, after the separation between them, Poan experienced the recent
tetraploidization ∼6.4 Mya. The WGD introduced the gene family expansion and contraction. We found that the expanded gene families are involved in a lot of GO terms, some of which are related to the environmental adaption such as “respond to hormone” and “reproductive shoot system development”. The phytohormone biosynthesis and signaling may also play key roles in tuberous root development [62]. The rapid growth of stolons laid a solid foundation for reproduction and increase in tuberous root. The identified Poan-specific gene families are enriched in GO terms such as “response to light density” and “response to UV-B”. Light quality and intensity is one of the important factors of tuberous root formation [63]. It is important for Poan to adapt to the light density. Furthermore, Poan may have evolved the adaptability to survive under the strong ultraviolet light in QTP.

We adopted three methods including cytological evidence, k-mer analysis and ploidy analysis to investigate the ploidy of Poan. All evidences suggested that Poan was an allotetraploid plant. Through gene synteny and genetic distance analyses, we successfully identified the sub-genome structure of Poan. Each of them contained seven chromosomes, consistent with basic chromosomal number of Rosaceae species. The approaches used in this study provide a feasible strategy for investigating the allopolyploid plant genome.

Moreover, some studies had reported that Poan also has other varieties such as: pentaploid and hexaploid [64]. Therefore, our genome assembly of Poan represents the finite genetic variation of this species. More materials should be covered for identifying the more detailed genetic variations through re-sequencing or pan-genome assembling. However, based on our research, the typical features of Poan had been elucidated both in the genomic and transcriptomic aspects. The A sub-genome of Poan, genetically close to Pomi, was larger in size and evolved differently to the B sub-genome. The homoeologous genes had tissue-specific expression bias despite lacking significant genome-wide expression dominance. The overall R gene numbers in Poan and Pomi were obviously less than Frve, indicating the gene contraction in Poan and Pomi, which is probably be due to environmental adaption. The variation of R genes should lead to different disease resistance among these species. We further investigated the key genes involved in tannin, flavonoid and triterpene metabolism in Poan (Supplementary Table S12). Given that Poan are rich in these compounds, it is conducive to molecular research in the future. Furthermore, compared to other herbaceous plants, the key genes involved in starch biosynthesis were significantly expanded. It could be inferred that the copy number variation in the Poan genome was caused by tetraploidization. In addition, these key genes were highly expressed in the root tissue, and they evenly were distributed in the A and B sub-genomes (Supplementary Figure S9). These observed results strongly suggested that copy number variation caused by allotetraploidization and high expression level of key enzymes involved in starch biosynthesis lead to the tuberous root formation in Poan.

In conclusion, owing to the high-speed development of sequencing techniques, we assembled a reference genome of Poan, which shed light on the tuber formation and genome evolution. The data generated from this research will be significant resources for genetic studies and genomics-assisted breeding programs of Poan.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/genes12121993/s1, Figure S1: Morphological characteristics of P. anserina, Figure S2: Karyotype of Poan genome, Figure S3: The chromosomal contact matrix heatmap in Poan, Figure S4: Functional annotation of Poan gene set, Figure S5. LTR-RT insertion time of Poan, Pomi and Frve, Figure S6. The syntenic depth of all homologous gene pairs between Pomi and Poan genome, Figure S7. The syntenic blocks between top 10 contigs of Pomi and Poan genome, Figure S8. Overall homoeologous gene expression levels between A and B sub-genomes in Poan, Figure S9. Distribution of key genes involved in starch biosynthesis, Table S1. Summary of sequencing data, Table S2. Genome size analysis by k-mer (k = 17), Table S3. Illumina PE reads mapping result, Table S4. Statistics of completeness validation of genome assembly, Table S5. Summary of gene set of Poan and Pomi, Table S6. Significantly enriched GO terms of species-specific gene families, Table S7. Significantly enriched KEGG terms of species-specific gene families, Table S8. Orthogroups shared by woody plants and herbaceous plants, Table S9. Significantly enriched GO terms of expanded gene families,
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Table S10. Characterized R genes in F. vesca, P. anserina and P. micrantha, Table S11. The expression level (TPM) of key genes related to starch metabolism, Table S12. The key genes involved in tannin, flavonoid and triterpenes metabolism in Poan.

Author Contributions: H.Z. and B.L designed the study. X.G. and S.L. performed bioinformatics analysis. B.L. and X.G. wrote the paper. R.L., Y.L., S.C. and D.C. analyzed part of data. X.G. and Y.Z. collected the samples. All authors have read and approved the final manuscript.

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Data Availability Statement: This Whole Genome Shotgun project has been deposited at GenBank under the accession PRJNA640225. All sequencing data are available at NCBI Sequence Read Archive SRP267963. The genome assembly and annotation files are available at figshare (https://figshare.com/projects/The_genome_assembly_and_annotation_files_of_Potentilla_anserina/83771 (accessed on 25 November 2021)).

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References
1. Moritkawa, T.; Imura, K.; Akagi, Y.; Muraoka, O.; Ninomiya, K. Ellagic acid glycosides with hepatoprotective activity from traditional Tibetan medicine Potentilla anserina. J. Nat. Med. 2017, 72, 29018991. [CrossRef] [PubMed]
2. Northwest Plateau Institute of Biology, Chinese Academy of Sciences. Economic Flora of Qinghai; Qinghai Peoples Publishing House: Xining, China, 1987; pp. 270–273.
3. Xia, L.; You, J. The determination of amino acids composition of the traditional food Potentilla anserina L. root by high-performance liquid chromatography via fluorescent detection and mass spectrometry. Int. J. Food. Sci. Technol. 2011, 46, 1164–1170. [CrossRef]
4. Guo, T.; Wei, J.; Ma, J. Antittusive and expectorant activities of Potentilla anserina. Pharm. Biol. 2015, 54, 1–5. [CrossRef]
5. Schimner, O.; Lindenbaum, M. Tannins with Antimutagenic Properties in the Herb of Alchemilla Species and Potentilla anserina. Planta Med. 1995, 61, 141–145. [CrossRef]
6. Kombal, R.; Glasl, H. Flavan-3-Ols and flavonoids from Potentilla anserina. Planta Med. 1995, 61, 484–485. [CrossRef]
7. Zhao, Y.L.; Cai, G.M.; Hong, X.; Shan, L.M.; Xiao, X. Anti-hepatitis B virus activities of triterpenoid saponin compound from Potentilla anserina L. Phytotherapy 2008, 15, 253–258. [CrossRef]
8. Chen, J.R.; Yang, Z.Q.; Hu, T.J.; Yan, Z.T.; Niu, T.X.; Wang, L.; Cui, D.A.; Wang, M. Immunomodulatory activity in vitro and in vivo of polysaccharide from Potentilla anserina L. Fitoterapia 2010, 81, 1117–1124. [CrossRef]
9. Erikkson, T.; Donoghue, M.; Hibbs, M. Phylogenetic analysis of Potentilla using DNA sequences of nuclear ribosomal internal transcribed spacers (ITS), and implications for the classification of Rosoideae (Rosaceae). Plant Syst. Evol. 1998, 211, 155–179. [CrossRef]
10. Persson, N.; Toresen, I.; Andersen, H.; Smedmark, J.; Erikkson, T. Detecting destabilizing species in the phylogenetic backbone of Potentilla (Rosaceae) using low-copy nuclear markers. AoB Plants 2020, 12, pla017. [CrossRef] [PubMed]
11. Töpel, M.; Lundberg, M.; Erikkson, T.; Eriksen, B. Molecular data and ploidal levels indicate several putative allopolyploidization events in the genus Potentilla (Rosaceae). PLoS Curr. 2011, 3, RRN1237. [CrossRef]
12. Buti, M.; Moretto, M.; Barghini, E.; Mascagni, F.; Natali, L.; Brilli, M.; Lomsadze, A.; Sonogo, P.; Giongo, L.; Alonge, M.; et al. The genome sequence and transcriptome of Potentilla micrantha and their comparison to Fragaria vesca (the woodland strawberry). GigaScience 2018, 7, gix010. [CrossRef]
13. Ranallo-Benavidez, T.; Jaron, K.; Schatz, M. GenomeScope 2.0 and Smudgeplot for reference-free profiling of polyploid genomes. Nat. Commun. 2020, 11, 1432. [CrossRef] [PubMed]
14. Chin, C.S.; Peluso, P.; Sedlazeck, F.; Nattestad, M.; Concepcion, G.; Clum, A.; Dunn, C.; O’Malley, R.; Figueroa-Balderas, R.; Morales-Cruz, A.; et al. Phased diploid genome assembly with single-molecule real-time sequencing. Nat. Methods 2016, 13, 1050–1054. [CrossRef] [PubMed]
15. Walker, B.; Abeel, T.; Shea, T.; Priest, M.; Abouelliel, A.; Sakthikumar, S.; Cuomo, C.; Zeng, Q.; Wortman, J.; Young, S.; et al. Pilon: An Integrated Tool for Comprehensive Microbial Variant Detection and Genome Assembly Improvement. PLoS ONE 2014, 9, e112963. [CrossRef]
16. Servant, N.; Varoquaux, N.; Lajoie, B.; Viara, E.; Chen, C.J.; Vert, J.P.; Heard, E.; Dekker, J.; Barillot, E. HiC-Pro: An optimized and flexible pipeline for Hi-C data processing. Genome Biol. 2015, 16, 259. [CrossRef]
17. Dudchenko, O.; Batra, S.; Omer, A.; Nyquist, S.; Hoefer, M.; Durand, N.; Shamim, S.; Machol, I.; Lander, E.; Aiden, A.; et al. De novo assembly of the Aedes aegypti genome using Hi-C yields chromosome-length scaffolds. Science 2017, 356, eaal3327. [CrossRef]
18. Durand, N.; Shamim, S.; Machol, I.; Rao, S.; Huntley, M.; Lander, E.; Aiden, E. Juicer Provides a One-Click System for Analyzing Loop-Resolution Hi-C Experiments. Cell Syst. 2016, 3, 95–98. [CrossRef] [PubMed]
19. Li, H. Fast and Accurate Long-Read Alignment with Burrows-Wheeler Transform. *Bioinformatics* 2010, 26, 589–595. [CrossRef] [PubMed]
20. Simão, F.; Waterhouse, R.; Ioannidis, P.; Zdobnov, E. BUSCO: Assessing genome assembly and annotation completeness with single-copy orthologs. *Bioinformatics* 2015, 31, 3210–3212. [CrossRef]
21. Ou, S.; Chen, J.; Jiang, N. Assessing genome assembly quality using the LTR Assembly Index (LAI). *Nucleic Acids Res.* 2018, 46, e126. [CrossRef]
22. Ou, S.; Jiang, N. LTR_retriever: A Highly Accurate and Sensitive Program for Identification of Long Terminal Repeat Retrotransposons. *Plant Physiol.* 2017, 176, 1410–1422. [CrossRef] [PubMed]
23. Lowe, T.; Eddy, S. tRNAScan-SE: A program for improved detection of transfer RNA Genes in genomic sequence. *Nucleic Acids Res.* 1997, 25, 955–964. [CrossRef]
24. Camacho, C.; Coulouris, G.; Avagyan, V.; Ma, N.; Papadopoulos, J. BLAST+: architecture and applications. *BMC Bioinform.* 2009, 10, 1–9. [CrossRef] [PubMed]
25. Nawrocki, E.; Kolbe, D.; Eddy, S. Infernal 1.0: Inference of RNA Alignments. *Bioinformatics* 2009, 25, 1335–1337. [CrossRef] [PubMed]
26. Kalvari, I.; Nawrocki, E.; Ontiveros-Palacios, N.; Argasinska, J.; Lammikiewicz, K.; Marz, M.; Griffiths-Jones, S.; Toffano-Nicoche, C.; Gautheret, D.; Weinberg, Z.; et al. Rfam 14: Expanded coverage of metagenomic, viral and microRNA families. *Nucleic Acids Res.* 2020, 49, D192–D200. [CrossRef] [PubMed]
27. Flynn, J.; Hubley, R.; Goubert, C.; Rosen, J.; Clark, A.; Feschotte, C.; Smit, A. RepeatModeler2 for automated genomic discovery of transposable element families. *Proc. Natl. Acad. Sci. USA* 2020, 117, 201921046. [CrossRef]
28. Xu, Z.; Wang, H. LTR-FINDER: An efficient tool for the prediction of full-length LTR retrotransposons. *Nucleic Acids Res.* 2007, 35, W265–W268. [CrossRef]
29. Jurka, J.; Kapitonov, V.; Pavlícek, A.; Klöners, P.; Kohany, O.; Walichiewicz, J. RepBase Update, a database of eukaryotic repetitive elements. *Cytogenet. Genome Res.* 2005, 110, 462–467. [CrossRef]
30. Stanke, M.; Steinkamp, R.; Waack, S.; Morgenstern, B. AUGUSTUS: A web server for gene finding in eukaryotes. *Nucleic Acids Res.* 2004, 32, W309–W312. [CrossRef] [PubMed]
31. Korf, I. Gene finding in novel genomes. *BMC Bioinform.* 2004, 9, 1–9.
32. Goodstein, D.; Shu, S.; Howson, R.; Neupane, R.; Hayes, R.; Fazio, J.; Mitros, T.; Dirks, W.; Hellsten, U.; Putnam, N.; et al. Phytome: A comparative platform for green plant genomics. *Nucleic Acids Res.* 2011, 40, D1178–D1186. [CrossRef]
33. Birney, E.; Durbin, R. Using GeneWise in the Drosophila Annotation Experiment. *Genome Res.* 2000, 10, 547–548. [CrossRef] [PubMed]
34. Grabherr, M.; Haas, B.; Yassour, M.; Levin, J.; Thompson, D.; Amit, I.; Adiconis, X.; Fan, L.; Raychowdhury, R.; Zeng, Q.; et al. Full-Length transcriptome assembly from RNA-Seq data without a reference genome. *Nat. Biotechnol.* 2011, 29, 644–652. [CrossRef] [PubMed]
35. Haas, B.; Salzberg, S.; Zhu, W.; Pertea, M.; Allen, J.; Orvis, J.; White, O.; Buell, C.; Wortman, J. Automated eukaryotic gene structure annotation using EVidenceModeler and the Program to Assemble Spliced Alignments. *Genome Biol.* 2008, 9, R7. [CrossRef]
36. El-Gebali, S.; Mistry, J.; Bateman, A.; Eddy, S.; Luciani, A.; Potter, S.; Qureshi, M.; Richardson, L.; Salazar, G.; Smart, A.; et al. The Pfam protein families database in 2019. *Nucleic Acids Res.* 2018, 47, D427–D432. [CrossRef] [PubMed]
37. Cantalapiedra, C.; Hernández-Plaza, A.; Letunic, I.; Bork, P.; Huerta-Cepas, J. eggNOG-mapper v2: Functional Annotation, Orthology Assignments, and Domain Prediction at the Metagenomic Scale. *Mol. Biol. Evol.* 2021, 38, 5825–5829. [CrossRef] [PubMed]
38. Sug, S.; Lee, T.; Cheng, C.H.; Bublé, K.; Zheng, P.; Yu, J.; Humann, J.; Ficklin, S.; Gasic, K.; Scott, K.; et al. 15 years of GDR: New data and functionality in the Genome Database for Rosaceae. *Nucleic Acids Res.* 2018, 47, D1137–D1145. [CrossRef]
39. Emms, D.; Kelly, S. OrthoFinder: solving fundamental biases in whole genome comparisons dramatically improves orthogroup inference accuracy. *Genome Biol.* 2016, 16, 157. [CrossRef] [PubMed]
40. Dechoo, B.; Luo, H.; Huo, P.; Wang, Z.; Zhang, S.; He, Z.; Wu, Y.; Lianhe, Z.; Liu, J.; Guo, J.; et al. KOBAS-i: Intelligent prioritization and exploratory visualization of biological functions for gene enrichment analysis. *Nucleic Acids Res.* 2021, 49, W317–W325. [CrossRef]
41. Edgar, R. MUSCLE: Multiple Sequence Alignment with High Accuracy and High Throughput. *Nucleic Acids Res.* 2004, 32, 1792–1797. [CrossRef]
42. Stamatakis, A. RAxML Version 8: A Tool for Phylogenetic Analysis and Post-Analysis of Large Phylogenies. *Bioinformatics* 2014, 30, 1312–1313. [CrossRef] [PubMed]
43. Yang, Z.H. PAML 4: Phylogenetic analysis by maximum likelihood. *Mol. Biol. Evol.* 2007, 24, 1586–1591. [CrossRef] [PubMed]
44. Kumar, S.; Stecher, G.; Suleski, M.; Hedges, S. TimeTree: A Resource for Timelines, Timetrees, and Divergence Times. *Mol. Biol. Evol.* 2017, 34, 1812–1819. [CrossRef] [PubMed]
45. Wang, Y.; Tang, H.; Debarry, J.; Tan, X.; Li, J.; Wang, X.; Lee, T.H.; Jin, H.; Marler, B.; Guo, H.; et al. MCScanX: A toolkit for detection and evolutionary analysis of gene synteny and collinearity. *Nucleic Acids Res.* 2012, 40, e49. [CrossRef] [PubMed]
46. Finn, R.; Clements, J.; Eddy, S. HMMER web server: Interactive sequence similarity searching. *Nucleic Acids Res.* 2011, 39, W29–W37. [CrossRef]
47. Bolger, A.M.; Lohse, M.; Usadel, B. Trimmomatic: a flexible trimmer for Illumina sequence data. *Bioinformatics* **2014**, *30*, 2114–2120. [CrossRef] [PubMed]

48. Kim, D.; Paggi, J.; Park, C.; Bennett, C.; Salzberg, S. Graph-based genome alignment and genotyping with HISAT2 and HISAT-genotype. *Nat. Biotechnol.* **2019**, *37*, 1. [CrossRef]

49. Liao, Y.; Smyth, G.; Shi, W. FeatureCounts: An efficient general purpose program for assigning sequence reads to genomic features. *Bioinformatics* **2013**, *30*, 923–930. [CrossRef] [PubMed]

50. Jarvis, D.; Ho, Y.S.; Lightfoot, D.; Schmöckel, S.; Li, B.; Borm, T.; Ohyanagi, H.; Mineta, K.; Michell, C.; Saber, N.; et al. The genome of *Chenopodium quinoa*. *Nature* **2017**, *542*, 1–6. [CrossRef]

51. Yasui, Y.; Hirakawa, H.; Oikawa, T.; Toyoshima, M.; Matsuzaki, C.; Ueno, M.; Mizuno, N.; Nagatoshi, Y.; Imamura, T.; Miyago, M.; et al. Draft genome sequence of an inbred line of *Chenopodium quinoa*, an allotetraploid crop with great environmental adaptability and outstanding nutritional properties. *DNA Res.* **2016**, *23*, dsw037. [CrossRef]

52. Sahebi, M.; Hanafi, M.; Wijnen, A.; Rice, D.; Rafii, M.; Azizi, P.; Osman, M.; Taheri, S.; Abu Bakar, M.F.; Mat Isa, M.N.; et al. Contribution of transposable elements in the plant’s genome. *Genome Biol.* **2018**, *665*, 155–166. [CrossRef]

53. Zhang, Q.J.; Li, W.; Li, K.; Nan, H.; Shi, C.; Zhang, Y.; Dai, Z.Y.; Lin, Y.L.; Yang, X.L.; Tong, Y.; et al. The Chromosome-Level Reference Genome of Tea Tree Unveils Recent Bursts of Non-autonomous LTR Retrotransposons to Drive Genome Size Evolution. *Mol. Plant* **2020**, *13*, 935–938. [CrossRef]

54. Zhang, Q.J.; Gao, L.Z. Rapid and Recent Evolution of LTR Retrotransposons Drives Rice Genome Evolution During the Speciation of AA-Genome Oryza Species. *G3 Bethesda Md.* **2017**, *7*, 1875–1885. [CrossRef]

55. Panchy, N.; Lehti-Shiu, M.; Shiu, S.H. Evolution of Gene Duplication in Plants. *Plant Physiol.* **2016**, *171*, 2294–2316. [CrossRef] [PubMed]

56. Qiao, X.; Li, Q.; Yin, H.; Qi, K.; Li, L.T.; Wang, R.; Zhang, S.; Paterson, A. Gene duplication and evolution in recurring polyploidization-diploidization cycles in plants. *Genome Biol.* **2019**, *20*, 38. [CrossRef]

57. Li, M.; Wang, R.; Wu, X.; Wang, J. Homoeolog expression bias and expression level dominance (ELD) in four tissues of natural allotetraploid *Brassica napus*. *BMC Genom.* **2020**, *21*, 330. [CrossRef]

58. Utsumi, Y.; Tanaka, M.; Utsumi, C.; Takahashi, S.; Matsui, A.; Fukushima, A.; Kobayashi, M.; Sasaki, R.; Oikawa, A.; Kusano, M.; et al. Integrative omics approaches revealed a crosstalk among phytohormones during tuberous root development in cassava. *Plant Mol. Biol.* **2020**, *8*, 1–21. [CrossRef] [PubMed]

59. Zierer, W.; Rüscher, D.; Sonnewald, U.; Sonnewald, S. Tuber and Tuberous Root Development. *Annu. Rev. Plant Biol.* **2021**, *72*, 551–580. [PubMed]

60. Cobon, A.M.; Mattefield, M. Morphological and cytological studies on a hexaploid clone of *Potentilla anserina* L. *Watsonia* **1976**, *11*, 125–129.