

三个多倍体猕猴桃基因组 survey 分析及系统进化研究

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摘要: 【目的】全面了解多倍体猕猴桃种质资源的染色体倍性和基因组特征, 并分析其在猕猴桃属植物中的系统进化关系, 以期为多倍体猕猴桃全基因组组装提供参考。【方法】基于流式细胞术分析中华猕猴桃 AcD2301 (*Actinidia chinensis*)、软枣猕猴桃 AcD2302 (*Actinidia arguta*)、对萼猕猴桃 AcD2303 (*Actinidia valvata*) 染色体倍性, 利用 Illumina 二代测序平台开展基因组 survey 分析, 并基于 SNP 构建 15 种猕猴桃属植物系统进化树。【结果】中华猕猴桃 AcD2301、软枣猕猴桃 AcD2302、对萼猕猴桃 AcD2303 的染色体倍性分别为四倍体、四倍体、六倍体, 与 survey 分析结果一致。K-mer 分析预测中华猕猴桃 AcD2301、软枣猕猴桃 AcD2302、对萼猕猴桃 AcD2303 单套基因组大小分别约为 626 Mb、668 Mb、585 Mb, 杂合度为 3.00%、3.30%、8.06%, 重复序列比例 43.70%、45.30%、40.7%。系统进化树显示软枣猕猴桃与对萼猕猴桃亲缘关系较近, 且均与中华猕猴桃独立进化而来。【结论】分析了中华猕猴桃 AcD2301、软枣猕猴桃 AcD2302、对萼猕猴桃 AcD2303 的染色体倍性、基因组大小和系统进化关系, 为将来开展的多倍体猕猴桃全基因组测序提供参考, 也为深入研究猕猴桃多倍化和系统进化提供了理论支持。

关键词: 猕猴桃; 基因组 survey 分析; 基因组大小; 系统进化

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Survey analysis and phylogenetic study of three polyploid kiwifruit genomes

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Abstract: 【Purpose】Plant polyploidization is the evolution of adapting to environmental changes and protecting their own population development. The polyploidization of kiwifruit will double its chromosome number and affect the structure and function of its genome, thereby enriching the genetic diversity of kiwifruit. To comprehensively understand the chromosomal ploidy and genomic characteristics of polyploid kiwifruit germplasm resources, and analyze their systematic

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evolutionary relationships in kiwifruit. **【Methods】**This study analyzed the chromosome ploidy of AcD2301 (*Actinidia chinensis*), AcD2302 (*Actinidia arguta*) and AcD2303 (*Actinidia valvata*) with reference to the diploid *Actinidia chinensis* Hongyang. The samples were analyzed by flow cytometry on the CyFlow Space flow cytometer after being lysed by CyStar UV Precise P kit and dyed by DAPI fluorescent dye in the dark. The total genomic DNA of kiwifruit was extracted by CTAB method, and then electrophoresis was conducted by 0.8% agarose gel. DNA quality was detected with UV spectrophotometer. Use the second-generation sequencing technology Illumina NovaSeq sequencing platform to perform double end sequencing on the sample library. Use software such as FastP to view the distribution of base quality, average error rate distribution of reads, and base content distribution of reads sequencing. Filter the raw data with adapters and low-quality reads to obtain high-quality sequences, and compare them with nucleic acid databases. High quality sequencing data was generated using Jellyfish (version 2.3.0) software k-mer19 to generate K-mer frequency tables, and genome size, heterozygosity, and repeatability were estimated using the GenomeScope 2. The next-generation sequencing data of kiwifruit, which have been published in the NGDC and NCBI databases, were compared with the reference genome *Actinidia chinensis* 'Hongyang' v4.0. SNP calling was performed using GATK software, and the Maximum likelihood algorithm in fastTree software was used to construct phylogenetic trees of 15 kiwifruit species, including *Actinidia chinensis*, *Actinidia arguta* and *Actinidia valvata*.

【Result】 The samples were subjected to flow cytometry analysis, and the peak values of the diploid 'Hongyang' kiwifruit were compared with the reference species. The chromosome ploidy of AcD2301 and AcD2302 were both tetraploid, while the chromosome ploidy of AcD2303 was hexaploid. The subsequent genome survey analysis results were consistent with this. The total genomic DNA of the sample was extracted by CTAB. After being detected, the AcD2301, AcD2302 and AcD2303 gene DNA were sequenced by the Illumina NovaSeq sequencing platform. Sequencing yielded raw data of 162.91 Gb, 139.74 Gb, and 142.44 Gb, followed by filtering to obtain high-quality data of 160.64 Gb, 138.16 Gb, and 140.73 Gb. The sequencing quality assessment showed that the Q20 and Q30 values of AcD2301 were 96.95% and 91.91%, respectively. The Q20 and Q30 values of AcD2302 were 97.09% and 92.07%, respectively. The Q20 and Q30 values of AcD2303 were 96.80% and 91.43%, respectively; The GC contents are approximately 37.20%, 36.77%, and 36.15%, respectively. The sequencing data quality values are all greater than 35, and the base error rates are all less than 0.045, indicating that the genome reads have high quality and can be used for subsequent analysis. Randomly select reads from the sequencing data of AcD2301, AcD2302, and AcD2303 and compare them with the nucleic acid library (NT library). The results show that the randomly selected reads can all be compared with the genome of kiwifruit plants, indicating that there is no contamination in the sequencing data.

Through K-mer analysis of the kiwifruit genome data after quality control, AcD2301 estimated the genome size to be 626 Mb, heterozygosity to be 3.00%, and repeat sequence ratio to be 43.70%; The estimated size of the AcD2302 genome is 668 Mb, with a heterozygosity of 3.30% and a repeat sequence ratio of 45.30%; The estimated genome size of AcD2303 is 585Mb, with a heterozygosity of 8.06% and a repeat sequence ratio of 40.7%. In addition, the support rates for homologous tetraploids of AcD2301 and AcD2302 were 97% and 96.7%, respectively. To analyze the evolutionary relationship of kiwifruit plants, SNP sequences were screened from the second-generation sequencing data of 15 kiwifruit, including AcD2301, AcD2302, and AcD2303. The Maximum likelihood algorithm was used to construct a phylogenetic tree. The results showed that the 15 kiwifruit plants were divided into three major evolutionary branches, with *Actinidia chinensis* AcD2301 as an independent branch, *Actinidia chinensis* 'Donghong' as another independent branch, and the remaining 13 kiwifruit species forming one evolutionary branch. *Actinidia chinensis* var. *deliciosa* in the third evolutionary branch is a small evolutionary branch, while the other 12 kiwifruit species form a small evolutionary branch. For the latter, *Actinidia hubeiensis* is a separate group; The remaining 11 kiwifruit species are grouped together, with the 6 kiwifruit species in the net fruit group clustered into a small evolutionary branch, while the 9 kiwifruit species in the remaining branches are all part of the spotted fruit group. From this, it can be seen that AcD2302, which belongs to the net fruit group, is closely related to AcD2303, and both have evolved independently from AcD2301 in the spotted fruit group. 【Conclusion】 The chromosome ploidy, genome size, and phylogenetic relationships of AcD2301, AcD2302 and AcD2303 were analyzed, providing reference for future polyploid kiwifruit whole genome sequencing and theoretical support for further research on kiwifruit polyploidization and phylogenetic relationships.

Key words: kiwi; Genome survey analysis; Genome size; System evolution

猕猴桃 (*Actinidia* spp.) 是猕猴桃科 (Actinidiaceae) 猕猴桃属 (*Actinidia* Lindl.) 植物，是 20 世纪初开始人工驯化栽培的特色经济果树，由于果实风味独特、营养丰富、维生素 C 含量高等优点，被誉为水果之王且深受广大消费者青睐^[1-2]。2024 年联合国粮农组织 FAO (<https://www.fao.org/home/zh/>) 统计数据显示，截至 2022 年世界猕猴桃采收面积 28.61 万 hm²，产量 429.15 万 t，是全球性重要的水果产业。其中中国猕猴桃产量约占世界猕猴桃总产量的 2/3，已成为我国重要的特色水果产业之一。猕猴桃为功能性雌雄异株植物，起源和分布中心均在我国，是广大山区常见的一种水果，生长在路旁、林中、水沟边、灌丛中，自然状态下存在着广泛的种间和种内杂交现象，造成了猕猴桃属植物复杂的形态结构变异。在猕猴桃属植物中，多倍化现象普遍存在，例如已知的主要品种中华猕猴桃红阳、枣猕猴桃魁绿、味猕猴桃贵长分别为二、四、六倍体。此外，猕猴桃种内染色体倍性变异也较为常见，

不同倍性材料在生态适应^[3]、抗逆^[4]及果实品质^[5]方面存在显著差异。猕猴桃多倍化是适应环境变化保护自身种群发展的进化，不仅使猕猴桃的染色体数目加倍，还影响其基因组的结构和功能，从而丰富猕猴桃遗传多样性^[6]。

随着基因组学时代的到来和发展，测序成本不断降低，高通量测序已被广泛应用于植物基因组测序中。在猕猴桃属植物中，中华猕猴桃 (*Actinidia chinensis*)^[7]、毛花猕猴桃 (*Actinidia eriantha*)^[8]、阔叶猕猴桃 (*Actinidia latifolia*)^[9]、山梨猕猴桃 (*Actinidia rufa*)^[10]、软枣猕猴桃 (*Actinidia arguta*)^[11]、长叶猕猴桃 (*Actinidia hemsleyana*)^[12]、葛枣猕猴桃 (*Actinidia polygama*)^[10]等基因组已有报道，为其他猕猴桃属植物的全基因组测序、重要性状解析和遗传改良等工作奠定了基础^[13]。然而中华猕猴桃、美味猕猴桃、软枣猕猴桃等主要栽培利用的物种普遍存在多倍化的现象，尽管不同倍性种质的基因组信息有共性之处，但多倍体猕猴桃的全基因组信息仍有待解析^[14-15]。此外，对萼猕猴桃作为新型猕猴桃砧木，具备较好的抗涝、抗寒、抗病能力，在产区中也已经得到较好的推广^[16]，但缺乏其基因组信息，阻碍了对其重要抗逆性状解析的基础。因此，考察中华猕猴桃、软枣猕猴桃、对萼猕猴桃的倍性及基因组信息对后续指导多倍体基因组的组装和辅助其他相关研究具有十分重要的意义。

笔者在本研究中选取野生种质中华猕猴桃 AcD2301 (*Actinidia chinensis*)、软枣猕猴桃 AcD2302 (*Actinidia arguta*)、对萼猕猴桃 AcD2303 (*Actinidia valvata*) 进行多倍体猕猴桃基因组 survey 分析及系统进化研究，通过流式细胞术、K-mer 分析和系统进化树构建，进行染色体倍性、物种杂合率、基因组重复序列比例和基因组大小的评估及系统进化关系研究，以期为多倍体猕猴桃全基因组组装提供参考，也为深入研究猕猴桃多倍化和系统进化提供了理论支持。

1 材料和方法

1.1 试验材料

试验材料中华猕猴桃 AcD2301、软枣猕猴桃 AcD2302、对萼猕猴桃 AcD2303 均为野生资源（表 1），保存于贵州省农业科学院果树研究所百宜落叶果树试验基地。试验样品采集，剪取顶端幼嫩叶片，液氮速冻后置于-80 °C超低温冰箱保存备用。

表 1 样品采集信息

Table 1 Sample collection information

样品名称 Sample name	性别 Sexuality	采样时间 Sampling time	采样地点 Sampling site
中华猕猴桃 AcD2301 <i>Actinidia chinensis</i> AcD2301	雌 Female	2023 年 5 月 May, 2023	贵州省贵阳市 Guizhou, Guiyang
软枣猕猴桃 AcD2302 <i>Actinidia arguta</i> AcD2302	雌 Female	2023 年 5 月 May, 2023	贵州省贵阳市 Guizhou, Guiyang
对萼猕猴桃 AcD2303 <i>Actinidia valvata</i> AcD2303	两性花 Bisexual flower	2023 年 5 月 May, 2023	贵州省贵阳市 Guizhou, Guiyang

1.2 试验方法

1.2.1 染色体倍性检测

以二倍体红阳猕猴桃 (*Actinidia chinensis* var. Hongyang, 2X=58) 为内参, 采用流式细胞术进行染色体倍性检测^[17]。分别称取 AcD2301、AcD2302 和 AcD2303 新鲜顶端叶片 0.2 g, 置于培养皿中, 用 CyStain UV Precise P 试剂盒进行细胞核裂解, 提取完成后用 50 μm Celltrics 滤网过滤至样品管中, 加入 DAPI 荧光染液避光染色 2 min 后在 CyFlow Space 流式细胞仪上进行流式细胞术测试, 用 FloMax 软件分析核悬浮液。

1.2.2 DNA 提取及测序

采用 CTAB 法提取猕猴桃基因组总 DNA, 并通过 0.8% 琼脂糖凝胶电泳检测 DNA 提取质量, 同时采用紫外分光光度计对 DNA 进行定量。利用第二代测序技术 Illumina NovaSeq 测序平台对样本文库进行双末端测序。采用 fastp^[18]等软件查看碱基质量分布、Reads 平均错误率分布、Reads 测序碱基含量分布, 原始数据过滤接头和低质量 reads 获得高质量序列, 并与核酸库进行比对, 排除外源物种污染。

1.2.3 基因组 Survey 分析

高质量测序数据基于 jellyfish (version 2.3.0) 软件设置 K-mer 为 19 生成 K-mer 频数表和频率直方图, 统计总 K-mer 数、唯一 K-mer 数等, 并运用 GenomeScope 2 工具进行基因组大小、杂合度和重复序列比例的估计^[19-20]。

1.2.4 基于 SNP 的系统进化树构建

基于自测数据 (AcD2301、AcD2302、AcD2303) 和公共数据库 (NGDC、NCBI) 下载部分已公布的猕猴桃二代测序数据 (表 2), 在贵州果树所生物信息学分析平台进行系统进化分析, 与参考基因组红阳 v4.0^[21]进行比对, 利用 GATK 软件^[22]进行 SNP calling (仅保留二等位基因), 用 fastTree 软件中的 Maximum likelihood 算法构建系统进化树, 并将树文件进行可视化。

表 2 猕猴桃属植物种名及拉丁文名称

Table 2 Kiwifruit plant species name and Latin name

编号 numbering	种名 species name	拉丁文名称 Latin name	数据来源 data sources
AcD2301	中华猕猴桃	<i>Actinidia chinensis</i>	自测数据 Self-test data
AcD2302	软枣猕猴桃	<i>Actinidia arguta</i>	自测数据 Self-test data
AcD2303	对萼猕猴桃	<i>Actinidia valvata</i>	自测数据 Self-test data
CRR635714	中华猕猴桃东红	<i>Actinidia chinensis</i> var. Donghong	NGDC
CRR635715	阔叶猕猴桃	<i>Actinidia latifolia</i>	NGDC
SRR3543582	美味猕猴桃	<i>Actinidia chinensis</i> var. <i>deliciosa</i>	NCBI
SRR18177732	湖北猕猴桃	<i>Actinidia hubeiensis</i>	NCBI
SRR3723918	浙江猕猴桃	<i>Actinidia zhejiangensis</i>	NCBI
SRR3705798	黄毛猕猴桃	<i>Actinidia fulvicoma</i>	NCBI
SRR3705797	毛花猕猴桃	<i>Actinidia eriantha</i>	NCBI
SRR3723917	山梨猕猴桃	<i>Actinidia rufa</i>	NCBI

SRR3407085	软枣猕猴桃	<i>Actinidia arguta</i>	NCBI
SRR3474219	葛枣猕猴桃	<i>Actinidia polygama</i>	NCBI
SRR3474216	大籽猕猴桃	<i>Actinidia macrosperma</i>	NCBI
SRR3474220	对萼猕猴桃	<i>Actinidia valvata</i>	NCBI

注：国家生物信息中心（NGDC，中国）、National Center for Biotechnology Information（NCBI，美国）。

Note: National Center for Biotechnology Information (NGDC, China)、National Center for Biotechnology Information (NCBI, USA).

2 结果与分析

2.1 猕猴桃染色体倍性分析

以二倍体红阳猕猴桃 (*Actinidia chinensis* var. *Hongyang*, $2n=58$) 为内参，分析 3 份猕猴桃样品的倍性，图 1 展示为猕猴桃多倍体样品倍性的流式直方图。其流式直方图中横坐标代表荧光强度，纵坐标代表细胞数量，荧光强度与 DNA 含量成正比，即峰值的位置反应测试样品的倍性。根据内参物种二倍体红阳猕猴桃（图 1-A）的峰值比较，AcD2301（图 1-B）和 AcD2302（图 1-C）的染色体倍性均为四倍体，而 AcD2303（图 1-D）染色体倍性为六倍体，流式细胞术测得染色体倍性结果与后续全基因组测序结果一致，图中杂峰的原因可能为部分细胞核降解造成。

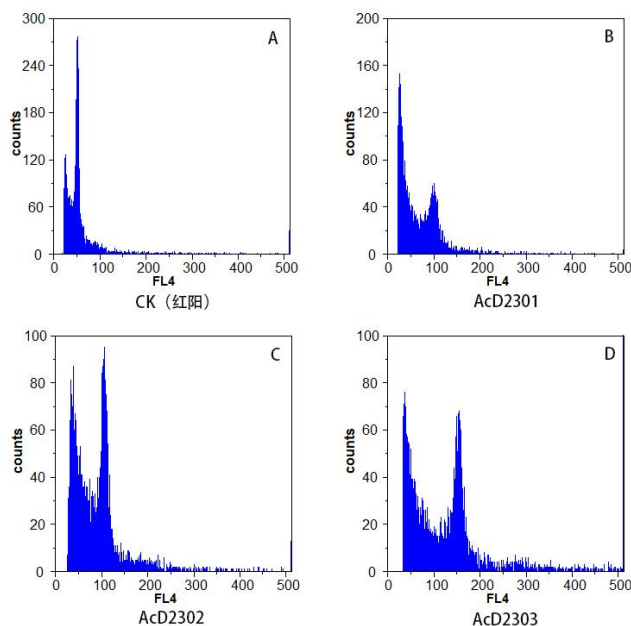


图 1 猕猴桃多倍体流式直方图

Fig. 1 Kiwi polyploid flow cytometry histogram

2.2 猕猴桃基因组测序及质控

通过二代 Illumina NovaSeq 测序平台对 AcD2301、AcD2302 和 AcD2303 基因 DNA 进行测序，分别获得 162.91 Gb、139.74 Gb 和 142.44 Gb 原始测序数据，经过过滤后分别获得 160.64Gb、138.16 Gb 和 140.73 Gb 高质量测序数据；测序的质量评估结果显示，AcD2301 的 Q20、Q30 值分别为 96.95%、91.91%，AcD2302 的 Q20、Q30 值分别为 97.09%、92.07%，AcD2303 的 Q20、Q30 值分别为 96.80%、91.43%，表明基因组数据良好，可用于后续分析。AcD2301、AcD2302 和 AcD2303 基因 GC 含量分别约为 37.20%，36.77% 和 36.15%（表 3）。

AcD2301(图2-A、图3-A)、AcD2302(图2-B、图3-B)和AcD2303(图2-C、图3-C)基因组中大部分测序数据质量值均大于35(图2),其碱基错误率均小于0.045(图3),表明其基因组测序的Reads质量较高,测序结果可信度较高。

表3 基因组测序数据统计表

Table 3 Statistical table of genome sequencing data

样品	AcD2301	AcD2302	AcD2303
总 Reads 数 The total number of reads	1078845806	925411346	943316252
总碱基数 Total number of bases /bp	162905716706	139737113246	142440754052
过滤后 Reads 数 Filtered reads number	1067883632	916913380	934022544
过滤后碱基/bp Filtered bases/bp	160639029587	138162662210	140726147170
Q20 rate/%	96.95	97.09	96.80
Q30 rate/%	91.91	92.07	91.43
GC/%	37.20	36.77	36.15

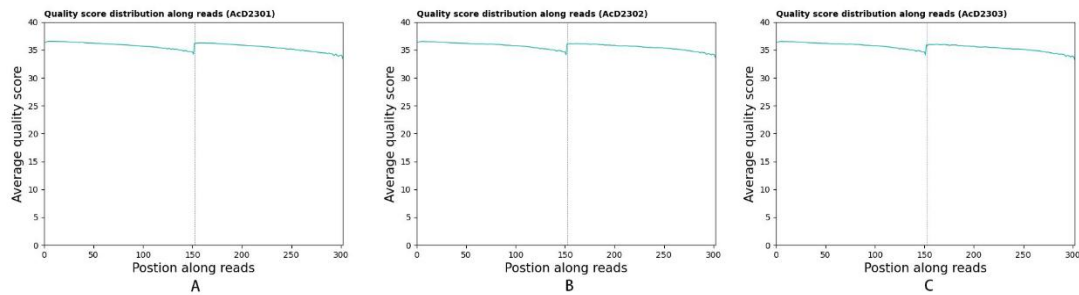


图2 单碱基质量分布

Fig. 2 Single base quality distribution

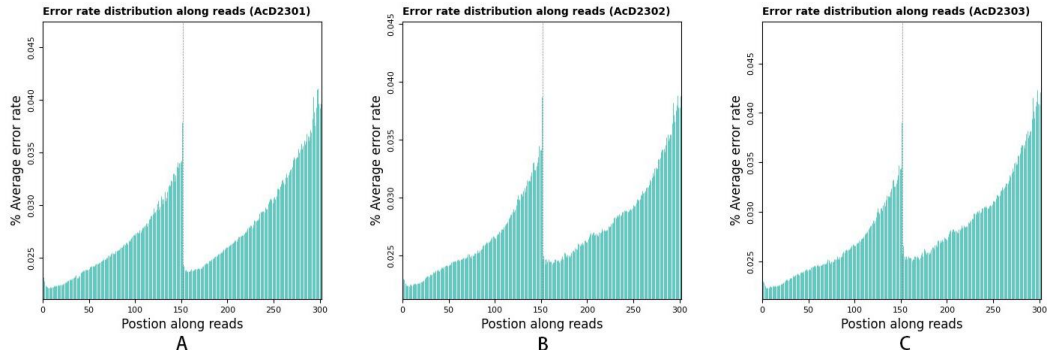


图3 Reads 平均错误率分布

Fig. 3 Distribution of average error rate of reads

2.3 猕猴桃基因组测序数据与NT数据库比对

进一步从AcD2301、AcD2302和AcD2303基因组测序数据中随机抽取10 000条Reads数据使用blast软件与核酸库(NT库)进行比对,挑选最优比对结果按物种统计(表4),结果显示随机选取Reads均能比对上猕猴桃属植物基因组,表明此次测序的基因组数据不存在

在污染，部分样本与核酸库比对率较低的原因与取样大小有关。

表 4 NT 比对结果统计

Table 4 NT comparison results statistics

物种名 Species name	AcD2301	AcD2302	AcD2303
	reads 数/比率	reads 数/比率	reads 数/比率
	The number of reads / ratio	The number of reads / ratio	The number of reads / ratio
中华猕猴桃 <i>Actinidia chinensis</i>	865/8.65	636/6.36	445/4.45
茶树 <i>Camellia sinensis</i>	214/2.14	188/1.88	183/1.83
软枣猕猴桃 <i>Actinidia arguta</i>	-	296/2.96	-
小叶猕猴桃 <i>Actinidia lanceolata</i>	138/1.38	144/1.44	108/1.08
黑蕊猕猴桃 <i>Actinidia melanandra</i>	85/0.85	-	-
对萼猕猴桃 <i>Actinidia valvata</i>	-	-	101/1.01
山梨猕猴桃 <i>Actinidia rufa</i>	68/0.68	101/1.01	67/0.67
君迁子 <i>Diospyros lotus</i>	45/0.45	46/0.46	-
葛枣猕猴桃 <i>Actinidia polygama</i>	-	-	61/0.61

2.4 猕猴桃基因组 survey 分析

高质量数据通过 K-mer 分析，预估物种基因组大小，并对物种的杂合度、重复情况进行分析。通过对质控后的猕猴桃基因组数据进行 K-mer 分析（图 4、表 5）得，AcD2301（图 4-A）预估单套基因组大小为 626 Mb，杂合度为 3.00%，重复序列比例 43.70%；AcD2302（图 4-B）预估单套基因组大小为 668 Mb，杂合度为 3.30%，重复序列比例 45.30%；AcD2303（图 4-C）预估单套基因组大小为 585 Mb，杂合度为 8.06%，重复序列比例 40.7%。基于猕猴桃基因组 survey 数据分析得 AcD2301（图 4-A）同源四倍体支持率为 97%，AcD2302（图 4-B）同源四倍体支持率为 96.7%。

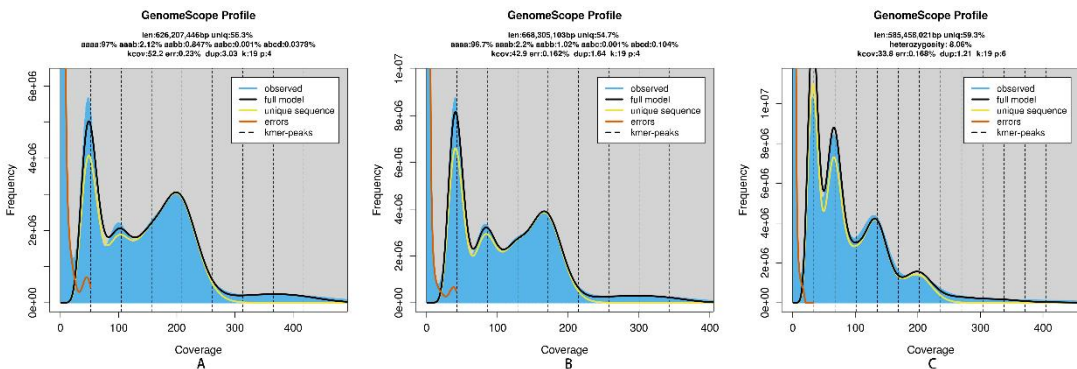


图 4 K-mer 分布曲线

Fig. 4 K-mer distribution curve diagram

表 5 基于 19-kmer 基因组信息估计统计表

Table 5 Estimated statistical table based on 19-kmer genome information

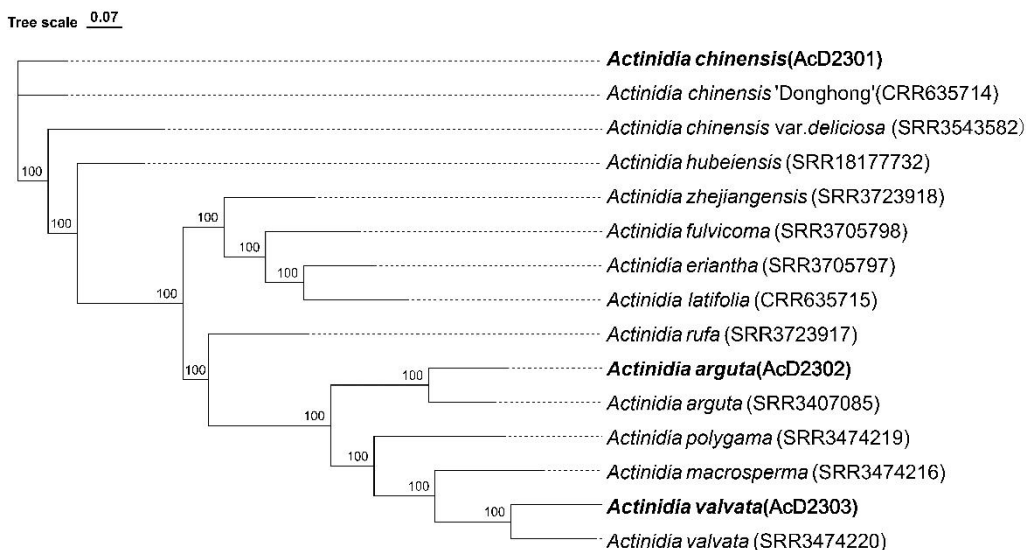
类型 Type	AcD2301	AcD2302	AcD2303
K 个碱基的序列 Sequence of K bases	19	19	19
K-mer 分布图的峰值 Frequency peaks of K-mer distribution chart	104	85.8	67.6

The peak of K-mer distribution map

杂合度 Heterozygosity/%	3.00	3.30	8.06
基因组大小 Genome size/Mb	626	668	585
重复序列比例 Repeat sequence ratio /%	43.70%	45.30%	40.7%
使用数据量 Amount of data used /G	160.6	138	141
数据量乘数 Quantity of data multiplier	256	206	240

2.5 猕猴桃属植物进化树分析

为分析猕猴桃属植物的进化关系，筛选了已报道的中华猕猴桃东红、美味猕猴桃和湖北猕猴桃等 15 种猕猴桃属植物的二代测序数据中的 SNP 序列，采用 Maximum likelihood 算法构建系统进化树。该系统进化树显示（图 5），15 种猕猴桃属植物分为三大进化分支，且均得到了较好的支持，其中中华猕猴桃 AcD2301 为一个独立进化枝，东红猕猴桃为另一个独立进化枝，其余 13 种猕猴桃组成一个进化枝。在第三个进化分支中的美味猕猴桃为一个小进化枝，其余 12 种猕猴桃组成一个小进化分支。对于后者湖北猕猴桃单独一组；其余 11 种猕猴桃为一组，其中净果组的 6 种猕猴桃（软枣猕猴桃 AcD2302、软枣猕猴桃、葛枣猕猴桃、大籽猕猴桃、对萼猕猴桃 AcD2303、对萼猕猴桃）聚为一个小进化分支，而其余分支的 9 种猕猴桃均为斑果组。



系统进化树构建中下载并使用已公布的猕猴桃二代测序数据有东红猕猴桃、阔叶猕猴桃（<https://ngdc.cncb.ac.cn>）；美味猕猴桃、湖北猕猴桃、浙江猕猴桃、黄毛猕猴桃、毛花猕猴桃、山梨猕猴桃、软枣猕猴桃、葛枣猕猴桃、大籽猕猴桃、对萼猕猴桃（<https://www.ncbi.nlm.nih.gov>）。

In the construction of phylogenetic tree, the published next-generation sequencing data of kiwifruit were downloaded and used, including *Actinidia chinensis* Donghong, *Actinidia latifolia* (<https://ngdc.cncb.ac.cn>); *Actinidia chinensis* var. *deliciosa*, *Actinidia hubeiensis*, *Actinidia zhejiangensis*, *Actinidia fulvicoma*, *Actinidia eriantha*, *Actinidia rufa*, *Actinidia arguta*, *Actinidia polygama*, *Actinidia macrosperma*, and *Actinidia valvata* (<https://www.ncbi.nlm.nih.gov>).

图 5 猕猴桃属植物系统进化树

Fig. 5 The phylogenetic tree of kiwifruit plants

3 讨 论

多倍化是推动植物遗传多样性和适应环境变化的重要机制之一，在植物中广泛存在，其中猕猴桃属植物中多倍化现象非常普遍。猕猴桃多倍化表现为体细胞均增大，果型更加圆润饱满，叶片颜色更深、表皮毛被明显增多、产量高、抗性强等特征^[3-5]。尽管以往的研究已经从很大程度上揭示了猕猴桃属物种的基因组信息以及该物种的主要倍性，但仍有部分物种尚未明确。本研究中就基于流式细胞术分析 AcD2301、AcD2302、AcD2303 的染色体倍性分别为四倍体、四倍体和六倍体。而基于猕猴桃基因组 survey 数据分析所得 AcD2301 同源四倍体支持率为 97%，AcD2302 同源四倍体支持率为 96.7% 的倍性，与上述结果基本一致。以上工作为进一步丰富猕猴桃物种基因组奠定了基础。

基于测序技术解析全基因组信息，为植物起源、进化、生殖、发育、抗性和性别调控等研究提供了基础。不同种类的植物基因组大小相差很大，根据目前已经公布的基因组数据中梅溪蕨（*Tmesipteris ob lanceolata*）的基因组大小约 160.45 Gb，而旋刺草（*Genlisea aurea*）的基因组大小仅为约 0.063 6 Gb，相差约 2500 倍^[23]。目前主要采用流式细胞术和高通量测序技术等方法评估植物的基因组大小，例如在四数九里香^[24]、白及^[25]、荆芥^[26]等多种植物基因组大小特征评估中都有应用。流式细胞术是通过测量细胞中 DNA 与荧光染料结合后发出的荧光信号强度，来间接预估基因组大小的相对值，而基因组 survey 分析是利用高通量测序技术对植物基因组进行测序和直接获取基因组大小等信息的测序技术，这两种技术结合起来评估基因组大小和特征相对可靠^[27]。已报道猕猴桃属植物的基因组大小通常在 600 Mb 左右，中华猕猴桃为 610.1 Mb^[7]，毛花猕猴桃为 619.3 Mb 和 611.7 Mb^[8]等，本研究结果所揭示的单套基因组大小较为相近，AcD2301 为 626 Mb、AcD2302 为 668 Mb、AcD2303 为 585 Mb，均都在 600 Mb 左右，但基因组具体大小又取决于不同的种质资源。

基因组学研究还可以揭示物种的遗传多样性、基因组演化历程以及基因功能等，通过构建系统进化树可以直观地展现亲缘关系和进化历程^[28]。已有研究通过 UPGMA 聚类分析得到星毛组的中华猕猴桃与净果组的软枣猕猴桃亲缘关系较远^[29]，并且与净果组的对萼猕猴桃亲缘关系也较远^[30]，由此推测同为净果组的软枣猕猴桃和对萼猕猴桃亲缘关系较近，并均与星毛组的中华猕猴桃亲缘关系较远。本研究中构建的猕猴桃属植物系统进化树，证明了软枣猕猴桃 AcD2302 与对萼猕猴桃 AcD2303 亲缘关系较近，且均与中华猕猴桃 AcD2301 独立进化而来的结果一致，为阐明物种进化关系及基因组的内在结构奠定基础。

4 结 论

中华猕猴桃 AcD2301、软枣猕猴桃 AcD2302、对萼猕猴桃 AcD2303 的染色体倍性分别为四倍体、四倍体和六倍体，与全基因组测序预估结果一致；基于全基因组 survey 分析预测基因组大小分别为 626Mb、668Mb、585Mb，杂合度为 3.00%、3.30%、8.06%，重复序列比例 43.70%、45.30%、40.7%。SNP 系统进化树发现软枣猕猴桃 AcD2302 与对萼猕猴桃 AcD2303 亲缘关系较近，且均与中华猕猴桃 AcD2301 独立进化而来。

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