

Analyzing genetic diversity of chloroplast genomes in Liliales

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Abstract

Liliales is a monocotyledonous order and contains both photosynthetic and mycoheterotrophic species that distribute locally or worldwide. In this study, the genetic diversity of chloroplast genomes in Liliales was explored regarding their nucleotide diversity and repeated composition. The analysis of nucleotide diversity revealed various hotspots in large and small single-copy regions whereas the IR regions had low sequence divergence. Although each family has specific hotspots, the *rps15-ycf1* region was commonly found as a highly variable area in the cpDNA of observed taxa. In the cpDNA of Liliales, mononucleotide simple sequence repeat (SSR) is the most common type. The majority of SSRs are located in non-coding regions. Similarly, more long repeats were found in non-coding areas than in coding sequences. Additionally, the complement repeat exceeds forward type in the cpDNA of Liliales. The highest number of long repeats was found in *Corsia dispar* whereas that of SSRs was detected in *Smilax china*. The results of nucleotide diversity and repeat analyses provided fundamental information for further studies on population genetics, molecular marker development and evolutionary history of Liliales.

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1 Introduction

Liliales, a monocotyledonous order of angiosperms, includes nine families of over 1500 species [1]. The families of Liliales distribute worldwide or locally. For example, Smilacaceae species are widespread from Australia, Europe, to Africa and Asia whereas the monotypic family, Petermanniaceae, can only be found in Australia. Additionally, there are two types of plants in Liliales, including mycoheterotrophic type in Corsiaceae and photosynthetic type in the remained families [2]. Because of their opposite patterns of lifestyle and distribution, Liliales is a good model to explore the evolutionary history of land plants. Previously, biogeography and divergent time estimation of Liliales were conducted [3]. The results showed a divergent time of 124 million years ago (mya) from other monocots and the families were

splitted approximately 113 mya. In addition to divergent time, the origin of Liliales was found in Australia where the ancestors of Liliales would then be widespread and evolved [3]. Beside order level, the divergent time estimation and biogeography of each family of Liliales were approached. Liliaceae originated from temperate Asia in the late Cretaceous (85 mya) to occupy the northern hemisphere [4]. Meanwhile, the members of Melanthiaceae used the Bering Land Bridge to migrate from North America to East Asia around 92.1 mya [5]. Colchicaceae arose in Australia 67 mya and migrated to Africa and North America [6]. Smilacaceae is an interesting family of Liliales that has many fossil records for elucidating the evolutionary history of Liliales [7-8]. These findings suggested an interesting evolutionary history of Liliales, especially at genomic level.

Chloroplast genome (cpDNA) is one of three existing genomes (including mitochondrial, nucleus and chloroplast genomes) in most land plants. Typically, cpDNA has a quadripartite structure which includes one large single copy (LSC) and a small single copy (SSC) separated by two inverted repeat (IR) regions [9]. Also, cpDNA contains 80 protein-coding genes, 30 tRNAs and four rRNAs and some of the protein-coding genes are related to photosynthesis. These genomic data are crucial for elucidating the phylogeny of land plants; therefore, the 1000 plant genomes project was conducted, followed by another 10 000 plant genomes project [10-11]. As a result, a billion years of evolutionary history of plants was explored [12]. Additionally, cpDNA is a useful source for mining molecular markers for population genetics and plant identification [13–17]. In Liliales, the cpDNA sequences from all families were reported [18-21]. These data provided essential information for elucidating the evolution of Liliales [4–8,22]. Although the complete cpDNA of Liliales have been reported, there has been no compilation of data for nucleotide diversity and repeat composition among Liliales families. Therefore, in this study, the available cpDNA data of Liliales were combined to locate the highly variable regions. Additionally, the simple sequence repeat (SSR) and long repeat were screened across cpDNA of Liliales. These new results will add insights into the evolutionary history of Liliales.

2 Materials and methods

2.1 Sampling chloroplast genome data

Complete chloroplast genome (cpDNA) sequences of Liliales were searched on NCBI (National Center for Biotechnology Information (<https://www.ncbi.nlm.nih.gov/>)) using the keywords “Liliales, chloroplast, complete genome”. The search results revealed all complete chloroplast genomes of Liliales, especially duplicated data for a species. Therefore, only one complete sequence (without unknown nucleotide in the genome) of a species was randomly selected as a representative because the similarity of the duplicated data is usually over 99.5 % (data not shown). Then, the selected complete genomes were downloaded under the GenBank (full) format which includes various information of chloroplast genomes such as gene content, gene

location, length, GC content, etc. All the data were imported to the Geneious Prime program for further analysis.

2.2 Nucleotide diversity analysis

To calculate the nucleotide diversity (Pi values, resulted from estimating the average number of nucleotide differences per site among DNA sequences) among chloroplast genomes of Liliales. The higher nucleotide diversity is, the higher genetic variation is detected in the genome. DnaSP 6 program was employed. First, the Pi values were estimated at familial level, making the complete cpDNA within each family of Liliales aligned using the MAUVE program embedded in Geneious Prime. The aligned sequences were then imported to DnaSP6 for Pi value calculation and sliding window analysis with the window size of 2 000 and the step size of 100. Among the families of Liliales, there is a monotypic family labeled as Petermanniaceae. Additionally, only one complete cpDNA was reported in Campynemataceae. The Corsiaceae includes heterotrophic species (i.e., *Corsia dispar* and *Arachnitis uniflora*) which exhibit extreme structural changes. Therefore, the nucleotide diversity analysis was not conducted for Petermanniaceae, Campynemataceae and Corsiaceae in this study.

2.3 Examination of repeat structure and microsatellites

For screening repeat number and location in cpDNA of Liliales, REPuter program was used with the minimum length of 20 bp for forward, reverse, and complement repeats. Meanwhile, Phobos program embedded in Geneious Prime was used for identifying microsatellites number and location with the minimum lengths of 10 bp for mono-, 12 bp for di-, 15 bp for tri-, 16 bp for tetra-, 20 bp for penta-, and 24 bp hexanucleotide repeats. A representative species of each genus in Liliales was selected randomly from available data for examining the repeat content in this study.

3 Results and discussion

3.1 Features of chloroplast genome in Liliales

On the NCBI database, 177 out of over 1500 species of Liliales have records of complete chloroplast genome (Table 1). Liliaceae family have the highest number of complete chloroplast genomes (105



species), followed by Melanthiaceae (49 species), Colchicaceae (9 species), Smilacaceae (4 species), Alstroemeriaceae (3 species), Philesiaceae (3 species), Corsiaceae (2 species) and one species each for Petermanniaceae and Campynemataceae. The lengths of chloroplast genomes range from 24 846 bp (*Arachnitis uniflora*, Corsiaceae) to 163 860 bp (*Paris liana*, Melanthiaceae). The GC content of Liliales is 37 % on average. Although *Corsia dispar* has a reduced size of cpDNA as found in *Arachnitis uniflora*, the GC content of the former is 30.8 %, which is lower than the latter (37.1 %) and other observed species (Table 1). Most of cpDNAs of Liliales encode 80 proteins, 30 tRNAs and four rRNAs (Table 1, Table 2). However, there are only 79 protein-coding genes in *Amana* species and *Chionographis japonica*, of which *infA* and *rps16* were lost, respectively. In contrast to other species, two members of Corsiaceae exhibited an extreme loss of protein-coding gene and tRNA (Table 2). Specifically, *Corsia dispar* has 30 protein-coding genes and 24 tRNAs whereas *Arachnitis uniflora* includes 16 protein-coding genes and 5 tRNAs.

Notably, these two species still have four rRNAs that are commonly found in other Liliales taxa (Table 2). There are two groups of species in Liliales according to cpDNA structure. The first group contains photosynthetic species that has typical structure of cpDNA including one large single copy (LSC), a small single copy (SSC) and two inverted repeat (IR) regions and contains approximately 80 protein-coding genes, 30 tRNA-coding genes and four rRNA-coding genes. The second group includes mycoheterotrophic species that exhibited an extreme loss of genes and significant changes of genome structure. Although *Arachnitis uniflora* has fewer genes than *Corsia dispar*, the former cpDNA has a typical quadripartite structure that was not found in the latter. This phenomenon suggested different stages of change in chloroplast genomes of mycoheterotrophic species. Previously, the plastid genomes of Ericaceae revealed the loss of genes related to photosynthesis whereas the other genes were remained [23]. Similarly, different numbers of gene loss were found in orchids that provided a scenario of 5 steps for the loss of plastid genes [24-25].

Table 1 Comparison of the features of plastid genomes from ten families of Liliales

Family	Species	Accession number	Length (bp)	GC content (%)	Gene content
					(Protein coding/ tRNA/rRNA)
Liliaceae (105 species)	<i>Amana anhuiensis</i>	KY101423	150 842	36.7	79/30/4
	<i>Amana baohuaensis</i>	MT898423	150 757	36.7	79/30/4
	<i>Amana edulis</i>	KY401425	151 136	36.7	79/30/4
	<i>Amana erytgronioides</i>	KY401421	150 858	36.7	79/30/4
	<i>Amana kuocangshanica</i>	KY401423	151 058	36.7	79/30/4
	<i>Amana wanzhensis</i>	KY401422	150 913	36.7	79/30/4
	<i>Calochortus uniflorus</i>	MK673754	155 794	37.4	80/30/4
	<i>Calochortus venustus</i>	MT261150	155 688	37.4	80/30/4
	<i>Cardiocrinum cathayanum</i>	KX575836	152 415	37.1	80/30/4
	<i>Cardiocrinum cordatum</i>	KX575837	152 410	37.1	80/30/4
	<i>Cardiocrinum giganteum</i>	KX528334	152 653	37.1	80/30/4
	<i>Clintonia udensis</i>	MT261153	156 214	37	80/30/4
	<i>Erythronium japonicum</i>	MT261155	151 416	36.6	80/30/4
	<i>Erythronium sibiricum</i>	KX644899	151 034	36.7	80/30/4
	<i>Fritillaria anhuiensis</i>	MH593363	152 119	37	80/30/4
	<i>Fritillaria cirrhosa</i>	KF769143	151 991	36.9	80/30/4
	<i>Fritillaria crassicaulis</i>	MK258147	151 852	37	80/30/4
	<i>Fritillaria dajinensis</i>	MH244913	151 991	36.9	80/30/4

<i>Fritillaria davidii</i>	MK158145	152 044	37	80/30/4
<i>Fritillaria delavayi</i>	MN480806	151 938	37	80/30/4
<i>Fritillaria eduardii</i>	MF947708	152 224	37	80/30/4
<i>Fritillaria hupehensis</i>	NC024736	152 145	37	80/30/4
<i>Fritillaria karelinii</i>	KX354691	152 118	36.9	80/30/4
<i>Fritillaria maximowiczii</i>	MK258138	152 434	37.1	80/30/4
<i>Fritillaria meleagroides</i>	MF947710	151 846	37	80/30/4
<i>Fritillaria pallidiflora</i>	MG211822	152 078	37	80/30/4
<i>Fritillaria persica</i>	MF947709	151 803	37	80/30/4
<i>Fritillaria przewalskii</i>	MH244908	151 983	36.9	80/30/4
<i>Fritillaria sichuanica</i>	MH244907	151 958	37	80/30/4
<i>Fritillaria sinica</i>	MH244912	152 064	36.9	80/30/4
<i>Fritillaria taipaiensis</i>	KC543997	151 693	37	80/30/4
<i>Fritillaria tortifolia</i>	MG211819	152 005	37	80/30/4
<i>Fritillaria thungergii</i>	MH244914	152 160	37	80/30/4
<i>Fritillaria unibracteata</i>	MH244909	151 058	37	80/30/4
<i>Fritillaria unibracteata</i> var. <i>wabuensis</i>	KF769142	151 009	37	80/30/4
<i>Fritillaria ussuriensis</i>	MT261156	152 156	37	80/30/4
<i>Fritillaria verticillata</i>	MG211823	151 959	37	80/30/4
<i>Fritillaria walujewii</i>	MG211820	151 920	36.9	80/30/4
<i>Fritillaria yuminensis</i>	MG200070	151 813	37	80/30/4
<i>Fritillaria yuzhongensis</i>	MK258139	151 652	37	80/30/4
<i>Gagea triflora</i>	MT261157	150 345	37	80/30/4
<i>Lilium bulbiferum</i>	MW465412	152 690	37	80/30/4
<i>Lilium amabile</i>	MT261159	152 614	37	80/30/4
<i>Lilium amoenum</i>	MT880912	152 280	37	80/30/4
<i>Lilium bakerianum</i>	KY748301	151 655	37.1	80/30/4
<i>Lilium brownii</i>	KY748296	152 677	37	80/30/4
<i>Lilium callosum</i>	MT261160	152 630	37	80/30/4
<i>Lilium candidum</i>	MK753244	152 101	37	80/30/4
<i>Lilium cernuum</i>	MT261161	152 553	37	80/30/4
<i>Lilium davidii</i> var. <i>unicolor</i>	MK954110	152 659	37	80/30/4
<i>Lilium distichum</i>	NC029937	152 598	37.1	80/30/4
<i>Lilium duchartei</i>	KY748300	152 287	37	80/30/4
<i>Lilium fargesii</i>	KX592156	153 235	36.0	80/30/4
<i>Lilium formosanum</i>	MT261162	152 610	37	80/30/4
<i>Lilium gongshanense</i>	MK493297	151 974	37	80/30/4
<i>Lilium hansonii</i>	MT261163	152 168	37	80/30/4
<i>Lilium henricii</i>	MH136807	152 784	37	80/30/4
<i>Lilium henryi</i>	KY748302	153 119	37	80/30/4
<i>Lilium japonicum</i>	MT261164	152 613	37.1	80/30/4
<i>Lilium lancifolium</i>	MH177880	152 479	37	80/30/4
<i>Lilium lankongense</i>	MK757466	152 611	37	80/30/4
<i>Lilium leichtlinii</i> var. <i>maximowiczii</i>	MK753242	152 604	37	80/30/4
<i>Lilium leucanthum</i>	KY748299	152 935	37	80/30/4

	<i>Lilium longiflorum</i>	KC968977	152 793	37.02	80/30/4
	<i>Lilium lophophorum</i>	MK493298	152 382	37	80/30/4
	<i>Lilium martagon</i> var. <i>pilosiusculum</i>	MF964219	152 816	37	80/30/4
	<i>Lilium matagense</i>	MN745201	152 402	37	80/30/4
	<i>Lilium meleagrinum</i>	MK493299	152 197	37	80/30/4
	<i>Lilium nanum</i>	MK493300	152 417	37	80/30/4
	<i>Lilium nepalense</i>	MK493301	152 316	37	80/30/4
	<i>Lilium pardalinum</i>	MH029495	151 969	37	80/30/4
	<i>Lilium pardanthinum</i>	MG704135	152 718	37	80/30/4
	<i>Lilium pensylvanicum</i>	MK493295	152 058	37.1	80/30/4
	<i>Lilium primulinum</i> var. <i>ochraceum</i>	KY7482988	152 036	37	80/30/4
	<i>Lilium pumilum</i>	MK954109	152 591	37	80/30/4
	<i>Lilium philadelphicum</i>	KY940847	152 175	37.1	80/30/4
	<i>Lilium regale</i>	MK493302	153 082	37	80/30/4
	<i>Lilium rosthornii</i>	MW136390	152 956	37	80/30/4
	<i>Lilium sargentiae</i>	MK493303	153 129	37	80/30/4
	<i>Lilium souliei</i>	MW007720	152 326	37	80/30/4
	<i>Lilium speciosum</i> var. <i>gloriosoides</i>	MN509267	152 912	37.02	80/30/4
	<i>Lilium sulphureum</i>	MK493304	153 107	37	80/30/4
	<i>Lilium superbum</i>	NC026787	152 069	37	80/30/4
	<i>Lilium taliense</i>	KY009938	153 055	36.9	80/30/4
	<i>Lilium tsingtauense</i>	KU230438	151 983	37	80/30/4
	<i>Lilium washintonianum</i>	MH590100	151 967	37.1	80/30/4
	<i>Lilium xanthellum</i>	MN745202	151 967	37.1	80/30/4
	<i>Lloydia tibetica</i>	MK673752	150 379	36.9	80/30/4
	<i>Medeola virginiana</i>	MK673752	153 914	37	80/30/4
	<i>Nomocharis aperta</i>	MK493293	152 042	37	80/30/4
	<i>Nomocharis pardanthina</i>	NC_038193	152 718	37	80/30/4
	<i>Notholirion bulbuliferum</i>	MN509268	153 019	37.1	80/30/4
	<i>Notholirion campanulatum</i>	MK673746	153 169	37	80/30/4
	<i>Notholirion macrophyllum</i>	MH011354	152 143	37.1	80/30/4
	<i>Prosartes lanuginosa</i>	MK673749	158 265	37	80/30/4
	<i>Scoliopus bigelovii</i>	MK673747	154 698	37.2	80/30/4
	<i>Streptopus ovalis</i>	MT261171	157 359	37.1	80/30/4
	<i>Tulipa altaica</i>	MK673755	146 887	37.1	80/30/4
	<i>Tulipa buhseana</i>	MT316022	152 062	36.6	80/30/4
	<i>Tulipa iliensis</i>	MW077740	152 073	36.6	80/30/4
	<i>Tulipa patens</i>	MT327740	152 050	36.7	80/30/4
	<i>Tulipa sylvestris</i>	MT261172	151 940	36.7	80/30/4
	<i>Tulipa thianschanica</i>	MT327741	152 122	36.6	80/30/4
	<i>Tricyrtis formosana</i>	MK673751	156 018	37.3	80/30/4
	<i>Tricyrtis macropoda</i>	MT261173	155 453	37.4	80/30/4
Smilacaceae	<i>Smilax china</i>	HM536959	157 878	37.25	80/30/4
	<i>Smilax glycyphylla</i>	MT261169	158 922	36.9	80/30/4
	<i>Smilax microphylla</i>	MW423607	158 246	37.1	80/30/4

	<i>Smilax nipponica</i>	MT261170	158 178	37.1	80/30/4
Philesiaceae	<i>Ripogonum scandens</i>	MT261167	160 287	37.6	80/30/4
	<i>Philesia magellanica</i>	MT261166	158 786	37.6	80/30/4
	<i>Lapageria rosea</i>	MT261158	160 054	37.5	80/30/4
Melanthiaceae (49 species)	<i>Chionographis japonica</i>	KF951065	154 646	37.7	79/30/4
	<i>Heloniopsis tubiflora</i>	KM078036	157 940	37.5	80/30/4
	<i>Paris axialis</i>	MN125591	156 821	37.4	80/30/4
	<i>Paris bashanensis</i>	MN125580	157 320	37.7	80/30/4
	<i>Paris birmanica</i>	MN125580	157 857	37.3	80/30/4
	<i>Paris caobangensis</i>	MN125593	158 256	37.2	80/30/4
	<i>Paris caojianensis</i>	MZ147601	163 853	37	80/30/4
	<i>Paris cronquistii</i>	KX784041	157 710	37.3	80/30/4
	<i>Paris daliensis</i>	MN125574	158 118	37.3	80/30/4
	<i>Paris delavayi</i>	MN125581	158 575	37.2	80/30/4
	<i>Paris dulongensis</i>	MN125566	157 342	37.4	80/30/4
	<i>Paris dunniana</i>	KX784042	157 984	37.2	80/30/4
	<i>Paris fargesii</i>	KX784043	157 518	37.3	80/30/4
	<i>Paris forrestii</i>	KX784044	158 345	37.3	80/30/4
	<i>Paris incompleta</i>	MN125572	157 610	37.7	80/30/4
	<i>Paris japonica</i>	MH796668	155 957	37.6	80/30/4
	<i>Paris liiana</i>	MT857225	163 860	37	80/30/4
	<i>Paris luquanensis</i>	KX784045	158 451	37.3	80/30/4
	<i>Paris marei</i>	KX784046	157 891	37.3	80/30/4
	<i>Paris marmorata</i>	KX784047	157 566	37.3	80/30/4
	<i>Paris polyphylla var chinensis</i>	KX784048	158 307	37.2	80/30/4
	<i>Paris polyphylla var yunnanensis</i>	KX784049	157 547	37.3	80/30/4
	<i>Paris qiliangiana</i>	MN125576	158 354	37.2	80/30/4
	<i>Paris quadrifolia</i>	KX784051	157 097	37.7	80/30/4
	<i>Paris rugosa</i>	MN125570	157 239	37.4	80/30/4
	<i>Paris stigmatosa</i>	MN125570	157 239	36.8	80/30/4
	<i>Paris tengchongensis</i>	MN125584	157 150	37.4	80/30/4
	<i>Paris tetraphylla</i>	MN125596	156 567	37.5	80/30/4
	<i>Paris tibetica</i>	MN125596	157 389	37.4	80/30/4
	<i>Paris undulata</i>	MN125586	158 286	37.2	80/30/4
	<i>Paris vaniotii</i>	MN125567	156 846	37.4	80/30/4
	<i>Paris verticillata</i>	KJ433485	157 379	37.6	80/30/4
	<i>Paris vietnamensis</i>	KX784050	158 224	37.2	80/30/4
<i>Paris xichouensis</i>	MN125585	158 225	37.3	80/30/4	
<i>Paris yanchii</i>	MN125582	157 918	37.3	80/30/4	
<i>Trillium camschatcense</i>	MN125568	156 139	37.5	80/30/4	
<i>Trillium cuneatum</i>	NC027185	156 610	37.5	80/30/4	
<i>Trillium decumbens</i>	NC027282	158 552	37.7	80/30/4	
<i>Trillium govanianum</i>	MH796670	157 379	37.7	80/30/4	
<i>Trillium maculatum</i>	KR780075	157 359	37.5	80/30/4	
<i>Trillium tschonoskii</i>	KR780076	156 852	37.5	80/30/4	

	<i>Veratrum japonicum</i>	MG940972	151 791	37.7	80/30/4
	<i>Veratrum mengtzeanum</i>	MW147219	153 705	37.8	80/30/4
	<i>Veratrum oxysepalum</i>	MW147219	153 705	37.7	80/30/4
	<i>Veratrum patulum</i>	KF437397	153 699	37.7	80/30/4
	<i>Veratrum taliense</i>	MN125578	151 909	37.8	80/30/4
	<i>Xerophyllum tenax</i>	KM078035	156 746	37.8	80/30/4
	<i>Ypsilandra thibetica</i>	MH796671	157 613	37.5	80/30/4
	<i>Ypsilandra yunnanensis</i>	MH796672	158 806	37.4	80/30/4
Alstroemeriaceae	<i>Alstroemeria aurea</i>	KC968976	155 510	37.26	80/30/4
	<i>Bomarea edulis</i>	KM233641	154 925	38.2	80/30/4
	<i>Luzuriaga radicans</i>	KM233640	157 885	38.1	80/30/4
Colchicaceae (9 species)	<i>Androcymbium greuterocymbium</i>	MT261148	154 804	37.6	80/30/4
	<i>Colchicum autumnale</i>	KP125337	156 462	37.6	80/30/4
	<i>Disporum cantoniense</i>	MW759302	156 688	37.6	80/30/4
	<i>Disporum sessile</i>	MN332241	159 102	37.3	80/30/4
	<i>Gloriosa superba</i>	KP125338	157 924	37.6	80/30/4
	<i>Iphgenia indica</i>	MT012417	158 319	37.4	80/30/4
	<i>Tripladenia cunninghamii</i>	MT261174	155 652	37.6	80/30/4
	<i>Uvularia grandiflora</i>	MT261175	157 025	37.6	80/30/4
	<i>Wurmbea burtii</i>	MT261176	155 297	37.7	80/30/4
Petermanniaceae	<i>Petermannia cirrhosa</i>	MT261165	156 852	38	80/30/4
Campynemataceae	<i>Campynema lineare</i>	MT261151	156 305	36.9	80/30/4
Corsiaceae	<i>Corsia dispar</i>	MT261154	63 172	30.8	30/24/4
	<i>Arachnitis uniflora</i>	MT261149	24 846	37.1	16/05/4

Table 2 Gene contents in the chloroplast genomes of Liliales taxa

Group of gene		Name of gene(common)
RNA genes	Ribosomal RNAs	rrn4.5(x2) xz, rrn5(x2) xz, rrn16(x2) xz, rrn23(x2) xz
	Transfer RNAs	trnA-UGCa(x2) x, trnC-GCAxz, trnD-GUCx, trnE-UUCxz, trnF-GAAx, trnM-CAUxz, trnG-GCC, trnG-UCCa, trnH-GUGx, trnI-CAU(x2) x, trnI-GAUa(x2), trnK-UUUax, trnL-CAA(x2) x, trnL-UAAax, trnL-UAGx, trnM-CAUx, trnN-GUU(x2) x, trnP-UGGx, trnQ-UUGxz, trnR-ACG(x2) x, trnR-UCU, trnS-GCUx, trnS-GGAX, trnS-UGAX, trnT-GGUx, trnT-UGUx, trnV-GAC(x2) x, trnV-UACa, trnW-CCAxz, trnY-GUAX
Protein genes	Photosystem I	psaAx, psaB, psaC, psaI, psaJ
	Photosystem II	psbA, psbB, psbC, psbD, psbE, psbF, psbH, psbI, psbJ, psbK, psbL, psbM, psbN, psbT, psbZ
	Cytochrome	petA, petB, petD, petG, petL, petN
	ATP synthase	atpA, atpBx, atpEx, atpFa, atpH, atpIx
	Rubisco	rbcL
	NADH dehydrogenase	ndhAa, ndhBa(x2)x, ndhC, ndhD, ndhE, ndhF, ndhG, ndhH, ndhI, ndhJ, ndhK
	ATP-dependent protease subunit P	clpPaxz

	Chloroplast envelope membrane protein	cemA
Ribosomal proteins	large units	rpl2a(x2) xz, rpl14xz, rpl16xz, rpl20xz, rpl22x, rpl23 (x2), rpl32x, rpl33x, rpl36x
	small units	rps2xz, rps3xz, rps4xz, rps7(x2) xz, rps8xz, rps11xz, rps12a(x2) xz, rps14xz, rps15x, rps16x, rps18x, rps19(x2) xz
Transcription /translation	RNA polymerase	rpoA, rpoB, rpoC1a,rpoC2
	Initiation factor	infA
	Miscellaneous proteins	accDxz, ccsA, matKx
	Hypothetical proteins & Conserved reading frame	ycf1x, ycf2 (x2) x, ycf3, ycf4x, ycf15
a: gene has intron; x2: gene has two copies; x : remained in Corsia dispar; z : remained in Arachnitis uniflora		

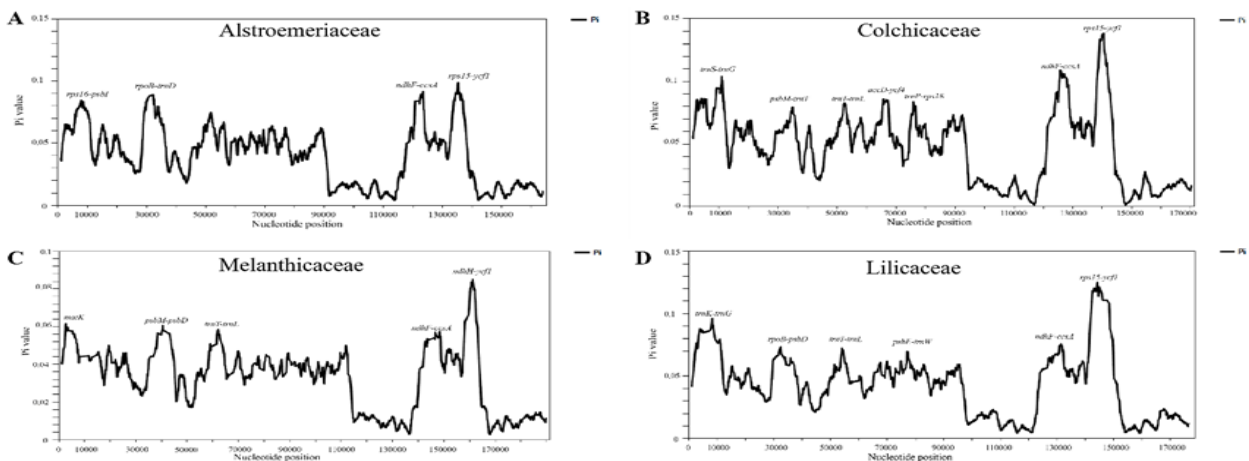
In the first stage, the *ndh* genes were lost, followed by the disappearance of photosynthetic genes. In the third stage, the genes for RNA polymerase were not found. In the fourth and fifth stages, genes for ATP synthase and other functions were lost, respectively. In Corsiaceae of Liliales, the cpDNA of *Arachnitis uniflora* and *Corsia dispar* are in the third and the fourth stages (Table 2). However, there are only two out of 26 species of Corsiaceae that have available cpDNA on NCBI data. Therefore, further studies that cover all species of Corsiaceae should be conducted to provide a better understanding of the evolutionary history of mycoheterotrophic species in Liliales.

Among photosynthetic species of Liliales, there are records of *infA* and *rps16* loss in cpDNA sequences of *Amana* and *Chionographis* species (Table 1). Previously, the loss of *infA* in cpDNA was detected in many angiosperms and the intact *infA* was found in nucleus [26]. Similarly, the loss of *rps16* was also found in other angiosperms that was compensated by a copy of *rps16* in the nucleus genome [27-28]. In

Liliales, the loss of gene was recorded but there is no study on the effect of that loss in cpDNA. Therefore, further studies on the impact of gene loss in photosynthetic as well as mycoheterotrophic species of Liliales should be conducted.

3.2 Nucleotide diversity patterns in chloroplast genomes of Liliales

The nucleotide diversity analysis revealed different P_i values among families of Liliales (Figure 1). The high P_i values (> 0.1) were recorded in Alstroemeriaceae, Colchicaceae and Liliaceae (Figure 1A, 1B, 1D) whereas Philesiaceae and Smilacaceae have smaller P_i values (< 0.04) (Figure 1E, 1F). In Melanthiaceae, the high P_i values range from 0.04 to 0.1 (Figure 1C). In Alstroemeriaceae, the high P_i values were found in *rps16-psbI*, *rpoB-trnD*, *ndhF-ccsA* and *rps15-ycf1* (Figure 1A). In Colchicaceae, *trnS-trnG*, *psbM-trnT*, *trnT-trnL*, *accD-ycf4*, *trnP-rps18*, *ndhF-ccsA* and *rps15-ycf1* exhibit high P_i values (Figure 1B). In Melanthiaceae, five regions including *matK*, *psbM-psbD*, *trnT-trnL*, *ndhF-ccsA* and *ndhH-ycf1* have high P_i values.



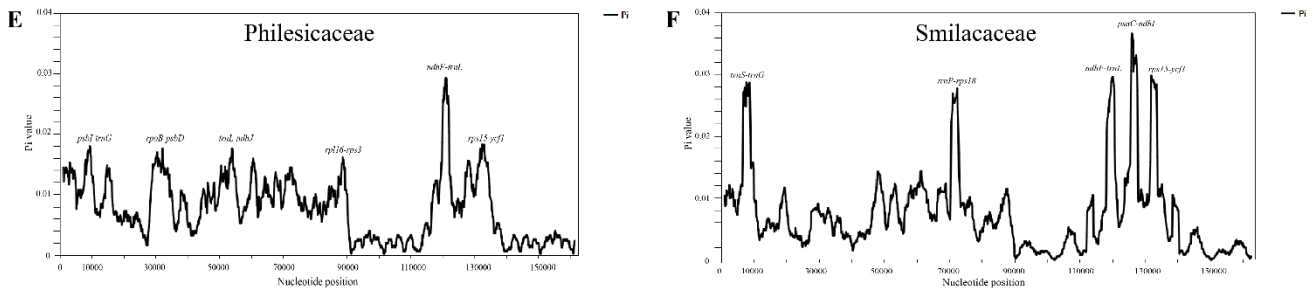


Figure 1 Sliding window analysis of the whole chloroplast genomes of Liliales species. (window length: 2 000 bp, step size: 100 bp). X-axis: position of nucleotide, Y-axis: Pi values of each window.

A. Alstroemeriaceae; B. Colchicaceae; C. Liliaceae; D. Melanthiaceae; E. Philesiaceae; F: Smilacaceae

In Liliaceae, *trnK-trnG*, *rpoB-psbD*, *trnT-trnL*, *psbE-trnW*, *ndhF-ccsA* and *rps15-ycf1* regions showed high Pi values. In Philesiaceae, high Pi values were found in *psbI-trnG*, *rpoB-psbD*, *trnL-ndhJ*, *rpl16-rps3*, *ndhF-trnL* and *rps15-ycf1* (Figure 1E). In Smilacaceae, five regions have high Pi values including *trnS-trnG*, *trnP-rps18*, *ndhF-trnL*, *psaC-ndhI* and *rps15-ycf1* (Figure 1F). Most of high Pi values were found in non-coding regions but some coding regions such as *matK* and *ycf1* also had high nucleotide diversity.

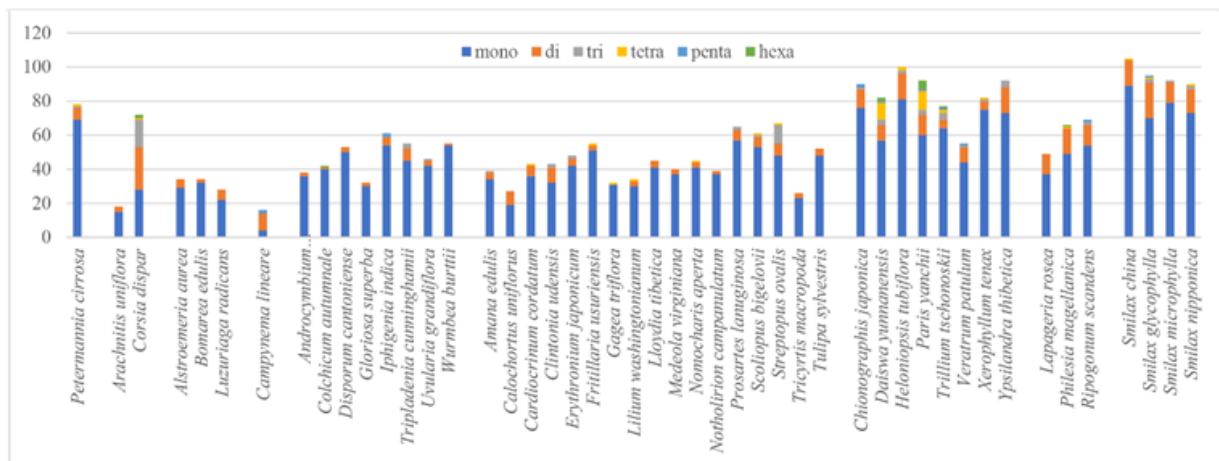
Similar to Liliales, nucleotide diversity has been explored in cpDNA of various angiosperms. For example, in *Paris* species (Melanthiaceae), divergent hotspots were found in both coding regions (*rpoC1* and *ycf2*) and non-coding regions (*trnS-trnG*, *rpl32-trnL*, etc.) [21]. In other land plants such as species of *Symplocos*, *Avena* and *Senecioneae*, various hotspots with high Pi values were located in different regions of cpDNA [29-31]. The information of hotspots is a useful source for developing molecular markers in

angiosperms [32]. In Liliales, highly variable regions were identified for each family, except Corsiaceae, Petermanniaceae and Campynemataceae due to the lack of data and mycoheterotrophic lifestyle. Among the hotspots, *ycf1* is the common region in observed families. However, this region should be verified with the lack of data from Campynemataceae and Petermanniaceae in further studies.

3.3 Comparison of Repeat composition in chloroplast genomes in Liliales

The analysis of SSRs in cpDNA of Liliales resulted in different numbers of repeats among families (Figure 2A). The highest number of SSRs was found in *Smilax china* (105 records) whereas *Campynema lineare* and *Arachnitis uniflora* have 16 and 18 SSRs in cpDNA, respectively. Among six types of SSR, the mononucleotide repeat (A/T) is the most abundant (2191 records) followed by dinucleotide (AT/TA/GC/CG) type (339 records).

A



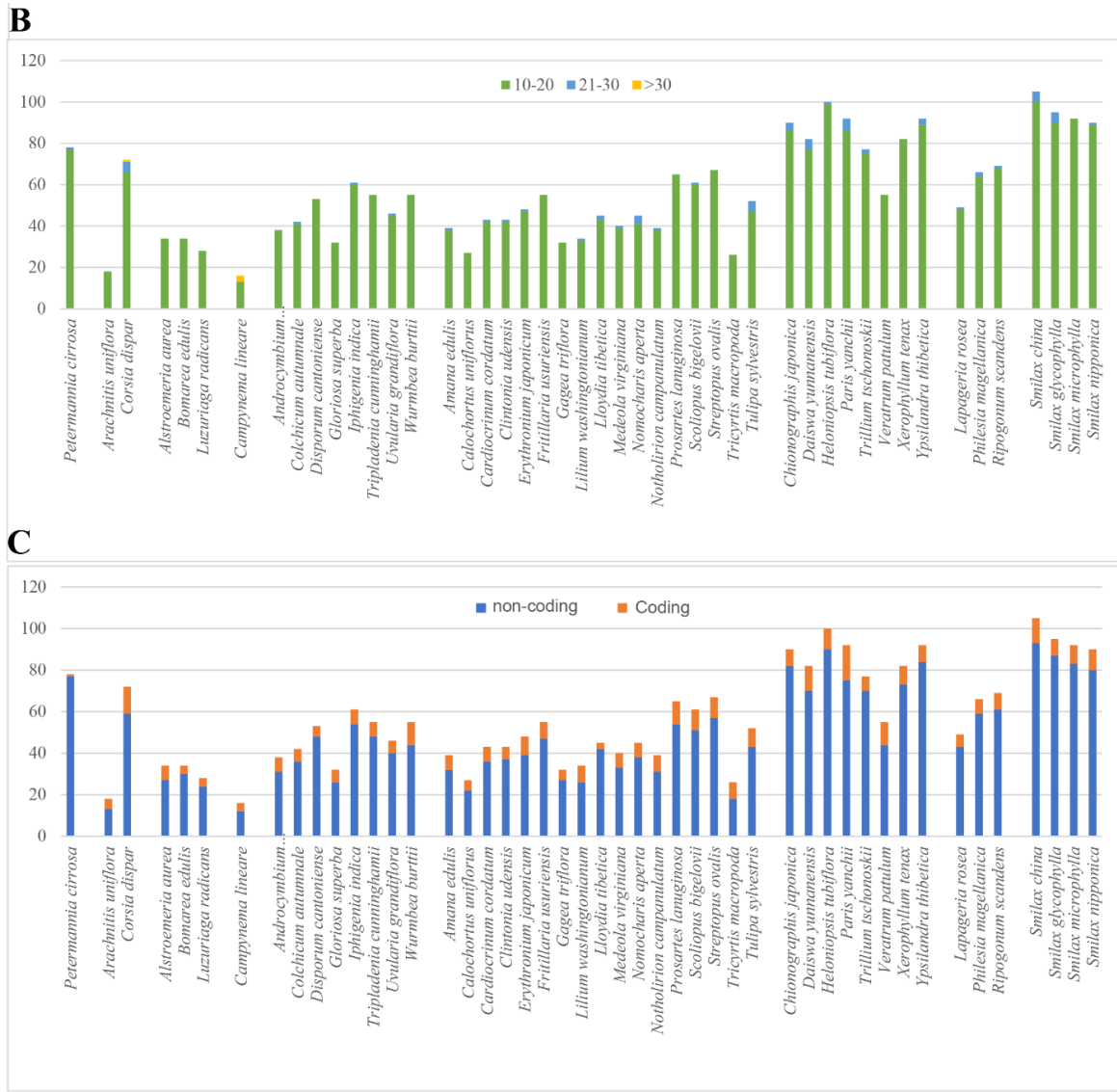
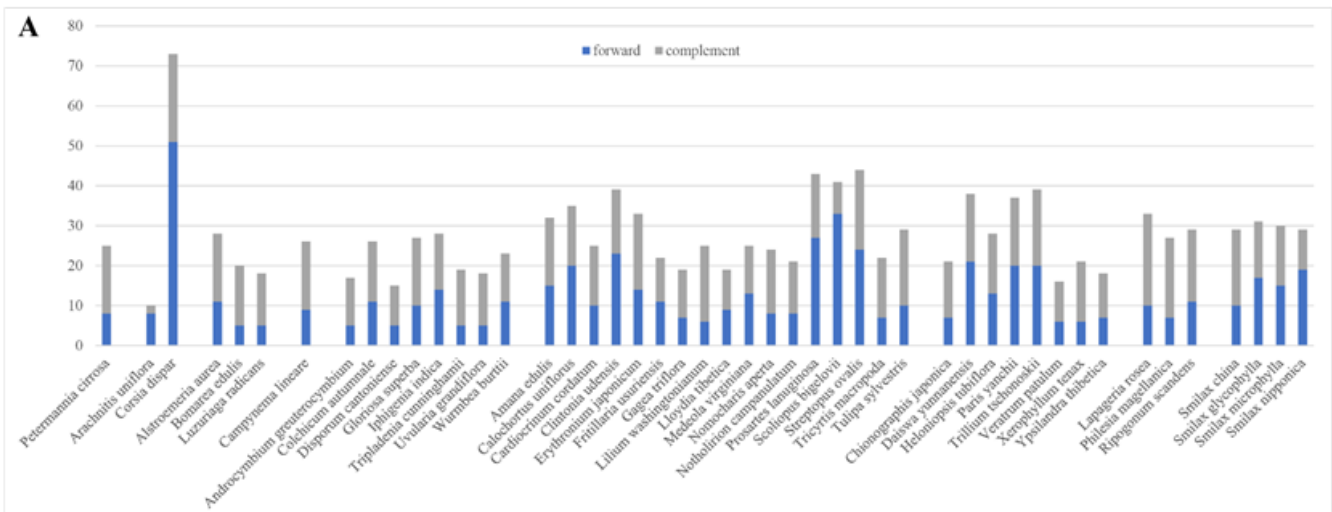


Figure 2 Quantity of SSR in chloroplast genomes of Liliales. A. Types of SSR; B. Length of SSR; C. Location of SSR.



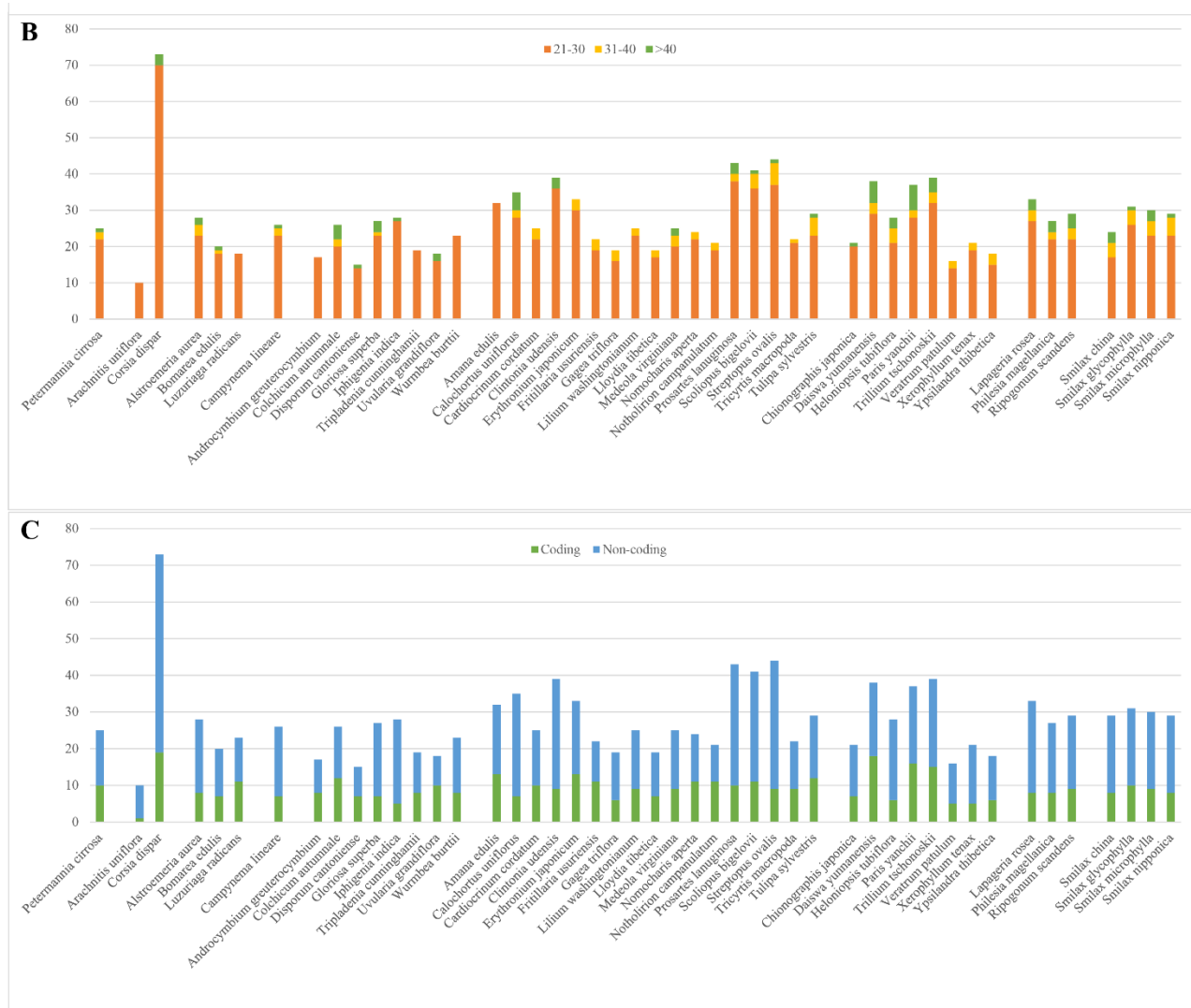


Figure 3 Quantity of long repeat in chloroplast genomes of Liliales. A. Types of repeat; B. Length of repeat; C. Location of repeat.

The types of tri-, tetra-, penta- and hexanucleotide are not common in Liliales, except Melanthiaceae members of which cpDNAs have a total of 59 records of these four types (AAT/ACAT/ATATC/AAAAT/AAAGAG). Although Melanthiaceae members possessed a larger number of repeats compared to other Liliales's families, the repeats in the chloroplast genome do not affect the morphological characteristics of Melanthiaceae, encoded by nuclear genes. The lengths of SSR varied across Liliales taxa (Figure 2B). Most of SSRs (2 591 units) have the lengths of up to 20 bp whereas only 64 SSRs have the lengths from 21 to 30 bp. Although *Campynema lineare* has the smallest number of SSRs in comparison to other taxa, it

contains three SSRs of which the length is over 30 bp. The location of SSRs is mainly in non-coding regions; however, 13.6 % of SSR was found in coding regions (Figure 2C). In Liliales cpDNAs, the coding regions containing SSRs are *rpoC1*, *rpoC2*, *rpoB*, *ycf1*, *cemA*, *psbD*, *psbC*, *psbF*, *accD*, *ndhF*, *ndhG*, *ndhI*, *rps2*, *rps7*, *rps14*, *rps19*, *rps3* and *atpB*. Among surveyed species of Liliales, there are 597 records of forward repeats and 700 complement repeats in cpDNA (Figure 3A). The highest number of repeats was detected in *Corsia dispar* (73 repeats) whereas *Arachnitis uniflora* only has 10 repeats including eight forward and two complement units. In most cpDNAs, the complement repeat exceeded; however, more forward repeats were recorded in

cpDNAs of *Corsia dispar*, *Arachnitis uniflora*, *Calochortus uniflorus*, *Clintonia udensis*, *Medeola virginiana*, *Prosartes lanuginosa*, *Scoliopus bigelovii*, *Streptopus ovalis*, *Daiswa yunnanensis*, *Paris yanchii*, *Trillium tschonoskii*, *Smilax glycyphylla* and *Smilax nipponica* (Figure 3A). The lengths of repeats are mostly shorter than 30 bp (Figure 3B). Only 13 % of repeat has the length over 30 bp. Similar to SSRs, the repeats are located mainly in non-coding regions (Figure 3C). In coding areas, the forward and complement repeats were found in *psaA*, *psaB*, *rpoC2*, *ycf1*, *ycf2*, *ndhF*, *ndhI*, *trnS*, *trnfM*, and *trnG*.

In chloroplast genomes, SSRs and repeats are useful information for tracking the evolution of the plants. The SSRs can be used to develop molecular markers for population genetics and identification of plants [17,33]. Additionally, SSR markers can be used for testing the breeding of plants [34,35]. Beside SSRs, the repeats are important factors affecting the structure of cpDNA during the evolutionary history [36,37]. Repeat is also the cause of new repeated generations in cpDNA [38]. In Liliales, *Corsia dispar*, a mycoheterotrophic species that is in the third stage of cpDNA structural change, has the highest number of repeats and does not have a typical quadripartite structure, suggesting the high impact of repeats on the plastid genome structure of *Corsia* species. The mycoheterotrophic lifestyle does not require photosynthesis; therefore, genes related to performing and controlling photosynthetic progress are not necessary in the plastid genome of *Corsia* species.

References

- 1.P. F. S. The Angiosperm Phylogeny Group, M. W. Chase, M. J. M. Christenhusz, M. F. Fay, J. W. Byng, W. S. Judd, D. E. Soltis, D. J. Mabberley, A. N. Sennikov, P. S. Soltis, "An update of the Angiosperm Phylogeny Group classification for the orders and families of flowering plants: APG IV," *Bot. J. Linn. Soc.*, vol. 181, no. 1, pp. 1–20, May 2016.
- 2.J. David L. and G. Bruce, "Corsia dispar D.L.Jones & B.Gray (Corsiaceae), a new species from Australia, and a new combination in Corsia for a New Guinea taxon," *Austrobaileya*, vol. 7, no. 4, pp. 717–722, 2008.
- 3.T. J. Givnish *et al.*, "Phylogenomics and historical biogeography of the monocot order Liliales: out of Australia and through Antarctica," *Cladistics*, vol. 32, no. 6, pp. 581–605, Dec. 2016.
- 4.J. S. Kim and J.-H. Kim, "Updated molecular phylogenetic analysis, dating and biogeographical history of the lily family (Liliaceae: Liliales)," *Bot. J. Linn. Soc.*, vol. 187, no. 4, pp. 579–593, Jul. 2018.
- 5.C. Kim, S.-C. Kim, and J.-H. Kim, "Historical Biogeography of Melanthiaceae: A Case of Out-of-North America Through the Bering Land Bridge," *Front. Plant Sci.*, vol. 10, Apr. 2019.
- 6.J. Chacón and S. S. Renner, "Assessing model sensitivity in ancestral area reconstruction using L <sc>agrange</sc> : a case study using the Colchicaceae family," *J. Biogeogr.*, vol. 41, no. 7, pp. 1414–1427, Jul. 2014.

Consequently, various genes in plastid were lost during evolutionary history. In the plastid genome, repeats initiated the deletion of genes as well as non-coding regions. At the present, only one complete plastid genome of *Corsia* has been reported. Therefore, more samples of *Corsia* should be sampled to investigate the effectiveness of repeats in the structural change of Corsiaceae of which *Arachnitis uniflora* has smallest number of repeats and remains the typical quadripartite structure.

4 Conclusions

Complete chloroplast genomes of Liliales were surveyed and the analysis of nucleotide diversity revealed various hotspots among the families of Liliales in both coding and non-coding regions. Additionally, various types of repeats were identified in representative species of Liliales that are crucial sources for further studies on population genetics and development of molecular markers. Last but not least, more samples of Corsiaceae, Campynemataceae and Colchicaceae should be collected to cover the gaps within those families for fulfilling the complete evolutionary history of the chloroplast genome in Liliales.

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7. Z. Qi *et al.*, “Phylogenetics, character evolution, and distribution patterns of the greenbriers, Smilacaceae (Liliales), a near-cosmopolitan family of monocots,” *Bot. J. Linn. Soc.*, vol. 173, no. 4, pp. 535–548, Dec. 2013.
8. C. Chen *et al.*, “Understanding the formation of Mediterranean–African–Asian disjunctions: evidence for Miocene climate-driven vicariance and recent long-distance dispersal in the Tertiary relict *Smilax aspera* (Smilacaceae),” *New Phytol.*, vol. 204, no. 1, pp. 243–255, Oct. 2014.
9. H. Daniell, C.-S. Lin, M. Yu, and W.-J. Chang, “Chloroplast genomes: diversity, evolution, and applications in genetic engineering,” *Genome Biol.*, vol. 17, no. 1, p. 134, Dec. 2016.
10. E. J. Carpenter *et al.*, “Access to RNA-sequencing data from 1,173 plant species: The 1000 Plant transcriptomes initiative (1KP),” *Gigascience*, vol. 8, no. 10, Oct. 2019.
11. S. Cheng *et al.*, “10KP: A phylodiverse genome sequencing plan,” *Gigascience*, vol. 7, no. 3, Mar. 2018.
12. M. A. Gitzendanner, P. S. Soltis, G. K.-S. Wong, B. R. Ruhfel, and D. E. Soltis, “Plastid phylogenomic analysis of green plants: A billion years of evolutionary history,” *Am. J. Bot.*, vol. 105, no. 3, pp. 291–301, Mar. 2018.
13. H. Robert J, *Molecular Markers in Plants*. Oxford, UK: Blackwell Publishing Ltd., 2012.
14. K. Semagn, Å Bjørnstad, and M. N. Ndjiondjop, “An overview of molecular marker methods for plants,” *African J. Biotechnol.*, vol. 5, no. 25, pp. 2540–2568, 2006.
15. T. N. Vu *et al.*, “Molecular markers for analysis of plant genetic diversity,” *Vietnam J. Biotechnol.*, vol. 18, no. 4, pp. 589–608, May 2021.
16. J. Hyun, H. D. K. Do, J. Jung, and J.-H. Kim, “Development of molecular markers for invasive alien plants in Korea: a case study of a toxic weed, *Cenchrus longispinus* L., based on next generation sequencing data,” *PeerJ*, vol. 7, p. e7965, Nov. 2019.
17. M. L. C. Vieira, L. Santini, A. L. Diniz, and C. de F. Munhoz, “Microsatellite markers: what they mean and why they are so useful,” *Genet. Mol. Biol.*, vol. 39, no. 3, pp. 312–328, Aug. 2016.
18. H. D. K. Do, C. Kim, M. W. Chase, and J. Kim, “Implications of plastome evolution in the true lilies (monocot order Liliales),” *Mol. Phylogenet. Evol.*, vol. 148, p. 106818, Jul. 2020.
19. H. D. K. Do, J. S. Kim, and J.-H. Kim, “Comparative genomics of four Liliales families inferred from the complete chloroplast genome sequence of *Veratrum patulum* O. Loes. (Melanthiaceae),” *Gene*, vol. 530, no. 2, 2013.
20. S.-C. Kim, J. S. Kim, and J.-H. Kim, “Insight into infrageneric circumscription through complete chloroplast genome sequences of two *Trillium* species,” *AoB Plants*, vol. 8, p. plw015, 2016.
21. Y. Song *et al.*, “Chloroplast Genomic Resource of Paris for Species Discrimination,” *Sci. Rep.*, vol. 7, no. 1, p. 3427, Dec. 2017.
22. H. D. K. Do and J.-H. Kim, “The implication of plastid transcriptome analysis in petaloid monocotyledons: A case study of *Lilium lancifolium* (Liliaceae, Liliales),” *Sci. Rep.*, vol. 9, no. 1, 2019.
23. T. Braukmann and S. Stefanović, “Plastid genome evolution in mycoheterotrophic Ericaceae,” *Plant Mol. Biol.*, vol. 79, no. 1–2, pp. 5–20, May 2012.
24. S. W. Graham, V. K. Y. Lam, and V. S. F. T. Merckx, “Plastomes on the edge: the evolutionary breakdown of mycoheterotroph plastid genomes,” *New Phytol.*, vol. 214, no. 1, pp. 48–55, Apr. 2017.
25. C. F. Barrett and J. I. Davis, “The plastid genome of the mycoheterotrophic *Corallorhiza striata* (Orchidaceae) is in the relatively early stages of degradation,” *Am. J. Bot.*, vol. 99, no. 9, pp. 1513–1523, Sep. 2012.
26. R. S. Millen *et al.*, “Many Parallel Losses of *infA* from Chloroplast DNA during Angiosperm Evolution with Multiple Independent Transfers to the Nucleus,” *Plant Cell*, vol. 13, no. 3, pp. 645–658, Mar. 2001.
27. A. A. Alqahtani and R. K. Jansen, “The evolutionary fate of *rpl32* and *rps16* losses in the *Euphorbia schimperii* (Euphorbiaceae) plastome,” *Sci. Rep.*, vol. 11, no. 1, p. 7466, Dec. 2021.
28. M. Ueda *et al.*, “Substitution of the Gene for Chloroplast RPS16 Was Assisted by Generation of a Dual Targeting Signal,” *Mol. Biol. Evol.*, vol. 25, no. 8, pp. 1566–1575, Apr. 2008.
29. S.-C. Kim, J.-W. Lee, and B.-K. Choi, “Seven Complete Chloroplast Genomes from *Symplocos*: Genome Organization and Comparative Analysis,” *Forests*, vol. 12, no. 5, p. 608, May 2021.
30. Q. Liu, X. Li, M. Li, W. Xu, T. Schwarzacher, and J. S. Heslop-Harrison, “Comparative chloroplast genome

- analyses of *Avena*: insights into evolutionary dynamics and phylogeny,” *BMC Plant Biol.*, vol. 20, no. 1, p. 406, Dec. 2020.
31. A. W. Gichira, S. Avoga, Z. Li, G. Hu, Q. Wang, and J. Chen, “Comparative genomics of 11 complete chloroplast genomes of Senecioneae (Asteraceae) species: DNA barcodes and phylogenetics,” *Bot. Stud.*, vol. 60, no. 1, p. 17, Dec. 2019.
32. W. Dong, J. Liu, J. Yu, L. Wang, and S. Zhou, “Highly Variable Chloroplast Markers for Evaluating Plant Phylogeny at Low Taxonomic Levels and for DNA Barcoding,” *PLoS One*, vol. 7, no. 4, p. e35071, Apr. 2012.
33. C. Li, Y. Zheng, and P. Huang, “Molecular markers from the chloroplast genome of rose provide a complementary tool for variety discrimination and profiling,” *Sci. Rep.*, vol. 10, no. 1, p. 12188, Dec. 2020.
34. A.-H. Yang, J.-J. Zhang, X.-H. Yao, and H.-W. Huang, “Chloroplast microsatellite markers in *Liriodendron tulipifera* (Magnoliaceae) and cross-species amplification in *L. chinense*,” *Am. J. Bot.*, vol. 98, no. 5, pp. e123–e126, May 2011.
35. M. Yang *et al.*, “Genetic linkage maps for Asian and American lotus constructed using novel SSR markers derived from the genome of sequenced cultivar,” *BMC Genomics*, vol. 13, no. 1, p. 653, Dec. 2012.
36. F. Yue, L. Cui, C. W. DePamphilis, B. M. Moret, and J. Tang, “Gene rearrangement analysis and ancestral order inference from chloroplast genomes with inverted repeat,” *BMC Genomics*, vol. 9, no. S1, p. S25, Mar. 2008.
37. J. D. Palmer, B. Osorio, J. Aldrich, and W. F. Thompson, “Chloroplast DNA evolution among legumes: Loss of a large inverted repeat occurred prior to other sequence rearrangements,” *Curr. Genet.*, vol. 11, no. 4, pp. 275–286, Jan. 1987.
38. H. D. K. Do and J.-H. Kim, “A dynamic tandem repeat in monocotyledons inferred from a comparative analysis of chloroplast genomes in melanthiaceae,” *Front. Plant Sci.*, vol. 8, 2017.

Phân tích đa dạng di truyền bộ gen lục lạp ở bộ Loa kèn

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Tóm tắt Bộ Loa kèn là một bộ thực vật một lá mầm và bao gồm cả loài thực vật tự dưỡng và dị dưỡng cộng sinh với nấm; phân bố rộng khắp hoặc cục bộ tại một số vùng nhất định. Trong nghiên cứu này, đa dạng di truyền của bộ Loa kèn được khảo sát thông qua phân tích đa dạng nucleotide và thành phần các loại trình tự lặp trong bộ gen lục lạp. Kết quả phân tích đa dạng nucleotide cho thấy rất nhiều trình tự có biến động cao trong vùng trình tự đơn lớn (LSC) và vùng trình tự đơn nhỏ (SSC) trong khi vùng trình tự lặp đảo thì có mức biến động thấp. Mặc dù từng họ trong bộ Loa kèn có các trình tự biến động đặc trưng nhưng vùng trình tự *rps15-ycf1* có biến động cao được tìm thấy hầu hết trong các bộ gen lục lạp. Trong bộ gen lục lạp của bộ Loa kèn, các trình tự lặp đơn giản (SSR) loại nucleotide đơn là loại phổ biến và hầu hết các trình tự SSR nằm ở vùng không mã hóa. Tương tự như vậy, các trình tự lặp dài cũng chủ yếu được tìm thấy ở vùng không mã hóa. Ngoài ra, trình tự lặp đảo là trình tự lặp phổ biến hơn so với trình tự lặp liên tục trong bộ gen lục lạp của bộ Loa kèn. Số lượng trình tự lặp dài cao nhất được tìm thấy trong bộ gen lục lạp của loài *Corsia dispar* trong khi trình tự lặp đơn giản được xác định nhiều nhất trong loài *Smilax china*. Các kết quả nghiên cứu đa dạng nucleotide và trình tự lặp sẽ cung cấp các thông tin nền tảng cho các nghiên cứu tiếp theo trong lĩnh vực di truyền quần thể, chỉ thị phân tử và lịch sử tiến hóa của bộ Loa kèn.

Từ khóa bộ Loa kèn, bộ gen lục lạp, đa dạng nucleotide, giá trị Pi, trình tự lặp.

