Comparison of intraspecific, interspecific and intergeneric chloroplast diversity in Cycads

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Cycads are among the most threatened plant species. Increasing the availability of genomic information by adding whole chloroplast data is a fundamental step in supporting phylogenetic studies and conservation efforts. Here, we assemble a dataset encompassing three taxonomic levels in cycads, including ten genera, three species in the genus Cycas and two individuals of C. debaoensis. Repeated sequences, SSRs and variations of the chloroplast were analyzed at the intraspecific, interspecific and intergeneric scale, and using our sequence data, we reconstruct a phylogenomic tree for cycads. The chloroplast was 162,094 bp in length, with 133 genes annotated, including 87 protein-coding, 37 tRNA and 8 rRNA genes. We found 7 repeated sequences and 39 SSRs. Seven loci showed promising levels of variations for application in DNA-barcoding. The chloroplast phylogeny confirmed the division of Cycadales in two suborders, each of them being monophyletic, revealing a contradiction with the current family circumscription and its evolution. Finally, 10 intraspecific SNPs were found. Our results showed that despite the extremely restricted distribution range of C. debaoensis, using complete chloroplast data is useful not only in intraspecific studies, but also to improve our understanding of cycad evolution and in defining conservation strategies for this emblematic group.

Cycads are iconic relict species, regarded as “living fossils” because of their recognizable intermediate morphological traits between angiosperms and gymnosperms. Cycads dominated the Mesozoic but their origin can be dated to the late Paleozoic (~265–290 Ma). However, molecular dating studies indicate that living cycad species could be not much older than ~12 Ma, rejecting both the hypothesized role of dinosaurs in generating extant diversity and the use of “living fossils” to describe current cycad species. Cycads are distributed in tropical and subtropical regions of Africa, Asia, Oceania and America. Ten genera and 344 species are currently accepted, with Cycas containing roughly 40% of the species in the Near Threatened and Vulnerable categories in the IUCN red list.

The genus Cycas L., in the monotypic family Cycadaceae, is the oldest genus of cycads, holding about 113 species. More than 20 species are found in China, with most of them endemic. Cycas debaoensis Y. C. Zhong & C. J. Chen, a critically endangered cycad species endemic to southwest China, only occurs in 11 small populations near the border of Guangxi province and Yunnan province. Previous studies have assessed genetic diversity in C. debaoensis using inter simple sequence repeat (ISSR) markers or nuclear microsatellites, showing limited gene flow among populations and low within-population diversity.

Chloroplasts (cps) are present in photosynthetically active green tissues and generally develop from proplastids in meristems or etioplasts after illumination of dark-grown tissues, and display a conserved structure of two inverted repeats (IR) separated by small (SSC) and large (LSC) single-copy regions. Due to their natural abundance in plant cells (∼3–5% of the cell DNA content comparing to nuclear DNA), cp sequences are a versatile tool for plant identification (DNA-barcoding) and evolutionary studies. They have been used at small and large temporal scales in plants. The use of cps is a very powerful tool to reconstruct plant phylogenies and infer historical biogeographic patterns of diversification. However, only a limited number of regions in the chloroplast genome have been used to address evolutionary, taxonomic and biodiversity questions in Cycas. With the rapid development of Next Generation Sequencing (NGS), it is now feasible to obtain the entire sequence of
the chloroplast using a genome skimming approach, resulting in high resolution phylogenies and allowing for estimations of timing of historical diversification, biodiversity and extent of genomic divergence\textsuperscript{14,20,21}.

It is well known that genetic diversity can greatly vary between taxa, due to either different intrinsic characteristics (e.g. reproductive system, genome size and organization) or to extrinsic features (e.g. endemic vs. widespread species, young vs. old species)\textsuperscript{17,18,22}. In addition, the hypothesis of a linear accumulation of mutations in sequences across time (i.e. a molecular clock) has been refuted in many groups\textsuperscript{23–25}.

In this study, we analyse the \textit{C. debaoensis} chloroplast as a reference together with molecular data of other \textit{Cycas} species to identify potential DNA-barcode loci, and compared generic-level chloroplast features in cycads, to highlight the evolutionary history of this group. We also compared the chloroplast features of two individuals of \textit{C. debaoensis} to provide new resources for marker development in this endangered species.

## Results and Discussion

### Genome size and features.

Using genome skimming and reference-guided assembly, we reconstructed the 162,094 bp long chloroplast genome \textit{C. debaoensis\_Jiang\_DB-2015}. The complete cp genome was submitted...
to GenBank under accession number KU743927. It was 2 bp longer than C. deboeaeosis (KM459003) due to two 1 bp indels in the LSC (Fig. 1). The two C. deboeaeosis individuals exhibited the typical composition of LSC, SSC regions and two IR copies of 88,854 bp, 23,088 bp, and 25,076 bp (Table 1). The overall GC content of C. deboeaeosis_Jiang_DB-2015 was 39.4%, and 38.7%, 36.6% and 42.0% in the LSC, SSC and IR regions, respectively. These values are similar to C. deboeaeosis KM459003. C. deboeaeosis (KM459003) and C. revoluta showed similar GC content (39.4%), slightly lower than 392 tattuungensis (39.5%) (Table 1). In total, 133 genes were annotated, including 87 protein coding genes, 37 tRNA genes and 8 rRNA genes in C. deboeaeosis_Jiang_DB-2015, while 156 and 169 genes were annotated in C. revoluta and C. tattuungensis, respectively. Twelve genes (atpF, rpoC1, rpl2, clpP, ycf3, trnG-UCC, trnK-UUU, trnL-UAA in LSC; ndhA locate in SSC; ndhB, trnA-UGC, trnG-GAU in the IRs regions) contain 1–2 introns, respectively; while fourteen genes (3 protein coding 7 tRNA and 4 rRNA) were duplicated in the IR regions (Table 2). The tufA gene, found in gymnosperms and hornworts and inherited from green algae is coding for a nonfunctional protein synthesis elongation factor (723-bp long in C. tattuungensis and Ginkgo)9. Interestingly, this gene was 1 bp longer in C. deboeaeosis_Jiang_DB-2015 than in other Cycas (Table 2).

**Repeat and SSR analysis.** Using REPuter, seven repeats were found in the chloroplast of C. deboeaeosis_Jiang_DB-2015, which were three forward (F) and four palindrome (P), with no reverse and complement repeats discovered (Table 3). The repeats were mainly distributed in the intergenic spacers of transfer RNA genes, some of them being located in transfer RNA itself (Table 3). Interspecific comparison and analysis in broader Cycadaeae showed that C. revoluta had the highest number of repeats (24), while C. deboeaeosis contained the fewest (7), and C. tattuungensis contained an intermediate number of repeats (16) (Supplementary Table S1, Fig. 1). In contrast, the comparison of simple sequence repeats (SSRs) revealed a relative conservatism in their numbers, with congeneric species showing similarities in both numbers and spatial patterns of SSRs occurrence (Fig. 3). This is of particular value in Cycadaeae, where species are usually poorly defined or absent. These generic molecular biomarkers have the potential to provide useful diagnostic data in redefining complex paraphyletic and polyphyletic species groups in the family, as previously highlighted for C. revoluta species delineation 22,29–31, as previously highlighted for C. deboeaeosis10,12. There were 39 SSRs in the chloroplast of C. deboeaeosis, 34 (87%) and 5 (13%) mono- and di-nucleotides SSRs, respectively (Table 4). These SSRs were mainly distributed in the IGS region (29; 74%), and the other 26% were distributed in CDS genes (Table 4). C. revoluta and C. tattuungensis had similar SSRs numbers and locations than C. deboeaeosis, most of them being mononucleotides and distributed in the IGS (Fig. 3). Interestingly, C. revoluta lacked some SSR patterns (G and GA mono- and di- nucleotides, respectively) (Table S2). These diagnostic SSRs can be used in combination with nuclear SSRs developed in the genus for Cycadaeae conservation or reintroduction, species biodiversity assessments and phylogenetic studies in native or introduced areas12,30–33.

**Cycads phylogenetice reconstruction and comparison.** In the maximum likelihood (ML) phylogenetic tree, all but two nodes were highly supported (bootstrap support ≥95), with the accessions of C. deboeaeosis closely related to the other Cycas species (Fig. 4A). Cycas spp. diverged first in the Cycadaeae, followed by Dioon, a clade containing Zamia, Ceratozamia and Stangeria. Bowenia diverged from the remaining Zamiaceae with relatively high support (bootstrap support 80%), Macrozamia being as sister to a clade containing both Encephalartos and Lepidozamia (Fig. 4A). This chloroplast phylogeny confirms the division of Cycadaeae into two suborders, each of them being polyphyletic in our analyses (Fig. 4B), but contradict the current family delimitations9, with the family Stangeriaceae being polyphyletic with high support, in agreement with the most recent phylogenetic
work in cycads35. However, we found each genus in Stangeriaceae grouping with one of the clade in Zamiaceae, contrary to Salas-Leiva et al.35, in which *Bowenia* diverged in a basal position for all cycads but *Cycas* and *Dioon*, and *Stangeria* group with *Zamia* (*Microcycas* being absent from our dataset). *C. debaoensis* diverged in a basal position of the genus, with *C. revoluta* and *C. taitungensis* being closely related. In addition, the branch leading to *Cycas* is longer than the branches leading to the other genera, in accordance with the hypothesis of a recent diversification during the Miocene19. These results demonstrate the suitability and efficiency of using complete chloroplast sequences in reconstructing the evolutionary history of cycads, as previously demonstrated for other groups36,37.

Although the clear delineation of the genera in cycads are mostly due to the lengths of the sequences provided by complete chloroplast sequencing, the variability was unevenly distributed along the circular molecule (Fig. 5). Indeed, the ribosomal RNA genes as well as the region between *psbA* and *rpoC1* genes (0–25 kb) and between *ndhC* and *rpl20* (50–72 kb) of the cp sequences exhibited relatively few variations among cycads (Fig. 5). A cluster of four *ndh* genes (120–124.5 kb) appeared to be strikingly conserved. Overall, the level of variation increased with taxonomic distance, meaning that regions showing polymorphisms among *Cycas* species, exhibited higher variation among different cycad genera. However, some notable well-defined (<500 bp long) polymorphic

| Repeat1 start (location) | Repeat2 start (location) | Size (bp) | Type | Region |
|--------------------------|--------------------------|-----------|------|--------|
| 1 88,852 (*trnL-CAU*)     | 137,037 (IGS *chlL-trnN-GUU*) | 20,257    | P    | LSC; IRb |
| 2 28,396 (IGS *ropB-trnC-GCA*) | 28,470 (IGS *ropB-trnC-GCA*) | 39        | F    | LSC    |
| 3 55,346 (IGS *trnM-CAU*)  | 55,346 (IGS *trnM-CAU-atpE*) | 32        | P    | LSC    |
| 4 48,629 (IGS *trnF-GAA-ndhf*) | 48,629 (IGS *trnF-GAA-ndhf*) | 31        | F    | LSC    |
| 5 113,680 (IGS *trnN-GUU-ndhf*) | 113,680 (IGS *trnN-GUU-ndhf*) | 30        | P    | IRa    |
| 6 113,680 (IGS *trnN-GUU-ndhf*) | 137,236 (IGS *chlL-trnN-GUU*) | 30        | F    | IRa; IRb |
| 7 137,236 (IGS *chlL-trnN-GUU*) | 137,236 (IGS *chlL-trnN-GUU*) | 30        | P    | IRb    |

Table 3. Repeat sequences and their distribution found by REPuter in the *C. debaoensis* (KU743927) chloroplast genome. IGS: Intergenic spacer.

Figure 2. Repeat sequences in four chloroplast genomes of *Cycas*. REPuter was used to identify repeat sequences with length ≥30 bp and sequence identity ≥90% in the chloroplast genomes. F and P indicate the repeat type F (forward) and P (palindrome), respectively. Repeats with different lengths are indicated in different patterns.

Figure 3. Number of simple sequence repeats in four chloroplast genomes of *Cycas*, classified by repeat type. mono-: mononucleotide SSRs; di-: dinucleotide SSRs.

work in cycads35. However, we found each genus in Stangeriaceae grouping with one of the clade in Zamiaceae, contrary to Salas-Leiva et al.35, in which *Bowenia* diverged in a basal position for all cycads but *Cycas* and *Dioon*, and *Stangeria* group with *Zamia* (*Microcycas* being absent from our dataset). *C. debaoensis* diverged in a basal position of the genus, with *C. revoluta* and *C. taitungensis* being closely related. In addition, the branch leading to *Cycas* is longer than the branches leading to the other genera, in accordance with the hypothesis of a recent diversification during the Miocene19. These results demonstrate the suitability and efficiency of using complete chloroplast sequences in reconstructing the evolutionary history of cycads, as previously demonstrated for other groups36,37.

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regions departed from this assertion in \textit{ycf1} and \textit{ycf2} IRa. Indeed, the polymorphisms in these regions were higher among \textit{Cycas} than among cycads as a whole. Interestingly, three SNPs were located in other regions (\textit{trnL}-\textit{trnF}, \textit{clpP} intron 2 and \textit{ycf2} IRb) that exhibited a continuous increase in polymorphism levels across the family. \textit{ycf1} has been recently proposed as a barcode locus\textsuperscript{38}, despite it not being present in all genera\textsuperscript{39}, and was identified with \textit{clpP} among the most variable loci in \textit{Parthenium} spp.\textsuperscript{40}. Finally, \textit{trnL}-\textit{trnF} was previously used in \textit{Cycas} phylogenetic studies\textsuperscript{41}, but also in species identification of trees and ferns\textsuperscript{42,43}. Here, we stress the need to further assess these loci as potential more informative substitutes to the official barcode loci\textsuperscript{44}.

**Comparative interspecific chloroplast genomic analysis.** Focusing on the three \textit{Cycas} species available in GenBank, the mVISTA results showed that the four chloroplasts were highly conserved; however, the coding regions appeared to be globally more variable than the non-coding regions (Fig. 5 and Suppl. Fig. 1). Furthermore, the coding regions, e.g. \textit{rpoB}, \textit{psbC}, \textit{clpP} (intron), \textit{ycf1}, and \textit{ycf2}; \textit{psbA-trnH} and \textit{trnL-trnF} intergenic spacers showed promising levels of variations for further development in applications such as DNA-barcoding or phylogenetic reconstruction.

\textit{Cycas} have been considered as a difficult group for DNA-barcoding\textsuperscript{45}. Previously, it was reported that the \textit{psbA-trnH} spacer was highly variable in Cycadales except \textit{Cycas}\textsuperscript{45}, but \textit{trnL-trnF} was used in \textit{Cycas} for phylogenetic studies\textsuperscript{41}. Although \textit{rbcL} + \textit{matK} were chosen as a two-locus DNA-barcode for their universality and efficacy in land plant\textsuperscript{46}, it was not variable enough in \textit{Cycas} (Fig. 5 and Suppl. Fig. 1). \textit{rpoB}, \textit{rpoC1}, and non-coding

### Table 4. Simple sequence repeats in the \textit{C. debaoensis} (KU743927) chloroplast genome.

| No. | SSR type | SSR size | start | SSR-containing region |
|-----|----------|----------|-------|-----------------------|
| 2   | p1       | 20       | 8367  | IGS (\textit{trnQ}–\textit{UUG}–\textit{psbK}); LSC |
| 1   | p1       | 20       | 73873 | clpP; LSC |
| 1   | p1       | 19       | 123912| IGS (\textit{ndhG}–\textit{ndhI}); SSC |
| 1   | p1       | 14       | 32963 | IGS (\textit{trnE}–\textit{UUC}–\textit{trnH}–\textit{GUG}); LSC |
| 1   | p1       | 13       | 68694 | IGS (\textit{rufA}–\textit{trnH}–\textit{GUG}); LSC |
| 2   | p1       | 11       | 101911| IGS (\textit{rps7}–\textit{rps12}); IRa |
| 1   | p1       | 11       | 148078| IGS (\textit{trnV}–\textit{GAC}–\textit{rps12}); IRb |
| 2   | p1       | 10       | 1872  | \textit{tmK}–\textit{UUU}; LSC |
| 1   | p1       | 10       | 11723 | IGS (\textit{trnG}–\textit{UCC}–\textit{trnG}–\textit{UCC}); LSC |
| 1   | p1       | 19       | 15487 | IGS (\textit{atpH}–\textit{atpI}); LSC |
| 1   | p1       | 18       | 10133 | IGS (\textit{trnS}–\textit{GCU}–\textit{ycf12}); LSC |
| 1   | p1       | 16       | 57648 | IGS (\textit{rpl16}); LSC |
| 3   | p1       | 15       | 54643 | IGS (\textit{trnM}–\textit{CAU}–\textit{ndhC}); LSC |
| 3   | p1       | 15       | 84795 | \textit{rpl16}; LSC |
| 3   | p1       | 15       | 134345| IGS (\textit{ycf1}–\textit{chlN}); SSC |
| 2   | p1       | 13       | 74064 | \textit{clpP}; LSC |
| 2   | p1       | 12       | 5951  | \textit{rps16}; LSC |
| 1   | p1       | 12       | 118369| IGS (\textit{rpl32}–\textit{trnP}–\textit{GGG}); SSC |
| 4   | p1       | 11       | 83222 | \textit{rps8}; LSC |
| 1   | p1       | 11       | 88618 | IGS (\textit{rps23}–\textit{trnL}–\textit{CAU}); LSC |
| 1   | p1       | 11       | 102859| IGS (\textit{rps12}–\textit{trnV}–\textit{GAC}); IRa |
| 1   | p1       | 11       | 149026| IGS (\textit{rps7}–\textit{rps12}); IRb |
| 3   | p1       | 10       | 63350 | IGS (\textit{ycf4}–\textit{cemA}); LSC |
| 1   | p1       | 10       | 69919 | IGS (\textit{trnP}–\textit{UGU}–\textit{psaF}); LSC |
| 1   | p1       | 10       | 125905| \textit{ndhA}; SSC |
| 1   | p1       | 14       | 70451 | IGS (\textit{psaJ}–\textit{trnH}–\textit{GUG}); LSC |
| 1   | p1       | 11       | 52764 | IGS (\textit{ndhC}–\textit{trnH}–\textit{GUG}); LSC |
| 2   | p1       | 10       | 70921 | IGS (\textit{rpl33}–\textit{trnH}–\textit{GUG}); LSC |
| 1   | p1       | 10       | 149995| IGS (\textit{rps7}–\textit{ndhB}); IRb |
| 1   | p1       | 14       | 44747 | \textit{ycf3}; LSC |
| 1   | p1       | 11       | 5229  | \textit{rps16}; LSC |
| 2   | p1       | 10       | 17681 | IGS (\textit{rps2}–\textit{rpoC2}); LSC |
| 1   | p1       | 10       | 100943| IGS (\textit{ndhB}–\textit{rps7}); IRa |
| 3   | p2       | (TA)14   | 28     | 29832 | IGS (\textit{petN}–\textit{psbM}); LSC |
| 3   | p2       | (TA)9    | 18     | 1453  | IGS (\textit{psbA}–\textit{trnK}–\textit{UUU}); LSC |
| 2   | p2       | (TA)6    | 12     | 15670 | IGS (\textit{atpH}–\textit{atpI}); LSC |
| 1   | p2       | (GA)6    | 12     | 68377 | IGS (\textit{psbE}–\textit{petL}); LSC |
| 1   | p2       | (AT)6    | 12     | 15808 | IGS (\textit{atpH}–\textit{atpI}); LSC |
(e.g. $\text{atpF-}\text{atpH}$, $\text{psbK-psbL}$, and $\text{rpl32-trnL}$) regions have been shown to be variable enough at higher taxonomic levels\(^{46}\), but also within cycads\(^{47,48}\). In light of this, $\text{psbC}$, $\text{clpP}$ (intron), $\text{ycf1}$, and $\text{ycf2}$ should be considered as candidates for future phylogenetic studies in $\text{Cycas}$.

**Intraspecific comparison.** Comparing the two individuals of $\text{C. debaoensis}$, we found 10 SNPs and 1 indel, plus one “N” position in our data that didn’t allow us to confirm its status (Table 5). Their genomic positions are indicated in Figs 1 and 5. Six and four SNPs were located in IGS and coding regions, respectively, and one indel in the $\text{clpC}$ intron. The genetic distance between the two individuals was 0.005553% (i.e. $\approx 1$ SNPs/indels per 20 kb). This result is consistent with previous studies showing low within-population diversity in addition to limited gene flow among populations in $\text{C. debaoensis}$\(^{10,11}\). Whittall et al.\(^{49}\) found a comparable level of intraspecific divergence in pines, irrespective of the rarity of the considered species. However, in the pest species $\text{Jacobaea vulgaris}$ (Asteraceae), the intraspecific divergence was four times higher\(^{50}\), perhaps due to its short generation times as opposed to those prevailing in slow growing and long-lived trees. Further studies are still needed to determine whether or not intraspecific genetic diversity is linked to geographic ranges or the intrinsic characteristics of the taxonomic group.

**Conclusions**

Comparing genomic diversity at different taxonomical, but also spatial and temporal scales, we were able to reconstruct a robust phylogeny for cycads, and to identify regions showing promising levels of variation at three levels (familial, generic and intraspecific rank). These regions can provide useful and alternative loci for species identification and population-based studies for conservation, ecology and evolution. Despite their restricted geographic ranges, we showed that several, potentially diagnostic intraspecific variations can be found in the chloroplasts of different individuals $\text{C. debaoensis}$, including 10 SNPs and 1 indel in as of yet unstudied regions.

Comparing results from the three scales, four regions appeared to be variable at the three considered taxonomic scales, namely $\text{ycf3}$, $\text{clpP}$, $\text{psbD}$ and the $\text{trnL-trnF}$ IGS. Therefore, we recommend future studies in cycads further evaluate these loci in details.

We expect that by providing and highlighting these new resources to the plant research community, it will allow for development of new diagnostic markers and innovative conservation strategies in this iconic, but highly threatened taxonomic group, especially in the case of $\text{C. debaoensis}$.

**Materials and Methods**

**DNA sequencing and genome assembly.** Total genomic DNA was extracted from 0.1 g of frozen fresh leaves, from an individual collected in Guangxi ($23°69′40″N, 106°15′83″E$) in 2015 (voucher deposited at our research group herbarium, Jiang_C2) according to the manufacturer instructions with the Plant Genomic DNA Kit (Tiangen Biotech Co., Ltd). A 350-bp paired-end library was then constructed using NEBNext Ultra II DNA Library Prep Kit (Ipswich, Massachusetts, USA) and sequenced by Novogene (Beijing, China). About 1 Gb of raw data were obtained on an Illumina HiSeq2500 platform (San Diego, California, USA), with a paired-end read length of $2 \times 150$ bp. The raw reads were submitted to the SRA under the accession number SRR3407155.

The raw data were imported in Geneious R9 (Biomatters Ltd, Auckland, New Zealand), and a cp genome was assembled according to Hinsinger and Strijk\(^{25}\). Raw reads were trimmed according to their $S'$ and $S''$-end quality, then a reference-guided assembly was performed, using the available cp of $\text{Cycas debaoensis}$ (KM459003) as a
reference for the mapping step. The cp genome annotation was transferred from *C. debaoensis* using the implemented function in Geneious R9 and their boundaries were manually checked. The circular cp genome map was generated using the Organellar Genome Draw program (OGDRAW).

**Figure 5.** mVISTA percent identity plot of available cycad chloroplasts, using *C. debaoensis* as a reference. Vertical scale indicates the percentage of identity ranging from 50% to 100%. Coding regions are in blue and noncoding regions are in pink. Cladogram redrawn from Fig. 5B; branch lengths are not representative of evolutionary changes; bootstrap support is indicated on the nodes. Black arrows indicate intraspecific variations in *C. debaoensis.*
CIPRES55, and set the substitution model accordingly. We built a maximum likelihood tree using PHYML56, with
a 012310 genera were compared using mVISTA57, with all positions. In addition, to identify regions with substantial variability, the complete cp genomes of nine cycad

Table 5. Intraspecific SNPs between two individuals of *C. debaoensis* (KU743927).

| Position | Type | Nucleotide | location | location type |
|----------|------|------------|----------|---------------|
| 8432     | A/T  | trnQ pufK  | IGS      |               |
| 9889     | C/T  | trnS ycf12 | IGS      |               |
| 33707    | G/T  | pfdD       | coding   |               |
| 44878    | C/G  | ycf3       | exon 1   |               |
| 48140    | A/C  | trnL trnF  | IGS      |               |
| 60132    | G/T  | trnR accF  | IGS      |               |
| 62218    | A/C  | psaI ycf4  | IGS      |               |
| 73073    | A/.  | clpP       | intron 2 |               |
| 86371    | C/T  | rps19 rpl2 | IGS      |               |
| 90307    | G/T  | ycf2       | coding (IRa) |         |
| 160640   | A/C  | ycf2       | coding (IRb) |         |

Intergeneric comparisons and phylogenetic reconstruction. Following the recommended best practices for complete organellar sequencing63, we performed a phylogenetic analysis to confirm the accuracy of our reconstructed plastid and sample identification. We retrieved all the Cycadales available in GenBank (accessed 2016/02/15), representing all the ten genera in the order except Microcycas. To this Cycadales dataset, we added the sequences of *Gingko biloba* (NC016986), *Pinus strobus* (NC026302), *Araucaria heterophylla* (NC026450), *Taxus mairei* (NC020321) and *Cupressus sempervirens* (NC026296) as outgroups.

Sequences were aligned using MAFFT53 with default options. We used the jModelTest54 implementation in CIPRES55, and set the substitution model accordingly. We built a maximum likelihood tree using PHYML55, with a 012310 + F model using four gamma categories and 1000 bootstrap replicates. The ML tree was built using all positions. In addition, to identify regions with substantial variability, the complete cp genomes of nine cycad genera were compared using mVISTA57, with *C. debaoensis* (KM459003) as a reference for the annotations.

Sequence divergences among cycads were estimated using the Kimura 2-parameter model 58, implemented in MEGA59. Codon positions included were 1st + 2nd + 3rd + Noncoding. All positions containing gaps and missing data were excluded prior to analyses.

Interspecific comparisons. The complete cp genomes of *C. debaoensis* and two other species in cycads (*C. revoluta* and *C. taitungensis*) were compared using mVISTA57, as described above.

For each species, repeats (forward, palindrome, reverse and complement sequences) were identified using REPuter60 with 30 bp and sequence identity greater than 90%. Simple sequence repeats (SSRs) of *C. debaoensis* and the two other species were detected using MISA53 by setting the minimum number of repeats to 10, 5, 4, 3, 3 and 3 for mono-, di-, tri-, tetra-, penta- and hexa nucleotides, respectively. We recorded substitutions and indels separately, as well as their location in the chloroplast genome (e.g. SSRs/repeats, coding region/rRNA/tRNA/IGS). Sequence divergence extent between the two individuals was estimated as described above.

Intraspecific genome comparison. The two complete chloroplasts of *C. debaoensis* (KU743927, KM459003) were aligned in Geneious R9 (Biomatters Ltd, Auckland, New Zealand) using the MAFFT algorithm 53, and differences were identified using the “Find Variations/SNPs” function and checked individually. We recorded substitutions and indels separately, as well as their location in the chloroplast genome (e.g. SSRs/repeats, coding region/rRNA/tRNA/IGS). Sequence divergence extent between the two individuals was estimated as described above.

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**Author Contributions**

G.-F. J. designed the study, collected the plant materials, extracted DNA, assembled and analyzed the data, and wrote the paper; D.D.H. designed the study, assembled and analyzed the data and wrote the paper; J.S.S. wrote the paper. All authors have read and approved the final manuscript.

**Additional Information**

Accession codes: Raw reads and assembled chloroplast of *C. debaoensis* are available under the accession numbers SRR3407155 and KU743927, respectively.

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