Unprecedented intraindividual structural heteroplasmy in
Eleocharis (Cyperaceae, Poales) plastomes

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Data deposition:
Abstract

Plastid genomes (plastomes) of land plants have a conserved quadripartite structure in a gene dense unit-genome consisting of a large inverted repeat (IR) that separates two single copy regions. Recently, alternative plastome structures were suggested in Geraniaceae and in some conifers and *Medicago* the co-existence of inversion isomers has been noted. In this study, plastome sequences of two Cyperaceae, *Eleocharis dulcis* (water chestnut) and *E. cellulosa* (gulf coast spikerush), were completed. Unlike the conserved plastomes in basal groups of Poales, these *Eleocharis* plastomes have remarkably divergent features, including large plastome sizes, high rates of sequence rearrangements, low GC content and gene density, gene duplications and losses, and increased repetitive DNA sequences. A novel finding among these features was the unprecedented level of heteroplasmy with the presence of multiple plastome structural types within a single individual. Illumina paired-end assemblies combined with PacBio single-molecule real-time sequencing, long-range PCR and Sanger sequencing data, identified at least four different plastome structural types in both *Eleocharis* species. PacBio long read data suggested that one of the four *E. dulcis* plastome types predominates.
Key words: chloroplast, plastid genome, rearrangement, homologous recombination, RDR, repeat
Introduction

Plastids are one of the most essential features in photosynthetic eukaryotes. Plastid genomes (plastomes) of land plants have a conserved quadripartite structure in a gene dense unit-genome consisting of large and small single copy (LSC and SSC; ~80 kb and 20 kb) regions separated by a large inverted repeat (IR\textsubscript{A} and IR\textsubscript{B}; ~25 kb) (Wicke et al. 2011; Ruhlman and Jansen 2014). Plastomes of most land plants range from 120-170 kb in size with a median size and GC content of 154 kb and 37.6 %, respectively (Wicke et al. 2011; Weng et al. 2014; Park et al. 2018). After massive transfer of plastid DNA to the host nucleus during symbiogenesis, most plastomes retain genes encoding approximately 80 proteins, 30 tRNAs and four rRNAs. Although variation in plastome size, structure and gene content is uncommon in most photosynthetic taxa, extensive variation has been documented in several angiosperm lineages (Ruhlman and Jansen 2018).

Expansion, contraction and loss of the IR are the main factors causing plastome size variation and considerable variation in IR extent has been observed in diverse lineages, including Geraniaceae (Guisinger et al. 2011; Blazier, et al. 2016a; Zhang et al. 2016; Weng et al. 2017; Ruhlman and Jansen 2018), Fabaceae (Anon 2015; Schwarz et al. 2015), Ericaceae (Fajardo et al. 2013; Martínez-Alberola et al. 2013),
Berberidaceae (Ma et al. 2013), Trochodendraceae (Sun et al. 2013), Plantaginaceae (Zhu et al. 2016), Cactaceae (Sanderson et al. 2015; Solórzano et al. 2019), Annonaceae (Blazier, et al. 2016b), Campanulaceae (Knox 2014; Cheon et al. 2017), Papaveraceae (Park et al. 2018) and Passifloraceae (Rabah et al. 2019; Shrestha et al. 2019). The plastome of *Pelargonium transvaalense* is ~243 kb in size due to a remarkably expanded IR (~88 kb), leaving relatively small LSC (~60 kb) and SSC (~7 kb) regions (Weng et al. 2017). Similar to plastome size variation, examples of gene and intron losses have been documented in several angiosperm families. The loss of the entire suite of *ndh* genes in *Erodium* (Blazier et al. 2011), Orchidaceae (Chang et al. 2006; Wu et al. 2010; Delannoy et al. 2011; Barrett et al. 2014; Kim et al. 2015; Ruhlman et al. 2015), Alismatales (Peredo et al. 2013), and Carnegiea (Sanderson et al. 2015) and loss of *accD, clpP, rpl20, rps7, and ycf1* genes and the introns of *atpF, rpoC1, and clpP* in *Passiflora* (Jansen et al. 2007; Rabah et al. 2019; Shrestha et al. 2019) are examples. The monocot family Poaceae (Poales) also experienced gene and intron losses, such as *accD, ycf1*, and *ycf2* genes and the introns of *clpP* and *rpoC1* (Hiratsuka et al. 1989; Maier et al. 1995; Morris and Duvall 2010). These changes do not occur in basal lineages of Poales (Bromeliaceae and Typhaceae) (Stevens 2001), which retain the structure and
content of conserved plastomes (Guisinger et al. 2010; Redwan et al. 2015; Poczai and Hyvönen 2017).

Although most plastomes contain very small amounts of repetitive DNA apart from the canonical IR, large numbers of repeats have been identified in several eudicot and monocot lineages with highly rearranged plastomes (Lee et al. 2007; Cai et al. 2008; Haberle et al. 2008; Guisinger et al. 2011; Zhang et al. 2011; Weng et al. 2014). In some of these a significant positive correlation between the degree of genomic rearrangements and the number of repeats was detected (Guisinger et al. 2011; Weng et al. 2014). Repetitive DNA in plastomes plays a central role in genomic rearrangements and sequence divergence via illegitimate recombination and slipped-strand mispairing (Asano et al. 2004; Rogalski et al. 2006; Timme et al. 2007; Gray et al. 2009; Zhang et al. 2011). While the large canonical IR is well known to reverse the polarity of SC regions through recombination (Kolodner and Tewari 1979; Bendich 2004), other inversions have been associated with short inverted repeats (Kim and Lee 2005; Schwarz et al. 2015; Rabah et al. 2019). The typical IR does not occur in conifer plastomes but lineage-specific, short inverted repeats are associated with large inversions. Since Tsumura et al. (2000) discovered inversion isomers associated with ~1.2 kb inverted repeat in Pinaceae, examples of repeat associated isomeric plastomes
within a single species or individual have been documented in several gymnosperm lineages including Pinaceae, Cupressaceae, Podocarpaceae and Taxaceae (Wu et al. 2011; Guo et al. 2014; do Nascimento Vieira et al. 2016; Hsu et al. 2016; Qu et al. 2017; Fu et al. 2019). In angiosperms, two distinct plastome configurations that differed in the orientation of a ~45 kb segment were detected among accessions of the legume Medicago truncatula with verification using DNA gel blot analysis (Gurdon and Maliga 2014). Recently, with the aid of PacBio long reads, alternative plastome arrangements were identified within the plastome of a single individual of Monsonia emarginata (Geraniaceae) and these structural variants were associated with large repeats (Ruhlman et al. 2017).

Approximately 4,000 land plant plastomes have been sequenced and are publicly available (accessed on 2 Oct 2019 from NCBI). However, taxon sampling is biased in favor of certain major groups, limiting the knowledge of plastome organization and evolution in many lineages. For example, despite 616 sequenced plastomes of the order Poales available in NCBI, just eight belong to the non-Poaceae lineages Bromeliaceae, Typhaceae, Eriocaulaceae, Cyperaceae and Joinvilleaceae (Guisinger et al. 2010; Redwan et al. 2015; Wysocki et al. 2016). Among 14 Poales families (APG IV, 2016), Cyperaceae (sedges) is the second largest with more than 5,500 species and a
cosmopolitan distribution. However, little is known about plastome organization and evolution within the family, which is represented by three plastomes in NCBI that have not yet been published. In this study plastome sequences of two *Eleocharis* species were completed. In contrast to basal Poales, *Eleocharis* plastomes exhibited multiple structural types within a single individual, unprecedented levels of genome rearrangement and repetitive DNA, and extensive gene loss and duplication.

**Materials and Methods**

**Plant materials and DNA extraction**

Corms of water chestnut (*Eleocharis dulcis*, accession # PI 106274) were provided by the U.S. National Plant Germplasm System in the United States Department of Agriculture and Gulf coast spikerush, *Eleocharis cellulosa*, was collected in west Texas (supplementary table S1, Supplementary Material online). Corms of *E. dulcis* and live plants of *E. cellulosa* were grown in the University of Texas at Austin (UT-Austin) greenhouses. Since culms (a hollow or pithy stalk or stem of grasses, sedges and rushes) play the primary role in photosynthesis due to highly reduced leaves modified into a tubular sheath at the base of culms, total genomic DNA was extracted from newly emergent culms using the cetyl trimethylammonium bromide (CTAB) method of
Doyle & Doyle (1987) with modifications that included the addition of 3% PVP and 3% beta-mercaptoethanol (Sigma, St. Louis MO). Following treatment with RNase A (ThermoScientific, Lafayette, CO) and phase separation with chloroform, DNA was recovered by ethanol precipitation, resuspended in DNase-free water and stored at -20 ºC for genome sequencing. DNA of *E. dulcis* but not *E. cellulosa* was extracted from a single individual. Voucher specimens of *E. dulcis* and *E. cellulosa* were deposited in the Billie L. Turner Plant Resources Center (TEX-LL) at UT-Austin (supplementary table S1, Supplementary Material online).

**Genome sequencing, assembly and annotation**

Genomic DNAs were sent to the Beijing Genomics Institute (BGI) for library preparation and Illumina sequencing. Approximately 40 million 150-bp, paired-end (PE) reads were generated on the Illumina HiSeq X-Ten sequencing platform (Illumina, San Diego, CA) and reads were assembled *de novo* using Velvet version 1.2.08 (Zerbino and Birney 2008) at the Texas Advanced Computing Center (TACC). Multiple *de novo* assemblies were performed with a range of *k*-mer sizes (intervals of four from 61 to 117), three minimum depths of coverage (200X, 500X and 1000X), default insert size estimation and scaffolding off. Plastid contigs from all assemblies were imported into Geneious v.9.1.8
and all protein-coding genes were identified by BLAST against reference plastomes of *Nicotiana tabacum* (NC_001879), *Typha latifolia* (NC_013823), *Ananas comosus* (NC_026220) and *Hypolytrum nemorum* (NC_036036) and by GeSeq with MPI-MP chloroplast references (Tillich et al. 2017). Once draft plastomes were assembled, all plastid contigs were mapped to drafts to identify the inverted repeat (IR) boundaries and mismatches among contigs. To validate any ambiguities and determine the IR boundaries Bowtie2 v.2 (Langmead and Salzberg 2012) was used to map the Illumina reads to draft plastomes from which IRa had been removed. Verification of protein-coding genes was manually performed in Geneious by aligning genes with their homologs from the plastome sequences listed above as well as from *Spinacia oleracea* (NC_002202) and *Arabidopsis thaliana* (NC_000932). Since the *H. nemorum* plastome is publicly available in NCBI but has not yet been published, annotations were carefully examined and manually modified following the method described above prior to use in comparative analyses. All transfer RNA genes were identified using tRNAscan-SE v.2.0 (Lowe and Chan 2016). In *Eleocharis* and *H. nemorum* plastomes, genes that were annotated with low similarity (25 - 50%) to reference genes were further examined and regarded as putatively missing or nonfunctional based on the following criteria used in Rabah et al. (2019): absence of an
open-reading frame (ORF) with a complete conserved domain as searched in conserved domain database (CDD, www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi) and presence of internal stop codons that interrupted the reading frame or the conserved domain. Linear plastome maps were drawn with OGDRAW v. 1.2 (Lohse et al. 2013). For repeat visualization, circular maps were drawn using Circoletto (Darzentas 2010).

Whole-genome alignments were performed to examine the arrangement of locally colinear blocks (LCBs) of different plastome structural types of E. dulcis and E. cellulosa and to estimate the genomic rearrangements of Eleocharis plastomes compared with the basal Poales, T. latifolia, as a reference (supplementary fig. S1, Supplementary Material online) using progressiveMauve 2.3.1 in Geneious (Darling et al. 2010) with default parameters. One copy of the IR (IRa) was removed from plastomes prior to the Mauve alignment. The LCBs of E. dulcis, E. cellulosa, H. nemorum, A. comosus and T. latifolia plastomes were manually numbered and annotated for genes within each LCB. Strand orientation was indicated with ‘–’, if the LCB was in the reverse orientation relative to the reference. Breakpoint (BP) and reversal distances were estimated using the web-based application CREx (Common Interval Rearrangement Explorer) (Bernt et al. 2007) with T. latifolia as a reference.
Long-range PCR and Sanger sequencing

Nine oligonucleotide primers were designed for long-range PCR using Primer 3 (supplementary table S2, Supplementary Material online) (Untergasser et al. 2012). Primer sequences were located in genes (petN, petD, rps2, rpoC2, rpoB, psbK, rpl20, rps8 and ndhJ) situated at the ends of syntenic blocks to confirm plastome assemblies. In addition, eleven primers were designed to perform nested PCR using amplicons of targeted regions (supplementary table S2, Supplementary Material online).

Long-range PCR with multiple primer combinations was performed using the high-fidelity TaKaRa PrimeStar GXL DNA Polymerase (Takara Bio USA, INC., Mountain View, CA, USA). PCR reactions were performed in 12.5 µl, including 7 µl distilled water, 2.5 µl 5X PrimeSTAR GXL Buffer, 1 µl dNTP mixture, 0.5 µl DNA polymerase, 0.5 µl each primer and 0.5 µl genomic DNA (~50 ng). Each target was amplified using variable conditions depending on the size: initial denaturation at 98 °C for 1 min, followed by 30 cycles of denaturation at 98 °C for 10 sec, annealing at 58-62 °C for 15 sec, and extension at 68 °C for 50 to 150 sec. PCR products were treated with Exonuclease I (New England BioLabs, Ipswich, MA, USA) and Shrimp Alkaline Phosphatase (Fermentas, Glen Burnie, MD, USA) to remove residual primers and PCR
amplicons were Sanger sequenced at the Genome Sequencing and Analysis Facility (GSAF) at UT-Austin.

**PacBio sequencing**

Genomic DNA of *E. dulcis* with a total mass of 13.1 ug was sent to BGI for 20 kb PacBio library construction and single-molecule real-time (SMRT) sequencing (Pacific Biosciences, Menlo Park, CA). PacBio long reads were generated from one SMRT cell on RSII system. Error correction of long reads was performed using the LSC v.2.0 tool (Au et al. 2012) with approximately 46 million Illumina PE reads generated from DNA of the same *E. dulcis* individual. Error-corrected reads were mapped to complete plastome monomers and to genes at the end of each Illumina/Velvet-assembled syntenic block to detect reads that may support the presence of multiple structural types.

**Repetitive DNA sequence analysis**

Repeat analyses were performed to calculate the content of tandem and dispersed repeats. One copy of the IR (IRa) was removed from plastomes prior to repeat analyses. Dispersed repeats were detected by using the command line version BLAST v.2.8.1+ (Altschul et al. 1990) with a word size of 16 and percent identity of 80.0 % and each *Eleocharis, H. nemorum, A. comosus* and *T. latifolia* plastomes as both query and subject.
BLAST results were carefully examined and duplicate repeats were eliminated. Tandem repeats were identified by using the web version of Tandem Repeats Finder v.4.09 with default parameters (Benson 1999). Total repeats were categorized into several size classes based on the length of BLAST hits. The proportion of total repeats and repeats by each size class was calculated for each plastome (length of repeats/length of plastome (without IRa) X 100).

Results

Multiple structural types in Eleocharis plastomes

Plastome contigs assembled de novo from Eleocharis dulcis and E. cellulosa Illumina PE reads were very complex and suggested multiple arrangements of syntenic blocks (fig. 1B; supplementary figs. S2-3, Supplementary Material online). All possible arrangements of the syntenic blocks were assembled to yield the greatest number of unique draft plastome monomers. Careful examination of draft assemblies suggested four structural types that vary with respect to the arrangement of syntenic blocks in the LSC region.

Although the Illumina PE reads mapped to the four plastome types with approximately 1900X and 1080X depth of coverage for E. dulcis and E. cellulosa (supplementary table S3, Supplementary
Material online), respectively, these data were generated from 300-350 bp insert libraries and therefore could not confirm the presence of different plastome types. To address this and to confirm the putative multiple types, long-range PCR was performed for both species.

Long-range PCR amplification provided strong evidence for multiple structural conformations of *Eleocharis* plastomes (fig. 1B; table 1, supplementary figs. S4-5, Supplementary Material online). All but three of the 20 unique junction regions among syntenic blocks 2 to 6 for each plastome type were amplified using multiple combinations of primers designed to anneal in nine genes located at the boundaries of each block (fig. 1B; supplementary table S2, Supplementary Material online). For *E. dulcis* all junction regions were amplified and most of PCR amplifications yielded fragments of expected sizes, which ranged from ~4.4 to ~11 kb, confirming the presence of four plastome structural types (table 1). Although PCR of the 2-5 and (−3)−4 junctions exhibited two amplicons of different sizes, those with the stronger intensity were consistent with the expected sizes. Three fragments that were amplified from the 2-3 junction showed a similar intensity and one of them was congruent with the expected size. For *E. cellulosa* all junctions for plastome types 1 and 4 were amplified with fragments of expected sizes except for the 5-6 and 4-6 junctions (table 1). Plastome types
2 and 3 were supported with PCR amplifications for two and three junctions. Unexpectedly, (-4)-5 and 5-6 junctions found only in *E. cellulosa* plastome types 1 and 4 were also amplified in *E. dulcis* (supplementary fig. S6 and table S4, Supplementary Material online). The sizes of each PCR amplicon from *E. dulcis* (~8 kb and ~4.8 / ~3 kb) were very close to those from *E. cellulosa* (~8.5 kb and ~4.5 kb).

To further examine the *E. dulcis* plastome, the PCR amplicons for several junction regions were Sanger sequenced (table 1) and boundary sequences were compared with draft plastome assemblies from Illumina data. DNA sequences of seven junction boundaries consistently mapped to the expected regions in each plastome type. Among seven junctions, three, 3-(-4), (-4)-(-5) and 5-(-3), were fully sequenced by nested PCR using the PCR amplicons as templates with primers designed to anneal near the 3’ end (supplementary table S2, Supplementary Material online). One other region ((-3)-6) was almost fully sequenced.

Long read PacBio SMRT sequencing was performed for *E. dulcis*. A total of 68,167 raw PacBio reads with the mean size of 10,035 bp was generated. Following correction with Illumina sequences, the read count was reduced to 61,379 with the mean size of 9,139 bp and mean GC content of 35.7 %. Mapping of long reads to each plastome type, emphasizing reads containing genes situated at the end of each syntenic block, showed that the
majority of reads were consistent with the plastome type 2 except for four reads. Specifically, when corrected long reads were mapped to one end of syntenic block 2, where petN gene is located, 119 of 123 mapped reads were consistent with 2-5 syntenic block junction, but four reads that showed partial inconsistency with plastome type 2 had the sequences of 2-3 junction (adjacency pair of petN – petD genes) in plastome type 1, confirming the presence of multiple structural types in a single individual of Eleocharis dulcis.

**Plastome features**

Minimal size variation was detected among the four plastome types in E. dulcis and E. cellulosa, ranging from 196,668 to 199,561 bp and 192,023 to 193,234 bp, respectively (supplementary table S5, Supplementary Material online), all of which were larger than the median plastome size (152 kb) for 3,656 angiosperms (accessed on 2 Oct 2019 from NCBI). Hypolytrum nemorum (NC_036036), representing a basal lineage of Cyperaceae, had a plastome size of 180,648 bp with the IR expanded to ~ 38 kb and was also larger than the median angiosperm plastome size but smaller than Eleocharis. The IR and SSC regions of both Eleocharis species were similar in size (IR: ~36 kb, E. dulcis and ~35 kb, E. cellulosa; SSC: ~10 kb) with only seven genes (ndhF, trnL-UAG, rpl32, trnT-UGU, ndhD,
psaC and ndhE genes) in the SSC (table 2). However, since the size variation in the IR and SSC among different plastome types could not be identified with the current data, minor size variation in LSC region (115,003 – 117,896 bp, E. dulcis and 111,670 – 112,881 bp, E. cellulosa) accounted for the overall size variation among four types identified within each species.

Plastomes of both Eleocharis species contained the same number of unique genes (105; 72 protein coding genes, 29 tRNA and 4 rRNA genes), which was fewer than H. nemorum (110; 76 protein coding genes, 30 tRNA and 4 rRNA genes) and the highly conserved basal Poales plastomes (113; 79 protein coding genes, 30 tRNA and 4 rRNA genes) due to putative gene losses. However, the total number of genes in Eleocharis was greater than basal Poales due to gene duplication events (table 2). The total number of genes was higher in E. dulcis (139) than E. cellulosa (132) due to more extensive duplications (table 2). The plastomes of both species experienced several putative gene losses including accD, clpP, rpl23, rps18, ycf1, ycf2, infA and trnT-GGU. The plastome of H. nemorum also lacked accD, rps18 and infA and experienced duplication of trnV-GAC (table 2). Gene density (the number of genes per kb) in E. dulcis, E. cellulosa and H. nemorum plastomes was 0.70, 0.68 and 0.75 and the percentage of the genome that contained coding regions was 38.2 %, 38.5 % and 54.8 %, respectively, substantially lower.
than basal Poales (0.82 and 57.2 % for *A. comosus* and 0.81 and 56.4 % for *T. latifolia*) (table 2).

The GC content of *E. dulcis* and *E. cellulosa* plastomes was low at 32.6 % and 32.9 %, respectively, compared to the earliest diverging taxon of Cyperaceae, *H. nemorum* (34.9 %). All Cyperaceae had GC values less than the basal Poales *T. latifolia* (36.6 %) and *A. comosus* (37.4 %) and the mean value (37.5 %) of 3,656 angiosperm plastomes (fig. 2, table 2). The IR GC content of *E. dulcis* (37.6 %) and *E. cellulosa* (37.8 %) plastomes was higher than the mean for angiosperm plastome IRs, whereas the GC content of LSC and SSC regions for *E. dulcis* (30.2 and 25.5 %) and *E. cellulosa* (30.4 and 25.8 %) was lower (table 2).

Overall, the organization of *Eleocharis* plastomes was substantially different from the highly conserved plastomes of basal Poales, Bromeliaceae and Typhaceae, whereas the basal taxon in Cyperaceae, *H. nemorum*, exhibited features intermediate between those of *Eleocharis* and basal Poales.

Whole-genome alignment using progressiveMauve (fig. 3, supplementary table S6, Supplementary Material online) identified 25 LCBs from five complete plastomes (*E. dulcis, E. cellulosa, H. nemorum, A. comosus* and *T. latifolia*). The two basal Poales plastomes were collinear, whereas *Hypolytrum* had breakpoint (BP) and reversal distances of 11 and 7, respectively compared to basal Poales plastomes (table 3). The BP and
reversal distances were 16 and 13 (E. dulcis) or 16 and 14 (E. cellulosa) between Hypolytrum and Eleocharis plastomes, respectively. The largest BP distance (25) was between E. dulcis and basal Poales plastomes, whereas the largest reversal distance (21) was between E. cellulosa and basal Poales. The BP and reversal distances were 5 and 4 between the two Eleocharis species, respectively. Among the four plastome types in each Eleocharis species, six LCBs (1-6 syntenic blocks) were identified excluding small LCBs of large repeats (supplementary fig. S3, Supplementary Material online). Plastomes of both Eleocharis species had different orders of LCBs (3, 4 and 5 syntenic blocks) in the LSC region and in E. cellulosa LCB 2 was reversed compared to E. dulcis.

**Repetitive DNA in Eleocharis plastomes**

Repeat analyses revealed that Eleocharis plastomes contain abundant repetitive DNA with minor variation among different structural types (supplementary tables S7-8, Supplementary Material online). Along with Eleocharis, basal Cyperaceae and Poales taxa were included in repeat analyses. The amount and proportion of dispersed repeats in Eleocharis plastomes (39,752 bp, 24.3% for E. dulcis and 31,118 bp, 19.7 % for E. cellulosa) were substantially greater than in H. nemorum (8.8 %, 12,520 bp),
which in turn was greater than \textit{A. comosus} (1.1\%, 1,495 bp) and \textit{T. latifolia} (0.9\%, 1,210 bp) (table 4).

Dispersed repeats were grouped into five size classes (fig. 4). The \textit{Eleocharis dulcis} plastome contained the largest number of dispersed repeats (321) ranging from 17 bp to ~5 kb followed by \textit{E. cellulosa} (238; 17 bp to ~3 kb) and \textit{H. nemorum} (157; 17 to ~850 bp), while basal Poales had 57 and 47 repeats smaller than 60 bp (fig. 4, supplementary table S8, Supplementary Material online). Repeats >1000 bp were only detected in \textit{Eleocharis} plastomes where they represented more than 10\% and 8\% of the \textit{E. dulcis} and \textit{E. cellulosa} genomes, respectively (fig. 4). The three Cyperaceae plastomes contain larger repeats in the range of 61 to 1000 bp relative to other Poales, and these larger repeats constitute a substantial proportion of the plastome (fig. 4). In \textit{Eleocharis} many of the larger repeats in the >201 bp size range were concentrated between syntenic blocks 2 to 6 (fig. 1A).

The greatest proportion of tandem repeats were identified in \textit{H. nemorum} (6,638 bp; 4.7\%) followed by \textit{E. dulcis} (5,864 bp; 3.6\%) (table 4). Unlike dispersed repeats, only \textit{E. dulcis} plastome contained unusually large amounts of tandem repeats. \textit{Eleocharis cellulosa} had fewer tandem repeats (2,833 bp; 1.8\%) than \textit{T. latifolia} (3,270 bp; 2.4\%). Some tandem repeats were embedded in large dispersed repeats in \textit{Eleocharis} plastomes, and
there were instances where small dispersed repeats were located within tandem repeats. In the *E. dulcis* plastome, three copies of the largest tandem repeat (278 bp) with 99% sequence identity included *trnfM-CAU* in the IR. Most tandem repeats were found in intergenic spacers (IGS) except for occurrences in the introns of several genes (*rps16, ycf3, rpoC1, rpl16* and *ndhB*), including 16 copies of a 48 bp tandem repeat that share 84% sequence identity within the intron of *rpoC2* causing an expansion of its size to 5,097 bp. Similar to *E. dulcis*, most tandem repeats in *E. cellulosa* were found in IGS regions. Three introns included tandem repeats, the introns of *rps16, ndhB* and *rpoC2*. The *rpoC2* gene (4,707 bp) was expanded by 12 copies of a 48 bp tandem repeat with 84% sequence identity within its intron. Three and seven copies of a 48 and 45 bp tandem repeat were identified within *rpoC2* (4,707 bp) in *H. nemorum*. Overall, the exceptionally abundant repeats that enlarged IGS regions contributed to plastome size expansion in *Eleocharis*.

Repetitive DNA also resulted in gene duplications in *Eleocharis* plastomes. Three copies of the *trnfM-CAU* were located in the 278 bp tandem repeat of the *E. dulcis* IR, resulting in six total copies of this sequence. Several other genes were multiplied, including four copies of *rpl36* and two copies of *rpoA* in the LSC, and three copies of *trnH-GUG, trnV-GAC* and *trnP-UGG* (two copies in the IRs). In *E. cellulosa,
while *rpl36* was duplicated in a 593 bp dispersed repeat only in the LSC of plastome type 4, all four structural types had three copies of *trnF-M-CAU*, *trnH-GUG* and *trnV-GAC* (two copies in the IRs) located in various sized dispersed repeats (177-456 bp). In *H. nemorum*, one additional copy of *trnV-GAC* along with two in the IR was identified in the LSC within an 800 bp dispersed repeat with 98.5 % sequence identity.

**Discussion**

Despite being the second largest family of Poales, plastome organization and evolution of Cyperaceae has not been examined. In this study, plastomes of two *Eleocharis* species, *E. dulcis* and *E. cellulosa*, were atypical with respect to size, gene content, GC content, gene order and gene density compared to the highly conserved basal Poales plastomes. The most unusual feature was the presence of multiple plastome structural types within each *Eleocharis* species. Furthermore, plastome structural heteroplasmy was observed within single individuals and was associated with a remarkable abundance of repetitive DNA. The discussion mainly focuses on possible mechanisms that could generate different structural types in plastomes and comparisons to alternative plastome structures (isomeric forms) reported in a few distantly related seed plant lineages. The plastome
organization of *Eleocharis* is also compared with basal Cyperaceae and other Poales.

**Structural heteroplasmy in *Eleocharis* plastomes**

In two species of *Eleocharis*, at least four different plastome structural types are present, and in one species, *E. dulcis*, this unprecedented structural heteroplasmy was identified in a single plant. Initial assemblies using Illumina short reads suggested multiple plastome types, a suggestion confirmed by long-range PCR, Sanger sequencing and PacBio SMRT data. A few previous studies in conifers and *Medicago* (Fabaceae) detected isomeric plastomes associated with inversions mediated by lineage-specific short repeats (only for conifers) using a variety of approaches ranging from Southern hybridization to Illumina sequencing and PCR (Tsumura et al. 2000; Wu et al. 2011; Guo et al. 2014; do Nascimento Vieira et al. 2016; Hsu et al. 2016; Qu et al. 2017; Fu et al. 2019). Recently, PacBio long read data combined with Illumina read assembly demonstrated alternative gene order arrangements for the *Monsonia emarginata* (Geraniaceae) plastome (Ruhlman et al. 2017).

Although four major plastome structural types were clearly evident in *Eleocharis*, long-range PCR data suggested the possibility of a higher degree of heteroplasmy as shown in supplementary table S4, Supplementary Material online. A
similar suggestion was made by Ruhlman et al. (2017) for Monsonia. The possibility that the unexpected PCR results may be the result of amplifications of plastid DNA that has been transferred to nuclear or mitochondrial genomes cannot be ruled out. It is also possible that PCR artifacts arising from PCR-mediated recombination (Lahr and Katz 2009; Alverson et al. 2011) could explain the unexpected amplicons. However, consistent sizes between PCR amplicons for most junctions, Sanger sequencing, Illumina assemblies and PacBio reads strongly support the unprecedented structural heteroplasmy in Eleocharis.

For decades plastomes were considered circular with limited recombination. The presence of isomeric, presumed circular plastomes with different orientations of the single copy regions was attributed to intramolecular recombination between IR copies within a single unit genome, and was referred to as “flip-flop” recombination (Palmer 1983; Palmer 1985; Brears et al. 1986; Stein et al. 1986). Two previous studies in Pinaceae adopted the flip-flop intramolecular recombination model to explain the different plastome isoforms (Tsumura et al. 2000; Wu et al. 2011), whereas other studies of conifers suggested a repeat-mediated homologous recombination (HR) mechanism based on a circular unit genome (Guo et al. 2014; do Nascimento Vieira et al. 2016; Hsu et al. 2016; Qu et al. 2017; Fu et al. 2019). These previous studies ignored the overwhelming evidence that
the plastome exists as linear, branched and occasionally circular forms (Deng et al. 1989; Lilly et al. 2001; Oldenburg and Bendich 2004; Scharff and Koop 2006; Shaver et al. 2006). In fact, it has been shown that the different isomers of the SC regions occur by a recombination-dependent-replication (RDR) mechanism involving different unit genomes (Oldenburg and Bendich 2004; Maréchal and Brisson 2010). Thus, any discussion of the mechanism for generating multiple structural types of plastomes must consider RDR as the primary cause.

Ruhlman et al. (2017) proposed that RDR between different linear copies of plastomes generated alternative plastome arrangements associated with large inverted repeats (> 2 kb) in Mononínia emarginata (see fig. 4 in Ruhlman et al. 2017). The unusually large number and size of repeats (up to ~5 kb in E. dulcis and ~3 kb in E. cellulosa) could account for the large number of plastome structural types by providing homologous sequence for the RDR pathway. Many of the large (> 1 kb) and intermediate (201 – 1000 bp) repeats in Eleocharis plastomes are located between syntenic blocks 3-5 and may be involved with repeat-mediated RDR within or between unit genomes. Structural heteroplasmy may be more prevalent than appreciated, especially in other lineages that have plastomes with highly rearranged gene orders and abundant larger repeats, such as Passiflora, Trifolium, Campanulaceae and Geraniaceae.
The phylogenetic distribution of this phenomenon in Cyperaceae or other Poales is unknown due to limited taxon sampling in the family and order. The publicly available plastome of one of the basal taxa in Cyperaceae, *Hypolytrum nemorum*, has not been published and no raw read data is available for analysis. Plastome sequences of two basal lineages, Bromeliaceae and Typhaceae, do not show structural heteroplasmy (Guisinger et al. 2010; Redwan et al. 2015; Poczai and Hyvönen 2017) and two other Poales lineages (Poaceae and Eriocaulaceae) do not exhibit multiple plastome types (Darshetkar et al. 2019; Orton et al. 2019). Additional sampling of plastome sequences in Cyperaceae and related families of cyperids is needed to explore the origin and extent of structural heteroplasmy in this clade.

**Plastome organization**

In addition to remarkable structural heteroplasmy, *Eleocharis* plastomes have a number of unusual genomic characteristics compared to basal Poales, including size expansion, abundant repetitive DNA, higher degree of genome rearrangement, IR expansion, increased IGS, low GC content, low gene density and numerous gene duplications and losses.

In most angiosperms, increases in plastome size are caused by IR expansion as reported in several unrelated lineages,
including *Pelargonium* (IR ~ 88 kb; Chumley et al. 2006; Weng et al. 2014; Weng et al. 2017), *Annona* (IR ~ 64.5 kb; Blazier, et al. 2016b), *Lamprocapnos* (IR ~ 51 kb; Park et al. 2018), *Asarum* (IR ~45 kb; Sinn et al. 2018) and *Passiflora* (IR ~35 kb; Shrestha et al. 2019). Plastome size expansion in *Eleocharis* is mainly due to increased number and size of repeats and to a lesser extent IR expansion (tables 2, 4). Increased plastome size in the basal member of Cyperaceae, *Hypolytrum nemorum*, is also caused by these two phenomena. This is in contrast to other members of the Poales (*Typha, Ananas* and *Eriocaulon*), which have typical IRs and much less repetitive DNA and size variation (Guisinger et al. 2010; Redwan et al. 2015; Darshetkar et al. 2019). *Eleocharis* plastomes have similar SSC/IR boundaries to *H. nemorum* but they have an independent IR expansion into the LSC. Despite IR expansion into both the SSC and LSC and increased number of repeats in the IGS regions of the IR in *Eleocharis* plastomes, the size of their IR is smaller than *H. nemorum* because they lack *ycf1* and *ycf2*. Repeat content and IR expansion have contributed to the increase in plastome size of *Eleocharis* and both of these factors affect GC content. The IR has higher GC content than SC regions (table 2) so IR expansion should result in higher overall GC content in plastomes. However, *Eleocharis* plastomes had the lowest GC values (~ 32.6 %) among photosynthetic angiosperms despite their
genome size expansion (fig. 2). Other Cyperaceae have a similar pattern, which is likely due to having more repeats in IGS regions, which are known to be GC poor (table 4, Cai et al. 2006). The prevalence of repeats in the IGS regions also contributed to the lower gene density in Eleocharis and Hypolytrum compared to basal Poales plastomes (table 2). Thus, unlike most other photosynthetic angiosperms, Eleocharis plastomes have experienced increased plastome size driven by IR expansion and accumulation of repeats, and low GC content and gene density.

Whole-genome alignment revealed extensive gene order changes in Eleocharis plastomes (fig. 3, table 3) relative to T. latifolia. The exceptional number of gene order changes is likely related to the abundance of repeats, which facilitate repeat-mediated homologous recombination. The IR was previously suggested to play a role in plastome structural stability based on the presence of extensive rearrangements in IR-lacking groups in legumes (Palmer and Thompson 1982) and Cryptomeria (Hirao et al. 2008). However, many studies failed to find a correlation between plastome stability and the presence of the IR in several angiosperm lineages, including Campanulaceae (Haberle et al. 2008; Knox 2014), Oleaceae (Lee et al. 2007), Ericaceae (Fajardo et al. 2013; Martínez-Alberola et al. 2013), Plantago (Zhu et al. 2016), Geraniaceae (Chumley et al. 2006; Guisinger et al. 2011;
Weng et al. 2014; Blazier, et al. 2016a) and Passifloraceae (Rabah et al. 2019; Shrestha et al. 2019). In Eleocharis and other lineages the presence of dispersed repeats was more likely the primary factor facilitating plastome rearrangements.

Plastomes of basal Poales, Typha (Guisinger et al. 2010) and Ananas (Redwan et al. 2015), as well as Eriocaulon (Darshetkar et al. 2019) have a conserved gene content, similar to the ancestral angiosperm plastome (Ruhlman and Jansen 2014). However, Cyperaceae plastomes have numerous gene duplications and losses (table 2). Many gene duplications are the result of IR expansion into the SSC and LSC. The other duplicated genes are associated with the dispersed repeats. Gene losses are more extensive in Eleocharis than in H. nemorum (table 2). The missing Eleocharis genes have been pseudogenized or lost in several eudicot lineages, including Campanulaceae (Cosner et al. 1997; Haberle et al. 2008; Knox 2014; Cheon et al. 2017), Trifolium (Cai et al. 2008; Magee et al. 2010; Sabir et al. 2014), Ericaceae (Fajardo et al. 2013; Martínez-Alberola et al. 2013), Carnegiea (Sanderson et al. 2015), Silene (Erixon and Oxelman 2008; Sloan et al. 2012), Geraniaceae (Chumley et al. 2006; Guisinger et al. 2011; Weng et al. 2014; Blazier, et al. 2016a; Park et al. 2017; Weng et al. 2017) and Passifloraceae (Rabah et al. 2019; Shrestha et al. 2019). Plastomes of these groups have abundant repetitive DNA and highly rearranged gene
order, a syndrome that is shared with Eleocharis. In monocots, Poaceae plastomes are structurally similar to basal Poales except for three inversions and plastome size reduction caused by several gene losses (Quigley and Weil 1985; Hiratsuka et al. 1989; Doyle et al. 1992; Maier et al. 1995; Michelangeli et al. 2003; Morris and Duvall 2010; Harris et al. 2013). The loss of accD, ycf1, ycf2, clpP, and rpl23 is homoplastic in Poales as the same losses have occurred independently in the distantly related families Poaceae and Cyperaceae (supplementary fig. S1, Supplementary Material online). More plastome sequencing in Cyperaceae lineages is needed to elucidate the evolution of clpP, ycf1, ycf2 and rpl23 in the family. Moreover, transcriptome data would allow the determination of the fate of missing genes in Eleocharis and other related Cyperaceae.

Conclusion

Analyses utilizing Illumina and PacBio SMRT sequences, long-range PCR and Sanger sequencing data demonstrated unprecedented structural heteroplasmy within a single Eleocharis dulcis individual. At least four different plastome structural types are present in both Eleocharis species and it is likely that this phenomenon is much more widespread in the family. The long-held notion that plastomes are highly conserved in structure within individuals and species must be reconsidered,
especially for lineages with a prevalence of large dispersed repeats that are likely to exhibit structural heteroplasmy in plastomes.

Acknowledgements

This work was supported by Texas Ecolab program to CL and the S. F. Blake Centennial Professorship to RKJ and TAR. The authors thank the Billie L. Turner Plant Resources Center (TEX-LL) at UT-Austin for serving as a repository for voucher specimens, United States Department of Agriculture Germplasm Resources Information Network for providing corms and the Texas Advanced Computing Center (TACC) at UT-Austin for access to supercomputers.
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Figure legends

Figure 1: Schematic representations of *Eleocharis dulcis* and *E. cellulosa* plastomes. (A) Unit-genome maps and repetitive DNA content of *E. dulcis* and *E. cellulosa* plastomes. Completed *Eleocharis* plastome sequences were submitted to OGDRAW (Lohse:2013hc) to generate physical maps and Circoletto (Darzentas:2010jw) to visualize repetitive DNA. Structural type and plastome size (in parentheses) are shown below the species name. Syntenic blocks (numbered 1 – 6) detected by progressiveMauve (Darling:2010iu) are depicted by open boxes; the negative symbol (−) indicates reverse-oriented strands relative to the reference (*Typha*). Red boxes indicate blocks that that vary in order among different structural types while blocks encompassed by black boxes maintain their positions across types. Genes indicated in red font are located near the end of each syntenic block and were employed as annealing sites for PCR confirmation of plastome arrangements. The vertical black bar at the top of each linear map is provided for scale. The circular representation of each species is below each linear map with syntenic block numbers shown. Dispersed repeats and IR are shown within the circular map in blue and red, respectively. (B) Each syntenic block is illustrated with a different color and follow the same numbering convention as in (A). Representative genes near the end of each syntenic block are
shown at the top. Gene symbols in white font indicate those in long range PCR and correspond to the red gene symbols in (A). Different syntenic block arrangements are highlighted in red boxes for both Eleocharis species. IR, inverted repeat; SSC, small single copy region.

**Figure 2:** *Eleocharis* plastomes exhibit atypical size and GC content. Plastome size and GC content of 3,656 angiosperms in the NCBI Genome database were plotted. Parasitic (yellow) and autotrophic (teal) species are indicated with different colors. The two *Eleocharis* plastomes are labeled with red font. Other Cyperaceae and larger plastomes, including *Pelargonium* and *Annona cherimola*, are indicated with teal font. The *Erodium* plastomes with high GC content are also represented with a teal label. bp, basepairs.

**Figure 3:** Whole-plastome alignment of five Poales species. Newly completed plastomes of *Eleocharis dulcis* and *E. cellulosa* and publicly available basal Cyperaceae and Poales plastomes from NCBI (*Hypolytrum nemorum*, *Ananas comosus* and *Typha latifolia*) were analyzed by progressiveMauve to identify locally collinear blocks (LCBs) with the *Typha* plastome as a reference. One copy of the inverted repeat was removed before the analysis and numerals at top indicate size in kilobases (kb). The
corresponding LCBs among five plastomes are shaded and connected with a line of the same color. The histogram inside each block shows pairwise nucleotide sequence identity. LCBs that are flipped across the plane indicate an inverted strand.

Figure 4: Repetitive DNA content in five Poales plastomes. (A) The number of dispersed repeats in different size classes. (B) The proportion of plastome that represents dispersed repeats in different size classes. bp, basepairs.
Supplemental figures

Supplemental figure S1: Cladogram of 14 Poales families.
Phylogenetic relationships Poales families are based on APG IV (APG IV, 2016; Stevens 2001).

Supplemental figure S2: Schematic representations of four plastome structural types in Eleocharis. Unit-genome maps of Eleocharis dulcis (A) and E. cellulosa (B) are presented for four structural types in each species. Each plastome type was submitted to OGDRAW {Lohse:2013hc} to generate a schematic map. Syntenic blocks detected by progressiveMauve {Darling:2010iu} are numbered from 1 to 5, along with the inverted repeat (IR) and small single copy (SSC) region (representing block 6) and enclosed by open boxes. Syntenic block numbers with ‘-’ indicates reverse-oriented strands. Red boxes indicate syntenic blocks that vary in order among different structural types. Gene names below red boxes are located near the end of each syntenic block and used in primer design. Bars at the bottom of A and top of B indicate scale.

Supplemental figure S3: Whole-plastome alignment of four plastome structural types in Eleocharis. Four plastome types of Eleocharis dulcis (A) and E. cellulosa (B) were analyzed by progressiveMauve to identify locally collinear blocks (LCBs)
with type 1 as the reference. One copy of the inverted repeat was removed prior to analysis. Type identifiers are shown at the left below each plastome. Corresponding LCBs are numbered, colored and connected with a line of the same color. Histograms inside each block shows pairwise nucleotide sequence identity. LCBs that are flipped across the plane indicate inversions. Numbers at top indicate size in kilobases (kb).

**Supplemental figure S4:** PCR confirmation of adjacency among syntenic blocks in four structural types of the *Eleocharis dulcis* plastome. Genes used to design PCR primers were located near the end of each syntenic block and are given at the top of each lane with the numbered-order of adjacency pair from 1 to 4 below in yellow font. The pair of numbers at the bottom of each lane denotes the adjacency of syntenic blocks as shown in Figure 1B. The approximate size of PCR amplicons in each lane is inferred using the 1 kb DNA ladder on either the left or right side of each photo. The unlabeled lanes without any numerals are not relevant to the current study. kb, kilobases.

**Supplemental figure S5:** PCR confirmation of adjacency among syntenic blocks in four structural types of the *Eleocharis cellulosa* plastome. Genes located near the end of each syntenic block used in primer design to confirm the adjacency pairs are
indicated at the top of each lane with the numbered-order of adjacency pair from 1 to 4 below in yellow font. The pair of numbers at the bottom of each lane denotes the adjacency of syntenic blocks as shown in Figure 1B. The approximate size of PCR amplicons in each lane is inferred relative to the 1 kb DNA ladder on the either left or right side of each photo. The unlabeled lanes without any numerals are not relevant to the current study. Kb, kilobases.

**Supplemental figure S6:** PCR confirmation of two unexpected adjacency pairs of syntenic blocks in *Eleocharis dulcis* plastome. These pairs were not originally identified in *E. dulcis* plastome types but in *E. cellulosa* plastome types 1 and 4. Genes used in primer design were located near the end of each syntenic block and are represented at the top of each lane. The pair of numbers at the bottom of each lane denotes the adjacency pair of syntenic blocks as shown in Figure 1B. The approximate size of PCR amplicons in each lane is inferred using the 1 kb DNA ladder on either the left or right side of each photo. The unlabeled lanes without any numerals are not relevant to the current study. Kb, kilobases.
Supplemental tables

Supplemental table S1: Newly sequenced Eleocharis plastomes.

Supplemental table S2: List of oligonucleotide primers used for PCR and sequencing to confirm plastome assemblies.

Supplemental table S3: Summary of Illumina sequencing data for Eleocharis plastomes.

Supplemental table S4: Long-range PCR results for unexpected adjacency pairs of syntenic blocks in Eleocharis dulcis.

Supplemental table S5: Summary of major features of four types of Eleocharis dulcis and E. cellulosa plastomes.

Supplemental table S6: Genes and intergenic regions included in locally collinear blocks (LCB) identified by progressiveMauve alignment of Eleocharis, Hypolytrum and basal Poales plastomes.

Supplemental table S7: Statistics of dispersed and tandem repeats in four plastome types of two Eleocharis species.

Supplemental table S8: The number of dispersed repeats in size classes in four plastome types of two Eleocharis species.
### Table 1: Verification of multiple structural types in *Eleocharis* plastomes.

| Species          | Type | Syntenic block | Primer F | Primer R | Size (bp) | PCR Band size (bp) | Sanger Seq. | PacBio Seq. |
|------------------|------|----------------|----------|----------|-----------|--------------------|-------------|-------------|
| 1                |      | 2 – 3          | E_petN   | E_petD   | 8,188     | 5,500 / 6,200 / 8,100 | +           |             |
|                  |      | 3 – (-4)       | E_rps2   | E_rpoB   | 4,842     | ~5,000            | +           | +           |
|                  |      | (-4) – (-5)    | E_rpoC2  | E_rp120  | 4,368     | ~4,300            | +           | +           |
|                  |      | (-5) – 6       | E_psbK   | E_rps8   | 8,550     | ~8,500            |             |             |
|                  |      |                |          |          |           |                   |             |             |
| Eleocharis dulcis|      | 2 – 5          | E_petN   | E_psbK   | 11,127    | ~11,000* / ~6,200 | +           |             |
|                  |      | 5 – (-3)       | E_rp120  | E_rps2   | 4,381     | ~4,300            | +           | +           |
|                  |      | (-3) – 4       | E_petD   | E_rpoC2  | 5,261     | ~3,000            | +           | +           |
|                  |      | 4 – 6          | E_rpoB   | E_rps8   | 5,181     | ~5,000            | +           | +           |
|                  |      |                |          |          |           |                   |             |             |
| 3                |      | 2 – (-4)       | E_petN   | E_rpoB   | 4,869     | ~4,000            | +           | +           |
|                  |      | (-4) – 3       | E_rpoC2  | E_petD   | 5,257     | ~4,300            | +           | +           |
|                  |      | 3 – (-5)       | E_rps2   | E_rp120  | 4,381     | ~4,300            | +           | +           |
|                  |      | (-5) – 6       | E_psbK   | E_rps8   | 8,550     | ~8,500            | +           | +           |
|                  |      |                |          |          |           |                   |             |             |
| Eleocharis cellulosa|    | (-2) – 3       | E_ndhJ   | E_petD   | 5,387     | ~5,400            | n/a         | n/a         |
|                  |      | (-3) – (-4)    | E_rps2   | E_rpoB   | 4,816     | ~5,000            | n/a         | n/a         |
|                  |      | (-4) – 5       | E_rpoC2  | E_psbK   | 9,000     | ~8,500            | n/a         | n/a         |
|                  |      | 5 – 6          | E_rp120  | E_rps8   | 3,025     | ~4,500            | n/a         | n/a         |
|                  |      |                |          |          |           |                   |             |             |
| 3                |      | (-2) – 5       | E_ndhJ   | E_psbK   | 8,326     |                  | n/a         | n/a         |
|                  |      | (-3) – 5       | E_rps2   | E_rpoB   | 2,685     | ~2,700            | n/a         | n/a         |
|                  |      | (-4) – (-5)    | E_rpoC2  | E_rp120  | 2,685     | ~2,700            | n/a         | n/a         |
|                  |      | (-5) – 6       | E_psbK   | E_rps8   | 9,339     |                  | n/a         | n/a         |
|                  |      |                |          |          |           |                   |             |             |
| 4                |      | (-2) – 3       | E_ndhJ   | E_petD   | 5,387     | ~5,000            | n/a         | n/a         |
|                  |      | (-3) – (-5)    | E_rps2   | E_rp120  | 4,346     | ~4,300            | n/a         | n/a         |
|                  |      | (-5) – 4       | E_psbK   | E_rpoC2  | 9,001     | ~8,000            | n/a         | n/a         |
|                  |      | 4 – 6          | E_rpoB   | E_rps8   | 3,495 / 2,312 | ~5,000   | n/a         | n/a         |

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+ indicates that adjacencies of syntenic blocks were confirmed.
* Band size in bold indicates the one with higher intensity.
a Almost entire junction was sequenced.
b Boundaries of junction were sequenced.
c Entire junction was sequenced.
F = forward. R = reverse. n/a = not available.
Table 2: Summary of major features of *Eleocharis*, *Hypolytrum* and basal Poales plastid genomes.

| Family          | Taxon                      | *Eleocharis dulcis* (Type 2) | *Eleocharis cellulosa* (Type 2) | *Hypolytrum nemorum* | *Ananas comosus* | *Typha latifolia* |
|-----------------|----------------------------|------------------------------|---------------------------------|----------------------|------------------|-------------------|
|                 | Genome size (bp)          | 199,561                      | 193,234                         | 180,648              | 159,636          | 161,572           |
|                 | LSC (% of genome)         | 117,896 (59.1)               | 112,881 (58.4)                  | 95,644 (52.9)        | 87,482 (54.8)    | 89,140 (55.2)     |
|                 | SSC (% of genome)         | 9,601 (4.8)                  | 10,311 (5.3)                    | 8,150 (4.5)          | 18,622 (11.7)    | 19,652 (12.2)     |
|                 | IR (% of genome)          | 36,032 (18.1)                | 35,021 (18.1)                   | 38,427 (21.3)        | 26,766 (16.8)    | 26,390 (16.3)     |
|                 | Total number of genes     | 139                          | 132                             | 135                  | 131              | 131               |
|                 | Number of unique genes    | 105                          | 105                             | 110                  | 113              | 113               |
|                 | Number of unique protein-coding genes (duplicated in IR) | 72 (14)                      | 72 (12)                         | 76 (12)              | 79 (6)           | 79 (6)            |
|                 | Number of unique tRNA genes (duplicated in IR) | 29 (10)                      | 29 (9)                          | 30 (8)               | 30 (8)           | 30 (8)            |
|                 | Number of unique rRNA genes (duplicated in IR) | 4 (4)                        | 4 (4)                           | 4 (4)                | 4 (4)            | 4 (4)             |
|                 | Number of genes with introns | 17                            | 17                             | 18                   | 18               | 18                |
|                 | GC content (%)            | 32.6                         | 32.8                            | 34.9                 | 37.4             | 36.6              |
|                 | GC content of IR/ LSC/ SSC (%) | 37.6/ 30.2/ 25.5             | 37.8/ 30.4/ 25.8                | 38.5 / 32.6/ 28.1    | 42.7/ 35.4/ 31.4 | 42.4 / 34.4/ 30.5 |
|                 | Genic DNA (% of genome, GC %) | 76,207 (38.2, 38.4)         | 74,394 (38.5, 38.5)             | 98,905 (54.8; 38.1)  | 91,317 (57.2, 40.2) | 91,212 (56.4, 39.9) |
|                 | Intergenic spacers (% of genome, GC %) | 123,354 (61.8, 29.1) | 118,840 (61.5, 29.3)          | 81,743 (45.2; 31.1)  | 68,611 (42.8, 33.7) | 70,541 (43.6, 32.4) |
|                 | Gene density              | 0.70                         | 0.68                            | 0.75                 | 0.82             | 0.81              |
|                 | Putative gene losses      | accD, clpP, rpl23, rps18, ycf1, ycf2, infA and trnT-GGU | accD, clpP, rpl23, rps18, ycf1, ycf2, infA and trnT-GGU | accD, rps18 and infA | –                 | –                 |

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| Putative gene duplications | rpoA (2) rpl36 (4), trnM-CAU (6), trnH-GUG (3), trnP-UGG (3) and trnV-GAC (3) | rpl36 (2)* trnM-CAU (3), trnH-GUG (3) and trnV-GAC (3) | trnV-GAC (3) | - | - |

The number in () in putative gene duplications indicates the number of copies. Asterisk (*) on rpl36 gene indicates gene duplication in only plastome type 4 of *E. cellulos*
Table 3. Pairwise comparison of breakpoint and reversal distances for Eleocharis, Hypolytrum and basal Poales plastomes.

|              | Typha latifolia | Ananas comosus | Hypolytrum nemorum | Eleocharis dulcis | Eleocharis cellulosa |
|--------------|----------------|----------------|---------------------|-------------------|---------------------|
| Typha latifolia | -              |                |                     |                   |                     |
| Ananas comosus | 0/0            | -              |                     |                   |                     |
| Hypolytrum nemorum | 11/7          | 11/7          | -                   |                   |                     |
| Eleocharis dulcis | 25/20         | 25/20          | 16/13               | -                 |                     |
| Eleocharis cellulosa | 24/21         | 24/21          | 16/14               | 5/4               | -                   |
Table 4: Statistics of dispersed and tandem repeats in *Eleocharis*, *Hypolytrum*, and basal Poales plastomes.

| Family       | Cyperaceae | Bromeliaceae | Typhaceae |
|--------------|------------|--------------|-----------|
| Species      | *Eleocharis dulcis* Type 2 | *Eleocharis cellulosa* Type 2 | *Hypolytrum nemorum* | *Ananas comosus* | *Typha latifolia* |
| Genome size (IRa excluded) | 163,529 | 158,213 | 142,221 | 132,862 | 134,642 |
| GC %         | 31.6       | 31.8       | 34        | 36.3     | 35.5     |

**Dispersed Repeats (DR)**

| Length of DR | 39,752 | 31,118 | 12,520 | 1,495 | 1,210 |
| GC % of DR   | 30.9   | 30.2   | 32.6   | 35.9   | 33.7   |
| GC % without DR | 31.8 | 32     | 33.9   | 36.3   | 35.5   |
| % of DR in genome | 24.3 | 19.7   | 8.8    | 1.1    | 0.9    |

**Tandem Repeats (TR)**

| Length of TR  | 5,864 | 2,833 | 6,638 | 2,057 | 3,270 |
| GC % of TR    | 25.9  | 25.5  | 27.8  | 18.4  | 13.5  |
| GC % without TR | 31.8 | 31.9  | 34.3  | 36.6  | 36    |
| % of TR in genome | 3.6  | 1.8   | 4.7   | 1.5   | 2.4   |

**Total Repeats**

| Length of total repeats | 42,216 | 32,695 | 16,718 | 3,552 | 4,436 |
| GC % of total repeats  | 30.4   | 29.9   | 31.1   | 25.7   | 19.1   |
| GC % without total repeats | 32    | 32.1   | 34.3   | 36.6   | 36    |
| % of total repeats in genome | 25.8 | 20.7   | 11.8   | 2.7    | 3.3    |
Eleocharis dulcis

Type 2
(199,561 bp)

Eleocharis cellulosa

Type 2
(193,234 bp)

A

B

Syntenic blocks

Eleocharis dulcis

Type 1

Type 2

Type 3

Type 4

Eleocharis cellulosa

Type 1

Type 2

Type 3

Type 4

http://mc.manuscriptcentral.com/gbe/article-abstract/doi/10.1093/gbe/evaa076/5819553
Typha latifolia (Typhaceae, reference)

Ananas comosus (Bromeliaceae)

Hypolytrum nemorum (Cyperaceae)

Eleocharis dulcis

Eleocharis cellulosa
