The C-Fern (*Ceratopteris richardii*) genome: insights into plant genome evolution with the first partial homosporous fern genome assembly

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Ferns are notorious for possessing large genomes and numerous chromosomes. Despite decades of speculation, the processes underlying the expansive genomes of ferns are unclear, largely due to the absence of a sequenced homosporous fern genome. The lack of this crucial resource has not only hindered investigations of evolutionary processes responsible for the unusual genome characteristics of homosporous ferns, but also impeded synthesis of genome evolution across land plants. Here, we used the model fern species *Ceratopteris richardii* to address the processes (e.g., polyploidy, spread of repeat elements) by which the large genomes and high chromosome numbers typical of homosporous ferns may have evolved and have been maintained. We directly compared repeat compositions in species spanning the green plant tree of life and a diversity of genome sizes, as well as both short- and long-read-based assemblies of *Ceratopteris*. We found evidence consistent with a single ancient polyploidy event in the evolutionary history of *Ceratopteris* based on both genomic and cytogenetic data, and on repeat proportions similar to those found in large flowering plant genomes. This study provides a major stepping-stone in the understanding of land plant evolutionary genomics by providing the first homosporous fern reference genome, as well as insights into the processes underlying the formation of these massive genomes.

There are estimated to be over 400,000 species of extant land plants¹, encompassing an enormous array of morphological, physiological, and ecological diversity. Accompanying this diversity is extraordinary variation in genome size²⁻⁴, spanning a 2,500-fold range from the bladderwort *Genlisea aurea* (~60 Mbp)⁴ to that of the monocot *Paris japonica* (150 Gbp)⁵. How these genomes are chromosomally partitioned also varies immensely, as land plants span a 360-fold range in chromosome number, from 2n = 4 in *Haplopappus gracilis*, *Brachychlorella dichroosomatica*, *Zingieria biebersteiniana*, and *Colpodium versicola* to 2n = 1,440 in the fern *Ophioglossum reticulatum*, the highest number reported for any eukaryote³⁻⁷. Understanding the processes underlying this enormous breadth in both genome size and chromosome number has long been a major area of interest among evolutionary biologists. However, sampling biases towards smaller, less complex genomes (e.g., *Arabidopsis*: 135 Mbp, n = 5) and crops have pervaded plant genome projects. Fortunately, recent technological advances have enabled the assembly and analysis of large genomes, such as those of conifers⁸⁻¹⁰, providing novel insights into the processes underlying genome and chromosomal composition. Despite this progress, the large genomes typical of homosporous ferns remain uninvestigated.

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Polyploidy, or whole-genome duplication (WGD), is the traditional explanation for the large genomes and numerous chromosomes found in many plants, as WGD results in the complete doubling of the genome\(^{13,15}\). Among flowering plants, phylogenetic and genomic studies have identified WGD events preceding key radiations, such as those of the core eudicots (~70% of the flowering plants)\(^{11}\), monocots\(^{14}\), and the entirety of flowering plants\(^{14-17}\). In addition, it was demonstrated that even species with minute genomes, such as the carnivorous plant \textit{Utricularia gibba}, with \(n = 13\) and a genome size of 80 Mbp, have experienced multiple WGD events; \textit{U. gibba} has undergone at least three ancient WGD events in the last 80 million years\(^{14}\). Thus, genome size and chromosome number may not be reliable indicators of WGD. This disparity between genome size and chromosomal composition relative to WGD frequency has altered our understanding of genome evolution as the question has changed from whether or not an organism is polyploid, to how many rounds of polyploidy an organism or lineage has experienced in its evolutionary history.

Although a few fern genomes, such as those of the heterosporous water ferns (<1% of fern diversity), are less than 250 Mbp\(^{15}\), the average homosporous fern genome is 12 Gbp, nearly five times the size of the genome of maize (2.5 Gbp) and over 80 times that of \textit{Arabidopsis}. In addition, homosporous ferns typically have substantially more chromosomes than seed plants, with an average haploid chromosome number of 59 compared to 16 in flowering plants or 12 in gymnosperms\(^{20}\). As a result, longstanding hypotheses have proposed that multiple, repeated WGD events were the major factor contributing to the high chromosome numbers and large genomes of ferns\(^{20-22}\).

It was originally hypothesized that homosporous ferns undergo intense selection favoring polyploidy to buffer against a putatively high rate of inbreeding that results from their unique life history\(^{23,24}\). The homosporous fern life cycle includes a free-living haploid gametophyte phase with the potential for intragametophytic selfing (IGS)\(^{15}\), or gametophytic selfing sensu Haufler et al.\(^{24}\) – a process that can produce a completely homozygous diploid plant in a single generation and thus expose any deleterious mutations. However, numerous isozyme analyses demonstrated that fern species with the lowest chromosome numbers within a given genus (ranging from \(n = 27\) to 52) were functionally diploid, producing typical diploid numbers of isozyme loci rather than multiple loci as seen in truly polyploid species with multiples of these low chromosome numbers\(^{15-27}\). Despite the lack of isozyme evidence for repeated polyploidy in diploid fern species, multiple copies of chlorophyll a/b-binding protein genes were discovered in the diploid fern \textit{Polystichum munitum}, but the duplicated genes were nonfunctional\(^{28}\). Furthermore, early population genetic investigations showed that homosporous ferns have highly variable mating systems and are typically outcrossing, refuting the hypothesized force (intense inbreeding depression via IGS) driving selection for polyploidy\(^{29}\). More recently, a genetic linkage map showed that \textit{Ceratopteris richardii} has one of the highest proportions of duplicated loci among plants (76%) yet lacks large, duplicated blocks that would be indicative of polyploidy\(^{30}\). In addition, a paralog-age distribution analysis of \textit{Ceratopteris} estimated an ancient polyploidy event over 180 million years ago (mya); however, the data used for this analysis were from a shallow EST library\(^{31}\). Despite little evidence for ancient polyploidy in ferns, chromosome count models suggest that 31% of fern and lycophtye speciation events involve WGD, compared to 15% in flowering plants\(^{32}\). However, these estimates of WGD refer to relatively recent polyploidy events (neopolyploidy) evident from chromosome numbers rather than ancient (paleopolyploidy) events deep in evolutionary history.

Ferns are also the only major lineage of land plants with a significant positive relationship between genome size and chromosome number, suggesting that fern chromosomes are relatively static compared to those of angiosperms and gymnosperms for which no such correlation exists\(^{33,34}\). While repeated episodes of WGD followed by extensive silencing and rearrangement cannot be discounted as an explanation for the paradoxical genomic, genetic, and chromosomal composition of ferns\(^{35}\), alternative processes underlying their large genomes and high chromosome numbers must be explored. Most notable among these alternative explanations for the large genomes of ferns is the impact of transposable elements (TEs) on genome size, as TEs make up the majority of genome space in a variety of eukaryotic lineages. For example, TEs are responsible for the difference in genome size between cultivated rice (\textit{Oryza sativa}, 390 Mbp) and wild rice (\textit{O. australiensis}, 965 Mbp)\(^{36}\). Phylogenetic reconstructions of major TE families in various plant lineages suggest that bursts of TE insertion result in inflated genome size\(^{38-40}\). However, genome inflation does not seem to be a one-way street, because unequal homologous recombination can eliminate repetitive regions, such as those produced by TEs\(^{40,41}\). Analysis of three conifer "giga-genomes" (20–30 Gbp) showed that these large genomes were derived not through WGD, but rather via extensive expansion of ancient TEs (especially retrotransposons) and an apparent inability to shed these repetitive regions via unequal recombination\(^{9,42}\). While TEs provide a possible alternative explanation for the large genome sizes of ferns as demonstrated in conifers, they cannot explain the high chromosome numbers of ferns. It is possible that ferns have ancestrally high chromosome numbers and a relatively low rate of WGD, yet this begs the question of how the high chromosome numbers were initially obtained. Aneuploidy or chromosomal fission are also possible explanations for the high chromosomal complement of most ferns\(^{22,26,43}\). There are now hundreds of published flowering plant, gymnosperm, lycophtye, and lycophtye genomes, alongside the recent publication of two heterosporous water fern genomes\(^{19}\). While these water fern genomes, \textit{Azolla} and \textit{Salvinia}, are much-needed references within the fern clade, they are atypical of 99% of ferns, in that these species are heterosporous and have very small genomes with few chromosomes (1 C = 0.25–1.76 Gbp, \(n = 9–22\))\(^{19}\). To date, no sequenced genome is yet available for any homosporous fern\(^{44}\). This major information gap is made more startling when the high species diversity (>10,000 species), significant ecological roles, and economic importance of homosporous ferns are considered\(^{45-50}\). Due to their crucial phylogenetic position as sister to seed plants, ferns are key for investigating an array of both genomic and non-genomic traits and will permit a synthesis of genome evolution across seed plants\(^{51,52}\).

Here we investigated the genome of the homosporous fern \textit{Ceratopteris richardii} (C-fern; 11.25 Gbp, \(n = 39\)), characterizing and classifying TE composition and assessing the extent of WGD. Our genomic data for C-fern, together with the recently published heterosporous water fern genomes\(^{19}\), help provide a genome evolutionary
context not just for ferns, but also for all vascular plants. Collectively, these data will permit deductions about ancestral genome characteristics of seed plants and ferns, as in studies of other phylogenetically pivotal lineages. Specifically, this Ceratopteris genome provides critical insights into the evolutionary genomics and paradoxes of the genomically long-neglected fern clade, in addition to serving as a valuable reference for future investigations into land plant genome composition and dynamics.

**Results**

**Genome sequencing and assembly.** Here we present the first sequence of a homosporous fern genome, providing a new resource for plant and evolutionary biology. The ability of homosporous ferns to undergo IGS (see above) partially simplified the assembly of this complex genome, as it made the sporophyte completely homozygous so that heterozygosity was not an issue in assembly. However, the quality of the Ceratopteris genome assembly and the computational resources required to assemble and analyze the genome reflect the technological difficulties of working with such a large and complicated genome with no closely related reference genome.

With paired-end short-read libraries totaling ~24X coverage from 1.8 billion cleaned reads, we assembled the 11.25 Gbp Ceratopteris genome into ~15 million contigs (>100 bp) or 988,403 scaffolds (>1,000 bp) (Table 1). We then combined and reduced the number of scaffolds using 8–10 Kbp mate-pair reads (13X coverage), producing a genome assembly (CFern v1.1) of 626,576 scaffolds with an N50 of 16 Kbp and total length of 4.25 Gbp, representing about 38% of the Ceratopteris genome.

We also sampled a smaller portion of the Ceratopteris genome using long-read sequencing of 32 bacterial artificial chromosomes (BACs) of Ceratopteris. This subsample assembly (BAC.SubSample) only totaled 3 Mbp of the Ceratopteris genome (0.03%), but had an N50 of 97 Kbp, providing a small, but more accurate and contiguous sampling of the 11.25 Gbp genome as long-read technology is less biased by repeat elements and mis-assemblies. The GC content of Ceratopteris was 37.7%, very similar to that of both the gymnosperm Picea abies (Norway spruce) (37.6%) and the flowering plant Amborella trichocarpa (37.5%), yet lower than that of maize (46.9%), the liverwort Marchantia polymorpha (42.0%), and the lycophyte Selaginella moellendorfii (45.3%).

**Transcriptome assembly.** From 12 PacBio SMRT cells, we obtained ~850,000 reads from which we produced 97,084 full-length, high-quality, cleaned transcripts (IsoSeq.HQ) ranging from 285 to 11,353 bp in length. When mapped onto the CFern v1.1 assembly at 98% identity and 98% coverage, the IsoSeq.HQ transcripts were collapsed into 4,620 genes and 10,043 isoforms; however, when coverage was reduced to 50%, there were 11,924 genes and 23,278 isoforms. The 2.5-fold increase in identified genes and isoforms via reduced coverage shows that our scaffolds do not span entire genes in the majority of cases. To overcome this fragmentation and provide

| Cytometric Genome Size | 11.25 Gbp |
|------------------------|-----------|
| Chromosome number | 39 |
| **Assembly V1.0** | |
| Total Size | 4.21 Gbp |
| N50% | 300 bp |
| Gaps | 0 |
| % GC | 36 |
| CFern v1.0 (>1,000 bp) | 988,403 scaffolds |
| Total Size | 2.69 Gbp |
| N50 | 3,376 bp |
| % Gaps | 0.5 |
| % GC | 36 |
| CFern v1.1 (>1,000 bp) | 626,576 scaffolds |
| Total Size | 4.25 Gbp |
| N50 | 16,289 bp |
| % Gaps | 37 |
| % GC | 38 |
| CFern v1.1A (>10,000 bp) | 133,755 scaffolds |
| Total Size | 2.79 Gbp |
| N50 | 22,401 bp |
| % Gaps | 44 |
| % GC | 38 |
| BAC.SubSample | 35 scaffolds |
| Total Size | 3.03 Mbp |
| N50 | 97,182 bp |
| % Gaps | 0 |
| % GC | 39 |

Table 1. Ceratopteris genome assembly statistics.
a set of high-confidence gene models, we implemented the Cogent genome-free protocol\(^\text{54}\) to produce 18,179 gene models (\textit{UniCFernModels}) from the \textit{IsoSeq.HQ} transcripts. Searching for 1,440 embryophyte single-copy orthologs\(^\text{55}\), we found 53% complete, 4.5% fragmented, and 42.6% missing.

### Polyploidy

To address the decades-old question of how common ancient polyploidy is in ferns, we employed sequence-based and cytogenetic approaches, which assessed three different temporal scopes of evolutionary history. Using paralog-age distribution analyses, we identified 1,800 paralogous gene pairs in the \textit{UniCFernModels} with a \(K_S\) value between 0.1 and 2.1. A minor peak around \(K_S = 0.3\) was detected; however, such small, “recent” peaks are often a result of small-scale gene duplications, not WGD\(^\text{56}\). In contrast, a single major peak was revealed in the synonymous distance plot of \textit{Ceratopteris}, similar to those observed in \textit{Azolla} and \textit{Equisetum} (Fig. 1). Based on the significant transition from positive to negative in the SiZER plot, the \textit{Ceratopteris} peak was at \(K_S = 1.1\), compared to 0.8 in \textit{Azolla} and 0.75 in \textit{Equisetum}, similar to the original results found by Vanneste \textit{et al.}\(^\text{57}\).

To determine whether the peaks found in these three ferns (\textit{Ceratopteris}, \textit{Azolla}, \textit{Equisetum}) represent a shared WGD rather than three distinct WGD events, we used the Multi-taxon Paleopolyploidy Search (MAPS)\(^\text{58}\). We first recovered 10,182 orthogroups from the clustered amino acid sequences of \textit{Ceratopteris}, \textit{Azolla}, \textit{Equisetum}, \textit{Amborella}, \textit{Selaginella}, and \textit{Physcomitrella}. We isolated 4,836 orthogroups with amino acid sequences from all six species and estimated gene family trees for each orthogroup. Of the subtrees that fit the known fern topology, ((\textit{Ceratopteris}, \textit{Azolla}), \textit{Equisetum}), 34% supported a gene duplication in the most recent common ancestor (MRCA) of \textit{Ceratopteris} and \textit{Azolla}, and 19% of subtrees fitting the ((\textit{Ceratopteris}/\textit{Azolla}, \textit{Equisetum}), \textit{Amborella}) topology supported a gene duplication shared across the three fern species (Fig. 1) – relatively low proportions compared to similar studies that identified shared WGD events\(^\text{59}\). These low proportions suggest three lineage-specific WGD events rather than one or two shared events between the three fern taxa.

### Figure 1

**Polyploidy analyses of three fern species.** (A) Paralog-age distribution analyses and associated SiZER plots of three fern species. Upper panels are \(K_S\)-based histograms (0.05 bins) of paralogs in \textit{Ceratopteris richardii}, \textit{Azolla filiculoides}, and \textit{Equisetum giganteum}. Lower panels are SiZER plots of the above paralog-age distribution data and associated smoothing functions where blue indicates significant (\(\alpha = 0.05\)) increases, red significant decreases, purple insignificance, and gray too few data points to determine. The white lines show the effective window widths for each bandwidth. Both upper and lower panels are on the same x-axis. (B) MAPS analysis across land plants and the associated WGD events (shown as stars). The percentages of subtrees that contain gene duplications shared by the descendent species of a given node are above the phylogeny (connected by dotted lines). Dates are based on Testo and Sundue\(^\text{60}\) and Morris \textit{et al.}\(^\text{61}\).
While the previously described methods of data analysis for assessing WGD are appropriate at deeper time scales, both are susceptible to missing more recent WGD events. As mentioned above, relatively recent (close to zero along the x-axis) WGD in the $K_s$ plots may be mistakenly attributed to small-scale duplications, while MAPS can only identify WGD events that have occurred prior to the MRCA of the next closest taxon included in the analysis. In the case of *Ceratopteris*, that would be 280 million years to the divergence of *Ceratopteris* and *Azolla*, thus only events older than that can be identified by MAPS.

Our cytogenetic approach using FISH suggests the ploidy of an organism by localizing 125–150 Kbp BAC DNA fragments to the chromosomes where the DNA fragment is found. If the organism is diploid, only two localizations will be apparent, while a polyploid should have more than two localizations. BAC-FISH evidence of polyploidy is relatively short-lived as studies of *Nicotiana* allopolyploids found that five million years after the WGD event, the two parental genomes in the polyploid were no longer distinguishable due to genome turnover, mutations, and small-scale duplications. However, our BAC-FISH results further corroborated our sequencing results in demonstrating a lack of evidence for recent WGD in *Ceratopteris*. Significantly, we detected only two primary localizations of each BAC probe we exposed to the *Ceratopteris* chromosome preparations (Fig. 2), suggesting diploidy. In a few cases, weak secondary localizations, or “ghost bands,” were found on multiple chromosomes; however, these are likely a result of repeat elements that are distributed throughout the numerous chromosomes.

Repeat diversity. In total, ~42% of the *CFern v1.1 A* assembly was composed of repeat elements (Fig. 3a). The *Copia* LTR RTs were the most prolific with over 800,000 elements making up 16.5% of the assembled genome, followed by the *Gypsy* LTR RT superfamily with 330,000 elements and accounting for 7.5% of the genome (Table 2). In comparison, Class II DNA transposons include members of 17 different super-families, yet only totaled 52,000 elements and <1% of the genome. The LINE RTs similarly covered 1.6% of the genome across 64,000 elements. Low-complexity, satellite, and simple repeats all covered <0.5% of the genome.

The repeat content and percent coverage were considerably higher in the long-read BAC.SubSample assembly (63%). Nearly 26% of the subsample assembly was made up of *Gypsy* LTR RTs and 21.8% was *Copia* LTR RTs.
while the LINE RTs and DNA transposons represented 3.2% and 0.16%, respectively. Low-complexity and simple repeats made up 0.2% and 2.2% of the BAC.SubSample, respectively. The mean lengths of all of the repeat types in the BAC.SubSample, with the exception of the DNA TEs, were more than double those of the CFern v1.1A assembly, and the Copia and Gypsy elements were nearly three times as large in the subsample compared to those of the CFern v1.1A assembly.

Similar to the BAC.SubSample results, read-based analyses of Ceratopteris also estimated that ~60% of the Ceratopteris genome is repetitive with 17% in Gypsy elements and 30% in Copia elements (Fig. 3c). The read-based analysis and BAC.SubSample characterization analysis are more accurate for assessing general genome composition compared to repeat characterization of the CFern v1.1A assembly, as they are not biased by the short-read assembly which can have trouble assembling repetitive regions beyond the length of the reads. This assembly bias is probably the reason the CFern v1.1A assembly was limited in low-complexity, satellite, and simple repeat elements. This limitation is also apparent in comparing the mean lengths of the CFern v1.1A LTR RTs to those of the BAC.SubSample (Fig. 3b), as the latter assembly could likely span those longer repetitive regions via long-read technology.

We directly compared the repeat content of Ceratopteris to that of other land plants by applying the same assembly-based repeat characterization protocol to Amborella trichopoda, a monocot (Zea mays), a liverwort (Marchantia polymorpha), a lycophyte (Selaginella moellendorffii), a conifer (Picea abies), and a moss (Physcomitrella patens). We chose to run our own analyses on these genome assemblies rather than comparing our results to those of previously published results due to the wide variation in repeat characterization analyses utilized. In addition, we ran the read-based clustering analysis on the above taxa, with the exception of Selaginella, using short reads from these assemblies covering 0.5X of each genome. Due to the previously mentioned limitations of the CFern v1.1A assembly, we largely focused on the BAC.SubSample and read-based results for comparing relative proportions of repeats to other taxa.

Compared to six other land plant genome assemblies, the BAC.SubSample was second in repeat proportion behind only Picea (Fig. 3a). The BAC.SubSample had proportions of Copia elements similar to those of Zea, substantially higher than any of the other genomes analyzed. The other super-family of LTR RTs, the Gypsy elements, represented 25.7% of the BAC.SubSample with a mean length of 1,660 bp. In contrast, 35% of the Zea and Physcomitrella genomes were made up of Gypsy elements with mean lengths of 2,755 and 1,574 bp, respectively. The read-based analyses generally agreed with the repeat proportions of the six analyzed taxa with the exceptions of Amborella and Picea (Fig. 3c). These two taxa had lower overall repeat proportions in the read-based analyses (58% vs. 25% in Amborella, 80% vs. 50% in Picea), matching previous similar studies.

We assessed LTR RT richness by comparing recent (>90% LTR similarity) LTR RT exemplars among the seven species compared here (Table 3). Zea was by far the most diverse with 4,561 distinct LTR RT exemplars, followed by Physcomitrella at 1,217 exemplars and Picea with 509. The CFern v1.1A assembly was low in recent
LTR RT diversity with only 22 exemplars, similar to that of Amborella and Marchantia, which had 11 and 30, respectively. Ancient (75–90% LTR similarity) LTR RT richness differed greatly from recent LTR RT diversity in Zea and Physcomitrella, which only had 45 and 16 ancient exemplars, respectively (Table 3). Picea also had fewer ancient exemplars than recent LTR RTs with 276, but CFern v1.1 and Amborella both had more ancient than recent LTR RT exemplars with 82 and 55, respectively.

The quality of the genome assembly could have had a large effect on these interspecific comparisons of repeat diversity, number, and size, as well as genome size, as earlier demonstrated with the BAC.SubSample. For example, these eight genome assemblies spanned a 27,000-fold difference in scaffold N50 lengths between that of Picea (8 Kbp) and maize (217,960 Kbp). Thus, in addition to the genomes of CFern v1.1 and Picea being many times larger than that of maize, they are much more fragmented, making the identification of repeat elements more difficult and biased for identifying those repeats with smaller lengths.

To investigate LTR RT insertion timing, we identified 62 full-length, high-confidence LTR RT elements in the CFern v1.1 and BAC.SubSample assemblies. The insertion timing of these LTR RTs was relatively uniform.

Table 2. Ceratopteris repeat diversity and composition.

| Class          | Order       | Superfamily | Element count | Length (bp) | % Genome |
|----------------|-------------|-------------|---------------|-------------|----------|
| Retrotransposon| Uncategorized| 1458        | 523887        | 0.02        |
|                | RTE-BovB    | 421         | 128815        | 0.00        |
|                | Jockey      | 434         | 127827        | 0.00        |
|                | R1          | 2565        | 1241869       | 0.04        |
|                | RTE-X       | 8733        | 3937956       | 0.14        |
|                | L2          | 19047       | 3652063       | 0.13        |
|                | Li-Tx1      | 23635       | 1552256       | 0.56        |
|                | L1          | 47105       | 20464731      | 0.73        |
| LTR            | Uncategorized| 23507       | 7859618       | 0.28        |
|                | DIRS        | 361         | 45453         | 0.00        |
|                | Pao         | 1494        | 462814        | 0.02        |
|                | Gypsy-Troyka| 5331        | 3393861       | 0.12        |
|                | ERV1        | 8083        | 6191165       | 0.22        |
|                | Gypsy       | 329706      | 207014935     | 7.42        |
|                | Copia       | 812470      | 460237954     | 16.50       |
| DNA Transposon | Uncategorized| 3289        | 693293        | 0.02        |
|                | hAT-Tip100  | 374         | 251768        | 0.01        |
|                | CMC-Mirage  | 416         | 153347        | 0.01        |
|                | MULE-MuDR   | 425         | 41133         | 0.00        |
|                | TcMar       | 530         | 83430         | 0.00        |
|                | hAT-hATw    | 622         | 343180        | 0.01        |
|                | Harbinger   | 627         | 224160        | 0.01        |
|                | PiggyBac    | 1230        | 122417        | 0.00        |
|                | Dada        | 1981        | 1130399       | 0.04        |
|                | CMC-EnSpm   | 2339        | 1276711       | 0.05        |
|                | Sola        | 2739        | 1565090       | 0.06        |
|                | hAT         | 2774        | 819873        | 0.03        |
|                | Maverick    | 4082        | 2443709       | 0.09        |
|                | hAT-Ac      | 4625        | 1727566       | 0.06        |
|                | PIF-Harbinger| 4982       | 1214259       | 0.04        |
|                | En-Spm      | 10115       | 5487072       | 0.20        |
|                | hAT-Tag1    | 15720       | 6076071       | 0.22        |
|                | Helitron    | 4260        | 812693        | 0.03        |

Table 3. Genome composition and LTR-RT statistics in sampled land plant genomes.

| Genus            | Selaginella | Marchantia | Physcomitrella | Amborella | Zea | Ceratopteris | Picea |
|------------------|-------------|------------|----------------|-----------|-----|--------------|------|
| Genome Size      | 0.11        | 0.2        | 0.48           | 0.87      | 2.1 | 11.25        | 20   |
| % GC             | 45.3        | 42         | 33.7           | 37.5      | 46.9| 37.7         | 37.6 |
| N50 (Kbp)        | 1750        | 1366       | 17435          | 4927      | 217959| 22           | 8    |
| Recent LTR RT    | 166         | 30         | 1217           | 11        | 4561| 22           | 509  |
| Ancient LTR RT   | 33          | 24         | 16             | 55        | 45  | 8            | 276  |
over the past 7 million years (Fig. 3d). However, we found considerable differences in the LTR RT identification of these two assemblies as the majority of the identified LTR RTs in the BAC.SubSample originated within the last million years, while the CFern v1.1A assembly did not identify a single LTR RT within the past million years and instead had largely older (>4 mya) LTR RTs. In addition, we note that the BAC.SubSample had 28 full-length, high-confidence LTR RTs, while the CFern v1.1A assembly had 34, despite nearly a 1,000-fold difference in assembly length. These results suggest that the long-read sequencing of the BAC.SubSample was able to span and properly assemble these repetitive elements, while the short reads of CFern v1.1 could only assemble older, more heterogeneous repetitive elements.

Discussion

Ferns are the second most species-rich clade of vascular plants, with over 10,000 species. In addition, ferns are the sister lineage to all seed plants and first appeared approximately 430 million years ago according to fossil-calibrated phylogenies. The oldest unequivocal fossil fern is Ibyka amphikoma with a minimum age of 384 million years. However, most extant fern diversity arose within the last 40–60 million years during the Cenozoic Era following the rise to dominance of the angiosperms. Despite their substantial morphological diversity, sister relationship to seed plants, and lengthy evolutionary history, ferns represent the final frontier of land plant genomics.

Here we provide the first draft genome assembly of the 11.25 Gbp Ceratopteris genome, as well as a high-confidence set of gene models. We assessed the role of WGD in the evolutionary history of Ceratopteris at three distinct temporal scales. Despite a genome size five times that of classically "large-genome" flowering plants (e.g., maize) and with eight times more chromosomes than Arabidopsis, which has undergone at least five WGD events, we found evidence consistent with only one ancient WGD event in Ceratopteris. The diploid signal localizations of our BAC-FISH approach refute any recent WGD events that may have been ambiguous in the paralog-age distribution analyses. The three peaks in the paralog-age distribution analyses of Ceratopteris, Azolla, and Equisetum overlap and thus could potentially be a shared event before the divergence of these three ferns (Fig. 1). However, MAPS analysis indicates that only a minority of subtrees support shared duplications among these three taxa, suggesting three lineage-specific WGD events rather than a single shared event. Based on our analyses and the timing of the WGD event in Equisetum, another lineage belonging to the broader fern (monilophyte) clade, the WGD of Ceratopteris is likely older than that of Equisetum (92 mya) yet younger than the most recent common ancestor of Ceratopteris and Azolla (~280 mya) (Fig. 1b). The approaches used here are those standardly used for inferring ancient WGD events from transcriptomic and genomic data. Even complete transcriptomes or chromosome-level genomes can underestimate the true number of WGDs as pseudogenes and gene fragments may be filtered out during analyses. As such, all paralog-age distribution analyses and the resulting inferences regarding WGDs are considered minimum estimates; however, similar analyses of diploid and triploid Ceratopteris thalictroides found nearly identical Ks frequency distributions to the results presented in this study. If additional WGD events were obscured from the Ks analyses due to incomplete gene sampling, it is unlikely that transcriptome samples from a congenic species would hide the same events. The authors of the above-mentioned study dated the WGD event to 52 ± 1 mya due to their use of a relatively rapid synonymous substitution rate (11.64 × 10−9). Synonymous substitution rates are highly variable among plant lineages and across time; thus, we are more confident in our relative dating of the Ceratopteris WGD as occurring between that of Equisetum and the most recent common ancestor of Ceratopteris and Azolla due to our MAPS results.

Our results do not support hypotheses of frequent WGD in ferns followed by massive gene silencing and the slow loss of genetic material and instead lend credence to the hypothesis that ferns had ancestrally high chromosome numbers and underwent WGD rarely yet were unable or very slow to lose genetic material. This conclusion is in agreement with past studies based on isozymes, transcripts of Equisetum giganteum, a genetic linkage map of Ceratopteris, as well as ancestral reconstructions that found ancient WGD events to be rare in the evolutionary history of ferns despite the presence of many neopolyploids. While genomic analyses in flowering plants have shown that even very small genomes, such as that of Arabidopsis, have undergone numerous rounds of polyploidy, yet still have a low number of chromosomes, we find that ferns are much less dynamic, having undergone relatively few WGD events, yet retaining a high number of chromosomes. Ancestral reconstructions of chromosome numbers across ferns have suggested that the common ancestor of all ferns had a haploid chromosome number of 22, while many of the more diverse fern lineages had higher ancestral chromosome numbers, such as n = 30 in Pteridaceae. If chromosome numbers were ancestrally high in ferns, only a single WGD event would therefore be needed to reach a chromosome number of n = 39 in Ceratopteris (or n = 59, the average across all ferns) since the divergence of the common ancestor of ferns from that of seed plants 400 million years ago. Alternatively, the high chromosome numbers of ferns could be a result of aneuploidy or chromosomal fission. To better understand the cause of the high chromosome numbers of homosporous ferns, comparative syntenic and phylogenetic analyses will have to be applied across multiple fern taxa based on complete genome assemblies.

Similar to other large plant genomes, a large proportion of the Ceratopteris genome is composed of LTR RTs and other transposable elements (Fig. 3). This “genome obesity” is the likely result of a steady accumulation of transposable elements and an inability to discard them, as found in smaller flowering plant genomes. Importantly, Ceratopteris had a very low diversity of recent LTR RT exemplars when compared to other large-genome species such as maize or Picea. While this finding could be indicative of low LTR RT richness and high abundance, given that the counts of the LTR RTs were considerably higher in Ceratopteris compared to the other genomes, it is also possible that we are unable to identify the majority of full-length LTR RTs due to low scaffold contiguity with the CFern v1.1 assembly. The BAC.SubSample assembly and read-based analyses provide a more accurate representation of the general repeat composition of Ceratopteris; however, these results provide a much smaller representation of the genome and are limited in their resolution. Clearly, long-read technology will
be necessary to overcome and fully analyze a genome of this size, as short-read sequencing simply cannot span and assemble the repetitive structures found in *Ceratopteris*. However, expense must be taken into account in any sequencing project, and here the deep sequencing of an 11.25 Gbp genome using long-read technology would take much longer and cost much more than similar sequencing with short reads.

This study provides a major stepping-stone in the understanding of plant evolutionary genomics by providing the first homosporous fern reference genome, as well as unique insights into the processes underlying the formation of these massive genomes. Future efforts should focus on long-read technology to provide a complete assembly of multiple homosporous ferns—thus permitting more extensive comparisons of genome evolution and synteny across green plants.

**Methods**

**Tissue samples.** *Ceratopteris richardii* (Pteridaceae) is a fast-growing tropical fern, used globally in research laboratories as well as in K-12 and undergraduate biology courses for studying alternation of generations in plants. Inbred lines and single-gene mutants are commercially available and readily produced. For this study, spores from the Hn-n inbred line were kindly donated by Dr. Leslie Hickok (University of Tennessee). The spores were germinated on nutrient media\(^{77,78}\) and grown following the recommended conditions in the C-Fern Manual ([www.c-fern.org](http://www.c-fern.org)). We isolated the germinated gametophytes to individual petri dishes and growth media. Given that *C. richardii* is homosporous, the gametophytes are typically bisexual and produce both antheridia and archegonia. By isolating the gametophytes prior to sexual maturity, we ensured that any sporophytes that did develop were a product of gametes from a single gametophyte and thus completely homozygous (doubled haploid).

**Library construction and sequencing.** We extracted genomic DNA (gDNA) from *Ceratopteris* using a modified CTAB protocol\(^{79}\) and quality checked and quantified the gDNA using a Qubit fluorometer (Invitrogen, Carlsbad, CA, USA) and NanoDrop spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA). Genomic short-read library preparation and sequencing for *Ceratopteris* were completed by the University of Florida's Interdisciplinary Center for Biotechnology Research (UF ICBR). The gDNA was fragmented and size-selected for ~300 base pair (bp) inserts, and the sequencing of 150 bp paired-end (PE) reads was conducted on two runs of the Illumina NextSeq platform (Illumina, San Diego, CA, USA). Mate-pair (MP) libraries (125 bp PE, 8–10 Kbp inserts) were prepared and sequenced at the DNA Genome Sequencing and Analysis Core on two lanes of Illumina HiSeq 2000 (Illumina, San Diego, CA, USA).

We also subsampled the *Ceratopteris* genome with long-read technology to avoid the assembly biases inherent in short-read technology by sequencing BAC clones (Plate CR_Ba #624, Green Plant BAC Library Project, provided by Clemson University Genomics Institute). We selected 34 *Ceratopteris* Hn-n BAC clones to be grown, pooled, purified, and sequenced using the RS II platform (Pacific Biosciences, Menlo Park, CA, USA) at the Arizona Genomics Institute. The reads were cleaned and de novo assembled using the Hierarchical Genome Assembly Process (HGAP) in the SMRT Analysis software package (Pacific Biosciences, Menlo Park, CA, USA) to produce the BAC.SubSample assembly.

Long-read technology was also used to acquire a high-confidence set of gene models from sporophyte tissue. We extracted total RNA from sexually mature leaf tissue using the RNeasy Plant Mini kit (Qiagen, Hilden, Germany). The total RNA was size-selected for 0.8–2, 2–3, 3–5, and >5 Kbp with the SageELF (Sage Science, Beverly, MA, USA) at the UF ICBR. The libraries were prepared following the SMRTbell library protocol, and each library was sequenced on three PacBio SMRT cells (Pacific Biosciences, Menlo Park, CA, USA) to produce the BAC.SubSample assembly.

**Genome assembly.** The raw genomic PE reads were trimmed of adapters and then quality-filtered with Trimmomatic\(^{80}\), while the raw MP reads were trimmed of adapters and separated into MP, PE, and unknown reads with NxTrim\(^{81}\). All libraries were quality-checked before and after cleaning with FastQC ([www.bioinformatics.babraham.ac.uk/projects/fastqc/](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/)). We divided the cleaned PE reads into 24-mers with Jellyfish\(^{82}\) and plotted their frequencies with KAT\(^{83}\) to assess environmental contamination, organellar genome content, nuclear genome size, and repeat content.

The PE reads were assembled using Meraculous\(^{284}\) and a k-mer size of 61 based on the results of KmerGenie\(^{85}\) to produce assembly *CFern* v1.0. The scaffolds from the *CFern* v1.0 assembly were further scaffolded with the MP reads using the SSPACE assembler\(^{86}\) to produce the genome assembly *CFern* v1.1. To compare the content of *CFern* v1.1 with the overall content of the cleaned reads, we divided the assembly into 24-mers with Jellyfish\(^{82}\) and compared the resulting frequencies to those of the cleaned PE reads using the compare feature of KAT\(^{83}\). For subsequent analyses, only scaffolds over 10 Kbp were used (*CFern* v1.1A).

**Transcriptome assembly.** We cleaned and processed the long reads following the IsoSeq protocol\(^{34}\) in which the circular consensus sequences (CCS) were acquired from the raw reads and then classified and clustered. Only full-length, high-quality (accuracy > 99%), polished sequences (*IsoSeq.HQ*) were used for analysis following the Iterative Clustering and Error correction (ICE)/Quiver algorithm. The *IsoSeq.HQ* sequences were further collapsed into unique isoforms and genes using both genome-based and sequence-based protocols (see below).

For the genome-based method, the *IsoSeq.HQ* sequences were mapped to the *CFern* v1.1A assembly using GMAP (parameters: -f samse –n 0 –z sense_force\(^{87}\). The sam file output was sorted (parameters: -k 3,3 –k 4,4n), and transcripts were collapsed together (collapse_isoforms_by_sam.py, [https://github.com/Magdoll/cDNA_Cupcake](https://github.com/Magdoll/cDNA_Cupcake)). We used both 98% coverage and 98% identity as our full-length mapping cutoff and then searched for incomplete genes with 50% coverage and 98% identity.
Due to the fragmented state of the CFern v1.1A assembly, many transcripts did not map. Thus, we also used CD-Hit v4.6.4 (parameters: –c 0.99 –G 0 –al 0.90 –AL 100 –as 0.99 –AS 30) to cluster and collapse highly similar transcripts into putative isoforms without a reference genome. We then used those sequences with the Coding Genome reconstruction tool98 for genome-free isoform collapse and gene identification. This pipeline divided the sequences into 30-mers and then grouped those kmers into clusters based on pairwise distances. De Brujin graphs of the sequences for each cluster were then used to resolve sequencing errors and alternative splicing events and output putative genes. Due to the high accuracy, full-length, de novo nature of IsoSeq and subsequent cleaning protocols, these genes served as our reference gene models for Ceratopteris (referred to as UniCFernModels).

Polyploidy. The UniCFernModels data set was used in the DupPipe pipeline80,91 to estimate the relative age of gene duplications. DupPipe finds duplicate gene pairs and then estimates the divergence of these genes using the number of substitutions per synonymous site (Ks). The frequency of duplicate genes corresponding to a given level of divergence, as a substitute for timing, was plotted as a histogram, and peaks were inferred to represent synchronous gene duplications, indicative of ancient polyploidy events80,92. Genes from two other ferns, Equisetum giganteum97 and Azolla filiculoides95, were similarly analyzed and plotted for comparison. To reduce the subjectivity of smoothing based on varying bin sizes, we analyzed the Ks values of these three ferns using the SiZer (Significance of Zero Crossings of the Derivative)93 package in R v3.4.294. This analysis determines whether an increase or decrease in a scatterplot or histogram is significant at α = 0.05 and plots the changes along the original x-axis with blue coloration indicating a significant increase, red a significant decrease, purple insignificance, and gray too few data points to determine.

To determine whether the three ferns examined here (Ceratopteris, Equisetum, and Azolla), spanning over 400 million years since their most recent common ancestor78, share any ancient polyploidy events, we clustered the predicted proteins of Ceratopteris, Equisetum, Amborella, Selaginella, and Physcomitrella into orthogroups using OrthoFinder95. Only orthogroups with gene representatives from all six species were retained. The protein sequences of each orthogroup were aligned with MAFFT98, and the alignments were converted to nucleotide alignments using the pxxa2cdn tool in Phyx97. The alignments were stripped of highly ambiguous (>90% missing data) columns, and genes were produced with RAXML using 100 rapid bootstrap searches and the GTR+GA model of evolution98. These gene family trees were entered into the Multi-tAxon Paleopolyploidy Search (MAPS) package98. This package first filters all of the gene family trees for subtaxa that match the known species tree [here (Physcomitrella, (Selaginella, (Amborella, (Equisetum, (Ceratopteris, Azolla))))]. It then counts the number of subtaxa with gene duplications at a specific node in the species tree relative to the number of available subtaxa. A node with a high proportion of gene duplications is presumed to have a shared polyploidy event.

We also used a cytogenetic approach to assess more recent WGD. We conducted fluorescent in situ hybridization (FISH) using the previously described BAC clones as probes following Chester et al.99 and Champal et al.100. To produce the probes, the BAC DNA was extracted from the Escherichia coli culture and amplified by rolling circle amplification (RCA)105. The RCA product was labeled by nick translation with Cy5-dUTP and purified with a QiAquick Nucleotide Removal kit (Qiagen, Venlo, Netherlands).

Root tips for chromosome preparations were collected in the mornings and immediately treated with pressurized nitrous oxide for 1 hour before being fixed in 3:1 ethanol (EtOH): glacial acetic acid overnight at room temperature and transferred to 70% EtOH at −20 °C for long-term storage. The root tips were then treated and chromosome spreads prepared to produce slides for in situ hybridization with the fluorescently labeled probes99. The BAC FISH images were taken on an AxioImager M2 microscope with an AxioCam MR camera (Carl Zeiss AG, Oberkochen, Germany).

Repeat characterization. We took both structural- and homology-based approaches to repeat characterization following Campbell et al.102. As long terminal repeat retrotransposons (LTR RTs) comprise a sizable proportion of most plant genomes, a variety of tools was used to characterize these repeats in the CFern v1.1A assembly. Recent LTR RTs were clustered based on 90% LTR similarity usingLTRharvest (parameters: -minlenltr 100 -maxlenltr 6000 -mindistltr 1500 -maxdistltr 25000 -mintsd 5 -maxtsd 5 -motif tgca -similar 90 -vic 10)103 from the GenomeFerns package101. LTRdigest was then used to find elements with poly purine tracts (PPT) or primer binding sites (PBS) using the Genomic tRNA database102. Those elements were identified and further filtered for false positive elements by removing gappy elements (>50 Ns), recent gene duplications where the flanking regions of the LTRs are alignable, and nested RT insertions using custom scripts. LTR RTs with nested DNA transposons were also identified by searching DNA transposase protein sequences with BLASTx106. LTR RT exemplars were then identified based on 80% identity and 90% coverage from the filtered elements based on the internal sequences of the LTR RTs and then based on the LTR sequences. Older LTR RTs were similarly collected but with 75% similarity among the LTR sequences and lacking the TGCA motif. To exclude more recent LTR RTs from the older LTR RT library, the younger LTR RT exemplars were used to mask and exclude elements found in the older LTR RT library with RepeatMasker107. The two LTR RT libraries were combined (allLTR.lib) and used as the reference library to mask the CFern v1.1A assembly with RepeatMasker107.

The unmasked remainder of the assembly was inputted into RepeatModeler to identify repeat families de novo108. The RepeatModeler library and LTR RT library were combined, and unidentified repeats were searched against a transposase database107,109 using BLASTx and identified to superfamily when possible108. To ensure that fragmented plant genes were not included in the final repeat library, we queried all of our repeats with the SwissProt plant protein110 and NCBI RefSeq plant protein databases using BLASTx106. With our clean, final repeat library, we used RepeatMasker to quantify the repeat elements throughout CFern v1.1A.

To make direct comparisons with other plant genome assemblies of varying sizes, qualities, and lineages, we followed the same repeat annotation protocol for the genomes of Amborella trichopoda111, a monocot (Zea mays)112, a liverwort (Marchantia polymorpha)113, a lycophyte (Selaginella moellendorffii)114, a conifer (Picea abies)115,
and a moss (*Physcomitrella patens*) [63]. We also ran the same protocol on the BAC.SubSample assembly. To remove assembly biases, we used RepeatExplorer 2111 on cleaned reads for the above-mentioned taxa, with the exception of Selaginella which only used Sanger sequencing. Raw reads were downloaded from the NCBI Sequence Read Archive and EMBL European Nucleotide Archive, cleaned up to reduced to 0.5X coverage of their respective genomes, then run in RepeatExplorer2111 via the Elixir CZ Galaxy portal under default parameters against the Viridiplantae version 3.0 transposable element protein domain database.

### Dating repeat insertion events.
We used the highly accurate but conservative LTR_Retriever package [68] to identify full-length LTR RTs and date their insertion using both the CFern v1.1A and BAC.SubSample assemblies. We provided candidate LTR RTs from LTR_harvest and LTR_finder using a 90% similarity minimum threshold between LTRs and the presence of the TGCA motifs. The candidate LTR RTs were filtered, removing non-LTR RT repeat elements or those with large amounts of tandem repeats or gaps. Especially in fragmented genome assemblies, such as the CFern v1.1A, these requirements hugely reduce the number of LTR RT candidates but ensure that only full-length LTR RTs are analyzed. Following filtering, the long terminal repeat regions of each transposable element were aligned, and the Jukes-Cantor model was used to estimate the divergence time of the two LTR regions. We used a mutation rate of $6.5 \times 10^{-9}$ per site per year to estimate the years since insertion [16]. This mutation rate is half that of rice [69] and is a general estimate; therefore, the insertion times should only be used in reference to the relative timing of insertion, rather than as exact dates.

### Data availability
All of the raw reads and associated genomic and transcriptomic assemblies can be found in the NCBI BioProjects under PRJNA511033. All of the tissue used for sequencing came from one doubled haploid genotype (Voucher: M. Whitten #5841, University of Florida Herbarium, FLAS).

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Author contributions
D.B.M., P.S.S. and D.E.S. designed the study with advice from E.B.S., P.G.W. and W.B.B. K.H. optimized the BAC-FISH protocol for Ceratopterus. D.B.M. carried out the study, analyzed the data, and wrote the manuscript. All authors reviewed and edited the manuscript.

Competing interests
The authors declare no competing interests.

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