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Craig F. Barrett
Michael R. McKain
Brandon T. Sinn
Xue Jun Ge
Yuqu Zhang

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Authors
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Craig F. Barrett1,*, Michael R. McKain2, Brandon T. Sinn1, Xue-Jun Ge3, Yuqu Zhang3, Alexandre Antonelli4,5,6, and Christine D. Bacon4,5

1Department of Biology, West Virginia University
2Department of Biological Sciences, University of Alabama
3Key Laboratory of Plant Resources Conservation and Sustainable Utilization, South China Botanical Garden, Chinese Academy of Sciences, Guangzhou, PR China
4Department of Biological and Environmental Sciences, University of Gothenburg, Sweden
5Gothenburg Global Biodiversity Centre, Göteborg, Sweden
6Royal Botanical Gardens Kew, Richmond, United Kingdom

*Corresponding author: E-mail: cfb0001@mail.wvu.edu.

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Abstract

Mechanisms of genome evolution are fundamental to our understanding of adaptation and the generation and maintenance of biodiversity, yet genome dynamics are still poorly characterized in many clades. Strong correlations between variation in genomic attributes and species diversity across the plant tree of life suggest that polyploidy or other mechanisms of genome size change confer selective advantages due to the introduction of genomic novelty. Palms (order Arecales, family Arecaceae) are diverse, widespread, and dominant in tropical ecosystems, yet little is known about genome evolution in this ecologically and economically important clade. Here, we take a phylogenetic comparative approach to investigate palm genome dynamics using genomic and transcriptomic data in combination with a recent, densely sampled, phylogenetic tree. We find conclusive evidence of a paleopolyploid event shared by the ancestor of palms but not with the sister clade, Dasypogonales. We find evidence of incremental chromosome number change in the palms as opposed to one of recurrent polyploidy. We find strong phylogenetic signal in chromosome number, but no signal in genome size, and further no correlation between the two when correcting for phylogenetic relationships. Palms thus add to a growing number of diverse, ecologically successful clades with evidence of whole-genome duplication, sister to a species-poor clade with no evidence of such an event. Disentangling the causes of genome size variation in palms moves us closer to understanding the genomic conditions facilitating adaptive radiation and ecological dominance in an evolutionarily successful, emblematic tropical clade.

Key words: Arecaceae, Arecales, chromosome, dysploidy, genome duplication, genome size.

Introduction

Genomic studies across the eukaryotic tree of life reveal that genome size is not indicative of organismal complexity, known as the “C-value paradox,” or “C-value enigma” (e.g., Thomas 1971; Cavalier-Smith 1978; Lewin 1983; Gregory 2001, 2005). For example, the genomes of some simple chlorophyte algae are orders of magnitude larger than the genomes of many flowering plants, despite multicellularity and the extensive differentiation of tissues found in the latter. Genome size and complexity have been hypothesized to correlate with or even drive rates of speciation, but evidence is equivocal (reviewed by Kraaijeveld [2010]). Instead, polyploidy and genome size variation may be more strongly correlated with species richness among major plant clades (e.g., Soltis et al. 2009; Wood et al. 2009; Jiao et al. 2011; Puttick et al. 2015). Plant genomes vary immensely in size (2,400-fold), from 61 megabases (Mb) in the carnivorous Genlisea (Fleischmann et al. 2014) to the lilioid species Paris japonica, at 148.8 gigabases (Gb; Pellicer et al. 2010).

What causes such drastic genome size variation in plants? Genome expansion in plants occurs by well-characterized mechanisms, and polyploidy, including both autoploidy...
and allopolyplody, is often implicated (e.g., Hawkins et al. 2008; Soltis et al. 2009; Grover and Wendel 2010; Wendel 2015; Kellogg 2016). Genome expansion may also occur via tandem or segmental duplication of chromosomal regions (e.g., Zhang 2003). Genome size reduction is less well understood. From a mechanistic perspective, genomes can decrease in size via fractionation and diploidization following a polyploidy event, wherein chromosomes undergo purging of many duplicated regions and structural rearrangements; illegitimate recombination between chromosomes, where misalignment during synthesis leads to large chromosomal deletions; intrastrand recombination, where misalignments occur within a single chromosome leading to large deletions; homologous recombination during meiosis; and chromosomal inversions, particularly those that expose formerly pericentric regions to the distal, telomeric ends of chromosomes where they can be more easily be deleted (Devos et al. 2002; Bennetzen et al. 2005; Hawkins et al. 2008; Zenil-Ferguson et al. 2016; Ren et al. 2018). Transposable elements also play a major role in both genomic growth via bursts of, for example, copy-paste transposition, and in genomic downsizing by causing misalignments during synthesis (e.g., Bennetzen et al. 2005).

Monocots, comprising nearly one fourth of all flowering plant species, display the largest range of genome size variation among flowering plants (Leitch et al. 2010). Among monocots, the palms (family Arecaceae, with >2,500 accepted species) represent a diverse and ancient clade >100-Myr old (Couvreur et al. 2011; Givnish et al. 2018) and comprise major ecological components of all tropical ecosystems on Earth, especially in Southeast Asia and the Neotropics, where they are particularly diverse and abundant (Uhl and Dransfield 1987; Dransfield et al. 2008; ter Steege et al. 2006; Dransfield et al. 2008; Baker and Dransfield 2010; Wendel 2015; Kellogg 2016). Genome expansion may also occur via past whole-genome duplications (WGDs) (Al-Mssallem et al. 2011; Bacon et al. 2012). Observations based on comparing silent substitutions among duplicate gene pairs (Ks plots) suggest at least that oil and date palms (Elaeis guineensis and Phoenix dactylifera, respectively) show evidence of past whole-genome duplications (WGDS) (Al-Mssallem et al. 2013; Singh et al. 2013). The only formal phylogenomic analyzes to include more than one palm species are those of D’Hont et al. (2012) and McKain et al. (2016), providing more conclusive evidence of a shared WGD event among the two model palms, which represent subfamilies Arecocoeae and Coryphoideae, respectively.

Several questions remain with respect to genome evolution in the palms. Did WGD events influence genome evolution across the palms and close relatives, and if so, how and at what point in their evolutionary history? Does variation in genome size and chromosome number carry phylogenetic signal across palms and relatives? Here, we use publicly available and newly generated transcriptomic and genomic data, a densely sampled phylogenetic tree, and published data on genome size and chromosome number to address the above questions. Our specific objectives are to 1) reconstruct the evolution of genome size and chromosome number and 2) detect and place the hypothesized WGD event(s), both within a phylogenetic context.

Palms are a model lineage in which to test relationships among trait evolution, biogeography, paleoenvironments, and tropical biodiversity (e.g., Eiserhardt et al. 2011, 2013; Kissling, Baker, et al. 2012; Kissling, Eiserhardt, et al. 2012; Baker and Couvreur 2013; Couvreur and Baker 2013; Bacon et al. 2018). Analyses in palms will help to elucidate patterns of genome size evolution in long-lived monocots, which are typically understudied in the world of evolutionary genomics. Ultimately, our aim is to generate a framework in which to
integrate genome evolutionary dynamics, biogeography, and trait evolution to elucidate the drivers of palm biodiversity.

Materials and Methods

Phylogenetic Trees

Two recently published trees include dense taxon sampling for the palms (Faurby et al. 2016; Antonelli et al. 2017). The “SUPERSMART” tree (Antonelli et al. 2017) was chosen because it has the best taxonomic representation that matches the available genome size, chromosome number, and genome skim data (see below). The tree contains 733 species and 293 genera and is based on all publicly available data from 37 loci (see Antonelli et al. 2017 for details).

Transcriptomic Data

Data were compiled from previously published RNA-seq data sets across monocots from the Sequence Read Archive (https://www.ncbi.nlm.nih.gov/sra; last accessed December 12, 2018), and the OneKP Project (https://sites.google.com/a/ualberta.ca/onekp/; last accessed December 12, 2018). Complete genomes were downloaded from GenBank (https://www.ncbi.nlm.nih.gov/genbank; last accessed December 12, 2018). Additional RNA-seq data sets were generated for Chamaedora seifrizii (Arecales: Arecoideae) and one representative species from four genera in family Dasypogonaceae: Baxteria australis, Calectasia narragara, Dasypogon bromeliifolius, and Kingia australis (supplementary table S1, Supplementary Material online).

Genome Size and Chromosome Numbers

Genome sizes and chromosome numbers were obtained from the Royal Botanic Gardens, Kew Angiosperm DNA C-values database (Bennett and Leitch 2012; http://data.kew.org/cvalues) and Dransfield et al. (2008), using only “prime” estimates (i.e., excluding those with low confidence). Data and trees were pruned in the “APE” package of R (Paradis and Schliep 2019) to match sampled tips from the SUPERSMART tree at the species level.

Data and Tree Articulation

We attempted to maximize the match of each data set (tree, chromosome number, and genome size) at the species level (supplementary fig. S1 and table S2, Supplementary Material online). In cases where genome size, chromosome number, or genome skim data did not match at the species level, and there were multiple genome size estimates represented by different species within a genus, we used another species of the same genus for the genome size estimate. Although ideally, we would prefer only data from the same species for genome size (further, even from the same individuals per species), using a congener is unlikely to bias our results, because the focus of this analysis is on large-scale relationships among repeat fractions and genome sizes.

Transcriptome Assembly and Gene Tree Reconstruction

RNA-seq data were assembled using Trinity v.2.2.0 (Grabherr et al. 2011; Haas et al. 2013) as described in McKain et al. (2016). Reads were cleaned using Trimmomatic v.0.32 (Bogler et al. 2014) with adapter trimming for TruSeq adapter sequence using one seed mismatch, a palindrome threshold of 30, and a simple clip threshold of 10. After adapter trimming, a sliding window of 10 base pairs a minimum threshold average Phred score of 20 was used to trim reads based on quality. Finally, reads <40 bp in length were discarded. Once assembled, reads were mapped back to transcripts using bowtie v.1.0.0 (Langmead et al. 2009), and read abundance per transcript was estimated using RSEM v.1.2.29 (Li and Dewey 2011) using the “align_and_estimate_abundance.pl” script packaged with Trinity. FPKM (fragments per kilobase of exon per million fragments mapped) was estimated for each gene identified by Trinity. The percentage of mapped fragments per isoform was estimated and transcripts with a value of <1% were removed from further analysis. FPKM filtered transcripts were translated using the RefTrans pipeline (McKain et al. 2016, https://github.com/mrmckain/RefTrans).

Transcripts were aligned to gene models from the Ananas comosum v.1.0 (Ming et al. 2015), Asparagus officinalis v.1.0 (Harkess et al. 2017), E. guineensis v.1.0 (Singh et al. 2013), Oryza sativa v.7.0 (Kawahara et al. 2013), Phalaenopsis equestris v.1.0 (Cai et al. 2015), and P. dactylifera v.1.0 (Al-Mssallam et al. 2013) genomes using BlastX with an e-value cutoff of 1.0 × e−10 (Camacho et al. 2009). BLAST results were filtered to identify best hits as defined by transcript and gene model pairs with the lowest e-value and at least 85% bidirectional overlap. Best hit gene models were used to translate transcripts using GeneWise 2.2.0 (Birney et al. 2004). The longest translation for each transcript was estimated, and transcripts with a value of <12 taxa were removed from assemblies.

OrthoFinder v.2.2.1 (Emms and Kelly 2015) under default settings was used to circumscribe putative gene families. Diamond v.0.9.19 (Buchfink et al. 2014) with an e-value cut off of 0.001 and the BlastP algorithm was used to align sequences to each other for the initial steps of OrthoFinder. In addition to transcriptomes, gene models from genome sequences for P. dactylifera v.1.0, E. guineensis v.1.0, Musa acuminate v.1.0 (D’Hont et al. 2012), A. comosum v.1.0, O. sativa v.7.0, Pha. equestris v.1.0, and As. officinalis v.1.0 were used in gene family estimation. Orthogroups were filtered to remove those with sequences from <12 taxa. Amino acid sequences for each orthogroup were aligned using MAFFT v.7.313 with
automatic alignment algorithm selection (Katoh and Standley 2013). Aligned amino acid sequences were used to create a codon alignment of the nucleotide sequences using PAL2NAL v.13 (Suyama et al. 2006) under default parameters. Gene trees were reconstructed using RAxML v.8.2.4 (Stamatakis 2014) under a GTR+gamma evolutionary model and 500 standard bootstrap replicates.

Gene trees and accompanying codon alignments were passed to the perl script clone_reducer (Estep et al. 2014; https://github.com/mrmckain/clone_reducer; last accessed December 12, 2018) to identify putative single copy gene families. This script identifies clades with a bootstrap value of 50 or more that comprise a single species. The longest sequence in this clade is then used to represent the clade as a whole. From these reduced alignments, a set of 1,102 gene families were identified as single copy. It is possible that these are not truly single copy but appear single copy due to the incomplete sampling of the genome by transcriptomes. New gene trees were reconstructed for these reduced alignments as described above. The most likely tree for each of these gene families was used to estimate a coalescence-based species tree using ASTRAL—III v.4.xx (Mirabab et al. 2014) using default parameters. Due to its low total transcripts, Calectasia grandiflora was not included in the estimation of this species tree. We placed Calectasia in the position identified by Barrett et al. 2016, which had a congruent topology to the estimated relationships presented here.

Identification and Phylogenetic Placement of WGD Events
After filtering for a minimum number of 12 taxa per tree, a total of 6,242 gene trees were used to identify and phylogenetically place putative WGD events. The software PUG (McKain et al. 2016) was used to identify putative gene duplications that coincide with the topology of the reconstructed coalescence-based species phylogeny. We ran PUG with the “estimate_paralogs” parameter flag, which has PUG identify all possible paralogs in a given gene tree by identifying all possible transcript pairs derived from the same taxon in a single gene tree. Each multilabeled gene tree was rerooted to a non-Arecaceae and Dasypogonaceae outgroup with preference given in the order: Acorus americanus, As. officinalis, Pha. equestris, Typha latifolia, A. comosum, Neoregalia carolinae, O. sativa, Hanguana malayana, Costus pulverulentus, Musa acuminate, and Tradascantia paludosa. With PUG, each putative paralog pair was queried to identify the most recent common ancestor node in the gene tree. The taxon composition of the subtree identified by the most recent common ancestor node was used to identify the equivalent node in the species tree. A placement of the duplication on the species tree was considered acceptable if taxa above the node match those in the gene tree and at least one species sister to this clade in the species tree was found sister to the equivalent clade in the gene tree. For all gene trees and paralogs, we ran PUG to identify both unique duplications (the default) and all duplications (flag “all_pairs”) to identify support for putative WGD events.

Ancient State Reconstruction, Shifts, and Phylogenetic Signal
We reconstructed ancestral genome sizes and chromosome numbers initially in the “APE” and “PHYTOOLS” (“contmap” function, Revell 2012) under a Brownian Motion Model. We further applied an Ornstein–Uhlenbeck model to investigate evidence of significant shifts in trait values over time and across the tree using the R package “1ou” (Khabbazian et al. 2016), which requires no a priori assumptions on the locations of trait shifts. We additionally analyzed evolutionary changes in chromosome number across the tree with ChromEvol (Glick and Mayrose 2014). This software compares explicit models of chromosome evolution by parameterizing ascending and descending dysploidy (where the current number of chromosomes, $j = i + 1$ or $i − 1$, respectively, where “$i$” represents the ancestral chromosome number); WGD ($j = 2i$); demipolyploidy ($j = 1.5i$); chromosome number changes involving a “base” haploid chromosome number ($x$); and linear versus constant rates of change, where linear changes in chromosome number are dependent upon the current chromosome number. We removed Voanioala gerardi (2n = 596) from the analysis because the sampling in that clade is inadequate to reconstruct such a drastic change in chromosome number. We tested ten models of chromosome evolution under the same set of dysploidy parameters as above. We compared the fit of alternative models via the Akaike Information Criterion (Akaike 1974) and Akaike Weights (Wagenmakers and Farrell 2004). We tested for correlation between log-transformed genome size and chromosome number using phylogenetically independent contrasts (Felsenstein 1985; Garland et al. 1992).

Results
Evidence of WGD in Palms
We found unequivocal evidence for an ancient WGD event shared by all representatives of the palms included here, but not shared with the sister clade, Dasypogonales (fig. 1). Coalescent analysis of relationships based on 1,102 single copy nuclear loci yields a tree with representative Arecoidae sister to Coryphoideae, which together are sister to the monotypic Nypoideae, with Mauritia, representing the Calamoideae, the subfamily sister to rest of Arecaceae (fig. 1). Ceroxyloideae were not sampled here. We analyzed a total of 6,242 gene families and detected 2,685 unique gene duplications supporting the species tree topology with a minimal
Bootstrap value of 80, representing 31.5% of all sampled gene families. The palms shared 278 unique gene duplications (3,321 paralog pairs), representing 4.6% of all gene families analyzed.

Genome Size

We found a lack of phylogenetic signal for genome size (fig. 2, \( n = 54 \) species; Pagel’s \( \lambda = 7.97 \times 10^{-10}, P = 1.0; \) Blomberg’s \( K = 0.47, P = 0.6 \)). The ancestral genome size for palms under a BM model is \( 3.6 \text{ Gb} \) (95% confidence interval, or CI = 0.74 to 8.0 Gb). We found limited evidence for significant shifts in genome size; all of these shifts are increases relative to inferred ancestral values, in Borassus, Coccothrinax, Pinanga, Iriartea, and Voanioala (fig. 2). Comparison of chromosome number and genome size via phylogenetically independent contrasts yields no significant correlation (\( F = 0.3832, P = 0.54 \)).

Chromosome Number

Ancestral state reconstruction of diploid chromosome number as a continuous character under a BM model yielded a pattern of phylogenetic signal (supplementary fig. S2A, Supplementary Material online). The ancestral 2\( n \) value under BM is 32 for palms (\( n = 195 \) species). There is a reduction to 2\( n = 26 \) in Calamus (subfamily Calamoideae), and a general increase to 2\( n = 36 \) in subfamily Coryphoideae. Chromosome number is unchanged at the crown nodes of subfamilies Ceroxylidoideae and Arecoideae, and a reduction to 2\( n = 26 \) is again observed in many species of Chamaedorea, for which there is dense sampling relative to other genera. A putative chromosome doubling is observed in Arenga caudata relative to all other members of this genus (ancestral 2\( n = 32 \)), but few other such events are observed in our data set. Voanioala gerardi, with 596 chromosomes, was removed as an outlier. We found evidence for 77 shifts in chromosome number across the palms sampled (OU model, BIC = -5,739.041; supplementary fig. S3B, Supplementary Material online). We also found significant phylogenetic signal for chromosome number (Pagel’s \( \kappa = 0.41, P = 2.5 \times 10^{-10}; \) Blomberg’s \( K = 0.29, P = 0.001 \)).

A model of linear dependency had the best fit to our data among ten different models of chromosome evolution in ChromEvol (AIC weight = 0.264; supplementary table S4, Supplementary Material online). The maximum-likelihood estimate for ancestral chromosome number was 2\( n = 30 \), though posterior probability estimates were low for the deepest nodes of the tree (i.e., PP < 0.7; fig. 3). ChromEvol detected 34 changes in chromosome number in contrast to the 77 shifts identified under an OU model. Most changes in chromosome number were ascending dysploidy, that is, increases in chromosome number of \( n \geq 1 \) (fig. 3C), and there was one possible case of WGD in Arenga caudata (2\( n = 32 \rightarrow 64 \)).
Discussion

Our principal objective was to investigate the evolutionary history of genome evolution in the palms. We found unequivocal evidence of a WGD event shared by all palm subfamilies but not with the sister clade, the monocot order Dasypogonales. We also found evidence of phylogenetic signal for chromosome numbers, evolving predominantly via a linear model of dysploid change.

Shared WGD in the Palms

We found evidence for a shared WGD event across all palm subfamilies, suggesting that polyploidy likely played a role in the diversification and evolutionary success of this emblematic tropical clade (fig. 1). With our data it may be impossible to infer whether this was the result of auto- versus allo-polyploidy: coupled with extinction, accumulation of substitutions among retained duplicates over long temporal scales has likely saturated any patterns that could be used to distinguish between these two processes. Methods used to detect ancient allopolyploidy mostly center around deep reticulate patterns or preferential paralog retention from one parental species, but all these methods require at least some knowledge of the potential donor lineages (see Clark and Donoghue 2017). The palm WGD event must have occurred between ~119 and 85 Ma, that is, after the estimated split of orders Arecales and Dasypogonales but before the first divergence of subfamily Calamoideae from the rest of the palms (Couvreur et al. 2011; Givnish et al. 2018). Although previous studies have alluded to a palm WGD, the hypothesis was only based on divergence comparisons of paralogs within genomes and limited taxon sampling (Al-Mssallem et al. 2013; Singh et al. 2013; He et al. 2015; but see D’Hont et al. 2012; McKain et al. 2016 for explicit phylogenomic comparisons). Such “Ks” comparisons, in which frequency distributions of divergence among paralogs are compared within individual genomes or transcriptomes, are informative for evidence of WGD within a particular genome, but they do not provide a rigorous, phylogenetic, comparative test of shared WGD among taxa. In the present analysis, we definitively and precisely confirm the phylogenetic placement of a palm WGD event, moreover indicating that the palm event is older than has been recently hypothesized (70–75 Ma; e.g., van de Peer et al. 2017).

The fact that this WGD event is not shared with the sister clade of palms, order Dasypogonales, is of high significance in terms of potential implications for palm diversification. A growing number of examples like that of Arecales–Dasypogonales is being revealed with the expansion of phylogenomic studies (e.g., see Soltis et al. 2009; Renny-Byfield and Wendel 2014; Panchy et al. 2016). The most comprehensive analysis to date across angiosperms, using RNA-seq data from the 1KP project, revealed that 70 of 99 WGD events are associated with increases in species richness of one clade relative to a species-poor sister clade (Landis et al. 2018). Here, we present a scenario of a species-rich, evolutionarily successful, ecologically dominant, widespread clade with evidence of ancient WGD prior to or coincident with an adaptive
radiation. In contrast, its sister clade is relatively species-poor, geographically restricted, ecologically marginal, and lacking evidence of WGD. Although the relationship among ancient WGD and subsequent adaptive radiation (Arecales, vs. a lack thereof in Dasyypogonales) may be anecdotal, there are many other diverse plant clades with a history of WGD (van de Peer et al. 2017; Landis et al. 2018). These notably include the order Poales and family Poaceae (Paterson et al. 2009; Tang et al. 2010; McKain et al. 2016), Orchidaceae (Zhang et al. 2017; Unruh et al. 2018), Brassicaceae (Edger et al. 2015), Fabaceae (Lavin et al. 2005; Pfeil et al. 2005; Cannon et al. 2015), and Solanaceae (Schlueter et al. 2004). Thus, it is likely that the ancient WGD identified in this study contributed to palm diversification and ecological dominance in tropical and subtropical ecosystems globally. There is only limited evidence of WGD in palms from the RNA-seq or genome data included in this study (at the base of subfamily Arecoideae, see fig. 1), there are several interesting candidates based on chromosomal information, including, for example, Arenga, Jubaeopsis, Rhapis, and Voanioala (Röser et al. 1997; Leitch et al. 2010). It is unclear in Voanioala whether repeated rounds of WGD have led to the remarkable proliferation of chromosomes and large genome size, or if another mechanism is responsible, for example, rampant TE accumulation and chromosomal fissions.

Our results naturally prompt a further question: What are the functions of retained paralogs, after post-WGD diploidization has largely purged the duplicated remainder of the genome? We are currently limited in terms of our use of RNA-seq data, as these were taken from a single tissue type (young, developing leaves; e.g., Matasci et al. 2014). Thus, analysis of such a “snapshot” of gene expression may severely limit, or even bias, an assessment of retained duplicate gene function in palms from a whole-organismal perspective. Such an analysis would require more inclusive transcriptomes, sampling multiple tissues both spatially and temporally, as well as complete or draft genomes. This would provide crucial information related to the question of whether WGD did in fact contribute to genetic novelty and thus adaptive radiation in the palms relative to the sister clade, for example, as in the retention of duplicated glucosinolate pathway genes as novel herbivore defense mechanisms in Brassicaceae (Edger et al. 2015).

A second putative palm WGD was found prior to the divergence of the Areceae and Cocoseae tribes in the subfamily Arecoideae (fig. 1). This event is supported by 94 unique gene duplications (285 paralog pairs) with a bootstrap support value of 80% or more. Further investigation is needed to

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**Fig. 3.**—Maximum-likelihood estimate of ancestral state reconstruction of chromosome number based on a model of linear rate dependency in ChromEvol (i.e., chromosome number changes depend on the current chromosome number), allowing both ascending and descending dysploid changes (\(j + 1, j - 1\)) and WGD (2j). Colors correspond to 2n chromosome numbers, and numbers in brackets indicate the ML estimate of chromosome number for that node. Asterisks refer to the posterior probability for the highest-likelihood reconstruction of chromosome number. Black dots correspond to four of the five palms subfamilies sampled.
verify this WGD event through increased sampling of Arecoideae, and based on the low support values and the putative paraphyly of Howea in the coalescence tree, this may be an artifact. We detected both the sigma (228 unique duplication, Bootstrap = 80) and tau (731 unique duplications, Bootstrap = 80) events described in earlier analyses (McKain et al. 2016), with tau after the divergence of Acorus, and sigma prior to the diversification of Poales. We also confirmed previously identified events in Bromeliaceae (196 unique duplications, Bootstrap = 80; McKain et al. 2016), Commelinales + Zingiberales (283 unique duplications Bootstrap = 80, D’Hont et al. 2012), and Zingiberales (538 unique duplications, Bootstrap = 80, D’Hont et al. 2012) (supplementary table S3, Supplementary Material online). There was also signal for a commelinid WGD event occurring after the divergence of Asparagales from the remainder of the monocots but is likely an artifact of sampling.

Evolution of Genome Size and Chromosome Number

Genome size is not correlated with chromosome number in the palms when accounting for phylogenetic relationships, nor does it carry phylogenetic signal based on our current sampling. Gene space varies among palms, from over 35,000 genes in oil palm to over 40,000 in date palm (Al-Mssallem et al. 2013; Singh et al. 2013). Further, the oil palm genome reveals evidence for a role for segmental gene duplications in gene space expansion (Singh et al. 2013). An estimate based on a recently published transcriptome of N. fruticans, a monotypic species of mangrove-growing palms, reveals that up to 45,000 genes may be present (>32,000 were identified via BLAST searches), but these numbers carry great uncertainty as only leaf tissue was sampled (He et al. 2015). Repeat content is known to be a major driver of genome size in plants (e.g., Pellicer et al. 2018). The estimated total repeat content from the date palm genome (transposons, satellite DNA) is ~38%, whereas this number is greater in oil palm, at an estimated 57% (Al-Mssallem et al. 2013; Singh et al. 2013). It is highly unlikely that increases in gene content alone explain the most drastic examples of genome size expansion in palms (e.g., Voanioala), and thus these were likely due to rampant increases in repetitive elements.

Genome size increases appear to be associated with high species diversity in some palm genera but not in others (fig. 2 and supplementary table S2, Supplementary Material online). For example, Coccothrinax (up to 7.27 Gb) and Pinanga (up to 8.66 Gb) are both relatively species-rich genera (>50 and 100 spp., respectively) with genomes much larger than the ancestral size estimated for palms (3.6 Gb). By contrast, three other genera with large genomes are relatively species poor: (12.01 Gb, one sp., I. deltoidea), Borassus (8.41, five spp.), and Voanioala (38.24 Gb, one sp., Voanioala gerardi, but possibly up to four spp.; see Gunn 2004). Clearly more sampling of genome sizes is needed across the palms, especially at—or even below—the species level, allowing a test of the hypothesis that genome size variation and not genome size per se is associated with species diversity (e.g., see Puttick et al. 2015). Ideally, such comprehensive sampling of genome sizes should be paired with phylogenomic information for all species to allow phylogenetically informed comparisons. Moreover, intrageneric and even intraspecific variation in genome size can be substantial (e.g., in Dypsis, Phoenix, Pinanga; summarized in Dransfield et al. [2008] with references therein) necessitating population-level sampling.

We identified major trends in chromosome number evolution across the palms, even with only 195 species sampled for chromosome number and phylogenetic tree information. By explicitly modeling chromosome number across the tree, we detected ~34 changes in chromosome number, which is fewer than the number of significant shifts detected under an OU model (fig. 3). The treatment of chromosome number as a continuous character may be misleading, and thus explicit models of changes in chromosome number are necessary to effectively capture the evolutionary dynamics of changes across the tree. A linear model of chromosome evolution had the best fit out of ten alternative models (supplementary table S4, Supplementary Material online). This is a state-dependent model in which chromosome number changes are dependent upon the current chromosome number (Glick and Mayrose 2014). Although <10% of the ~2,500 palm species were sampled in this study, this suggests that sampling was enough to track a linear mode of evolution across many clades (fig. 3). Large sampling gaps would be expected to obscure the pattern of chromosome number changes; for example, a linear dysploid transition from $2n = 30 \rightarrow 32 \rightarrow 34 \rightarrow 36 \rightarrow 34 \rightarrow 32$ within a lineage or clade might be observed as $2n = 30 \rightarrow 36 \rightarrow 32$ if the taxa with $2n = 32, 34$, and 34 are not sampled, respectively.

Specifically, ascending dysploidy appears to be the predominant mode of chromosomal change in palms based on the data available, suggesting an overall net trend to more chromosomes. The only information on chromosome number available for the sister clade of palms, Dasypogonaceae, is that of Dasypogon hookeri, which contains less than half the ancestral chromosome number of palms ($2n = 14$ vs. $2n = 30$, fig. 3; Röser 2000; Leitch et al. 2010). It is plausible that a WGD event in the palm ancestor not shared with Dasypogonaceae may be responsible for this difference. It would be surprising to observe such a conspicuous pattern of chromosome number “doubling” given the propensity for idiosyncratic chromosomal number change post-WGD; such a doubling after the split of ancestral Arecaceae and Dasypogonaceae would have had to persist for >100 my of evolution (based on the divergence time estimates in Givnish et al. [2018]). However, just as palms display some of the slowest substitution rates among monocots (see Barrett et al. 2016), plant taxa with relatively longer generation times...
generally experience slower rates of postpolyploid diploidization, and perhaps the same is true for descending dysploidy (Mandáková and Lysak 2018).

Our finding of no significant relationship between genome size and chromosome number corroborates an earlier analysis based on comparison of genome size across different categories of chromosome numbers (Leitch et al. 2010). Changes in chromosome number can, however, be an important evolutionary force involved in species diversification, often following a polyploidy event. During post-WGD diploidization and fractionation, dysploidy changes in chromosome number can result in reproductive isolation and thus cladogenesis (e.g., Dodsworth et al. 2016; Clark and Donoghue 2017; Mandáková and Lysak 2018). Although WGD (genome doubling or additive fusion) is an important factor in plant diversification in many clades, less study has been devoted to the evolutionary consequences of dysploidy, which appears to be the predominant mode of chromosomal evolution in palms. Additional sampling of both Dasypogonaceae and Arecaceae is needed, as is a more inclusive, phylogenetically comparative analysis of chromosome number across monosporic, for example, including both anagenetic and cladogenetic changes (e.g., chromoSSE; Freyman and Höhna 2018).

Conclusions and Future Directions
Here, we have unequivocally identified an ancient WGD event shared by all palms and characterized the predominant mode of chromosomal change in palms as dysploidy. Remaining questions include the role of repetitive elements in palm genome size evolution and how different genomic attributes have collectively influenced species diversification during the long evolutionary history of this ecologically dominant, evolutionarily successful clade. In the future, it will be critical to obtain whole-genome sequences for multiple representatives of each palm subfamily (including the genome of N. fruticans, the sole member of subfamily Nypoideae), along with each of the four genera of Dasypogonaceae. These genomic resources will allow 1) comparative analyses of genome architecture and synteny, 2) analysis of gene family expansion and contraction with respect to adaptive radiation of the palms, 3) ancestral reconstruction of genome content and architecture (i.e., gene family copy numbers, gene order along chromosomes, and repeat content), and 4) associations of genomic features, important phenotypic traits, ecology, biogeography, and species diversification rates. Such a densely sampled, integrative framework in the palms will advance our understanding of the evolution of tropical biodiversity.

Supplementary Material
Supplementary data are available at Genome Biology and Evolution online.

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