

Investigating the diversification of holocentromeric satellite DNA *Tyba* in *Rhynchospora* (Cyperaceae)

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- Background and Aims Satellite DNAs (satDNAs) are repetitive sequences composed by tandemly arranged, often highly homogenized units called monomers. Although satDNAs are usually fast evolving, some satDNA families can be conserved across species separated by several millions of years, probably because of their functional roles in the genomes. *Tyba* was the first centromere-specific satDNA described for a holocentric organism, until now being characterized for only eight species of the genus *Rhynchospora* Vahl. (Cyperaceae). Here, we characterized *Tyba* across a broad sampling of the genus, analysing and comparing its evolutionary patterns with other satDNAs.
- **Methods** We characterized the structure and sequence evolution of satDNAs across a robust dadated phylogeny based on Hybrid Target-Capture Sequencing (hyb-seq) of 70 species. We mined the repetitive fraction for *Tyba*-like satellites to compare its features with other satDNAs and to construct a *Tyba*-based phylogeny for the genus.
- **Key Results** Our results show that *Tyba* is present in the majority of examined species of the genus, spanning four of the five major clades and maintaining intrafamily pairwise identity of 70.9% over 31 Myr. In comparison, other satellite families presented higher intrafamily pairwise identity but are phylogenetically restricted. Furthermore, *Tyba* sequences could be divided into 12 variants grouped into three different cladespecific subfamilies, showing evidence of traditional models of satDNA evolution, such as the concerted evolution and library models. Besides, a *Tyba*-based phylogeny showed high congruence with the hyb-seq topology. Our results show structural indications of a possible relationship of *Tyba* with nucleosomes, given its high curvature peaks over conserved regions and overall high bendability values compared with other noncentromeric satellites.
- **Conclusions** Overall, *Tyba* shows a remarkable sequence conservation and phylogenetic significance across the genus *Rhynchospora*, which suggests that functional roles might lead to long-term stability and conservation for satDNAs in the genome.

Key words: Holocentromere, repetitive DNA, satellite DNA, phylogenetic signal, Rhynchospora Vahl.

INTRODUCTION

Repetitive DNA has become a key element in genomic studies since it was discovered that these sequences compose a large fraction of most eukaryotic genomes (Gemmell, 2021). The satellite DNAs (satDNAs) are one of the most well-studied types of repetitive DNA, being composed by tandemly arranged, usually highly homogenized units called monomers (Garrido Ramos, 2017). Monomer consensus sequences are used to characterize similar satDNA into families (Novák et al., 2017; Oliveira et al., 2021). One of the key features of these sequences is the remarkably fast rates of sequence evolution, in terms of abundance and/or nucleotide sequence, resulting in accumulation of changes in short evolutionary times (Macas et al., 2002; Lower et al., 2018). This rapid diversification presents major challenges to the study of satDNA evolution (Plohl et al., 2012).

Although different theories have been proposed to explain satDNA evolution (Lower et al., 2018), the most prominent

models are the concerted evolution and library models, which are often complementary to each other (Plohl *et al.*, 2012; Garrido-Ramos, 2015; Camacho *et al.*, 2022). The concerted evolution model follows the idea that the monomers of a satDNA array accumulate mutations in an independent manner and that these mutations are homogenized and fixed along the array following a molecular drive process (Dover, 2002; Plohl *et al.*, 2012; Garrido-Ramos, 2015). The library model postulates that related species share a library of satDNA families of varying abundances caused by random expansion or contraction of arrays of these different satDNAs (Salser *et al.*, 1976; Fry and Salser, 1977).

These two models try to explain the fast changes of satDNA repeats in sequence and abundance, although there are cases of long-term satDNA sequence conservation, usually observed in different taxonomic levels, such as families (Mravinac *et al.*, 2002), orders (Robles *et al.*, 2004) and even phyla (Petraccioli *et al.*, 2015). The maintenance of 'relic' satDNA families might

be indicative of functional roles for these sequences in eukaryotic genomes (Plohl *et al.*, 2012). This functionality might be
related to expression of non-coding RNA sequences, associated
with nucleotypic and/or structural effects (Mravinac *et al.*, 2005;
Plohl *et al.*, 2008). Sequence-based nucleosome-prediction
models using conserved satDNA monomers suggest that they
might have a role in nucleosome formation (Tsoumani *et al.*,
2013; Escudero *et al.*, 2019). These models analyse sequence
properties such as dinucleotide periodicity and curvature patterns to predict the 'bendability' values of a DNA sequence,
which can be an indicator of interaction with histones (Liu *et al.*, 2011; Zhang *et al.*, 2013).

Satellite DNAs are usually an integral part of the heterochromatin, frequently being found in functional regions such as the telomere and the centromere (Achrem et al., 2020). With regard to the centromere, most eukaryotes present monocentric chromosomes, in which the centromere is restricted to a single region, usually composed by long arrays of repetitive sequences, especially satDNAs (Plohl et al., 2014). These centromere-specific satDNAs present varying degrees of sequence conservation (Melters et al., 2013). For example, within the Poaceae family, the CentO centromeric satDNA found in Oryza was later shown to have high similarity with CentC, present in the centromeres of maize (Cheng et al., 2002; Zhong et al., 2002). In contrast, centromeric satDNAs of some Solanum species were shown to be chromosome specific (Gong et al., 2012; Zhang et al., 2014). In holocentric species, which present a dispersed centromere along each chromatid (Bureš et al., 2013), it was believed for a long time that centromere-specific repetitive sequences were not present (Marques and Pedrosa-Harand, 2016). For example, well-studied holocentric organisms, such as Luzula elegans and Caenorhabditi elegans, did not present repeats associated with CENH3, even after detailed genomic characterizations (Subirana and Messeguer, 2013; Heckmann et al., 2011). This changed with the discovery of the 172-bp satDNA Tyba in the sedge species Rhynchospora pubera (Cyperaceae). Tyba showed a line-like distribution along a groove positioned at the outer part of chromatids in R. pubera chromosomes co-localizing with the CENH3 protein, being the first centromere-specific satDNA described for a holocentric species (Marques et al., 2015). Tyba sequences were later confirmed to be present in eight other Rhynchospora species (Rhynchospora alba, R. breviuscula, R. cephalotes, R. ciliata, R. colorata, R. exaltata, R. tenerrima and R. tenuis), while being absent in Rhynchospora globosa (Rocha et al., 2016; Ribeiro et al., 2017; Costa et al., 2021; Hofstatter et al., 2022). This already suggests an old origin for this satellite DNA, given the estimated distance (~30 Myr; Buddenhagen, 2016) between the clade containing R. cephalotes + R. exaltata and the clade containing the other species that presented Tyba. Whole-genome sequencing of *Rhynchospora* species revealed that Tyba-based holocentromeres also impact genomic architecture, epigenome organization and karyotype evolution (Hofstatter et al., 2022). This suggests that the presence of Tyba at centromeres might have an adaptive role and might influence species diversification.

Rhynchospora is a cosmopolitan genus composed of ~400 species with a North American centre of diversity, with preliminary divergence time estimates placing the origin of the genus between 38 and 49 Mya (Buddenhagen, 2016; Thomas, 2020;

Silva Filho *et al.*, 2021). The presence of *Tyba* across evolutionarily distant species, such as *R. pubera* and *R. cephalotes*, coupled with its holocentromeric localization, makes this sequence an interesting case to study the evolution and dynamics of centromeric DNA in non-monocentric organisms in a macroevolutionary context. Here, we have performed a genus-wide investigation about the tempo and mode of *Tyba* evolution in *Rhynchospora*, comparing it with other satDNAs found in the genus. We mined repetitive DNA information from filtered offtarget next generation sequencing (NGS) reads (Costa *et al.*, 2021) from 70 species of *Rhynchospora* representing the major clades of the genus and used a robust phylogenetic framework to serve as the background for studying *Tyba* evolution. We aimed to answer the following questions.

- 1. How widely distributed is *Tyba* in *Rhynchospora* when compared with other satDNAs?
- 2. How conserved is *Tyba* across the whole genus?
- 3. What could be the reasons/mechanisms for spread of *Tyba* across the genus *Rhynchospora*?

MATERIALS AND METHODS

Sequence data acquisition and filtering

All target-capture sequencing data analysed here were obtained from the thesis by Buddenhagen (2016). Because we used offtarget reads from target-capture sequencing, we opted to exclude from our analysis the data of any species that showed a percentage of annotated repeats smaller than the one obtained for R. cephalotes by Costa et al. (2021), which was the species with the least amount of classified repeats (~5%) that still presented good correlation values with genome-skimming data. In total, 77 accessions, representing 70 Rhynchospora Vahl species (~20% of the genus), were selected for our satellite mining analysis. From the same dataset, we collected data from six Carex L. species, two Chorizandra R. Br. species, Exocarya sclerioides Benth., Hypolytrum nemorum (Vahl) Spreng. and Scirpodendron ghaeri (Gaertn.) Merr. to serve as the outgroup (Supplementary data Table S1). All sequences used were deposited in GenBank under project number PRJNA672127.

Phylogenetic analyses and molecular dating

To anchor our findings on a phylogenetic backbone, we used the robust RaxML topology constructed by Buddenhagen (2016) with 256 target loci obtained by hybrid target-capture sequencing (hyb-seq). Although the original sampling contained 115 *Rhynchospora* accessions, the reads of some of these were not sufficient for the RepeatExplorer analysis, yielding poor annotations. Therefore, we pruned the original tree, leaving only 77 *Rhynchospora* accessions and the 11 outgroup species. This was done with the *drop.tip* function implemented in the package *phytools* (Revell, 2012) in R (R Core Team, 2022). This pruned tree was then submitted to a molecular clock analysis. Divergence times were estimated on BEAST v.1.8.3 (Drummond and Rambaut, 2007) through the CIPRES Science Gateway, using the pruned tree as a fixed topology. For calibration, we used the same points defined by Buddenhagen (2016),

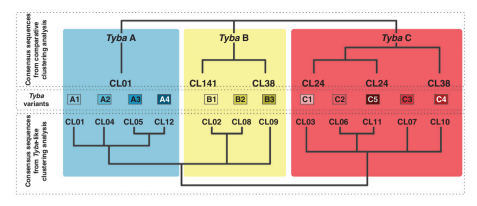


FIG. 1. Classification of satDNA family *Tyba* into subfamilies (uncovered in the species comparative clustering analysis) and variants [based on the *Tyba*-like (TL) reads in the clustering analysis]. Approximately maximum-likelihood (AML) trees show the phylogenetic relationships based on consensus sequence similarity. Coloured rectangles reflect the similarities between consensus sequences obtained in both analyses and are used to delimit sequences into subfamilies A (blue), B (yellow) and C (red). The clusters TL CL01, CL04, CL05 and CL12 formed a monophyletic clade and were all annotated as being similar to subfamily *Tyba* A (blue), being considered sequence variants A1–A4. Likewise, TL CL03, CL06, CL07, CL10 and CL11 formed a monophyletic clade that was annotated as being similar to *Tyba* C (red), thus being considered variants C1–C5. Although TL CL02, CL08 and CL09 were annotated as *Tyba* B, only CL02 and CL08 were monophyletic, with CL09 being in a polytomy with CL02 + CL08 and the clusters annotated as *Tyba* A. However, the most similar satellite to CL09 is CL02 (75% pairwise identity), which is further evidence that it should be considered a variant of *Tyba* B (yellow). Thus, CL02, CL08 and CL09 were considered to be variants B1–B3.

following a normal distribution with a 10% standard deviation. An uncorrelated relaxed lognormal clock (Drummond and Rambaut, 2007) and birth–death speciation model (Gernhard, 2008) were applied. Two independent runs of 100 000 000 generations were performed, sampling every 10 000 generations. After removing 25% of samples as burn-in, the independent runs were combined, and a maximum clade credibility (MCC) tree was constructed using TreeAnnotator v.1.8.2 (Rambaut and Drummond, 2013). To verify the effective sampling of all parameters and assess convergence of independent chains, we examined their posterior distributions in Tracer. The MCMC sampling was considered sufficient at effective sampling sizes (ESS) ≥ 200.

Satellite DNA mining

In order to prepare the target-capture sequencing data for satDNA mining, we first had to filter out all the reads containing the enriched targeted regions. For this, we followed the protocol presented by Costa et al. (2021) to acquire the unenriched off-target portion of the genomic libraries. The off-target datasets of each Rhynchospora species were uploaded to the RepeatExplorer pipeline (Novák et al., 2013) hosted at the webbased platform Galaxy (https://repeatexplorer-elixir.cerit-sc. cz/). RepeatExplorer uses a graph-based clustering algorithm to group sequences based on similarity, facilitating the identification of high-copy sequences of a genome. These clusters of sequences are then identified by cross-checking against repetitive element databases (Novák et al., 2013). Concurrently, the TAREAN (Tandem Repeat Analyzer) tool checks the clusters for predictions of tandem arrangement, building consensus sequences for these clusters (Novák et al., 2017).

Our dataset was submitted to three different run strategies in RepeatExplorer: (1) individual species clustering (ISC) analysis for each of the 77 *Rhynchospora* accessions in order to characterize the consensus of the satDNAs of each species; (2) a comparative clustering (CC) analysis with reads

from all the Rhynchospora accessions, in order to identify shared satellite DNAs; and (3) a Tyba-like clustering (TLC) analysis, a comparative analysis using only reads that were mapped to a database of previously published Tyba using the Geneious read mapper implemented in Geneious v.7.1.9 (low-sensitivity preset; Kearse et al., 2012). Given that the genome sizes of most analysed species were unknown, we input all reads left after the filtering of target regions (offtarget reads) for the ISC analysis, and the same number of reads (170 000) for each species was used to build the combined dataset for the CC analysis. This amount was decided based on the species that had the smallest number of reads analysed in the individual analysis (R. glaziovii, 173 544; Supplementary data Table S1). For the TLC analysis, we used all the 157 078 reads that were mapped to the Tyba database. The run parameters of the individual and comparative RepeatExplorer analysis were the same as those described by Costa et al. (2021).

The CC and TLC analysis were used to divide Tyba into subfamilies and variants, respectively (Fig. 1). The consensus sequences of clusters annotated as Tyba in the CC analysis were aligned using the Geneious alignment tool, with default settings (Kearse et al., 2012). The alignment was used to produce an 'approximately maximum-likelihood' (AML) phylogenetic tree using FastTree, as implemented in Geneious v.7.1.9 (Kearse et al., 2012), forming monophyletic clades that were considered different subfamilies of the satDNA (Fig. 1). The similarity between the shared satDNA found in the CC analysis (including non-Tyba satDNA families) was also assessed by a dotplot constructed with DOTTER (Sonnhammer and Durbin, 1995). The consensus sequences found in the TLC analysis were also used to construct an AML tree, and each of the clusters could be mapped to one of the subfamilies, being considered variants (Fig. 1). The consensus sequences found in the ISC analysis were mapped to the Tyba variants and non-Tyba shared satDNAs to estimate the sequence diversity of each shared satDNA. To assess the relationship between sequence divergence and age of the shared satDNAs, the pairwise identity (as a percentage) of sequences mapped to each shared satDNA was estimated with Geneious and plotted against the time of divergence between species that contained each of the shared satDNAs.

Satellite DNA-based phylogenetic analysis

We also performed a phylogenetic analysis based on the satDNA family Tyba and a separate analysis based on the other satDNAs found in the genus. For this, we separated all reads annotated as Tyba from the reads identified as other satDNAs ('non-Tyba' satDNAs) in the ISC analysis and adopted an Alignment and Assembly Free (AAF) methodology (Fan *et al.*, 2015). AAF constructs phylogenies directly from unassembled genome sequence data, bypassing both genome assembly and alignment. Thus, it calculates the statistical properties of the pairwise distances between genomes, allowing it to optimize parameter selection and perform bootstrapping.

We also used the individual Tyba consensus from the ISC analysis to estimate the phylogenetic signal (Pagel's λ ; Pagel, 1999) of monomer length and the GC content of Tyba sequences. For this, we use the phytools package (Revell, 2012) implemented in R (R Core Team, 2022). For species that had more than one Tyba variant, we used the consensus sequence of the most abundant Tyba cluster (Supplementary data Table S2). This analysis presents a measure of similarity between phylogenetically close species regarding the studied characteristics. In this case, a value of λ closer to one would mean that closely related species would have more similar Tyba variants, whereas λ closer to zero would mean that closely related species had less similar Tyba variants than expected (Pagel, 1999).

Sequence conservation

Given that the CC analysis included reads from all analysed species in a single run, it provides a broad view of satellite DNAs shared by two or more species. In contrast, with the ISC analysis each run contained reads from a single species, making it possible to find a greater number of satDNAs for each species. To determine whether these satDNAs were species specific or were part of the shared satDNAs, we mapped all the consensus sequences found in the ISC analysis to the consensus sequences of the shared satDNAs revealed in the CC analysis. For this, we uded the BowTie2 mapper (high-sensitivity preset, End-to-End, 60% pairwise identity) (Langmead and Salzberg, 2012). Given that we used a 60% pairwise identity threshold, ISC consensus sequences mapped to the same shared satDNA were considered to be from the same satDNA family. The alignments produced from the mapping were used to calculate the pairwise identity between all consensus sequences mapped to each shared satDNA. The age of shared satDNAs was estimated based on the age of the most common recent ancestor (MRCA) between the species that possessed the satDNAs, and a plot against pairwise identity was constructed using R.

Curvature/bendability analysis of shared satDNAs

Consensus sequences of the *Tyba* variants (mapped to their respective subfamily consensus sequences) and of the other shared satDNAs, including the non-centromeric satDNA previously found in *Rhynchospora globosa* (*RgSat*) (Ribeiro *et al.*, 2017)) were used to estimate bendability and curvature plots based on the DNA sequence using the bend.it server (http://pongor.itk.ppke.hu/dna/bend_it.html). It uses the DNase I-based bendability parameters of Brukner *et al.* (1995) and the consensus bendability scale (Gabrielan and Zohary, 2004). We also used the DNA curvature analysis website by Gohlke (https://www.lfd.uci.edu/~gohlke/dnacurve/) based on nucleosome positioning (Goodsell and Dickerson, 1994) to produce three-dimensional models of the *Tyba* subfamilies sequences by opening the Helix Coordinate PDB file in the Geneious software.

RESULTS

Phylogenetic framework

The phylogenetic reconstruction of *Rhynchospora* based on nuclear sequences showed five main clades (Fig. 2). According to our dating analysis, the crown age of the genus was estimated at 37.8 Mya [95% confidence interval (CI) = 33–42.1]. Clade V was the first to diverge, with crown age of ~25.6 Mya (95% CI = 22.2–28.8). Later, clade I (which comprises species traditionally classified as part of *Eurhynchosporae*; Gale, 1944) diverged from the remaining clades (31.8 Mya, 95% CI = 28–35.2), with clade IV diverging later from the rest at ~30.9 Mya (95% CI = 26.6–34.5). The remaining species split into clade II and clade III (species formerly recognized as *Pleurostachys*, recently synonymized as *Rhynchospora* section *Pleurostachys*; Thomas, 2020) at 27.6 Mya (95% CI = 24.5–31.2).

Tyba is a diverse satDNA family with several variants

The number of analysed reads of our ISC analyses ranged from 173 544 reads in R. glaziovii to 2 719 919 in R. pubera (Supplementary data Table S1). General satDNA abundance varied from 0.02% in R. emaciata to 9.14% in one of the accessions of R. corymbosa (Supplementary data Table S3). From the 537 satDNA consensus sequences found in the individual clustering analysis of all species, 133 were found to have ≥60% sequence similarity with one of the previously published Tyba sequences, with monomer length varying from 170 to 175 bp. These Tyba-like sequences were found in most species of clades I–IV, with only two species (R. biflora and R. racemosa) of clade II showing no trace of Tyba. Satellite DNAs from species of clade V did not show significant similarity with Tyba. We confirmed that these species from clades II and V did not present Tyba by being unable to find Tyba-like sequences in the full set of raw reads (Fig. 2).

In our CC analysis, we found 23 clusters that were annotated by TAREAN as satellite DNAs. From these, 17 clusters were found to have a significant amount of reads in more than one species, being considered shared satDNAs (Fig. 3;

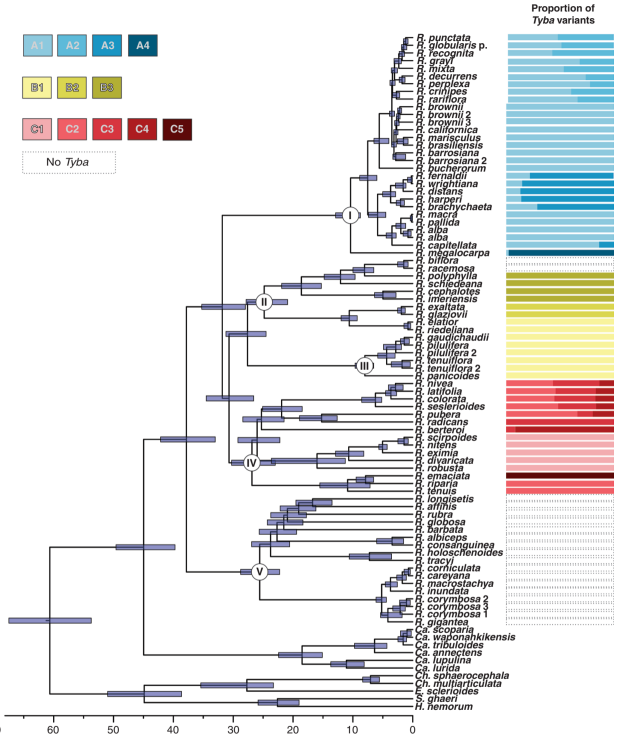


Fig. 2. Phylogenetic relationships and molecular dating of *Rhynchospora* species. Coloured bars (according to the key at the top left of the figure) represent the proportion of different *Tyba* variants found in the genome of each species. Bars at the nodes of the tree represent the 95% confidence interval of the molecular dating analysis, with the axis scale representing divergence time (in millions of years). Circles with roman numerals in the nodes delimit the five clades discussed.

Supplementary data Table S2). Six clusters of shared satDNAs (CL1, CL24, CL38, CL41, CL134 and CL141) were automatically annotated as having reads similar to *Tyba*, which was confirmed by the dotplot analysis (Supplementary data Fig. S1). The consensus sequences of these satDNAs were found to have

a 61.7% pairwise identity. Based on the high similarity and RepeatExplorer annotation, we determined that these satDNAs belonged to the same satDNA family. An AML phylogeny showed that these six *Tyba*-like satDNAs could be divided into three groups based on sequence similarity (subfamilies A, B

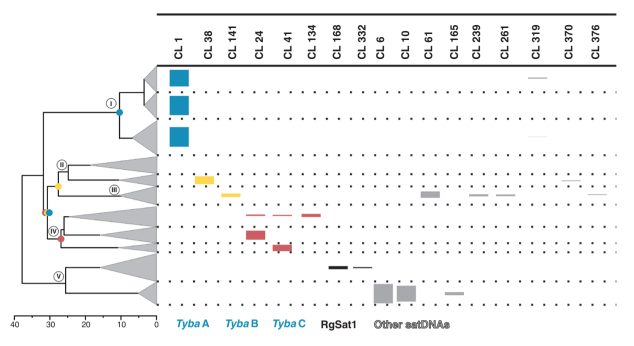


Fig. 3. Shared satDNAs in analysed *Rhynchospora*. Each line represents one of the clades of the simplified *Rhynchospora* phylogeny on the left (based on the tree presented in Fig. 1). Clusters annotated as satDNA by RepeatExplorer are presented in the upper part, and their relative abundance in a given species is represented by the height of the rectangles. Rectangle colours are according to the key below. Small coloured circles at the nodes of the simplified tree represent possible points of origin of the different *Tyba* subfamilies.

and C; see Fig. 1). Along with *Tyba*, we also annotated two further shared satDNA clusters that were found to be of the same family (Supplementary data Fig. S1) and presented high sequence similarity to RgSat, a non-centromeric satDNA found by Ribeiro *et al.* (2017). The clusters identified as RgSat also presented moderate similarity to CL06 (48.4% pairwise identity) and to the *Tyba* A subfamily sequences (46.0%), but these values were below our threshold for being considered to be from the same family.

In the TLC analysis, the 276 052 input reads were divided into 12 different clusters, each assigned to one of the main *Tyba* subfamilies by similarity hits to the custom repeat database (see Fig. 1; Supplementary data Table S2). An AML phylogenetic tree with the consensus sequence of the 12 *Tyba*-like clusters (TL-CLs) showed that these could also be divided into different clades. Each of these clades was assigned to one of the *Tyba* subfamilies discovered in the shared satDNA analysis based on the RepeatExplorer annotation. Thus, each of the TL-CLs was treated as one variant of a specific subfamily of *Tyba* (Fig. 1).

Tyba variants are clade specific within Rhynchospora

The CC analysis showed that *Tyba* was present among species of clades I, II, III and IV of *Rhynchospora*, with one of each *Tyba* subfamily being dominant in one specific clade, while subfamily B was present in both clades II and III (Fig. 3). No other satDNA was found in the CC analysis in more than one major clade. The RgSat satellite was found in clusters CL168 and CL332 (69.5% pairwise identity) and was present in only two species of clade V (*R. globosa* and *R. rubra*).

The TLC analysis mainly corroborated the results from the CC analysis (Fig. 3), with each clade presenting only one dominant *Tyba* subfamily (Fig. 2). Species of clades II and III showed only one variant from subfamily B per species. In contrast, in clades I and IV, a few subclades presented more than one variant from the same subfamily (A and C, respectively; Fig. 2).

Tyba sequences present high phylogenetic signal

Although discrepancies between AAF Tyba-like and hyb-seq topologies were observed at higher levels, we could retrieve a large number of monophyletic subclades congruent with the hyb-seq tree (Fig. 4). Most of the relationships inside clade I were well recovered in the AAF topology, with a few discrepancies, such as R. capitellata and R. bucherorum (Fig. 4). The only subclade from clade III that was not monophyletic in the AAF analysis was R. tenuis (R. emaciata + R. riparia). Support values of most nodes of the AAF tree were >95%, with a few lower supports at shallow levels and at the base of the clade containing species from clades II, III and IV (Fig. 4). In contrast, the AAF phylogeny based on the non-Tyba satDNAs had more incongruences when compared with the nuclear phylogeny (Supplementary data Fig. S2). Clades I and III were recovered as monophyletic, whereas species from the remaining clades were scattered throughout the phylogeny.

To investigate whether Tyba sequence variability reflected phylogenetic relationships, we also assessed Pagel's λ for the GC content (as a percentage) and monomer size (in base pairs) of the most abundant Tyba consensus sequence of each species. We found a high phylogenetic signal for both GC (λ = 0.92,

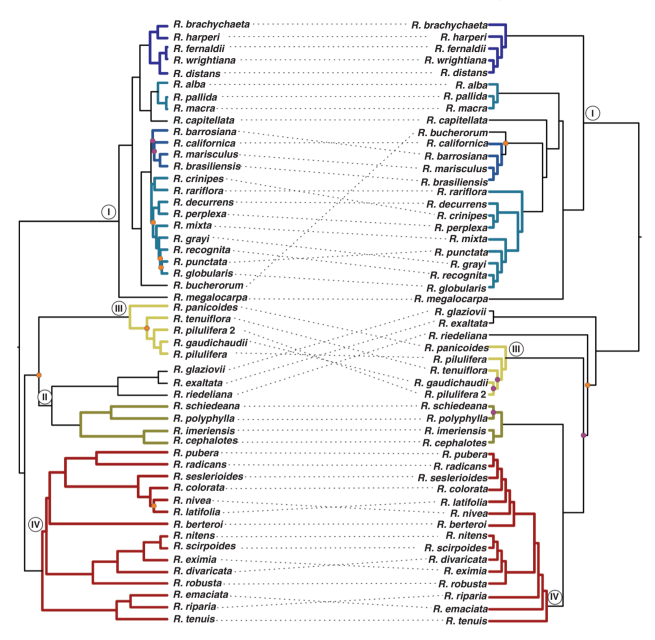


Fig. 4. Comparison of phylogenetic trees recovered from nuclear markers (left) and AAF of *Tyba*-like reads (right). Clades coloured in shades of blue (clade I), yellow (clade II) and red (clade III) are monophyletic in both trees. Circles at tree nodes refer to support values between 75 and 95% (orange) and <75% (purple).

Nodes without circles presented maximum support.

P < 0.001) content and monomer size ($\lambda = 0.93$, P < 0.001), indicating that Tyba sequences from closely related species have similar nucleotide composition and monomer size, whereas distant species do not share similar values of these traits.

Tyba sequences are remarkably different from other satDNAs

In order to determine sequence divergence within different satDNA families, we mapped all the 532 satDNA consensus sequences found in the ISC analysis to the 17 shared satDNA consensus sequences found in the CC analysis. In this way, we could identify whether/which satDNAs from each species could be considered to be from the same satDNA family. In

order to calculate intrafamily sequence divergence, we calculated the pairwise identity for all the resulting alignments. These identity values were plotted against the age of the MRCA of the species that shared that satDNA (Fig. 5). *Tyba* subfamilies presented lower pairwise identity than most of the other shared satDNAs (89.3, 76.4 and 82.3% for *Tyba* A, B and C, respectively), but were persistent through a longer evolutionary time (10, 28 and 27 Mya for *Tyba* A, B and C, respectively). Overall, all consensus satDNA sequences from the ISC analysis assigned to the *Tyba* family preserved a pairwise identity of 70.9%, with the MRCA of the species that contained *Tyba* having originated 31 Mya. The other shared satDNAs had high pairwise identity values (from 82.8% for CL10 to 100%

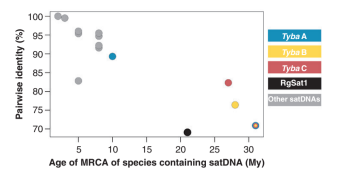


Fig. 5. Plot of pairwise identity through evolutionary time for the shared satDNAs recovered in the comparative clustering analysis. Dots represent the shared satDNAs, with colours reflecting the key on the right side. The multicoloured dot refers to the whole *Tyba* family.

for CL370), but were present only in species that diversified more recently, from 2 Mya (CL370) to 8 Mya (CL61) (Fig. 5). The only non-*Tyba* satDNA that was preserved through a relatively long evolutionary time was RgSat. Although it was present in only two species, those species shared a MRCA at ~21 Mya. However, the pairwise identity between the satDNAs mapped to the RgSat consensus was 69.1%. Overall, the *Tyba* subfamilies were older than most of the other satDNAs, while maintaining moderately high pairwise identity values (Fig. 5).

To check sequence conservation of the *Tyba* variants, the consensus sequences of the 12 variants were mapped to the consensus sequence of the three *Tyba* subfamilies, with identity values being calculated in a 2 bp sliding window. Overall, *Tyba* A had a smaller number of mutations among its variants, with a high identity throughout most of the sequence (Fig. 6). Both *Tyba* B and *Tyba* C variants showed a larger number of regions bearing low identity, but also presented highly conserved motifs, such as the ones highlighted by grey blocks in Fig. 6.

Curvature and bendability for the *Tyba* subfamilies and the other shared satDNAs were calculated based on DNase I parameters. Curvature peaks were found in regions that were fairly conserved in the variants of subfamilies A and C, which was not the case for the overall less-conserved variants of subfamily B (Fig. 6). Likewise, bendability values were higher at conserved regions separated by a less-conserved region between them, which presented the lowest bendability values (Fig. 6). The other shared satDNAs presented varied curvature/bendability profiles. Most sequences presented multiple curvature peaks, and bendability values did not follow the same pattern observed in *Tyba* (Supplementary data Fig. S2). The RgSat sequence, in contrast, presented a single thin curvature peak and overall low bendability values, with only one peak (Supplementary data Fig. S2).

DISCUSSION

The holocentromeric satDNA *Tyba* is conserved across the *Rhynchospora* phylogeny

We demonstrated that the holocentromeric satellite DNA *Tyba* is evolutionarily persistent and a relatively old family of satDNA in the genus *Rhynchospora*, especially when compared with the other shared satDNAs of the genus. Usually, satDNAs have a

fast rate of evolution and only species that are closely related share satellite families (Lower *et al.*, 2018). In more extreme cases, sequence divergence between centromeric satDNAs of different chromosomes in the same karyotypes have been reported (Zhang *et al.*, 2014).

The library model for satDNA evolution postulates that, depending on the homogenization rates (under concerted evolution), it is possible to find a low number of copies of a satDNA family in a phylogenetically distant species, whereas that family can be dominant on genomes in other closely related species (Plohl et al., 2012; Garrido-Ramos, 2015). Here, Tyba was found in four of the five main clades of Rhynchospora, and its monomer sequence could be classified into three subfamilies, with each clade presenting a single type. We could find different variants of Tyba in different abundances across closely related species, especially in clades I and IV. Although many species presented only one Tyba variant, it is important to note that we used reads in a way comparable to a genome-skimming analysis (Costa et al., 2021). This means that we had a small coverage for most of the analysed species, which, in turn, could mask low-abundant Tyba variants in the RepeatExporer analysis (Novák et al., 2017).

The propelling mechanisms of Tyba satDNA evolution

Different mechanisms seem to explain the diversification of *Tyba* holocentromeric satDNA in *Rhynchospora*. The high phylogenetic signal of GC content and monomer length and the moderate congruence between our *Tyba*-like AAF phylogeny and the hyb-seq phylogeny are evidence of efficient clade/ subclade-specific sequence homogenization of *Tyba*. Although the AAF phylogeny and the phylogenetic signal analysis used different types of data (raw NGS reads and TAREAN consensus sequences, respectively), both analyses showed that closely related species have more similar sequences of *Tyba* than distant species. Concerted evolution can lead to fast divergence between the satellites of reproductively isolated organisms, making these sequences potentially informative phylogenetic markers (Kuhn *et al.*, 2012; Lorite *et al.*, 2017; Belyayev *et al.*, 2019).

Our results suggest that both concerted evolution and the library model explain Tyba evolution in Rhynchospora. It has been demonstrated that mechanisms of concerted evolution can lead to the homogenization of different satDNA subfamilies that might have composed a genomic library of a group of species (Plohl et al., 2010; Ahmad et al., 2020). According to this view, all three subfamilies of Tyba might have existed in the satDNA library of the ancestor of all species that possess Tyba, as has been speculated for other phylogenetically conserved satDNA families (Quesada del Bosque et al., 2011, 2014; Lorite et al., 2017). However, ~30 Myr of evolutionary time might have led to the observed clade-specific homogenization. One of the results of this clade-specific homogenization is that the clade-specific homogenized sequences can become useful phylogenetic markers (Kuhn et al., 2012; Lorite et al., 2017; Belyayev et al., 2019). Although homogenization can occur independently from natural selection, it is possible that favourable characteristics could lead to evolutionary success of individuals presenting specific centromeric sequences, which could lead to the selection for particular sequence variants

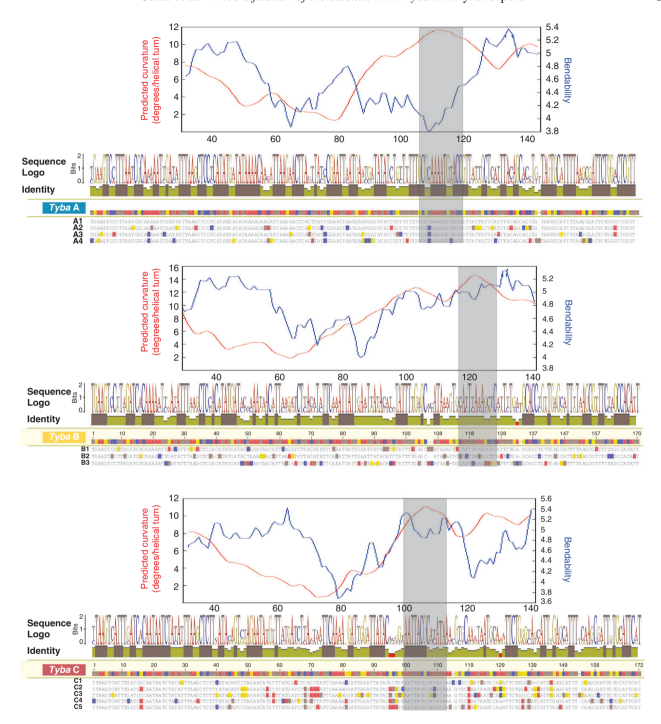


Fig. 6. Bendability (blue)/curvature (red) propensity, sequence logos, identity plot and alignment of *Tyba* variants assigned to each *Tyba* subfamily. Heights of nucleotides in the sequence logos are proportional to the number of times that base appeared in the alignment. Dark brown values in the identity plot represent high sequence identity, while yellow values represent moderate and red low sequence identity. Base pairs that are different from the consensus sequence are coloured in the alignment. Grey rectangles represent the approximate region where the curvature peak was estimated.

within a centromeric satDNA array (Pérez-Gutiérrez et al., 2012; Lower et al., 2018).

Upon its discovery, two variants of *Tyba* were identified in *R. pubera* (Marques *et al.*, 2015), and only one of these two was found in the closely related *R. tenuis* (Ribeiro *et al.*, 2017), both positioned in clade IV. In our analysis, we could observe a total of 12 different variants within three subfamilies in each of the

major *Rhynchospora* clades. Thus, patterns of both the library model and concerted evolution could be found in clade IV. Younger lineages present up to three different *Tyba* variants in their composition, whereas older lineages have only one dominant variant, which is congruent with the idea that concerted evolution is more efficient over long evolutionary times (Pérez-Gutiérrez *et al.*, 2012).

We found both variants previously discovered in R. pubera (Margues et al., 2015) and a third, new variant in this species. Comparatively, Tyba sequences from species of clades II and III appear to have had a faster homogenization process, with only a single variant being dominant in each species, but the same variant being shared with other species of the clade or even between clades suggests its older origin. This could lead to clade III and a subclade of clade II appearing as monophyletic in the AAF topology. A number of factors have been proposed to explain why homogenization rates might vary between satDNA sequences, such as array length, organization and genomic location (Navajas-Pérez et al., 2009; Kuhn et al., 2012; Lorite et al., 2017). The two variants of the oldest non-Tyba shared satDNA found in our CC analysis, RgSat, showed only 69% pairwise similarity. This satellite, first found in R. globosa, presents a traditional block-like distribution at the terminal regions of metaphase chromosomes and a clustered disposition at the chromocenters of interphasic nuclei (Ribeiro et al., 2017). In contrast, Tyba has been confirmed cytologically to be co-localized with the holocentromeres of species from clades II and IV (Marques et al., 2015; Rocha et al., 2016; Ribeiro et al., 2017; Costa et al., 2021) and was shown to have a dispersed distribution in interphasic nuclei (Marques et al., 2015; Ribeiro et al., 2017). Thus, the fact that a dispersed satDNA presents such high rates of homogenization, even when compared with a traditional block-like satDNA, is remarkable.

The *Eurhynchosporae* clade (formerly a tribe; Gale, 1944), corresponding to clade I in our phylogeny, presents an interesting case for satDNA evolution in a fast-diversifying group. This clade, which appears here as the first lineage to diverge among the clades that contain *Tyba*, was shown to present an increase in the diversification rate (Larridon *et al.*, 2021). In situations of rapid diversification, sequence evolution is often not rapid enough for sequences to diverge significantly between species (Giarla and Esselstyn, 2015). According to this view, species within a recent diversification event tend also to present with similar dominant satDNA variants (Quesada del Bosque *et al.*, 2013; Dogan *et al.*, 2021). Although the A1 variant is widespread among clade I in our analyses, the other three *Tyba* A variants appear in specific subclades at comparable abundance, in a classic library model display.

The evolutionary persistence of *Tyba* might be related to its centromeric distribution

Given its apparent phylogenetic conservation through long evolutionary times, what factors might be responsible for such long evolutionary reach of *Tyba*? It has been proposed in several studies that this long-term conservation of a satDNA sequence might suggest structural/nucleotypic functional roles for these elements (Mravinac *et al.*, 2005; Lower *et al.*, 2018). Immunofluorescence *in situ* hybridization experiments and chromatin immunoprecipitation sequencing (ChIP-seq) with the centromeric protein CENH3 have shown that *Tyba* is located at holocentromeres of *R. breviuscula*, *R. cephalotes*, *R. ciliata*, *R. pubera* and *R. tenuis* (Marques *et al.*, 2015; Rocha *et al.*, 2016; Ribeiro *et al.*, 2017; Costa *et al.*, 2021; Hofstatter *et al.*, 2022), which could indicate a functional association. Although we cannot ascertain that every satDNA identified here as *Tyba*, or derived from *Tyba*, has a holocentromeric

distribution, it might have a conserved chromosomal location because *Tyba* has been shown to be centromere specific in phylogenetically distant species, such as *R. cephalotes* (Costa *et al.*, 2021) and *R. pubera* (Marques *et al.*, 2015). Furthermore, the remarkably conserved sequence composition, monomer length and high abundance might be yet another indicator that all *Tyba* satDNAs found here are indeed centromeric, given that centromeric satDNAs are often the most abundant in most organisms (Melters *et al.*, 2013). More recently, sequencing of whole *Rhynchospora* genomes demonstrated the centromeric role of *Tyba*, which can impact evolutionary processes such as genomic architecture, epigenome organization and karyotype evolution (Hofstatter *et al.*, 2022).

Although satDNA is a major constituent of eukaryotic centromeres, sequence conservation is extremely low in most cases (Plohl *et al.*, 2014; Garrido-Ramos, 2017). In fact, centromeric satDNAs have been shown to differ among chromosomes of the same karyotype in some plant species (Gong *et al.*, 2012; Iwata *et al.*, 2013; Zhang *et al.*, 2014). An investigation of putative centromeric satDNA of 282 species across eukaryotes showed that sequence similarity rapidly declines through divergence, reaching background noise levels after 50 Myr of divergence (Melters *et al.*, 2013), which would make *Tyba*, at 31 Myr old, relatively old when compared with most centromeric satDNAs.

Our curvature/bendability analysis have shown the highest values of curvature and bendability at similar regions of the Tyba subfamilies sequences. In addition, this analysis also showed that *Tyba* sequences are more flexible than the sequence of RgSat, a confirmed non-centromeric satDNA. Other shared satDNAs presented variable patterns of curvature, with multiple peaks and mostly constant bendability values throughout the sequences. The curvature patterns of Tyba are similar to the proposed patterns of core DNA (tight sequences that wrap around a histone), in which large curvature at the ends of the sequence facilitate nucleosome formation (Liu et al., 2008). In the same way, high and stable bendability values for satellite sequences were also proposed to facilitate nucleosomal organization of centromeric proteins (Escudero et al., 2019). Our data are supported by genomic analyses that confirmed the centromeric role of Tyba satDNA (Hofstatter et al., 2022).

One important observation is that *Tyba* is not present in all *Rhynchospora* species, being absent from the whole of clade V and a few species from clade II, which emphasizes that *Tyba* is not necessary for holocentricity. For example, the clade V species *R. globosa*, which has meiotic evidence of its holocentricity (Arguelho *et al.*, 2012), presented only satDNAs disposed in block-like patterns (Ribeiro *et al.*, 2017). Although *Tyba* is probably not necessary for the holocentromere of *Rhynchospora* species, it is possible that its sequence confers advantages regarding nucleosome formation and positioning that might have resulted in its conservation for a long evolutionary time (Talbert and Henikoff, 2020).

It has also been shown that satDNAs can be associated with retrotransposons at functional parts of the centromere, as is the case with the satDNA *CentO* and retrotransposon CRR of rice (Cheng *et al.*, 2002). A similar association was found in *R. pubera*, where the *Ty3-Gypsy* retroelement CRRh was shown to have a similar pattern to *Tyba*, spread throughout the holocentromeres (Marques *et al.*, 2015). It has been proposed that an association with mobile elements could facilitate the

spread of centromeric satDNAs. Most notably, the association with mobile elements seemed to have facilitated the spread and persistence of relic satDNA BIV160 for >500 Myr of evolution (Plohl et al., 2010). In the bivalve mollusc Crassostrea gigas, a close association of a Helitron transposable element with centromeric regions seemed to facilitate the spread of abundant satDNAs to different chromosomes (Tunjić-Cvitanić et al., 2021). In Rhynchospora, Hofstatter et al. (2022) discovered that 468 Tyba loci associated highly with fragments from a non-autonomous Helitron (TCR1) in R. pubera, suggesting that Tyba satellite spread could have been facilitated by TCR1 transposition activity. Thus, the association with mobile elements could favour the conservation of Tyba sequences across Rhynchospora holocentric chromosomes, providing the mobility and means of fixation for this satDNA.

Here, 404 satDNA consensus sequences found in the individual species clustering analysis (mean of approximately five per accession) could not be mapped to the Tyba and non-Tyba shared satDNAs, meaning that those were probably species specific. The oldest non-Tyba shared satDNA (RgSat, previously found in R. globosa by Ribeiro et al., 2017) was present only in two distantly related species (R. rubra and R. globosa) that shared a common ancestor ~21 Mya. In contrast, clades containing Tyba shared a common ancestor 31.8 Mya and maintained a pairwise identity >70%. Although the concept of evolutionarily old satDNAs might be uncommon, it is not exactly rare (see Plohl et al., 2010; Quesada del Bosque et al., 2013). The long-term conservation of these 'relic' sequences might indicate that this type of repeat could provide evolutionary advantages by playing a functional role (Plohl et al., 2012; Lower et al., 2018).

Here, we have unravelled the evolutionary history of the holocentromere-specific satellite Tyba (Marques et al., 2015, 2016; Ribeiro et al., 2018) in what is one of the largest samplings for satDNA-related analysis to date. We observed a high sequence conservation and phylogenetic significance for Tyba, especially when compared with other satDNAs observed in the genus. Tyba seems to have evolved by a combination of satDNA library diversification and concerted evolution. The sequence has been shown to persist throughout a long evolutionary time, with clade-specific variants, and was present in most of the genus, in striking contrast to the other satDNAs found in Rhynchospora. Our results suggest that such conservation could be linked to a functional role within the holocentromeres. Although we present a thorough outline of Tyba evolution, future studies with long-read sequencing techniques might help us to gain a better understanding of the organization of Tyba arrays, which could provide further clues about its intimate association with the centromeres of Rhynchospora.

SUPPLEMENTARY DATA

Supplementary data are available online at https://academic.oup.com/aob and consist of the following.

Figure S1: dotplot of shared satDNAs found in the comparative clustering analysis of *Rhynchospora* species. Figure S2: bendability/curvature propensity plots of shared satDNAs RgSat, CL06, CL10, CL165, CL239, CL261 and CL319. Table S1: list of *Rhynchospora* species, voucher number, herbarium,

number of reads before and after filtering target reads and total number of reads analysed in the individual clustering analysis into RepeatExplorer. Table S2: name, monomer length, GC content and consensus sequence of shared satDNAs uncovered in the comparative clustering analysis, *Tyba* variants uncovered in the *Tyba*-like clustering analysis and of the *Tyba* consensus uncovered in the individual species clustering analysis. Table S3: proportion of different repetitive DNA classes per species identified in individual clustering analyses.

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CONFLICT OF INTEREST

All authors have declared that there are no conflicts of interest regarding this article.

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