Analysis of the small chromosomal *Prionium serratum* (Cyperid) demonstrates the importance of reliable methods to differentiate between mono- and holocentricity

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Abstract

For a long time, the Cyperid clade (Thurniceae-Juncaceae-Cyperaceae) was considered a group of species possessing holocentromeres exclusively. The basal phylogenetic position of *Prionium serratum* (Thunb.) Drège (Thurniceae) within Cyperids makes this species an important specimen to understand the centromere evolution within this clade. In contrast to the expectation, the chromosomal distribution of the centromere-specific histone H3 (CENH3), alpha-tubulin and different centromere-associated post-translational histone modifications (H3S10ph, H3S28ph and H2AT120ph) demonstrate a monocentromeric organisation of *P. serratum* chromosomes. Analysis of the high-copy repeat composition resulted in the identification of two centromere-localised satellite repeats. Hence, monocentricity was the ancestral condition for the Juncaceae-Cyperaceae-Thurniaceae Cyperid clade, and holocentricity in this clade has independently arisen at least twice after differentiation of the three families, once in Juncaceae and the other one in Cyperaceae. In this context, methods suitable for the identification of holocentromeres are discussed.

Keywords CENH3/CENPA · Centromere type · Holocentric chromosome · Evolution · Cyperids · Thurniceae

Introduction

Centromeres are essential for the segregation of chromosomes to the daughter cells during mitosis and meiosis. Most organisms contain one single size-restricted centromere per chromosome (monocentromere) visible as a primary constriction during metaphase. However, in independent eukaryotic taxa, species with chromosomes without distinct primary constrictions visible at metaphase exist, which are referred to as holocentric. Instead, the spindle fibres attach along almost the entire poleward surface of the chromatids (reviewed in Schubert et al. (2020)). Holocentricity evolved at least 19 times independently in various green algae, protozoans, invertebrates, and different higher plant families (Dernburg 2001; Escudero et al. 2016; Melters et al. 2012). A phylogenetic analysis of more than 50,000 species demonstrated that holocentric species are most likely derived from their monocentric ancestors rather than the other way around (Escudero et al. 2016). In total, ~1.5–2.0% of flowering plants are likely to have holocentric chromosomes (Bures et al. 2012). It is possible that holocentricity is even more common than reported so far, as the identification of the centromere type, especially in small-sized chromosomes, is challenging. Besides mono- and holocentric chromosomes, species with elongated monocentromeres were reported, such as *Pisum* and *Lathyrus* species (Neumann et al. 2012; Neumann et al. 2013).
2016) and the red fire ant Solenopsis invicta (Huang et al. 2016). Consequently, they were regarded as evolutionary intermediates via runaway expansion of their centromeres toward the development of holocentromeres (Huang et al. 2016).

One common explanation for the evolution of holocentric chromosomes is their putative advantage related to DNA double-strand breaks (Zedek and Bures 2018). The studies on artificial chromosomal rearrangements in various holocentric species showed that chromosome fragments retaining centromere activity are transmitted during mitosis and meiosis (Jankowska et al. 2015). Comparisons of diversification rates between monocentric and holocentric sister clades in animals and plants did not detect an increase in diversification in holocentric species (Marquez-Corro et al. 2018). Nevertheless, these analyses depend on the correct identification of the centromere type in a large number of lineages.

Because holocentric taxa are often embedded within broader phylogenetic lineages possessing monocentric chromosomes, it is thought that holocentric chromosome organisation originated from the monocentrics and that this transition occurred independently in multiple phylogenetic lineages (Melters et al. 2012). However, the factors that induced this transition and its mechanisms are currently unknown. Investigations of the changes associated with the transition from monocentric to holocentric chromosome organisation are, in theory, most informative when phylogenetically closely related species that differ in the centromere type are compared.

In angiosperms, holocentric chromosomes have been confirmed in some dicot species, e.g. in the genus Cuscuta L., subgenus Cuscuta (Convolvulaceae) (Oliveira et al. 2020) and in a few species within the genus Drosera L. (Droseraceae) (Sheikh et al. 1995). Also, in monocots, for example, in the genus Luzula DC (Juncaceae) (Heckmann et al. 2013) and Rhynchospora Vahl. (Cyperaceae) (Marques et al. 2015; Ribeiro et al. 2017) holocentricity occurs. These last two families belong to the Cyperid clade (Thurniceae-Juncaceae-Cyperaceae), which was originally considered to share holocentric chromosomes as a synapomorphic feature (Greilhuber 1995; Judd et al. 2016; Melters et al. 2012). However, exceptions have been reported in the genus Juncus L., in which four species exhibited primary constrictions (Guerra et al. 2019). It suggests that this synapomorphy of the Cyperid clade is uncertain.

Aiming to improve the understanding on the origin and evolution of the holocentricity within the Cyperid clade, we studied the centromere organisation of Prionium serratum (L.f.) Drège (Thurniceae), a species phylogenetically situated at the base of the Cyperid clade (Silva et al. 2020) (Suppl. Fig. 1, Hochbach et al. 2018; Semmouri et al. 2019). The South African monocotyledonous plant genus Prionium E. Mey is an old, species-poor lineage which split from its sister genus about 26.1 million years ago (Kumar et al. 2017). P. serratum is suspected to be holocentric, as it is closely related to the families Juncaceae and Cyperaceae. Supported was this assumption by the fact that this species has a low genomic GC content, as it is typically described for holocentric species (Smarda et al. 2014). Furthermore, Zedek et al. (2016) observed no significant increase in the proportion of G2 nuclei after gamma irradiation of P. serratum, differing from the situation found in monocentric species.

To ascertain the centromere type of P. serratum, we determined the chromosomal distribution of the centromere-specific histone H3 (CENH3) protein and alpha-tubulin fibres. In addition, antibodies specific for the cell cycle–dependent pericentromeric phosphorylation of histone H3 (H3S10ph, H3S28ph) and histone H2A (H2AT120ph) were employed to distinguish between a mono- or holocentric chromosome structure. In monocentric plants, immunostaining of mitotic chromosomes with antibodies against H3S10ph and H3S28ph typically results in a specific labelling of the pericentromere only. In contrast, in holocentric plants, immunolabelling with the same antibodies produces a uniform staining of condensed chromosomes, due to the chromosome-wide distribution of the pericentromere (Gernand et al. 2003). The cell cycle–dependent phosphorylation of histone H2A at position threonine 120 is associated with active centromeres (Demidov et al. 2014; Dong and Han 2012). Contrary to the expectation, a monocentromeric organisation of the chromosomes was found. The analysis of the high-copy repeat composition resulted in the identification of two centromere-localised satellite repeats. In addition, a DNA replication behaviour was found typical for small genome monocentric species. The data are discussed in the context of centromere evolution in Cyperids and concerning the suitability of available methods to identify holocentromeres.

Materials and methods

Plant material

Individuals of Prionium serratum (L.f.) Drège collected in western Cape (Cape Town, South Africa; TE2016_413) and provided by the Herrenhäuser Gardens (Hannover, Germany, IPK herbarium 70142) and the Botanical Garden Halle (Halle, Germany) were grown in a greenhouse of the Leibniz Institute of Plant Genetics and Crop Plant Research (IPK Gatersleben, Germany).

Flow cytometric genome size measurement

For nuclei isolation, roughly 0.5 cm² of fresh leaf tissue was chopped together with equivalent amounts of leaf tissue of one of the internal reference standards, Raphanus sativus.
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tively using the SeqGrapheR program (Novak et al. 2013).

Searches using databases (GenBank) were performed and
performed using the TAREAN (TAndem REpeat ANalyzer)
sequence of the tandem repeats, three independent runs were
corrected if necessary. The size of the annotated clusters was
manually checked, and their automated annotation was
All clusters representing at least 0.01% of the genome were
in total ~ 276 Mbp, corresponding to 0.82× genome coverage.

Domains in RepeatExplorer pipeline (Novak et al. 2013).

90%. Protein domains were identified using the tool Find RT
formed with a minimum overlap of 55% and a similarity of

quality cut-off value of 10 and interlaced. Clustering was per-
filtered by quality with 95% of bases equal to or above the

elixir.cerit-sc.cz/). Low-coverage genomic paired reads were
extracted from root meristems and prepared for paired-end
sequencing on Illumina HiSeqX (Illumina, CA) by
Novogene (Beijing, China).

DNA/RNA extraction and sequencing

Genomic DNA was extracted from P. serratum leaves using
the DNeasy Plant Mini Kit (Qiagen) and sequenced using the
HiSeq 2500 system (Illumina, CA) at low coverage. RNA was
extracted from root meristems and prepared for paired-end
sequencing on Illumina HiSeqX (Illumina, CA) by
Novogene (Beijing, China).

In silico repeat analysis

The repetitive proportion of the genome was analysed by the
RepeatExplorer pipeline (Novak et al. 2013), implemented
within the Galaxy/Elixir environment (https://repeatexplorer-
elixir.cerit-sc.cz/). Low-coverage genomic paired reads were
filtered by quality with 95% of bases equal to or above the
quality cut-off value of 10 and interlaced. Clustering was per-
formed with a minimum overlap of 55% and a similarity of
90%. Protein domains were identified using the tool Find RT
Domains in RepeatExplorer pipeline (Novak et al. 2013).

Searches using databases (GenBank) were performed and
graph layouts of individual clusters were examined interac-
sively using the SeqGrapheR program (Novak et al. 2013).

The number of analysed reads was 2,752,532 comprising
in total ~276 Mbp, corresponding to 0.82× genome coverage.
All clusters representing at least 0.01% of the genome were
manually checked, and their automated annotation was
corrected if necessary. The size of the annotated clusters was
used to characterise and quantify the genome proportion of the
high-copy repeats. To reconstruct the conserved monomer
sequence of the tandem repeats, three independent runs were
performed using the TAREAN (TAndem REPeat ANalyzer)
tool implanted in RepeatExplorer (Novak et al. 2017).

Repeat amplification, probe labelling and fluorescent
in situ hybridisation

Satellite DNAs (satDNA) were PCR amplified with primers
facing outwards of a repeat unit or directly synthesised as
oligonucleotides with 5′-labelled fluorescence. Primers and
oligonucleotides were designed from the most conserved re-
region of the consensus sequences (Table 1). Forty nanograms
of genomic DNA were used for all PCR reactions with 1×
PCR buffer, 2 mM MgCl2, 0.1 mM of each dNTP, 0.4 µM
of each primer, 0.025 U Taq polymerase (Qiagen) and water.
PCR conditions were 94 °C 3 min, 30× (94 °C 1 min, 55 °C 1
min, 72 °C 1 min) and 72 °C 10 min. Amplicons and plasmid
DNA of the 45S rDNA-containing clone pTa71 (Gerlach and
Bedbrook 1979) were labelled with either Cy3, Atto488 or
Atto550 fluorophores by a nick translation labelling kit (Jena
Bioscience).

Mitotic chromosomes were prepared from root tips, pre-
treated in 2 mM 8-hydroxyquinoline at 7 °C for 24 h and fixed
in ethanol: acetic acid (3:1 v/v) for 2 to 24 h at room temper-
ature and stored at −20 °C. Fixed root tips were digested with
2% cellulose, 2% pectinase, and 2% pectolyase in citrate buff-
er (0.01 M sodium citrate dihydrate and 0.01 M citric acid) for
90 min at 37 °C and squashed in a drop of 45% acetic acid.

Fluorescent in situ hybridisation was performed as described
by Aliyeva-Schnorr et al. (2015). The hybridisation mix
contained 50% (v/v) formamide, 10% (w/v) dextran sulphate,
2× SSC, and 5 ng/µl of each probe. Slides were denatured at
75 °C for 5 min, and the final stringency of hybridisation was
76%.

RNA sequence analysis

We generated a total 15.6 Gbp of paired-end reads of 150 bp
(around 52 million reads per end). Prior to mapping, all reads
were preprocessed for quality control with FastQC, Galaxy ver-
sion 0.72 (Andrews 2010). Subsequently, they were processed
with the Trimmomatic program, Galaxy version 0.36.6
(Bolger et al. 2014) to trim adaptor contamination and
low-quality sequences. As a result, 93.9% of high-
quality sequences from total number were used for de
novo transcriptome assembly with Trinity version 2.4.0.
To evaluate the quality of assembly, its completeness and
to remove poorly supported contigs, we applied Transrate
v1.0.3 (Smith-Unna et al. 2016). The resulting dataset
with 68,922 contigs was further processed by CD-HIT-
EST, v. 4.6.8 program, using -c 0.95 -n 10 as parame-
ters (Fu et al. 2012; Li and Godzik 2006)t oc l u s t e r
EST, v. 4.6.8 program, using -c 0.95 -n 10 as parame-
ters (Fu et al. 2012; Li and Godzik 2006)to cluster
highly homologous sequences and remove redundant
transcripts. Afterwards, the result file with 67,565
contigs was used to identify candidate coding regions
within the transcript sequences (Transdecoder v. 5.3.0;
http://transdecoder.github.io). RNAseq data are
deposited in the European Nucleotide Archive under
PRJEB39221 and genomic data are under NCBI
SRX8683442. To identify a CENH3 candidate in the
RNAseq data, we performed BLASTP, Galaxy Version
0.3.3 (Cock et al. 2015) using CENH3s from other
monocotyledonous plants.
Phylogenetic analysis

The CENH3 sequence selected from *P. serratum* transcriptome dataset and those of other species downloaded from NCBI GenBank (see Fig. 1, Suppl. Table 1) were aligned with ClustalW implanted in MEGA X, using the default setting (Kumar et al. 2018; Thompson et al. 1994). The evolutionary relationship was inferred using the maximum likelihood method by the IQ-Tree web server (http://iqtree.cibiv.univie.ac.at) (Trifinopoulos et al. 2016). The built tree was visualised, labelled and exported by Interactive Tree Of Life (iTOL, https://itol.embl.de/) (Letunic and Bork 2007, 2019).

Indirect immunostaining

The PsCENH3: RVKHFSNKAVSRTKKRIGSTR-c peptide was used for the production of polyclonal antibodies in rabbits. LifeTein (www.lifetein.com) performed the peptide synthesis, immunisation of rabbits and peptide affinity purification of antisera. Mitotic preparations were made from root meristems fixed in paraformaldehyde and Tris buffer (10 mM Tris, 10 mM EDTA, 100 mM NaCl, 0.1% Triton, pH 7.5) or 1 × MTSB buffer (50 mM PIPES, 5 mM MgSO₄, and 5 mM EGTA, pH 7.2) for 5 min on ice and for another 25 min only on ice. After washing twice in Tris buffer or 1 × MTSB buffer, the roots were chopped in LB01 lysis buffer (15 mM Tris, 2 mM Na₂EDTA, 0.5 mM spermine-4HCl, 80 mM KCl, 20 mM NaCl, 15 mM β-mercaptoethanol, 0.1% (v/v) Triton X-100, pH 7.5), filtered through a 50-μm filter (CellTrics, Sysmex) and diluted 1:10, and subsequently, 100 μl of the diluted suspension was centrifuged onto microscopic slides using a Cytospin3 (Shandon, Germany) as described (Jasencakova et al. 2001). Immunostaining was performed as described by Houben et al. (2007). The following primary antibodies were used: rabbit anti-PsCENH3 (diluted 1:300), mouse anti-alpha-tubulin (clone DM 1A, Sigma, diluted 1:200), mouse anti-histone H3S10ph (Abcam, 14966, diluted 1:200), mouse anti-histone H3S28ph (Millipore, 09_797, diluted 1:200) and rabbit anti-histone H2A120ph ((Demidov et al. 2014), diluted 1:200). As secondary antibodies, a Cy3-conjugated anti-rabbit IgG (Dianova) and a FITC-conjugated anti-mouse Alexa488 antibody (Molecular Probes) were used in a 1:500 dilution each. Slides were incubated overnight at 4 °C, washed 3 times in 1 × PBS or 1 × MTSB and then the secondary antibodies were applied. Immuno-FISH was performed, according to Ishii et al. (2015).

DNA replication analysis

Roots were treated for 2 h with 15 μM EdU (5-ethynyl-2′-deoxyuridine, baseclick GmbH), followed by water for 30 min. Preparation of slides was performed as described for immunostaining. The click reaction was performed to detect EdU according to the manual (baseclick GmbH).

Microscopy

Images were captured using an epifluorescence microscope BX61 (Olympus) equipped with a cooled CCD camera (Orca ER, Hamamatsu). To achieve super-resolution of ~120 nm (with a 488-nm laser excitation), we applied spatial...
structured illumination microscopy (3D-SIM) using a 63x/1.40 Oil Plan-Apochromat objective of an Elyra PS.1 microscope system (Carl Zeiss GmbH) (Weisshart et al. 2016).

**Results**

*Prionium serratum* is a monocentric species

*Prionium serratum* was chosen to test whether holocentricity occurs at the base of the Cyperid clade, since this species is phylogenetically positioned at the base of a group of species recognised as holocentrics. Since the roughly 1-μm long mitotic metaphase chromosomes did not allow an unambiguous identification of a monocentromere-typical primary constriction or a holocentromere-typical parallel configuration of anaphase sister chromatids, we generated a CENH3-specific antibody suitable for immunostaining. The centromere-specific histone variant CENH3 was shown to be essential for centromere function in many species (Allshire and Karpen 2008).

First, the root transcriptome of *P. serratum* was determined, and the assembled RNAseq reads were used to identify CENH3. Only one CENH3 gene was identified in the transcriptome dataset. After alignment of the corresponding amino acid sequence against CENH3s of other plant species, the evolutionary tree grouped *P. serratum* CENH3 together with other Cyperid sequences belonging to *Luzula* (Juncaceae), *Rhynchospora*, *Cyperus* and *Carex* (Cyperaceae), supporting the correct identification of the CENH3 gene (Fig. 1).

Next, antibodies (anti-PsCENH3) designed to recognise CENH3 of *P. serratum* were generated and used for immunostaining. Typical monocentromere dot-like signals were found at interphase and at early prophase (Fig. 2a, b). Additionally, an intense labelling of the nucleoli, likely representing unspecific immunosignals, was detected. The observed interaction of CENH3 with alpha-tubulin fibres at metaphase demonstrated the centromere specificity of the CENH3 signals (Fig. 2c). The application of super-resolution microscopy confirmed the close proximity of CENH3 and tubulin signals (Fig. 2d). Besides, the cell cycle–dependent, pericentromere-specific distribution of H3S10ph, H3S28ph and H2AT120ph approved a monocentric chromosome type (Fig. 3a–c). Hence, *P. serratum* is a monocentric species based on the results obtained by the application of different (peri)centromere-specific antibodies.
Identification of a centromere-localised repeat family in *P. serratum*

The genome size of *P. serratum* (*2n* = 46) is 335 Mbp/1C, estimated by flow cytometry. Next-generation sequence reads were generated to investigate the repetitive composition of the *P. serratum* genome based on the graph-based clustering analysis, resulting in the identification of high-copy satellite repeats and transposable elements. About 26.9% of the genome is composed of repetitive elements. The top first 329 clusters with at least 0.01% genome proportion, classified as 13 lineages of class I transposable elements (LTR retrotransposons and non-LTR LINE), six class II DNA transposons, satellite DNA (satDNA) and ribosomal DNA (rDNA) (Table 2). The LTR retrotransposons constituted ~9% of the genome, with the Ty1-Copia elements being more abundant than the Ty3-Gypsy elements, representing genome proportions of 5.36% and 3.63%, respectively.

The k-mer-based TAREAN analysis resulted in the identification of 19 different satDNA families. Out of these, the seven most abundant satDNAs were used for FISH to determine their chromosomal distribution. PsSat7, representing the

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**Fig. 2** Immunodetection of centromeric protein CENH3 (red) in *P. serratum* interphase nuclei (a), prophase (b) and its interaction with alpha-tubulin (green) in metaphase chromosomes (c, d). (d) Image taken by spatial structured illumination microscopy (SIM), enlargement (square) shows the interaction between CENH3 and alpha-tubulin.
Arabidopsis-type telomere sequence, hybridised to the terminal regions of all chromosomes. It is likely that the copy number of telomere repeats differs between the individual chromosome ends, as the intensity of the signals varied (Fig. 4a). PsSat41, PsSat311 and PsSat157 clustered on two, four and one chromosome pairs, respectively (Fig. 4b, c, d). PsSat306 colocalised with 45S rDNA signals (Fig. 4e).

Centromere-like signals were only found after FISH with the satDNA family PsSat156 (Fig. 4f). PsSat156a and PsSat156b possess a sequence similarity of 96% but with different abundance at chromosomes. Besides dot-like signals, both probes showed enlarged hybridisation signals on one but different chromosome pairs each. To confirm the centromeric position of PsSat156a, b, immuno-FISH with the CENH3-specific antibody was performed. Colocalisation of both signals in metaphase chromosomes and interphase nuclei demonstrated the centromere specificity of the repeat family PsSat156 (Fig. 4g, Suppl. Fig. 2). No sequence similarity was found between PsSat156 and centromeric repeats of other species.

Finally, we analysed the DNA replication behaviour of *P. serratum* by 5-ethynyl-2′-deoxyuridine (EdU) incorporation, a nucleoside analogue of thymidine. In general, different stages of the S phase are characterized by contrasting DNA replication patterns. The early S phase was characterized by dispersed EdU signals, and clustered EdU signals were typical for the late S phase (Costas et al. 2011; Němečková et al. 2020). Whether the replication behaviour of mono- and holocentric species differs is unknown yet. However, in the holocentric species *L. elegans*, the chromosomes are less clearly compartmentalised into distinguishable early- and late-replicating chromosome regions (Heckmann et al. 2013).

The nuclei of *P. serratum* revealed two major types of labelling patterns (Fig. 5). The majority of nuclei (85% of 500 nuclei) showed an almost uniform labelling (Fig. 5a), and 15% of nuclei showed a cluster-like distribution of EdU signals (Fig. 5b). Uniformly labelled nuclei are likely at early S phase, while nuclei with clustered signals undergoing late replication. Comparable replication patterns were found in other species with monocentric chromosomes like *Arabidopsis thaliana* (Dvorackova et al. 2018) and *Zea mays* (Bass et al. 2015).

**Discussion**

**Centromere evolution in Cyperids**

The analysis of the centromeres by immunostaining using CENH3, alpha-tubulin, histone H3S10ph, H3S28ph and H2AT120ph antibodies demonstrated a monocentric centromere type for the phylogenetically basal *P. serratum*. Therefore, these data suggest that monocentric chromosomes may be an ancestral condition for the Juncaceae-Cyperaceae-Thurniaceae Cyperid clade. As monocentricity was also reported in species within the *Juncus* genus (Guerra et al. 2019), holocentric chromosomes in the Cyperid clade have evolved at least twice independently: once in Juncaceae and once in Cyperaceae, after the divergence of the three families.
The phylogenetic close proximity of *P. serratum* CENH3 with species possessing holocentromere suggests that the sequence divergence of CENH3 does not correlate with its corresponding centromere type. A similar centromere-type independent CENH3 evolution was found for mono- and holocentric *Cuscuta* species (Oliveira et al. 2020). Hence, our data suggested that sequence modifications of CENH3 are not necessarily involved in the change of the centromere type in angiosperms.

The abundance of repetitive DNA in *P. serratum*

The small *P. serratum* genome contains a relatively low percentage of transposable elements, ~9% retrotransposons and ~3% DNA transposons, with 13 and 6 different lineages, respectively. Most plant genomes contain only a few satDNA families, mainly repeats associated with pericentromeric or subtelomeric regions (reviewed in Garrido-Ramos 2015). Here, we identified 19 different satDNA families (~3% of the genome). Unlike in other small genome–sized, monocentric species, like *A. thaliana* (Maluszynska and Heslop-Harrison 1991), sugar beet (Kubis et al. 1998) and rice (Cheng et al. 2002), the centromeric satDNA is not the most abundant satDNA. PsSat306, the most abundant satDNA family, displays colocalisation with the 45S rDNA. PsSat306 likely originated from the intergenic repeat spacer region as described for other satellite repeats (reviewed in Garrido-Ramos 2015). In addition, the 5-ethynyl-2′-deoxyuridine (EdU) detection in interphase showed comparable late replication patterns as those observed in species with monocentric chromosomes (Bass et al. 2015; Dvorackova et al. 2018).

A clustered distribution at one or only a few chromosome pairs was found for three satDNA families, similar to other satellite repeats in several species within the clade, as the holocentric *Luzula* and *Rhynchospora* genera, and outside the clade in typical monocentric species, as *Chenopodium quinoa* (Heckmann et al. 2013; Heitkam et al. 2020; Ribeiro et al. 2017). Two of these tandem repeats (PsSat156a and PsSat156b) share the same distribution at centromeric regions but with different signal intensities. Most likely, they evolved from the same ancestral centromeric repeat unit and underwent amplification or reduction at different chromosome pairs.

### How to identify holocentricity?

Results in *P. serratum* demonstrated that the characterisation of the centromere type, especially in species with small-sized chromosomes could be challenging. Which is the best method to identify holocentricity? As listed in Table 3, a range of different methods has been used to determine the centromere type in the past. However, no universal method amenable for all species exists, due to either the limitation in optical resolution, availability of specific antibodies, or required equipment.

Cytological methods, by observing the absence of a primary constriction in mitotic chromosomes, paralleled segregation of anaphase sister chromatids and the faithful transmission of induced chromosomal fragments, are the prime methods of choice to identify holocentrics. In large chromosome species like *L. elegans* and *R. pubera*, holocentromeres form at somatic pro- and metaphase a distinct longitudinal groove along each sister chromatid which is visible by standard (Heckmann et al. 2011; Nagaki et al. 2005), structured illumination and scanning electron microscopy (Marques et al. 2015; Wanner et al. 2015).

### Table 2  Repetitive families of *P. serratum*

| Repeat families | Genome proportion (in %) | Total (in %) |
|-----------------|--------------------------|--------------|
| LTR Retrotransposons | Ty1-Copia | |
| Ale | 1.46 | |
| SIRE | 1.91 | |
| Tork | 1.68 | |
| Alesia | 0.19 | |
| Ivana | 0.09 | |
| TAR | 0.02 | |
| Ikeros | 0.01 | |
| Ty3-Gypsy | Tat | 2.85 |
| Chromovirus Tekay | 0.42 |
| Chromovirus CRM | 0.20 |
| Chromovirus Galadriel | 0.14 |
| Chromovirus Reina | 0.02 |
| LINE | 0.28 | |
| DNA Transposon | TIR | 0.35 |
| CACTA | 0.24 |
| hAT | 0.35 |
| MuDR Mutator | 0.95 |
| PIF Harbinger | 0.87 |
| Helitron | 0.09 |
| Satellite | 3.09 | |
| rDNA | 45S | 2.68 |
| 5S | 0.15 | |
| Unclassified | 8.89 | |
| Total | 26.93 | |
However, the first two methods are not applicable for small chromosome species. The analysis of irradiation-induced chromosome fragments is still one of the best methods to verify holocentricity (Hughes-Schrader and Ris 1941;
reviewed in Mola and Papesci (2006)). While acentric fragments of monocentric chromosomes form micronuclei, induced holocentric fragments are stably transmitted into the next cell generation and do not form micronuclei. But the application of this method requires specialised equipment for the generation of ionising radiation.

The analysis of meiotic chromosome dynamics has been used to determine holocentricity in species with moderate to large chromosomes (reviewed in Cuacos et al. 2015; Marques and Pedrosa-Harand 2016). Three principle options exist to deal with holocentricity during meiosis: (i) ‘chromosome remodelling’, (ii) ‘functional monocentricity’ and (iii) ‘inverted meiosis’. In the case of inverted meiosis, in contrast to monopolar sister centromere orientation, the unfused holokinetic sister centromeres behave as two distinct functional units during meiosis I, resulting in sister chromatid separation. Homologous non-sister chromatids remain terminally linked by a hardly visible chromatin fibre. Then, they separate at anaphase II. Thus, an inverted sequence of meiotic sister chromatid segregation occurs.

An almost terminal position of 45S rDNA, adjacent to telomeres, has been linked to holocentricity. This observation was made in 42 species of seven genera with holokinetic chromosomes (Roa and Guerra 2012). A possible explanation is that a secondary constriction in the interstitial region would interrupt the kinetochore plate along the holokinetic chromosome establishing a condition similar to dicentric chromosomes, leading to errors in chromosome segregation (Heckmann et al. 2011). But in holocentric Lepidoptera species also interstitial 45S rDNA sites were detected (Nguyen et al. 2010). Thus, since the terminal 45S rDNA location is not universal in holocentrics, it is not a universal evidence for holocentricity. Also, a terminal position of 45S rDNA was found in monocentric species (Schubert and Wobus 1985).

Visualisation of kinetochore proteins, such as CENH3 or CENP-C, by immunodetection shows the centromere type directly (Marques et al. 2016; Nagaki et al. 2005). This strategy is less restricted by chromosome size. However, it is often limited by the availability of species-specific kinetochore antibodies, which are both time- and cost-consuming in production. However, the absence of CENH3 in some species (Drinnenberg et al. 2014) and the microtubule attachment at CENH3-free chromosome regions in some species (Oliveira et al. 2020) make the application of anti-CENH3 as a universal marker for centromeres questionable. Nevertheless, the analysis of kinetochore proteins could be complemented by combining the investigation of the spindle fibre attachment using alpha-tubulin-specific antibodies if the size of chromosomes allows the identification of the spindle fibre attachment site. In addition, the application of antibodies specific for the cell cycle–dependent pericentromeric phosphorylation of histone H3 (H3S10ph, H3S28ph) and H2A (H2AT120ph) resulted in the identification of holocentromere-specific immunostaining patterns (Demidov et al. 2014; Gernand et al. 2003). In monocentric plants, immunostaining with antibodies against H3S10ph and H3S28ph results in a specific labelling of the pericentromere in mitotic chromosomes. In contrast, in holocentric plants, immunolabelling with the same antibodies results in uniform staining of condensed chromosomes, due to the chromosome-wide distribution of the pericentromere (Gernand et al. 2003). The application of these antibodies in a wide range of species is possible due to the evolutionarily conserved amino acid sequence of histone H3. However, in some monocentric species, the application of anti-H2AT120ph resulted in additional non-pericentromeric signals (Baez et al. 2019; Sousa et al. 2016).

Transmission electron microscopy studies also showed differences between holo- and monocentric chromosomes in

![Fig. 5 Two types of DNA replication patterns in *P. serratum* shown by EdU labelling (red) and interphase nuclei counterstained with DAPI (blue). (a) Mainly uniform labelling and (b) clustered distribution of EdU signals.](image-url)
2. Microscopy-independent methods: flow cytometry and assessment of genomic content

Pending on the characteristics for each particular species, different techniques should be combined to determine holocentricity, dehydration and straightforward method exists, if possible, different techniques should be taken with care. Hence, as no universal method for identifying the centromere type (Smarda et al. 2014; Zedek et al. 2016). But, as our analysis of *P. serratum* showed, indirect methods should be taken with care. Hence, as no universal and straightforward method exists, if possible, different techniques should be combined to determine holocentricity, depending on the characteristics for each particular species.

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**Authors’ contributions** MB and YTK performed repeat analyses, FISH and immunostaining experiments and wrote the manuscript; YD performed high-resolution microscopy; APH and VS analysed data; JF measured the genome size; and AH designed the study, analysed data and wrote the manuscript. TB performed FISH and immunostaining experiments and wrote the manuscript; YD performed high-resolution microscopy. Funding Open Access funding enabled and organized by Projekt DEAL. This work has been supported by the Deutsche Forschungsgemeinschaft (HO1779/32-1), DAAD/CAPES (57517412; 88881.14406/2017-01) and Taiwan Ministry of Science and Technology (MOST, 106-2917-I-006-012 and 109-2917-I-564-022).

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| Methods | Examples | Exceptions |
|---------|----------|------------|
| 1.1 Chromosome morphology and dynamics | *Genus Euschistus* (Hughes-Schrader and Schrader 1961) | • To deal with holocentric during meiosis, chromosome remodelling and functional monocentricity exist in addition; e.g. temporary kinetochore activity in the end of chromosomes in kissing bug (Perez et al. 1997) |
| 1. Stable transmission of irradiation-induced chromosome fragments | *Genus Chionographis* (Tanaka and Tanaka 1977) |
| 1. Lack of a primary constriction in mitotic metaphase chromosomes and paralleled separation of mitotic anaphase chromatids. | *Bombyx* species (Murakami and Imai 1974) |
| 1. Subgenus Cuscuta (Garcia 2001) | *Centromere groove in L. elegans* (Nagaki et al. 2005; Wanner et al. 2015) |
| 2. In large holocentric chromosomes, sister chromatids form a distinct longitudinal centromere groove. | *L. echinata* (Braselton 1981), *L. nivea* (Bokhari and Godward 1980) (reviewed in Mola and Papeschi, 2006), Cabral et al. (2014) |
| 3. Electron microscopy to determine distribution of the kinetochore plate | • *Rhynchospora* species (Cabral et al. 2014) |
| 4. Electron microscopy to determine distribution of the kinetochore plate | • *L. elegans* (Heckmann et al. 2014; Kusana 1962; Nordenskiold 1962) |
| 5. Existence of inverted meiosis | • *R. pubera* (Cabral et al. 2014) |

**Table 3** Methods to identify holocentricity

**Authors’ contributions** MB and YTK performed repeat analyses, FISH and immunostaining experiments and wrote the manuscript; YD performed high-resolution microscopy; APH and VS analysed data; JF measured the genome size; and AH designed the study, analysed data and wrote the manuscript.

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