Comparative analyses of gibbon centromeres reveal dynamic genus specific shifts in repeat composition.

AUTHORS
Gabrielle A. Hartley†1, Mariam Okhovat†2, Rachel J. O’Neill*1,3,4 and Lucia Carbone*2,5,6,7

1 Department of Molecular and Cell Biology, University of Connecticut, Storrs, CT 06269
2 Department of Medicine, Knight Cardiovascular Institute, Oregon Health and Science University, Portland, OR 97239
3 Institute for Systems Genomics, University of Connecticut, Storrs, CT 06269
4 Department of Genomics and Genome Sciences, UConn Health, Farmington, CT 06030
5 Division of Genetics, Oregon National Primate Research Center, Beaverton, OR 97006
6 Department of Molecular and Medical Genetics, Oregon Health and Science University, Portland, OR 97239
7 Department of Medical Informatics and Clinical Epidemiology, Oregon Health and Science University, Portland, OR 97239

† Authors contributed equally.

*Corresponding Authors: E-mails: rachel.oneill@uconn.edu; carbone@ohsu.edu
Abstract

Centromeres are functionally conserved chromosomal loci essential for proper chromosome segregation during cell division, yet they show high sequence diversity across species. Despite their variation, a near universal feature of centromeres is the presence of repetitive sequences, such as DNA satellites and transposable elements (TEs). Because of their rapidly evolving karyotypes, gibbons represent a compelling model to investigate divergence of functional centromere sequences across short evolutionary timescales. In this study, we use ChIP-seq, RNA-seq and fluorescence in situ hybridization to comprehensively investigate the centromeric repeat content of the four extant gibbon genera (Hoolock, Hylobates, Nomascus and Siamang). In all gibbon genera, we find that CENP-A nucleosomes and the DNA-proteins that interface with the inner kinetochore preferentially bind retroelements of broad classes rather than satellite DNA. A previously identified, gibbon-specific composite retrotransposon, LAVA, known to be expanded within the centromere regions of one gibbon genus (Hoolock), displays centromere- and species-specific sequence differences, potentially as a result of its co-option to a centromeric function. When dissecting centromere satellite composition, we discovered the presence of the retroelement-derived macrosatellite SST1 in multiple centromeres of Hoolock, whereas alpha-satellites represent the predominate satellite in the other genera, further suggesting an independent evolutionary trajectory for Hoolock centromeres. Finally, using de novo assembly of centromere sequences, we determined that transcripts originating from gibbon centromeres recapitulate the species-specific TE composition. Combined, our data reveal dynamic shifts in the repeat content which define gibbon centromeres and coincide with the extensive karyotypic diversity within this lineage.

Keywords

Chromosome evolution, centromeres, gibbon, primate genomics, transposable elements, satellite DNA
Introduction

Centromeres are essential genomic loci that support the assembly of the kinetochore, thereby facilitating spindle attachment and faithful segregation of chromosomes to daughter cells during mitosis and meiosis. Comparative studies across deep branches of eukaryotic lineages have shown a stark contrast between the functional conservation of the components at the interface of the inner kinetochore and the rapid evolution of centromeric DNA sequences (Henikoff, et al. 2001). The centromeric DNA of humans and other great apes largely consists of ~171 bp long AT-rich alpha-satellite monomers that are tandemly repeated in blocks and further arranged into larger repetitive units, known as higher order repeat arrays (HORs) (Willard 1985; Waye and Willard 1987; Alexandrov, et al. 1993; Rudd, et al. 2003). While the presence of alpha-satellites is a predominantly conserved characteristic of primate centromeres, the identification of large-scale, variable centromere haplotypes in humans suggests the genomic landscape of such satellites is mutable (Langley, et al. 2019). In addition to centromeric satellites, recent work in phylogenetically divergent species, such as marsupials (Longo, et al. 2009; Ferreri, et al. 2011; Renfree, et al. 2011; Johnson, et al. 2018), *Drosophila* (Chang, et al. 2019), plants (Zhong, et al. 2002), and fungi (Yadav, et al. 2018) show that centromeric retroelements may be a defining element of functional centromeric chromatin. Retroelements, mobile elements that propagate via an RNA intermediate, have been proposed as major players in centromere evolution because they can provide the material for generation of new satellite families. Such phenomena have been observed in plants (Kapitonov and Jurka 1999; Cheng and Murata 2003), *Drosophila* (Heikkinen, et al. 1995), and cetaceans (Kapitonov, et al. 1998). Furthermore, centromeric retroelements have been implicated in facilitating chromosome evolution through the introduction of large-scale genomic rearrangements as specific classes of centromeric retroelement have been found to be enriched at evolutionary breakpoints (Longo, et al. 2009).

Gibbons, the endangered small apes in the Hominoidea superfamily, a taxonomic group co-occupied by both small and great apes (Cunningham and Mootnick 2009), present a unique opportunity to study the evolution of functional centromere sequences in the context of rapid karyotypic change. Gibbons diverged from the common Hominoidea ancestor only ~17 million years ago (mya), and a more recent (~5 mya) radiation event gave rise to the four extant genera. Despite their recent divergence and species radiation, gibbons have accumulated an
unusually high number of chromosomal rearrangements compared to other lineages, suggesting a rapid rate of karyotype evolution (Carbone, Vessere, et al. 2006; Roberto, et al. 2007; Carbone, et al. 2009; Girirajan, et al. 2009; Carbone, et al. 2012; Carbone, et al. 2014). Moreover, drastically different karyotypes are found in each of the four extant gibbon genera (Hoolock [2n=38], Nomascus [2n=52], Hylobates [2n=44], and Symphalangus [2n=50]), derived by translocations, fusions, fissions and inversions, often concomitant with centromere inactivation and formation of evolutionary new centromeres (Capozzi, et al. 2012). These rapid karyotypic changes, which have been extensively mapped in gibbons, offer a unique opportunity to study centromeres that are variable within the same species and across different genera.

In our previous investigations of the evolutionary history of gibbon chromosomes, we discovered a non-autonomous, gibbon-specific composite retrotransposon, named LAVA (after its three component elements: LINE-Alu-VNTR-Alu-like), that depends on the L1 machinery to retrotranspose in the host genome. Based on structural characterizations of LAVA subfamilies (Lupan, et al. 2015), the average length of LAVA elements ranges from 1,661 to 2,557. The VNTR (Variable Number Tandem Repeat) region of LAVA is the most variable portion of the LAVA, with an average length that can range from 546 to 1,377bp across LAVA sub families and is generally inversely correlated to the age of the element (Lupan, et al. 2015). While the LAVA element can be found in the genomes of all gibbon species examined to date, it has expanded in almost all centromeres of only two gibbon species belonging to the Hoolock genus (Carbone, et al. 2012; Hara, et al. 2012). Thus, expansion of the LAVA retroelement in gibbon centromeres likely occurred recently and after divergence of the four gibbon genera. In fact, the LAVA centromeric expansion in Hoolock has occurred recently enough not to have yet equally impacted all chromosomes, as four chromosomes lack LAVA expansions (Carbone, et al. 2012).

To determine if recently expanded repetitive sequences are linked to centromeric function, we completed an in-depth cross-genera characterization of the sequences binding three DNA-interacting centromeric proteins: CENP-A, the histone that demarcates centromeric chromatin (Yoda, et al. 2000; Van Hooser, et al. 2001); as well as CENP-B and CENP-C, two inner kinetochore proteins that lie at the interface between centromeric chromatin and kinetochore-spindle complex (Okada, et al. 2007; Klare, et al. 2015). Combining next
generation sequencing, *de novo* sequence assembly, repeat annotation methods and fluorescence *in situ* hybridization (FISH), the putatively functional centromeric repeat content was characterized and compared across the four gibbon genera. Instead of the homogeneous satellite arrays that typify primate centromeres, all four gibbon genera showed diverse transposable element content in CENP-A chromatin and the functional part of the centromere that forms the inner kinetochore (CENP-B and CENP-C). Across the *Hoolock* karyotype, LAVA and SST1 (a retroelement-derived macrosatellite found in primates) define highly divergent functional centromere forms compared to the other genera. Finally, a subset of transcriptionally active retroelements, including LAVA, define functional elements within centromeres and recapitulate the largest variation among centromeres of gibbons.

**Results**

**Centromeres of all four gibbon genera are enriched in SINEs and depleted in LINEs**

We investigated the repeat composition of gibbon centromeres using chromatin immunoprecipitation sequencing (ChIP-seq) targeting centromere proteins (CENPs) known to demarcate active centromeric chromatin, including the centromeric histone variant CENP-A, and two additional proteins, CENP-B and CENP-C (Yoda, et al. 2000; Van Hooser, et al. 2001; Okada, et al. 2007; Klare, et al. 2015). CENP-A is the centromere-specific H3 variant (Earnshaw and Rothfield 1985) that replaces conventional H3 in a subset of nucleosomes (Blower, et al. 2002) and is required for kinetochore assembly. Via its histone chaperone, HJURP, CENP-A is assembled into centromeric chromatin during late-telophase, early G1 in humans every cell cycle (Jansen, et al. 2007; Foltz, et al. 2009), demarcating the chromosomal site for kinetochore formation and spindle attachment. CENP-B is known to bind CENP-B box containing sequences within a subset of functional centromeric sequences (Amor, et al. 2004a; Henikoff, et al. 2015; Aldrup-MacDonald, et al. 2016) and is found at centromeres throughout the cell cycle (Earnshaw, et al. 1989). However, it is unknown when in the cell cycle loading of CENP-B actively occurs (Gamba and Fachinetti 2020). CENP-C is proposed to serve as a centromere marker that may target CENP-A as part of the constitutive centromere-associated network (CCAN), however it is distinguished from CENP-A in the timing and mode of deposition. While CENP-A deposition is limited to late telophase/early G1 (Jansen et al.,
2007), CENP-C targets centromeres throughout the cell cycle (Earnshaw, et al. 1989; Du, et al. 2010). Thus, each of these three CENPs can independently serve as a marker for centromere function, yet their predicted binding profiles are unknown.

ChIP-seq was carried out on gibbon lymphoblastoid cell lines (LCLs) established previously (Carbone, et al. 2014; Lazar, et al. 2018; Okhovat, et al. 2020) and for this study, from four different gibbons species: *Nomascus leucogenys* (NLE), *Hoolock leuconedys* (HLE), *Hylobates moloch* (HMO) and *Symphalangus syndactylus* (SSY), each representing one of the four extant gibbon genera (Supplementary Table S1). Alignments of CENP-A, CENP-B and CENP-C protein sequences between human and gibbons indicates high likelihood of cross-reactivity of human-specific antibodies with gibbon (95% identity across CENP-A with only 1 amino acid difference in the peptide used to generate the antibody, 98% across CENP-B, and 100% across CENP-C proteins). To independently validate successful immunoprecipitation of DNA bound to each CENP in each species, fluorescence in situ hybridization (FISH) was used to visualize each ChIP-seq library on mitotic chromosome spreads from corresponding LCLs. Each ChIP-seq library probe localized to centromeres in the four species examined, indicative of successful CENP immunoprecipitation for all three antibodies in each species (Figure 1A, Supplementary Figure 1). In addition to centromeric enrichment, pericentromeric and telomeric probe localization of variable intensity was observed in some species and chromosomes, similar to the hybridization patterns previously observed using a centromeric alpha-satellite probe (Cellmare, et al. 2009). Cross-hybridization of ChIP-seq DNA to non-centromeric loci, which was most easily visible in *Symphalangus* (SSY) (Figure 1A, Supplementary Figure 1), likely reflects occasional sequence similarities between centromeric and non-centromeric repeats that cannot be distinguished by FISH.

To characterize the repeat content of CENP-A, CENP-B, and CENP-C bound DNA, the repeat composition of 2.5 million randomly subsampled read pairs from each ChIP-seq library were annotated using RepeatMasker (Tempel 2012). The repeat content of corresponding control libraries constructed from chromatin that did not undergo immunoprecipitation (i.e. input libraries) was similarly annotated and used to estimate genome-wide repeat composition per species. The overall abundance and repeat composition of CENP-A, CENP-B and CENP-C libraries were broadly similar across gibbon genera, but largely different from their corresponding input libraries (Figure 1B, Supplementary Table S2). Consistent with the known
genetic structure of complex, regional centromeres [reviewed in (Hartley and O'Neill 2019)], gibbon ChIP-seq libraries had significantly higher repeat content compared to corresponding inputs. On average, 43% of trimmed read sequences were annotated as repeats across all ChIP libraries vs. 36% in input reads, indicating gibbon centromeres are enriched for, but not solely characterized by, repeats (paired t-test p<0.0001; Figure 1B, Supplementary Table S2).

Most repeat classes, including SINEs, LINEs, LTRs, and DNA elements, displayed relatively similar abundance in each ChIP-seq dataset across the four gibbon genera. Representing 52-64% of annotated repeats, SINEs were the most prevalent repeat class in CENP-A, CENP-B and CENP-C bound DNA pools. The percentage of SINE elements was approximately double those found in the corresponding input samples, indicating enrichment of SINEs in the functional compartments of gibbon centromeres. LINEs, the most abundant repeat class genome-wide, were depleted in CENP-bound DNA and constituted only 11-17% of the annotated repeats in CENP ChIP-seq libraries across all four genera (Figure 1B, Supplementary Table S2).

**The gibbon-specific LAVA retrotransposon is enriched in *Hoolock* centromeres**

The prevalence of the repeat class defined as “retrotransposon” by RepeatMasker (representing non-LINE/LTR/SINE retrotransposon elements) varied among gibbon genera, with HLE distinctly showing higher prevalence (Figure 1B, Supplementary Table S2). Indeed, the HLE input had ~10 times the relative abundance of retrotransposon repeats compared to the inputs of SSY, HMO, and NLE (1.9% of HLE input repeats, vs. 0.19± 0.04% [mean± stdev] of input repeats in SSY, HMO, NLE). HLE CENP-bound DNA showed an almost three-fold enrichment in the proportion of repeats annotated as retrotransposons (5.41 ± 1.33%) relative to input and ~11-fold enrichment relative to CENP-bound DNA from SSY, HMO and NLE (0.51 ± 0.17% of repeats; mean ± stdev).

Earlier cross-species FISH experiments showed centromeric expansion of the gibbon-specific retrotransposon LAVA exclusively in HLE (Carbone, et al. 2012), hence we wanted to determine if our CENP-bound DNA annotated as retrotransposons might include this element and if this element is enriched in gibbon centromeres. Using *in silico* simulations, we first verified that RepeatMasker could identify LAVA repeats from short-read sequences reliably and with low false negative and false positive rates (Supplementary Table S3). These analyses
revealed that ~50% of LAVA repeats may be mis-annotated as “SVA_A”, likely due to large regions of homology between LAVA and SVA, particularly at the VNTR region (Variable Number Tandem Repeat; Figure 2A, Supplementary Table S3). Given that gibbon genomes contain a negligible number of SVA element insertions (Wang, et al. 2005; Ianc, et al. 2014), gibbon repeats classified as “SVA_A” by RepeatMasker are likely LAVA elements. Thus, reads annotated as either LAVA or SVA_A were collectively classified as LAVA in this study.

Inter-genera comparison of retrotransposon annotations revealed higher abundance and enrichment of LAVA elements (relative to input) in CENP-A and CENP-B, but not CENP-C, ChIP-seq libraries from HLE compared to the three other genera (Figures 1B, 2B, 2C, Supplementary Figure 2A, Supplementary Table S2). To confirm these findings, CENP-A ChIP-seq and analyses were repeated as above on three additional HLE gibbons and revealed the same levels of LAVA enrichment, indicating that LAVA enrichment in CENP-A and CENP-B bound centromeric DNA is HLE-specific (Figure 2C). These observations, together with our previous findings using whole-genome shotgun sequences (Okhovat, et al. 2020), indicate that the LAVA element is overall more prevalent in the HLE genome compared to other genera, and that LAVA is distinctly enriched in HLE centromeric regions demarcated by CENP-A and CENP-B DNA binding.

**LAVA elements in the Hoolock genome are structurally distinct from other genera**

Motivated by the higher abundance of LAVA elements in HLE, we sought to determine if HLE LAVA elements are also structurally different from those found in NLE, HMO, and SSY. In lieu of centromeric LAVA sequences, we took advantage of LAVA repeat annotations in our short read sequences originating from centromeres (i.e. CENP ChIP-seq) versus reads from the rest of the genome (i.e. input library) across all four gibbon species. We used Repstat (Johnson, et al. 2018), which is an analysis tool that can compare short read data from a specific repeat to their consensus sequence and test for statistically significant deviations from this consensus, among different data sets. Compared to LAVA reads identified in the libraries of the three non-HLE genera, portions of reads annotated as LAVA in the CENP and input libraries from HLE were significantly longer and had more substitutions (i.e. higher percent divergence) relative to the consensus LAVA sequence (adjusted p<0.01) (Figure 3A, Supplementary Table S4). Pairwise comparisons among HMO, NLE, and SSY libraries revealed no difference in the
inferred length distribution and sequence of portions of reads annotated as LAVA repeats (adjusted p>0.05), with only a few exceptions (Figure 3A, Supplementary Table S4). Overall, these findings indicate that the sequence and structure of LAVA elements in the HLE genome and centromeres are distinct from those found in other gibbon species.

Repstat analyses of LAVA-annotated sequencing reads across HLE libraries revealed that portions of reads annotated as LAVA repeats in all three CENP protein bound DNA libraries were longer compared to those identified in the HLE input library (adjusted p<0.01) (Figure 3A, Supplementary Table S4). Moreover, compared to input, reads annotated as LAVA in the CENP-A and CENP-B, but not CENP-C, ChIP-seq datasets were significantly less diverged relative to the consensus LAVA sequence (adjusted p<0.05). Among the three HLE CENP-ChIP-seq libraries, LAVA annotated reads identified in the CENP-C library were shorter (adjusted p<0.01), and had significantly less sequence identity/more insertions relative to the LAVA consensus sequence compared to LAVA repeats annotated in either HLE CENP-B or CENP-A libraries (adjusted p<0.01) (Figure 3A, Supplementary Table S4). There were no significant differences in the length or sequence of reads annotated as LAVA between HLE CENP-A and CENP-B libraries (adjusted p>0.05) (Figure 3A, Supplementary Table S4).

Together, these results suggest that the structure and sequence composition of centromeric LAVA elements in HLE, particularly those putatively bound by CENP-A and CENP-B, differ significantly from LAVA sequences found elsewhere in the HLE genome. These patterns are likely due to differences in insertion preference of specific LAVA elements into centromeres, differences in the evolutionary trajectories of LAVA following centromeric insertion and CENP interaction, or both.

The structure and sequence of centromeric and non-centromeric LAVA repeats in NLE, SSY and HMO were compared to test if structurally distinct centromeric LAVA are present only in HLE, and as demonstrated in earlier FISH assays (Carbone, et al. 2012) and in our ChIP-seq analyses herein, are thus associated with centromeric LAVA expansion exclusively found in HLE. Compared to input libraries, no difference in the length or sequence divergence from consensus was detected in LAVA reads found in any CENP library from NLE and SSY gibbons (adjusted p>0.05). In contrast, within the HMO species, LAVA-annotated sequences in all three CENP-bound DNA pools were significantly shorter and more diverged from the LAVA consensus compared to sequences annotated as LAVA in the input library (adjusted p<0.05).
(Figure 3A, Supplementary Table S4). Altogether, while we observed distinct structures in centromeric LAVA across genera, the strongest differences were observed in HLE.

**Gibbon centromere proteins putatively bind LAVA elements at CENP-B box enriched VNTR sequences**

Since LAVA is a composite retrotransposon (Figure 2A), we examined whether centromere proteins preferentially bind specific repeat components within the LAVA sequence. Meta-summit analysis, previously optimized for LAVA (Fernandes, et al. 2020; Okhovat, et al. 2020), were completed for CENP ChIP-seq datasets across all four gibbon genera, since even non-HLE libraries showed >two-fold enrichment of LAVA elements in CENP libraries relative to input (Figure 3B, Supplementary Figure 2B). Briefly, CENP-A, CENP-B, and CENP-C ChIP-seq reads were aligned to the gibbon reference genome (Nleu3.0) (Carbone, et al. 2014) and both unique and multi-mapping reads were used to identify significant narrow peak summits that overlapped LAVA elements annotated in the reference genome. For each sample, the overlapping summits were mapped to the consensus LAVA sequence to generate a pileup of summit positions along the consensus element. Summits across this summit pileup, i.e. “meta-summits”, represent putative CENP binding sites within the composite LAVA element. In total, the number of significant CENP summits per gibbon CENP library ranged from 6,380 to 158,933, and 1-3% of all summits overlapped LAVA elements across samples (Supplementary Table S5). The high cross-species and cross-library variation in the total number of CENP peaks overlapping LAVA was reflected in the summit pileup height differences. Across all genera, the meta-summits (i.e. putative binding sites) of CENP-A, CENP-B and CENP-C were located within the VNTR region of LAVA (Figure 3B, Supplementary Figure 2A), indicating this repeat component may be putatively bound by CENPs.

We then mined ChIP-seq data for the presence of detectable CENP-B box motifs in gibbon centromeres. Human CENP-A ChIP-seq data (Hasson, et al. 2013) was used as a positive control to validate reliability of our pipeline. Briefly, broad CENP-A ChIP-seq peaks were identified within the human genome and sequences corresponding to these broad peaks showed overall enrichment of the CENP-B box motif (JASPAR database, MA0637.1) (Masumoto, et al. 1989; Jolma, et al. 2013) relative to shuffled sequences. Next, CENP peaks were sorted based on their fold-enrichment relative to input (i.e. peak height) and we
compared enrichment of CENP-B box motifs between high- and low-height peaks. Since ChIP-seq height may be a proxy for binding strength/frequency, the expectation was that higher peaks would be associated with higher prevalence of the CENP-B box motifs. As expected, a significant association was found between CENP-B box motifs and CENP-A peak height in human CENP-A bound DNA sequences (adjusted p <0.05; Supplementary Table S6).

Using a similar approach to investigate the presence of the CENP-B box motif in gibbon CENP-bound DNA, we first identified 139,315-383,872 broad peaks across all gibbon CENP ChIP-seq libraries. Overall, sequences of these broad peaks were not significantly enriched in the CENP-B box motif in any of the CENP libraries when compared to shuffled sequences (adjusted p >0.05). However, when peaks were sorted by height, we found higher peaks had significantly more CENP-B box motifs compared to weaker peaks in at least one CENP pull-down library in each genus (adjusted p <0.05; Supplementary Table S6). In order to test our hypothesis that LAVA was co-opted in gibbons to become a functional centromeric element, we aimed to also investigate CENP-B box enrichment in this element. Due to unavailability of centromere sequences in the reference genome, we used LAVA elements present elsewhere in the genome and found that among putative CENP binding sites overlapping LAVA elements (i.e. CENP ChIP-seq summits overlapping LAVA in meta-summit analysis above), 7-16% contained predicted CENP-B box motifs (Supplementary Table S6), and the majority of these motifs (~88% on average) were located in the VNTR region of LAVA (Supplementary Figure 2C). In addition, the DNA sequences of CENP peak summits within LAVA repeats were collectively strongly enriched in the CENP-B box motif (Benferroni adjusted p<0.01; Supplementary Table S6), linking CENP-DNA binding to a specific DNA recognition motif within the LAVA retroelement. While further studies and contiguous centromere sequence assemblies are required to fully characterize the presence, enrichment and functionality of the CENP-B box motif in gibbon centromeres, our analyses suggest that some gibbon centromeres may contain this motif, albeit likely at a much lower prevalence compared to human centromeres.

Centromeric alpha-satellite and SST1 repeats vary across gibbon genera, particularly in *Hoolock*
We observed high variation in satellite class abundance among ChIP-seq libraries across genera, with HLE libraries (which have the highest LAVA abundance) showing the lowest satellite prevalence (Figure 1B). In general, satellite repeats were less abundant among HLE and HMO CENP ChIP-seq and input repeats (0.6-3% of repeats), compared to SSY and NLE [14.8 ± 9.4% and 8.4 ± 4.6% of repeats, respectively; (mean ± stdev); Supplementary Table S2]. Except for HLE, which had equally low satellite DNA abundance in both input and CENP ChIP-seq libraries, all other gibbon genera showed around three-fold lower satellite repeat content in their CENP ChIP repeat composition compared to input, suggesting that despite being present in gibbon centromeres, satellites are not highly enriched in these loci compared to the rest of the genome. These data support previous FISH analyses showing a lack of satellite enrichment in HLE centromeres (Carbone, et al. 2012), but are in contrast to observations that centromeres of great apes are highly enriched in satellites, particularly alpha-satellite repeats (Alkan, et al. 2007).

Within the satellite class of repeats, the alpha-satellite family was responsible for most of the variation observed across gibbon genera and libraries (Figure 4A) and was the most prevalent family of satellites in the genomes and centromeres of SSY and NLE gibbons, representing 94-99% of all satellite repeats per library of these two genera. However, this repeat family was less abundant in the centromeres of HMO (constituting <58% of satellites in CENP ChIP-seq datasets), and the least abundant in the HLE both genome-wide and in the centromeres (representing <6% of satellites in CENP libraries; Figure 4A, Supplementary Table S2).

Based on sequence library annotations, the lack of alpha-satellites in HLE libraries is partially offset by the presence of a retroelement-derived macro-satellite family known as SST1 (also called MER22; Figure 4A) (Fatyol, et al. 2000). The SST1 satellite family accounted for 56-72% of satellite repeats across HLE CENP libraries, while making up less than 30% of satellites in HMO CENP libraries and less than 3% of CENP satellites in SSY and NLE (Figure 4A). The distinctly higher absolute and relative abundance of SST1 repeats in HLE CENP ChIP-seq datasets, particularly in CENP-A and CENP-B libraries, was validated in all three biological HLE CENP-A replicates (Figure 4B, Supplementary Figure 3A). While non-HLE gibbon genera have a smaller number of SST1 repeats genome-wide than HLE, each genera showed relative enrichment of SST1 repeats in CENP ChIP-seq datasets compared to their
input (Figure 4B, Supplementary Table S2). Thus, while SST1 elements are likely present in some centromeres across all gibbon genera, SST1 repeats are more prevalent in HLE centromeres.

FISH to metaphase preparations using genus-specific SST1 probes (Supplementary Table S7) confirmed the centromeric localization of SST1 on three chromosome pairs in HLE, but on only one chromosome pair in each of the other three species (Figure 5A, Supplementary Figure 3B). The three HLE chromosomes that contained SST1-rich centromeres were identified by inverse-DAPI banding and chromosome painting with HLE specific painting probes (Nie, et al. 2001) as HLE4, HLE15 and HLE18 (Figure 5B). SST1, found on the q arms of human chromosomes 19 and 4 (Tremblay, et al. 2010), is consistent with the predicted synteny between gibbons and human based on previous FISH (Roberto, et al. 2007; Capozzi, et al. 2012) (Figure 6A). We used FISH with an HSA19 chromosome paint to verify SST1 hybridization patterns and found overlap between the gibbon SST1 centromeric signal and HSA19 on gibbon chromosomes NLE10 (Figure 6B), and HLE15 and HLE18 (Figure 6C). The gibbon SST1 centromeric signal and the HSA19 paint also revealed an unknown synteny with HLE4 (Figures 6A, 6C). The SST1 signal is absent in HLE6, despite HLE6 having a non-centromeric syntenic block within the proximal long arm of HSA19 (Figures 6A, 6C). Thus, SST1 centromeric signal seems to be linked to centromere-specific rearrangements distinctive to each gibbon lineage. Of note, the three chromosomes containing SST1 repeats in HLE were previously shown to have weak or no centromeric LAVA expansion (Carbone, et al. 2012) confirming that an inverse relationship may exist between centromeric LAVA and satellite abundance.

Similar to our analysis of LAVA repeats, we used Repstat (Johnson, et al. 2018) to compare sequences and structural characteristics of reads annotated as SST1 repeats within and across species. Overall, very few differences were found in SST1 structure across libraries within each species, indicating that SST1 satellites have similar structures in both centromeric and non-centromeric regions within each genus. Of the few significant structural differences found across libraries, most were found in CENP-C libraries in HLE and NLE (Figure 7; Supplementary Table S4), while no significant differences in SST1 sequences were identified in HMO and SSY (adjusted p>0.05) (Figure 7; Supplementary Table S4), albeit low abundance of SST1 in the SSY genome may have affected significance of comparisons. When the
structure of SST1 repeats was compared across gibbon genera, SST1-annotated regions in HLE were identified as the most distinct. For example, SST1 read annotations across all HLE ChIP-seq datasets were significantly longer than those found in the respective NLE, HMO, and SSY ChIP-seq datasets (adjusted p<0.01). Very few differences were identified in SST1 structure and sequence among the non-HLE species (Figure 7, Supplementary Table S4).

**Patterns of expression of centromeric LAVA and SST1 differ across gibbon genera**

Our analyses indicated that both LAVA and SST1 repeats have expanded, and may be functional in gibbon centromeres, particularly in HLE. To test if putatively functional retrotransposons in centromeres are transcriptionally active, we characterized the repeat composition of the transcriptome across gibbon genera. Total RNA sequencing (RNA-seq) data was generated from lymphoblastoid cell lines of the same four gibbons that were used for our ChIP-seq assays (Supplementary Table S1). Consistent with findings in input and CENP ChIP-seq datasets, a significantly higher number of reads annotated as SVA_A/LAVA (one-sided z test, p<0.00001), with 1.3-1.6 times more LAVAs annotated per million reads in HLE than any other library (Figure 8A, Supplementary Table S8) was observed. Similarly, more SST1 elements were identified in the HLE RNA-seq library compared to the other genera (one-sided z test, p<0.00001), with 6.0-18.9 times more SST1 repeats annotated per million reads in HLE than any other library (Figure 8A, Supplementary Table S8). Thus, LAVA and SST1 transcript abundance varies across gibbon genera in a pattern consistent with centromeric repeat composition observed in the CENP ChIP-seq and FISH results.

To further delineate centromere-specific transcripts in the absence of centromere-spanning gibbon genome assemblies, *ab initio* contig assemblies were generated from each CENP-A, CENP-B, and CENP-C ChIP-seq dataset using RepARK, a program designed to generate *de novo* repeat libraries from whole-genome next generation sequencing reads (Koch, et al. 2014). Assemblies generated from CENP libraries had an average of 2,926 (+1,256; stdev) contigs and an average contig length of 163bp (+471; stdev) (Supplementary Table S9). We then identified RNA-seq reads unable to map to Nleu3.0 (4-5% of reads in each species), reasoning that, since the reference genome lacks centromere sequences, these unmapped reads would be enriched for centromere-specific transcripts. Unmapped reads were then aligned to the CENP ChIP-seq derived contigs, resulting in transcripts that can be
ascribed to centromeres in each gibbon species (Supplementary Figure 4, Supplementary Table S9). Across RNA-seq libraries from all four species, 0.27 ± 0.17% of the total RNA-seq reads did not align to the reference genome but did align to at least one of the assembled CENP ChIP-seq contigs, representing putative centromere-specific transcripts (Supplementary Table S9). The most common repeat families identified among the putative centromere-specific transcripts across species included: simple repeats, low complexity repeats, Alus, SVAs, LINE elements, and ERV-Ks (Figure 8B, Supplementary Table S10, Supplementary Figure 5). Notably, Alu, SVA, and LINE elements are all transposable elements that constitute the composite LAVA element. Consistent with our CENP ChIP-seq results (Figures 1B, 4), SST1-annotated transcripts mapped to HLE CENP-A, CENP-B and CENP-C ChIP-seq contigs, as well as HMO CENP-A and CENP-C ChIP-seq contigs, suggesting that this satellite element is actively transcribed in centromeres. While our approach does not provide a comprehensive annotation of the transcriptional landscape of gibbon centromeres, these results clearly indicate that centromere repeats are transcribed in species-specific patterns that reflect the underlying sequence and repeat composition.

Discussion

The scarcity of centromere sequences in genome assemblies has long hindered comparative studies on the function and evolution of centromere DNA. While advanced long-read sequencing technologies are reducing centromeric sequence deficiencies in model organism reference genome assemblies (Jain, et al. 2018; Johnson, et al. 2018; Chang, et al. 2019; Miga, et al. 2020), alternative strategies are still needed for cross-species centromere studies. To compare centromere repeats across the four extant genera of gibbons, a group of endangered small apes that has experienced rapid chromosome and centromere evolution (Carbone, Vessere, et al. 2006; Roberto, et al. 2007; Carbone, et al. 2009; Girirajan, et al. 2009; Carbone, et al. 2012; Carbone, et al. 2014), we integrated centromere-specific ChIP-seq, traditional cytogenetic, and centromere-focused RNA-seq analyses. Overall, these combined analyses highlighted high variability in abundance, sequences, and transcription of satellite and retroelement DNA among the gibbon genera.

Targeting DNA at the inner kinetochore of the centromere using ChIP-seq for CENP-A, -B, and -C, we found that centromeres of the Hoolock genus, represented by the Eastern
Hoolock (HLE), is unique among gibbons with respect to both retroelements and satellites. In addition to a higher prevalence of the gibbon-specific LAVA retroelement within HLE centromeres, we found significant differences in the length and sequence of centromeric LAVA read annotations compared to those found in chromosome arms. In particular, within the HLE genome, LAVA read annotations originating from regions putatively bound by CENPs were significantly longer than non-centromeric LAVA reads (Figure 3A). As the VNTR (Variable Number Tandem Repeats) portion of LAVA (Figure 2A) is found to be variable across insertions (Lupan, et al. 2015), it is likely that its expansions within HLE centromeres may have resulted in longer centromeric LAVA elements. Of note, our meta-summit analysis indicated that CENPs preferentially bind LAVA elements at the VNTR and at sequences enriched in the predicted CENP-B box motif (Figure 3B).

To date, CENP-B box motifs have been found in satellite sequences, linking this motif to centromere function. However, the evolution of the evolution of putative CENP-B box motifs in LAVA does not necessarily reflect the selective order of events by which sequences become involved in centromere identity. For example, a recent study in New World Monkeys indicates that the acquisition of CENP-B box binding capacity in centromeric sequences occurred independently in different lineages after the gain of both the satellite elements themselves and centromere function (Thongchum, et al. 2019). Thus, CENP-B is proposed to be a stabilizing factor for previously established centromeres (Gamba and Fachinetti 2020), increasing stability under stress in mitotic cells (Dumont, et al. 2020) or segregation efficiencies during asymmetrical segregation in meiosis (Chmatal, et al. 2017; Iwata-Otsubo, et al. 2017). In this context, it is likely that the acquisition of retroelements to support centromere function precedes the emergence of functional CENP-B boxes within the retroelement sequences themselves. The prediction of CENP-B boxes within LAVA elements and the capacity to bind CENP-B, as indicated based on our ChIP-seq data, support the observation that LAVA elements are centromeric and have acquired centromere function (as also evident by CENP-A and CENP-C binding). While our predictions do not indicate any mechanism by which LAVA elements initially acquired centromere function, the evolution of CENP-B box motifs in some centromeric LAVA sequences indicates selective pressure for stabilization of these elements within centromeres via CENP-B binding activity, analogous to the role alpha satellites play in centromere stabilization in humans.
It should be noted that in lieu of centromeric LAVA sequences, our analysis of the length and sequence of centromeric LAVA elements relied solely on short-read sequences. Moreover, our meta-summit analyses were carried out by aligning ChIP-seq reads to the current gibbon reference genome (Nleu3.0), and are thus based on the assumption that centromeric LAVA repeats bear high sequence similarity to LAVA repeats present in the reference genome. Deriving centromere-spanning genome assemblies in the future will allow for better characterization of centromeric LAVA sequences and identification centromere-specific variants, and thus may reveal additional CENP-B binding sites. Moreover, while the enrichment of the CENP-B box motif in LAVA sequences and enrichment of LAVA in CENP-B ChIP-seq data is intriguing, these findings are inconsistent with a previous study reporting that gibbon genomes lack CENP-B box motifs (Suntronpong, et al. 2016). Unlike centromeres characterized by highly homogenized satellite arrays, gibbon centromeres seem to be characterized by a diverse group of repeat types and this heterogeneity makes the detection of a statistically significant enrichment more challenging. Thus, further studies are required to confirm if the predicted CENP-B box motifs within centromeric LAVA elements are a site of CENP-B-DNA interaction.

In contrast to the alpha-satellite-rich centromere structures described for many other primates (Mahtani and Willard 1990; Warburton, et al. 1996; Alkan, et al. 2007; Sujiwattanarat, et al. 2015; Miga, et al. 2020), we observed that satellites, including alpha-satellites, comprise only a small percentage (0.7-12.0%) of CENP-bound repeats in all gibbon genera. Previous cytogenetics data gathered for HLE had shown that presence of centromeric LAVA expansions on a chromosome is mutually exclusive to centromeric alpha-satellites (Carbone, et al. 2012), suggesting a potentially conflictual relationship between the two repeats within gibbon centromeres. The exclusive relationship between satellites and retroelements may be linked to rapid chromosome change and centromere turnover as our observations in gibbon are similar to the finding of satellite-free, retroelement-enriched centromeres identified in Equus species (Nergadze, et al. 2018), another species group characterized by rapid karyotype and centromere evolution (Carbone, Nergadze, et al. 2006)

We show that while the most prevalent satellite repeat family in SSY, NLE and HMO gibbons are alpha-satellites, the vast majority of HLE satellites bound by centromeric proteins are annotated as the retroelement-derived macrosatellite SST1 (Fatyol, et al. 2000; Tremblay,
et al. 2010), underscoring, once again, differences in the prevalence and composition of centromeric satellite repeats between Hoolock and the other gibbon genera. While the sequence and structure of SST1 repeats appear relatively unchanged across SSY, NLE, and HMO, significant variation exists for SST1 repeats between these three species and HLE. Further, SST1 maps to three chromosomes in HLE (HLE 4, 15, and 18) but only one chromosome in the other gibbon species (NLE 10, HMO 14, and SSY 13); each of these gibbon chromosomes carries syntenic blocks homologous to segments of human chromosome 19 (Figures 5, 6), which span the centromeres (i.e. both p and q arm). We therefore infer that the SST1 sequence within the portion syntenic to human chromosome 19 was likely centromeric on the ancestral Hylobatidae chromosome (HyA9) (Capozzi et al. 2012). The SST1-containing segment of HSA19 then remained as a stable centromere on a single chromosome pair except in HLE, where it was involved in complex interchromosomal rearrangements, resulting in utilization of the repeat as both putative functional centromeric material and as evolutionary breakpoints on three chromosomes lacking LAVA expansions. In humans, SST1 forms a large, non-centromeric array on the q arm of HSA19 (Tremblay, et al. 2010), yet it is centromeric in more distantly related primate lineages (Fatyol, et al. 2000). It is therefore possible, given SST1 mediated rearrangements with centromeric locations in related species, that SST1 might represent an ancestral centromeric retroelement.

In addition to analyzing the DNA sequences binding to the inner kinetochore, we inquired about possible transcription of centromeric DNA. Active transcription of centromeric repeats has been detected in several organisms (reviewed in (Smurnova and de Wulf 2018)), with transcription of centromeric DNAs playing a pivotal role in the maintenance of proper centromere function by promoting CENP-A deposition (Bergmann, et al. 2011; Chan, et al. 2012). Implicated in the processes required for key centromere protein localization, interruption of centromeric satellite DNA transcription leads to chromosomal missegregation (Chueh, et al. 2009; Rošić, et al. 2014). Using de-novo ChIP-seq assemblies as centromere-proxies, we determined that among the centromere-derived transcripts in all gibbon genera, there is a spectrum of transcribed repeats comprised predominately of retroelements. Given the lack of species-specific, contiguous genomic sequence, our estimates are likely an under-representation of the centromeric transcription present in these species; however, the proportions of these centromere-transcript annotations are broadly reflecting those we defined
by ChIP-seq. Furthermore, using Nleu3.0 as a filter to remove shared among genera and non-centromeric repeats from our analyses, we were able to define a small number of species-specific centromere transcripts. For example, HLE has a high level of HLE-specific SINE-derived transcripts (predominantly Alu) within the RNA-seq data. Given their overall high abundance in ChIP-seq (Figure 1B) and RNA-seq (Figure 8B) data, we expect these SINEs to be found across several HLE centromeres. However, the location of these elements have not yet been assigned to centromeres of specific chromosomes. Ongoing long-read based genome assembly work in this species will afford discrete, chromosome specific centromeric maps to further refine chromosome-specific content and transcriptional activity. Although a specific role for centromere transcription remains to be established in gibbons, herein we show that repeats found in functional centromeric domains of gibbons across the karyotype are transcriptionally active, supporting the model that centromeric transcription is integral to centromere protein specification and function.

The historical belief that satellite DNAs are the predominant functional component of eukaryotic centromeres, with large tandem arrays present in animals, plants, and fungi (reviewed in McKinley and Cheeseman 2015), is challenged by the finding that neocentromeres devoid of satellites can successfully recruit CENP-A (Vouillaire, et al. 1993; du Sart, et al. 1997; Barry, et al. 1999; Amor, et al. 2004b). In fact, many species are now known to have some centromeres that lack satellite arrays altogether, including Equus species of horse, zebra, and donkey (Wade, et al. 2009; Nergadze, et al. 2018), orangutan (Locke, et al. 2011), chicken (Shang, et al. 2010), and potato (Gong, et al. 2012). Moreover, long-read genome assemblies spanning centromeric contigs in Drosophila (Chang, et al. 2019) and koala (Johnson, et al. 2018) has shown that retroelements, not satellites, are the predominate CENP-A binding sequences. A model has been proposed to explain these shifts in repeat content (Klein and O’Neill 2018), wherein satellite arrays, such as alpha-satellite, can become stabilized and homogenize across a karyotype (as observed for great apes). Destabilization events lead to the seeding or invasion of centromeres by transposable elements, which can be co-opted to become functional centromere sequences. Recent work has shown that retroelement insertions are common following neocentromere formation in maize (Schneider, et al. 2016), and evolutionary new centromeres in Equus asinus (Nergadze, et al. 2018) and wallabies (Longo, et al. 2009) are found in retroelement-rich regions. Combined, these
observations suggest that retroelement insertion might favor CENP-A deposition or, vice versa, CENP-A chromatin might provide a “safe heaven” for retroelements.

In line with the the observations made by others, our data suggest that satellites are neither the dominant nor exclusive functional centromeric sequences. Rather, our identification of retroelement-enriched gibbon centromeres in *Hoolock* and low abundance of satellite DNA in all four gibbon genera further supports the hypothesis that retroelements are powerful determinants of centromeric function and this might be particularly true in the context of rapid chromosome evolution. In the future, access to affordable ultra-long read sequencing technologies, as well as availability of high-quality genus- or species-specific genome assemblies, will allow for a more comprehensive view into the extraordinary DNA diversity that shapes functionally-stable centromeres in gibbons and other organisms.

**Methods**

**Establishment of lymphoblastoid cell lines**

We used previously established Epstein Barr Virus (EBV) transformed lymphoblastoid cell lines (LCL) from a *Nomascus leucogenys* (NLE) (Lazar, et al. 2018), and established new LCL for *Hoolock leuconedys* (HLE), *Hylobates moloch* (HMO) and *Symphalangus syndactylus* (SSY) gibbons according to previous protocols (Lazar, et al. 2018). Briefly, opportunistic whole blood samples were collected in sodium heparin tubes from each gibbon during routine check-ups at the Gibbon Conservation Center (Santa Clarita, CA). We used Ficoll-Paque PLUS (GE Healthcare) to isolate lymphocytes from the blood. Next, we transformed 3-9x10⁶ lymphocytes with EBV from the marmoset cell line B95-8 (ATCC CRL-1612), using a standard protocol. To do so, we incubated the cells with EBV for 2 hours at 37°C, then diluted them with RPMI-1640 (Corning cellgro) supplemented with 10% FBS (Hyclone), 1X MEM Non-essential Amino Acids Solution (Corning cellgro), 1mM Sodium pyruvate (Corning cellgro) 1% Pen-Strep (Corning cellgro) and 2mM L-glutamine (Hyclone). Finally, we grew cells undisturbed for 10-12 days and started feeding the cells with the same supplemented RPMI-1640 as soon as signs of transformation were observed under the microscope.

**Chromatin immunoprecipitation sequencing (ChIP-seq)**
We used LCLs (described above) from four gibbons each belonging to one of the four extant gibbon genera (*Hoolock, Nomascus, Hylobates, and Symphalangus*) and performed CENP-A, CENP-B and CENP-C ChIP-seq using the Magnify ChIP kit (ThermoFisher Scientific) according to manufacturer’s instructions, with minor modifications. Briefly, per ChIP assay we fixed 3x10^5 cells with 1% formaldehyde at room temperature for 10 min. Fixation was quenched with glycine, and cells were washed three times with cold 1X PBS. Cells were lysed for 10 min on ice using the Magnify Lysis buffer, in presence of Proteinase inhibitor cocktail. Lysed cells were then sonicated using the Bioruptor Pico sonicator (Diagenode) for 12 cycles (30 sec on/off) and spun down to remove cell debris. A 1% aliquot was taken from the chromatin as input, and the rest was incubated with the appropriate antibody [2ug of CENP-A (ab13939), 4ug of CENP-B (ab25734), or 2ug of CENP-C (ab50974)] at 4°C, with overnight rotation. The next day we incubated samples with Dynabead protein A/G rotating at 4°C for 2 hours, followed by bead washes, reverse-crosslink and DNA purification according to Magnify ChIP kit protocol. Concentrations of all ChIP and input samples were measured using the Qubit dsDNA High Sensitivity kit (ThermoFisher Scientific). ChIP replicates were pooled to 2ng and used to construct sequencing libraries using the NEBNext Ultra II library construction kit (New England BioLabs) without size selection. Libraries were quantified and QC’d using the Bioanalyzer High Sensitivity DNA kit (Agilent) and paired-end sequenced on the Illumina NextSeq and HiSeq platforms.

In order to test reproducibility of LAVA’s distribution in HLE centromeres we repeated CENP-A ChIP on LCL from three additional unrelated HLE gibbons (Supplementary Table S1).

**In silico validation of RepeatMasker LAVA annotations**

Since our study focused heavily on the complex composite LAVA retrotransposon, we decided to first validate that the RepeatMasker tool (Tempel 2012) is able to reliably annotate this element from short-read sequences. We used *in silico* simulated short-read datasets to estimate RepeatMasker’s false-negative and false-positive rates in identifying LAVA repeats. Briefly, to estimate false negative rates of our analysis we used wgsim (Li 2011) to simulate three datasets, each containing 2.5 million 75 bp paired-end reads (indel rate =0.02, single nucleotide mutation rate=0.05), from the sequences of 1,204 LAVA elements annotated during the construction of the gibbon reference genome (Nleu3.0) (Carbone, et al. 2014). These
reads, which are expected to only include LAVA repeats, were annotated using RepeatMasker. The \%counts of repeat not annotated as the SVA family (either LAVA or SVA_A) is an estimate of the false-negative rate of our analysis. To estimate the false positive rate of our analysis, we first used custom bash scripts and BEDtools (Quinlan and Hall 2010) to mask and remove all LAVA sequences present in the gibbon reference genome. We then used wgsim (Li 2011) to simulate three short-read datasets, each containing 2.5 million random 75 bp paired-end reads (indel rate =0.02, single nucleotide mutation rate=0.05) from the modified genome sequence. We annotated these reads, which are expected to lack LAVA repeats, using RepeatMasker and calculated percent of reads annotated as LAVA repeats. The false-positive rate is estimated by the mean \%count of repeats annotated as SVA family (including SVA_A or LAVA annotations) across simulated datasets.

**ChIP-seq data processing, repeat annotation and repeat structure comparison**

The quality of raw ChIP-seq datasets were assessed using FastQC (Andrews 2010) and reads were trimmed using Trimmomatic (Bolger, et al. 2014) to remove adaptor sequences, as well as low quality and short reads (2:30:10:26:TRUE SLIDINGWINDOW:4:20 TRAILING:20 LEADING:20 MINLEN:50). Next, we used Seqtk (Shen, et al. 2016) to randomly select 2.5 million trimmed read pairs from each ChIP sample. The random subsets of reads were converted to fasta format using manual bash scripts and their repeats were annotated using RepeatMasker version 4.0.3 and the gibbon (Hylobates sp.) 20150807 Repbase library (Jurka, et al. 2005). We used a custom script to group repeats in each library based on repeat class and family and to calculate the \%count and \%length (of total repeats) they constituted. Since overall results from \%count and \%length calculations were highly similar within datasets, we only report the \%count in the main text of this study, but both numbers are reported in Supplementary Table S2. Considering the high homology between SVA_A and LAVA and the fact that only few copies of SVA elements exist in the gibbon genome, repeats annotated as SVA_A using RepeatMasker are likely LAVA repeats and were annotated as such in this study.

To compare the sequence and structure of repeats across libraries, 50,000 reads were subsampled from HLE ChIP-seq libraries and 100,000 reads were subsampled from HMO, NLE, and SSY ChIP-seq libraries using Seqtk (Shen, et al. 2016). Fewer reads were required
to be sampled from HLE due to the enrichment of SST1 and LAVA. Subsampled libraries were annotated using RepeatMasker using the previously described annotation pipeline. We then used Repstat (Johnson, et al. 2018) with two-sided Wilcoxon signed-rank test, to identify significant differences in repeat structures annotated by RepeatMasker across libraries. Since the subsampled SSY input library only contained one repeat annotated as SST1, this library was excluded from downstream Repstat analysis of SST1 repeats.

**Identifying putative CENP binding sites within the LAVA element and examining CENP-B motif enrichment**

We used an approach originally described by Fernandes et al. 2020 (Fernandes, et al. 2020) and adapted for the LAVA element in Okhovat et al. 2020 (Okhovat et al. 2020) to identify putative CENP bindings sites in the consensus LAVA sequence. Briefly, we used Bowtie2 (Langmead and Salzberg 2012) with the --very-sensitive settings to align the trimmed reads from each input and CENP ChIP-seq library to the gibbon reference genome (Nleu3.0). We used these alignments, which consisted of both unique and multimapping reads, to identify significant ChIP-seq peaks using MACS2 (Zhang, et al. 2008) (-f BAMPE --keep-dup=2 --nomodel -q 0.01 -g 2.8e9). For each CENP library, we used BEDtools (Quinlan and Hall 2010) to identify all CENP peak summits overlapping LAVA elements annotated in the reference genome (Carbone et al. 2014). We extended each of these summits 50bp in each direction and mapping them to the consensus LAVA sequence (from the Repase library) using BLAT (Kent 2002) to generated a pileup of summits. The summit pileups generated were then used by MACS2 (Zhang, et al. 2008) (--keep-dup all --nomodel --call-summits --extsize 50) to identify summits of the summit pileups (“meta-summits”), which represent putative binding sites of CENP inside the LAVA element.

To validate our approach in detecting CENP-B box motifs, we first analyzed public human CENP-A ChIP-seq data as positive control. Human CENP-A ChIP-seq reads were aligned to Hg38 using Bowtie2 with the “--very-sensitive” paired-end settings (Langmead and Salzberg 2012). We identified significant (q<0.01) broad peaks using Model-based Analysis of ChIP-seq 2 (MACS2) (Zhang, et al. 2008), while allowing a maximum of two duplicate reads per dataset. We then used AME from the MEME suite (Machanick and Bailey 2011) to examine enrichment of CENP-B box motifs (JASPAR database, MA0637.1) (Jolma, et al.
2013) in these broad peaks using two slightly different approaches. In the first approach, we examined overall enrichment of the CENP-B box motif in sequences corresponding to all broad peak sequences, relative to shuffled versions of the same sequences. In the second approach, we sorted CENP peaks based on their fold-enrichment and compared enrichment of CENP-B box motifs between high- and low- enrichment peaks. A similar approach was then used to investigate presence of CENP-B box motifs in gibbon centromeres by aligning all libraries to the Nleu3.0 genome. P values calculated for enrichment analyses were adjusted for multiple testing using the Bonferroni method.

To characterize prevalence of the CENP-B-box motif at putative CENP binding sites within the LAVA element, we extracted 100bp sequences centered at CENP ChIP-seq summits overlapping LAVA (identified in the meta-summit analysis above) and used the Find Individual Motif Occurences (FIMO) tool of the MEME suite (Machanick and Bailey 2011) to annotate all significant (p<0.05) CENPB-box motifs. We used the predicted motifs to calculate the percentage of summits containing the CENPB-box motif in each CENP library. Next, we selected 20 of the LAVA elements with highest number of predicted CENPB-box motifs and determined the percentage of total CENP-B-box motifs that were located in the VNTR. Lastly, to examine statistical significance of enrichment of the CENP-B box motif in the putative binding sites of CENP within the LAVA element, we used the Motif Enrichment Analysis pipeline (AME) of the MEME suite (Machanick and Bailey 2011), and compared prevalence of the CENP-B box motif (MA0637.1) in the 100bp summit sequences (described above) relative to shuffled versions of the same sequences (scrambled while preserving the 2-mer frequencies).

**SST1 probe design and fluorescence in situ hybridization (FISH)**

We used our CENPA ChIP-seq data to generate genus-specific repeat consensus sequences for SST1. First, we extracted CENPA reads annotated as SST1 repeats by RepeatMasker, in each genus. R1 and R2 reads were combined within each genus and aligned to the reference SST1 consensus sequence (obtained from the RepeatMasker repeat library), using default single-end bwa (Li and Durbin 2009) settings. Next, we used a combination of SAMtools mpileup and bcftools (Li, et al. 2009) to characterize sequence variations in each genus. Lastly, we used bcftools (Li, et al. 2009) to modify the reference SST1 sequence based on
genus-specific sequence variations and construct a genus specific centromere SST1 consensus sequence. PCR primers used for probe development were designed based on genus-specific HLE SST1 consensus sequences. The consensus SST1 and primer sequences can be found in Supplementary Table S7.

Each 50 μL PCR reaction contained 0.2 μM forward and reverse primers, 0.125 mM deoxyribonucleotide triphosphate (dNTPs), 2x PCR buffer, Taq polymerase, and 100 ng genomic DNA isolated from each gibbon LCL. PCR settings consisted of an initial 3 min denaturation at 94˚C, followed by 35 cycles of 30 sec at 94˚C, 30 sec at 58˚C and 1 min at 72˚C with a final 5 min extension at 72˚C. PCR products were purified using the QIAquick PCR purification kit (Qiagen) and PCR labelled in 25 μL reactions containing 1 x GoTaq buffer (Promega), 0.2 mM dGTTPs/dATTPs/dCTTPs, 0.15 mM dTTPs, 0.2 nM forward and reverse primers, 0.625 U of GoTaq (Promega), 0.04 mM digoxigenin-deoxyuridine triphosphatase (dig-dUTPs, Enzo) and 100 ng PCR product. PCR labelling consisted of a 30 sec initial denaturation at 98˚C, followed by 25 cycles of 10 sec at 98˚C, 30 sec at 60˚C, and 30 sec at 72˚C with a final extension of 5 min at 72˚C.

FISH was carried out on metaphase spreads prepared per standard protocols [(Rooney and Czepulkowski 1992). Briefly, 200-500 ng of SST1 probe was rehydrated in Hybrisol VII (MP Biomedicals) and hybridized on slides overnight in a humid chamber at 37˚C following denaturation at 80˚C for 5 min. Prior to hybridization, slides were treated with RNase A (0.1 mg/mL in 2X SSC) for 15 min at 37˚C, 0.192 M HCl for 10 min, and denatured in 70% formamide/2X SSC at 72˚C for 2 min. Post hybridization washes were performed once in 2XSSC room temperature to remove the coverglass, 0.4X SSC/0.3% NP40 for 2 min at 72˚C and once in 2X SSC/0.1% NP40 at room temperature. The reaction was blocked in 4X SSC/0.2% Tween 20/5% BSA for 30 min at 37˚C and detection was using 1:500 anti-digoxigenin fluorescein following manufacturer's instructions for 30 min at 37˚C. Excess reagents were removed by rinsing slides three times in 4X SSC/0.2% Tween 20 for 5 min at 45˚C, rinsing in distilled water, and dehydrating in an ethanol row. Slides were counterstained with a 1:5 dilution of DAPI in Vectashield (Vector Laboratories, Inc.). Images were captured on an Olympus AX70 microscope using CytoVision software (Leica Biosystems Richmond, Inc.).

Chromosome painting was carried out on metaphase spreads using Aquarius® Whole Chromosome Painting probes (Cytocell Ltd). Probes were hybridized to RNase A treated
slides overnight in a humid chamber at 37°C following slide and probe co-denaturation at 72°C for 5 min on a Hybaid in-situ block. Post hybridization washes were performed once in 2XSSC RT to remove the coverslip, 0.4X SSC at 60°C for 2 min and once in 2X SSC/0.05% Tween 20 for 1 min at room temperature. Slides were rinsed in distilled water, dehydrated in ethanol and counterstained with a 1:5 dilution of DAPI in Vectashield (Vector Laboratories, Inc.). Images were captured using an Olympus AX70 microscope and CytoVision software (Leica Biosystems Richmond, Inc.).

**Generating ab initio CENP contigs from ChIP-seq reads**

To generate longer reads that may improve contig assembly, we used PANDAseq (Masella, et al. 2012) with default settings to merge read pairs that had >25 bp overlap with each other (quality threshold= 0.6). The combination of merged reads and unmerged read-pairs were then used by RepARK (Koch, et al. 2014) to generate de-novo assemblies according to developer’s instructions. A k-mer size of 37 bp was used for RepARK assemblies, after the optimal k-mer size was identified using the K-mer Analysis Toolkit (Mapleson, et al. 2017).

**RNA-sequencing and identification of centromere-specific reads**

Total RNA was extracted from fresh frozen cell pellets of the same four gibbons used for CENP ChIP assay, using the mirVana™ Total RNA Isolation kit (ThermoFisher Scientific). High-quality RNA was subjected to Illumina Tru-seq stranded total RNA library preparation, including Ribo-depletion. Libraries were sequenced on the NextSeq 500 v2, paired-end 75 bp (High Output kit), with a target depth of ~65M total paired-end reads/sample and an average insert length of 260 bp. RNA-seq reads were pre-processed similar to ChIP-seq reads described above, without subsequence subsampling. In order to broadly identify centromere RNA-seq reads, RNA-seq libraries was aligned to each respective RepARK assembly using Burrows-Wheeler Aligner’s default bwa-mem algorithm (Li and Durbin 2009). Aligned reads were separated and converted to FASTA format using Bedtools (Quinlan and Hall 2010) and Seqtk (Shen, et al. 2016). Repeats were annotated using RepeatMasker (described previously). In order to identify unique, centromere-derived reads from RNA-seq libraries, each RNA-seq library was aligned to Nleu3.0 using Burrows-Wheeler Aligner’s default bwa-mem algorithm (Li and Durbin 2009). Because this genome assembly does not contain reference
centromere sequences, unaligned reads were assumed to contain reads corresponding to centromere-specific transcripts. Unaligned reads were separated using SAMtools (Li, et al. 2009) and converted to FASTA format using Bedtools (Quinlan and Hall 2010) and Seqtk (Shen, et al. 2016).

To identify centromeric reads, unaligned reads were then mapped to CENP RepARK contig assemblies described above using the bwa-mem algorithm. Unaligned reads were removed from the remaining pool using SAMtools and the remaining aligned reads processed to FASTA format using Bedtools (Quinlan and Hall 2010) and Seqtk (Shen, et al. 2016). Duplicate reads produced during alignment were removed using the FASTX-Toolkit (Hannon 2010). Repeats in the remaining centromere-derived reads were annotated using RepeatMasker, as previously described.

Data Availability Statement
The ChIP-seq and RNA-seq data generated in this work are available at the Gene Expression Omnibus (GEO) database (accession number GSE161217). Scripts are available at https://github.com/gabriellehartley/CENP-Gibbon-Analysis/.

Acknowledgments and Funding Sources
Authors would like to thank Kimberly Nevonen, Christine McCann, Judy Brown and Nicole Pauloski for assistance with experiments, and the staff at the Gibbon Conservation Center (Santa Clarita, CA), especially the director Gabriella Skollar, who provided the opportunistic gibbon blood samples used for the establishment of gibbon LCLs. We also acknowledge the KCVI Epigenetics Consortium and the Exacloud cluster at OHSU, and the Center for Genome Innovation and Computational Biology Core at UConn. This work was supported by the National Science Foundation (grant number 1613856 to L.C. and R.O.) and the National Institutes of Health (grant 5R01GM123312-02 to R.O. and R01HG010333 to L.C.).

Figure Legends
Figure 1. Validation of CENP-A ChIP, and characterization of repeat composition of CENP-A, CENP-B, and CENP-C bound DNA
A) DNA fluorescence in situ hybridization (FISH) using CENP-A ChIP library as a probe shows signals (yellow) prevalently localized on centromeric regions on DAPI stained metaphase chromosome preparations for each gibbon genus. Pictures of gibbon species used in this study are shown. FISH validation of CENP-B and CENP-C ChIP libraries can be found in Supplementary Figure S1. B) The percentage of all repeats composed of each repeat family (as classified by RepeatMasker) is shown in CENP-A, CENP-B, and CENP-C ChIP-seq library for each genus. Schematic structures of repeat classes are shown below the x-axis.

Figure 2. Retrotransposons drive centromere-protein binding DNA variation across gibbon genera.

A) The SVA and the gibbon-specific composite LAVA element are depicted. Full-length LAVA (mean length 1.6-2.6KB) is comprised of portions of other repeats (CT-rich, Alu-like, and VNTR (mean length 0.5-1.4KB) all derived from SVA, as well as fractions of AluSz and L1ME5) and flanked by target site duplications. Region of high identity between SVA and LAVA is indicated by a grey box. B) Sankey diagram demonstrates the amount of retrotransposons annotated as SVA (SVA_A) and LAVA in each ChIP-seq library. C) LAVA frequency in CENP-A ChIP-seq datasets is plotted against LAVA enrichment relative to input. HLE individuals (circled) cluster separately from the other species indicating LAVA is more abundant genome-wide and at the centromeres in this species.

Figure 3. CENP-bound LAVAs in HLE differ from the the rest of the genome and from other genera.

A) Heatmap of structural variation in LAVA among different ChIP-seq libraries performed in pairwise comparisons among species (top) and between input and ChIP-IP libraries (bottom). As indicated, color corresponds to p value reported by Repstat. Comparisons in purple are significant under a p value of 0.05. B) CENP-A ChIP-seq summit pileups against the consensus LAVA sequence are shown for each species. The LAVA element scheme on top represents annotations of the consensus sequence. The position of the significant metasummits across libraries is marked with dashed vertical lines. CENP-B and CENP-C ChIP-seq summit pileups are in Supplementary Figure S2.
Figure 4. Centromere protein bound satellite distribution varies across gibbon genera. 
A) Overview of annotated satellite repeats in each species-specific CENP-A, CENP-B, and CENP-C ChIP library. Each satellite classification (shown in color-coded key at top) is shown as a percentage of total satellites annotated via RepeatMasker with (top) and without (bottom) alpha-satellite content. SSY, NLE, and HMO samples show high alpha-satellite content, while SST1 is the most abundant centromeric satellite in HLE. B) The higher absolute and relative abundance of SST1 repeats in HLE CENP-A ChIP-seq datasets was validated in three biological replicates (circled).

Figure 5. Characterization of SST1 localization on gibbon chromosomes by FISH. 
A) DNA FISH with SST1 as a probe on metaphase chromosomes from HLE (left) and NLE (right) and counterstained with DAPI. SST1 (yellow) signal is found on three chromosome pairs in HLE and one chromosome pair in NLE. B) FISH with chromosome paints HLE18 and HLE17 (pink) identified the small chromosome pair with centromeric SST1 signals (yellow) as HLE18 whereas HLE17 appears depleted of SST1.

Figure 6. SST1 associates with centromere-specific chromosome rearrangements homologous to human chromosome 19.
A) Predicted synteny of human chromosome 19 (purple) to HLE, NLE, SSY and HMO chromosomes is depicted. Phylogenetic relationship are as per (Shi and Yang 2018). Chromosome homologies to human chromosomes and Hylobates ancestral synteny are shown based on (Capozzi, et al. 2012). Both inferred ancestral chromosome heteromorphs are shown (HyA9a and HyA9b). SST1 localization is indicated with red bar. Whole-chromosome paints of human chromosome 19 (pink) on NLE (B) and HLE (C) metaphase chromosomes co-hybridized with SST1 (yellow); relevant chromosomes identified by inverted DAPI are indicated (white) in merged image (right).

Figure 7: Centromere proteins in gibbons bind similar SST1 elements across genera. 
Heatmap of structural variation analysis of SST1 among different ChIP-seq libraries are shown for pairwise comparisons among species (top) and between input and ChIP-IP (bottom).
Color corresponds to p values reported by Repstat as indicated. Comparisons in purple are significant under a p value of 0.05.

**Figure 8: Genome-wide and centromere-derived expression profiles for repeats vary across genera.**

**A)** Combined number of LAVA (*Left*) and SST1 (*Right*) repeats annotated per one million reads across the RNA-seq library of each species. **B)** Repeat annotation of centromere-derived RNA-seq reads for each genus. Each repeat classification is shown as a percentage of total annotated repeats mapped to CENP-A RepARK contigs (top of each pair) and as percentage of total annotated repeats mapped to RepARK contigs that did not map to the NLE3.0 assembly (bottom of each pair). Key for repeat annotations shown to the right.
References

Aldrup-MacDonald ME, Kuo ME, Sullivan LL, Chew K, Sullivan BA. 2016. Genomic variation within alpha satellite DNA influences centromere location on human chromosomes with metastable epialleles. Genome Research 26:1301-1311.

Alexandrov IA, Medvedev LI, Mashkova TD, Kisselev LL, Romanova LY, Yurov YB. 1993. Definition of a new alpha satellite suprachromosomal family characterized by monomeric organization. Nucleic Acids Research 21:2209-2215.

Alkan C, Ventura M, Archidiacono N, Rocchi M, Sahinalp SC, Eichler EE. 2007. Organization and evolution of primate centromeric DNA from whole-genome shotgun sequence data. PLoS Computational Biology 3.

Amor DJ, Bentley K, Ryan J, Perry J, Wong L, Slater H, Choo KH. 2004a. Human centromere repositioning "in progress". Proceedings of the National Academy of Sciences of the United States of America 101:6542-6547.

Amor DJ, Bentley K, Ryan J, Perry J, Wong L, Slater H, Choo KH. 2004b. Human centromere repositioning "in progress". Proceedings of the National Academy of Sciences of the United States of America 101:6542-6547.

Andrews S. 2010. FastQC: A quality control tool for high throughput sequence data. In. http://www.bioinformatics.babraham.ac.uk.

Barry AE, Howman EV, Cancilla MR, Saffery R, Choo KH. 1999. Sequence analysis of an 80 kb human neocentromere. Human Molecular Genetics 8:217-227.

Bergmann JH, Rodríguez MG, Martins NM, Kimura H, Kelly DA, Masumoto H, Larionov V, Jansen LE, Earnshaw WC. 2011. Epigenetic engineering shows H3K4me2 is required for HJURP targeting and CENP-A assembly on a synthetic human kinetochore. EMBO 30:328-340.

Blower MD, Sullivan BA, Karpen GH. 2002. Conserved organization of centromeric chromatin in flies and humans. Dev Cell 2:319-330.

Bolger AM, Lohse M, Usadel B. 2014. Trimmomatic: a flexible trimmer for Illumina sequence data. Bioinformatics 30:2114-2120.

Capozzi O, Carbone L, Stanyon RR, Marra A, Yang F, Whelan CW, de Jong PJ, Rocchi M, Archidiacono N. 2012. A comprehensive molecular cytogenetic analysis of chromosome rearrangements in gibbons. Genome Research 22:2520-2528.

Carbone L, Harris RA, Gnerre S, Veeramah KR, Lorente-Galdos B, Huddleston J, Meyer TJ, Herrero J, Roos C, Aken B, et al. 2014. Gibbon genome and the fast karyotype evolution of small apes. Nature 513:195-201.

Carbone L, Harris RA, Mootnick AR, Milosavljevic A, Martin DI, Rocchi M, Capozzi O, Archidiacono N, Konkel MK, Walker JA, et al. 2012. Centromere remodeling in Hoolock leuconedys (Hylobatidae) by a new transposable element unique to the gibbons. Genome Biology and Evolution 4:648-658.
Carbone L, Harris RA, Vessere GM, Mootnick AR, Humphray S, Rogers J, Kim SK, Wall JD, Martin D, Jurka J, et al. 2009. Evolutionary breakpoints in the gibbon suggest association between cytosine methylation and karyotype evolution. PLoS Genetics 5:e1000538.

Carbone L, Nergadze SG, Magnani E, Misceo D, Cardone MF, Roberto R, Bertoni L, Attolini C, Piras MF, de Jong P, et al. 2006. Evolutionary movement of centromeres in horse, donkey, and zebra. Genomics 87:777-782.

Carbone L, Vessere GM, ten Hallers BFH, Zhu B, Osoegawa K, Mootnick A, Kofler A, Wienberg J, Rogers J, Humphray S, et al. 2006. A high-resolution map of synteny disruptions in gibbon and human genomes. PLoS Genetics 2:e223.

Cellmare A, Catacchio CR, Alkan C, Giannuzzi G, Antonacci F, Cardone MF, Della Valle G, Malig M, Rocchi M, Eichler EE, et al. 2009. New insights into centromere organization and evolution from the white-cheeked gibbon and marmoset. Molecular Biology and Evolution 26:1889-1900.

Chan FL, Marshall OJ, Saffery R, Kim BW, Earle E, Choo KHA, Wong LH. 2012. Active transcription and essential role of RNA polymerase II at the centromere during mitosis. Proceedings of the National Academy of Sciences of the United States of America 109:1979-1984.

Chang C-H, Chavan A, Palladino J, Wei X, Martins NMC, Santinello B, Chen C-C, Erceg J, Beliveau BJ, Wu C-T, et al. 2019. Islands of retroelements are major components of Drosophila centromeres. PLOS Biology 17:e3000241.

Cheng ZJ, Murata M. 2003. A centromeric tandem repeat family originating from a part of Ty3/gypsy-retroelement in wheat and its relatives. Genetics 164:665-672.

Chmatal L, Schultz RM, Black BE, Lampson MA. 2017. Cell biology of cheating-transmission of centromeres and other selfish elements through asymmetric meiosis. Progress in Molecular and Subcellular Biology 56:377-396.

Chueh AC, Northrop EL, Brettingham-More KH, Choo KHA, Wong LH. 2009. LINE retrotransposon RNA is an essential structural and functional epigenetic component of a core neocentromeric chromatin. PLoS Genetics 5.

Cunningham C, Mootnick A. 2009. Gibbons. Curr Biol 19:R543-544.

du Sart D, Cancilla MR, Earle E, Mao JI, Saffery R, Tainton KM, Kalitsis P, Martyn J, Barry AE, Choo KH. 1997. A functional neo-centromere formed through activation of a latent human centromere and consisting of non alpha satellite DNA. Nature Genetics 16:144-153.

Du Y, Topp CN, Dawe RK. 2010. DNA binding of centromere protein C (CENPC) is stabilized by single-stranded RNA. PLoS Genetics 6:e1000835.

Dumont M, Gamba R, Gestraud P, Klaasen S, Worrall JT, De Vries SG, Boudreau V, Salinas-Luypaert C, Maddox PS, Lens SM, et al. 2020. Human chromosome-specific aneuploidy is influenced by DNA-dependent centromeric features. The EMBO Journal 39:e102924.

Earnshaw WC, Ratrie III H, Stetten G. 1989. Visualization of centromere proteins CENP-B and CENP-C on a stable dicentric chromosome in cytological spreads. Chromosoma 98:1-12.
Earnshaw WC, Rothfield N. 1985. Identification of a Family of Human Centromere Proteins Using Autoimmune Sera From Patients With Scleroderma. Chromosoma 91.

Fatyol K, Illes K, Diamond DC, Janish C, Szalay AA. 2000. Mer22-related sequence elements form pericentric repetitive DNA families in primates. Molecular and General Genetics 262:931-939.

Fernandes JD, Zamudio-Hurtado A, Clawson H, Kent WJ, Haussler D, Salama SR, Haeussler M. 2020. The UCSC repeat browser allows discovery and visualization of evolutionary conflict across repeat families. Mobile DNA 11:13.

Ferreri GC, Brown JD, Oberfell C, Jue N, Finn CE, O'Neill MJ, O'Neill RJ. 2011. Recent amplification of the kangaroo endogenous retrovirus, KERV, limited to the centromere. Journal of Virology 85:4761-4771.

Foltz DR, Jansen LE, Bailey AO, Yates JR, 3rd, Bassett EA, Wood S, Black BE, Cleveland DW. 2009. Centromere-specific assembly of CENP-a nucleosomes is mediated by HJURP. Cell 137:472-484.

Gamba R, Fachelletti D. 2020. From evolution to function: Two sides of the same CENP-B coin? Experimental Cell Research 390:111959.

Girirajan S, Chen L, Graves T, Marques-Bonet T, Ventura M, Fronick C, Fulton L, Rocchi M, Fulton RS, Wilson RK, et al. 2009. Sequencing human–gibbon breakpoints of synteny reveals mosaic new insertions at rearrangement sites. Genome Research 19:178-190.

Gong Z, Wu Y, Koblizkova A, Torres GA, Wang K, Iovene M, Neumann P, Zhang W, Novak P, Buell CR, et al. 2012. Repeatless and repeat-based centromeres in potato: implications for centromere evolution. Plant Cell 24:3559-3574.

FASTX-Toolkit [Internet]. 2010. Available from: http://hannonlab.cshl.edu/fastx_toolkit/

Hara T, Hirai Y, Jahan I, Hirai H, Koga A. 2012. Tandem repeat sequences evolutionarily related to SVA-type retrotransposons are expanded in the centromere region of the western hoolock gibbon, a small ape. Journal of Human Genetics 57:760-765.

Hartley G, O'Neill RJ. 2019. Centromere Repeats: Hidden Gems of the Genome. Genes 10.

Hasson D, Panchenko T, Salimian KJ, Salman MU, Sekulic N, Alonso A, Warburton PE, Black BE. 2013. The octamer is the major form of CENP-A nucleosomes at human centromeres. Nature Structure and Molecular Biology 20:687-695.

Heikkinen E, Launonen V, Muller E, Bachmann L. 1995. The pvB370 BamHI satellite DNA family of the Drosophila virilis group and its evolutionary relation to mobile dispersed genetic pDv elements. Journal of Molecular Evolution 41:604-614.

Henikoff JG, Thakur J, Kasinathan S, Henikoff S. 2015. A unique chromatin complex occupies young alpha-satellite arrays of human centromeres. Science Advances 1:e1400234.

Henikoff S, Ahmad K, Malik HS. 2001. The centromere paradox: stable inheritance with rapidly evolving DNA. Science 293:1098-1102.

Ianc B, Ochis C, Persch R, Popescu O, Damert A. 2014. Hominoid composite non-LTR retrotransposons-variety, assembly, evolution, and structural determinants of mobilization. Molecular Biology and Evolution 31:2847-2864.
Iwata-Otsubo A, Dawicki-McKenna JM, Akera T, Falk SJ, Chmatal L, Yang K, Sullivan BA, Schultz RM, Lampson MA, Black BE. 2017. Expanded satellite repeats amplify a discrete CENP-A nucleosome assembly site on chromosomes that drive in female meiosis. Current Biology 27:2365-2373.

Jain M, Olsen HE, Turner DJ, Stoddart D, Bulazel KV, Paten B, Haussler D, Willard HF, Akeson M, Miga KH. 2018. Linear assembly of a human centromere on the Y chromosome. Nature Biotechnology 36:321.

Jansen LE, Black BE, Foltz DR, Cleveland DW. 2007. Propagation of centromeric chromatin requires exit from mitosis. J Cell Biol 176:795-805.

Johnson RN, O'Meally D, Chen Z, Etherington GJ, Ho SYW, Nash WJ, Grueber CE, Cheng Y, Whittington CM, Dennison S, et al. 2018. Adaptation and conservation insights from the koala genome. Nature Genetics 50:1102-1111.

Jolma A, Yan J, Whittington T, Toivonen J, Nitta KR, Rastas P, Morgunova E, Enge M, Taipale M, Wei G, et al. 2013. DNA-binding specificities of human transcription factors. Cell 152:327-339.

Jurka J, Kapitonov VV, Pavlincek A, Klonowski P, Kohany O, Walichiewicz J. 2005. Repbase Update, a database of eukaryotic repetitive elements. Cytogenet Genome Res 110:462-467.

Kapitonov VV, Holmquist GP, Jurka J. 1998. L1 repeat is a basic unit of heterochromatin satellites in cetaceans. Molecular Biology and Evolution 15:611-612.

Kapitonov VV, Jurka J. 1999. Molecular paleontology of transposable elements from Arabidopsis thaliana. Genetica 107:27-37.

Kent WJ. 2002. BLAT - the BLAST-like alignment tool. Genome Research 12:656-664.

Klare K, Weir JR, Basilico F, Zimniak T, Massimiliano L, Ludwigs N, Herzog F, Musacchio A. 2015. CENP-C Is a Blueprint for Constitutive Centromere-Associated Network Assembly Within Human Kinetochores. Journal of Cell Biology 210.

Klein SJ, O'Neill RJ. 2018. Transposable elements: genome innovation, chromosome diversity, and centromere conflict. In. Chromosome Research. p. 5-23.

Koch P, Platzer M, Downie BR. 2014. RepARK—de novo creation of repeat libraries from whole-genome NGS reads. Nucleic Acids Research 42:e80.

Langley SA, Miga KH, Karpen GH, Langley CH. 2019. Haplotypes spanning centromeric regions reveal persistence of large blocks of archaic DNA.

Langmead B, Salzberg SL. 2012. Fast gapped-read alignment with Bowtie 2. Nature Methods 9:357-359.

Lazar NH, Nevonen KA, O'Connell B, McCann C, O'Neill RJ, Green RE, Meyer TJ, Okhovat M, Carbone L. 2018. Epigenetic maintenance of topological domains in the highly rearranged gibbon genome. Genome Research 28:983-997.

wgsim - read simulator for next generation sequencing [Internet]. 2011.

Li H, Durbin R. 2009. Fast and accurate short read alignment with Burrows-Wheeler transform. Bioinformatics 25:1754-1760.
Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G, Durbin R, Genome Project Data Processing S. 2009. The Sequence Alignment/Map format and SAMtools. Bioinformatics 25:2078-2079.

Locke DP, Hillier LW, Warren WC, Worley KC, Nazareth LV, Muzny DM, Yang S-P, Wang Z, Chinwalla AT, Minx P, et al. 2011. Comparative and demographic analysis of orangutan genomes. Nature 469:529-533.

Longo MS, Carone DM, Green ED, O’Neill MJ, O’Neill RJ. 2009. Distinct retroelement classes define evolutionary breakpoints demarcating sites of evolutionary novelty. In. BMC Genomics. p. 334.

Lupan I, Bulzu P, Popescu O, Damert A. 2015. Lineage specific evolution of the VNTR composite retrotransposon central domain and its role in retrotransposition of gibbon LAVA elements. BMC Genomics 16.

Machanick P, Bailey TL. 2011. MEME-ChIP: motif analysis of large DNA datasets. In. Bioinformatics. p. 1696-1697.

Mahtani MM, Willard HF. 1990. Pulsed-field gel analysis of alpha-satellite DNA at the human X chromosome centromere: high frequency polymorphisms and array size estimate. Genomics 7:607-613.

Mapleson D, Garcia Accinelli G, Kettleborough G, Wright J, Clavijo BJ. 2017. KAT: a K-mer analysis toolkit to quality control NGS datasets and genome assemblies. Bioinformatics 33:574-576.

Masella AP, Bartram AK, Truszkowski JM, Brown DG, J.D. N. 2012. PANDAseq: paired-end assembler for illumina sequences. BMC Bioinformatics 13:1-7.

Masumoto H, Masukata H, Muro Y, Nozaki N, Okazaki T. 1989. A human centromere antigen (CENP-B) interacts with a short specific sequence in alphoid DNA, a human centromeric satellite. Journal of Cell Biology 109:1963-1973.

McKinley KL, Cheeseman IM. 2015. The molecular basis for centromere identity and function. Nature Reviews Molecular Cell Biology 17:16-29.

Miga KH, Koren S, Rhie A, Vollger MR, Gershman A, Bzikadze A, Brooks S, Howe E, Porubsky D, Logsdon GA, et al. 2020. Telomere-to-telomere assembly of a complete human X chromosome. Nature 585:79-84.

Nergadze SG, Piras FM, Gamba R, Corbo M, Cerutti F, McCarter JGW, Cappelletti E, Gozzo F, Harman RM, Antczak DF, et al. 2018. Birth, evolution, and transmission of satellite-free mammalian centromeric domains. Genome Research 28:789-799.

Nie W, Rens W, Wang J, Yang F. 2001. Conserved chromosome segments in Hylobates hoolock revealed by human and H. leucogenys paint probes. Cytogenetics and Cell Genetics 92:248-253.

Okada T, Ohzeki J, Nakano M, Yoda K, Brinkley WR, Larionov V, Masumoto H. 2007. CENP-B Controls Centromere Formation Depending on the Chromatin Context. Cell 131.
Okhovat M, Nevonen KA, Davis BA, Michener P, Ward S, Milhaven M, Harshman L, Sohota A, Fernandes JD, Salama SR, et al. 2020. Co-option of the lineage-specific LAVA retrotransposon in the gibbon genome. PNAS 117:19328-19338.

Quinlan AR, Hall IM. 2010. BEDTools: a flexible suite of utilities for comparing genomic features. Bioinformatics 26:841-842.

Renfree MB, Papenfuss AT, Deakin JE, Lindsay J, Heider T, Belov K, Rens W, Waters PD, Pharo EA, Shaw G, et al. 2011. Genome sequence of an Australian kangaroo, Macropus eugenii, provides insight into the evolution of mammalian reproduction and development. Genome Biology 12:R81.

Roberto R, Capozzi O, Wilson RK, Mardis ER, Lomiento M, Tuzun E, Cheng Z, Mootnick AR, Archidiacono N, Rocchi M, et al. 2007. Molecular refinement of gibbon genome rearrangements. Genome Research 17:249-257.

Rooney DE, Czepulkowski BH. 1992. Human cytogenetics: a practical approach. Oxford: IRL Press.

Rošić S, Kohler F, Erhardt S. 2014. Repetitive centromeric satellite RNA is essential for kinetochore formation and cell division. Journal of Cell Biology 207:335-349.

Rudd MK, Schueler MG, Willard HF. 2003. Sequence organization and functional annotation of human centromeres. Cold Spring Harbor Symposia on Quantitative Biology 68:141-150.

Schneider KL, Xie Z, Wolfgruber TK, Presting GG. 2016. Inbreeding drives maize centromere evolution. Proceedings of the National Academy of Sciences of the United States of America 113:E987-E996.

Shang W-H, Hori T, Toyoda A, Kato J, Popendorf K, Sakakibara Y, Fujiyama A, Fukagawa T. 2010. Chickens possess centromeres with both extended tandem repeats and short non-tandem-repetitive sequences. Genome Research 20:1219-1228.

Shen W, Le S, Li Y, Hu F. 2016. SeqKit: A Cross-Platform and Ultrafast Toolkit for FASTA/Q File Manipulation. In. PLoS One.

Shi C-M, Yang Z. 2018. Coalescent-based analyses of genomic sequence data provide a robust resolution of phylogenetic relationships among major groups of gibbons. Molecular Biology and Evolution 35:159-179.

Smurnova K, de Wulf P. 2018. Centromere and pericentromere transcription: roles and regulation... in sickness and in health. Frontiers in Genetics 9:674.

Sujiwattanarat P, Thapana W, Srikulnath K, Hirai Y, Hirai H, Koga A. 2015. Higher-order repeat structure in alpha satellite DNA occurs in New World monkeys and is not confined to hominoids. Nature Scientific Reports 5.

Suntronpong A, Kugou K, Masumoto H, Srikulnath K, Ohshima K, Hirai H, Koga A. 2016. CENP-B box, a nucleotide motif involved in centromere formation, occurs in a New World monkey. Biology Letters 12:20150817.

Tempel S. 2012. Using and understanding RepeatMasker. Methods in Molecular Biology 859:29-51.
Thongchum R, Nishihara H, Srikulnath K, Hirai H, Koga A. 2019. The CENP-B box, a nucleotide motif involved in centromere formation, has multiple origins in New World monkeys. Genes & Genetic Systems 94:301-306.

Tremblay DC, Alexander Jr. G, Moseley S, Chadwick BP. 2010. Expression, tandem repeat copy number variation and stability of four macrosatellite arrays in the human genome. BMC Genomics 11:632.

Van Hooser AA, Ouspenski, II, Gregson HC, Starr DA, Yen TJ, Goldberg ML, Yokomori K, Earnshaw WC, Sullivan KF, Brinkley BR. 2001. Specification of kinetochore-forming chromatin by the histone H3 variant CENP-A. Journal of Cell Science 114:3529-3542.

Voullaire LE, Slater HR, Petrovic V, Choo KH. 1993. A functional marker centromere with no detectable alpha satellite, satellite III, or CENP-B protein: activation of a latent centromere? American Journal of Human Genetics 52:1153-1163.

Wade C, Giulotto E, Sigurdsson S, Zoli M, Gnerre S, Imsland F, Lear T, Adelson D, Bailey E, Bellone R, et al. 2009. Genome sequence, comparative analysis and population genetics of the domestic horse (Equus caballus). Science 326:865-867.

Wang H, Xing J, Grover D, Hedges DJ, Han K, Walker JA, Batzer MA. 2005. SVA elements: a hominid-specific retroposon family. Journal of Molecular Biology 354:994-1007.

Warburton PE, Haaf T, Gosden J, Lawson D, Willard HF. 1996. Characterization of a chromosome-specific chimpanzee alpha satellite subset: evolutionary relationship to subsets on human chromosomes. Genomics 33:220-228.

Waye JS, Willard HF. 1987. Nucleotide sequence heterogeneity of alpha satellite repetitive DNA: a survey of alphoid sequences from different human chromosomes. Nucleic Acids Research 15:7549-7569.

Willard HF. 1985. Chromosome-specific organization of human alpha satellite DNA. American Journal of Human Genetics 37:524-532.

Yadav V, Sheng Sun S, Billmyre RB, Thimmappa BC, Shea T, Lintner R, Bakkeren G, Cuomo CA, Heitman J, Sanyal K. 2018. RNAi is a critical determinant of centromere evolution in closely related fungi. Proceedings of the National Academy of Sciences of the United States of America 115:3108-3113.

Yoda K, Ando S, Morishita S, Hourmura K, Hashimoto K, Takeyasu K, Okazaki T. 2000. Human centromere protein A (CENP-A) can replace histone H3 in nucleosome reconstitution in vitro. Proceedings of the National Academy of Sciences of the United States of America 97:7266-7271.

Zhang Y, Liu T, Meyer CA, Eeckhoute J, Johnson DS, Bernstein BE, Nusbaum C, Myers RM, Brown M, Li W, et al. 2008. Model-based Analysis of ChIP-Seq (MACS). In. Genome Biology. p. R137.

Zhong CX, Marshall JB, Topp C, Mroczek R, Kato A, Nagaki K, Birchler JA, Jiang J, Dawe RK. 2002. Centromeric retroelements and satellites interact with maize kinetochore protein CENH3. Plant Cell 14:2825-2836.
A

B

Downloaded from https://academic.oup.com/mbe/advance-article/doi/10.1093/molbev/msab148/6275246 by guest on 16 May 2021
