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
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Linear assembly of a human centromere on the Y chromosome

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The human genome reference sequence remains incomplete owing to the challenge of assembling long tracts of near-identical tandem repeats in centromeres. We implemented a nanopore sequencing strategy to generate high-quality reads that span hundreds of kilobases of highly repetitive DNA in a human Y chromosome centromere. Combining these data with short-read variant validation, we assembled and characterized the centromeric region of a human Y chromosome.

Centromeres facilitate spindle attachment and ensure proper chromosome segregation during cell division. Normal human centromeres are enriched with AT-rich ~171-bp tandem repeats known as alpha satellite DNA¹. Most alpha satellite DNAs are organized into higher order repeats (HORs), in which chromosome-specific alpha satellite repeat units, or monomers, are reiterated as a single repeat structure hundreds or thousands of times with high (>99%) sequence conservation to form extensive arrays². Characterizing both the sequence composition of individual HOR structures and the extent of repeat variation is crucial to understanding kinetochore assembly and centromere identity^{3–5}. However, no sequencing technology (including single-molecule real-time (SMRT) sequencing or synthetic long-read technologies) or a combination of sequencing technologies has been able to assemble centromeric regions because extremely high-quality, long reads are needed to confidently traverse low-copy sequence variants. As a result, human centromeric regions remain absent from even the most complete chromosome assemblies.

Here we apply nanopore long-read sequencing to produce high-quality reads that span hundreds of kilobases of highly repetitive DNA (**Supplementary Fig. 1**). We focus on the haploid satellite array present on the Y centromere (DYZ3), as it is particularly suitable for assembly owing to its tractable size, well-characterized HOR structure, and previous physical mapping data^{6–8}.

We devised a transposase-based method that we named ‘longboard strategy’ to produce high-read coverage of full-length bacterial artificial chromosome (BAC) DNA with nanopore sequencing (MinION sequencing device, Mk1B, Oxford Nanopore Technologies). In our

longboard strategy, we linearize the circular BAC with a single cut site, then add sequencing adaptors (**Fig. 1a**). The BAC DNA passes through the pore, resulting in complete, end-to-end sequence coverage of the entire insert. Plots of read length versus megabase yield revealed an increase in megabase yield for full-length BAC DNA sequences (**Fig. 1b** and **Supplementary Fig. 2**). We present more than 3,500 full-length ‘1D’ reads (that is, one strand of the DNA is sequenced) from ten BACs (two control BACs from Xq24 and Yp11.2; eight BACs in the DYZ3 locus⁹; **Supplementary Table 1**).

Correct assembly across the centromeric locus requires overlap among a few sequence variants, meaning that accuracy of base-calls is important. Individual reads (MinION R9.4 chemistry, Albacore v1.1.1) provide insufficient sequence identity (median alignment identity of 84.8% for control BAC, RP11-482A22 reads) to ensure correct repeat assembly¹⁰. To improve overall base quality, we produced a consensus sequence from 10 iterations of 60 randomly sampled alignments of full-length 1D reads that spanned the full insert length for each BAC (**Fig. 1c**). To polish sequences, we realigned full-length nanopore reads to each BAC-derived consensus (99.2% observed for control BAC, RP11-482A22; and an observed range of 99.4–99.8% for vector sequences in DYZ3-containing BACs). To provide a truth set of array sequence variants and to evaluate any inherent nanopore sequence biases, we used Illumina BAC resequencing (Online Methods). We used eight BAC-polished sequences (e.g., 209 kb for RP11-718M18; **Fig. 1d**) to guide the ordered assembly of BACs from p-arm to q-arm, which includes an entire Y centromere.

We ordered the DYZ3-containing BACs using 16 Illumina-validated HOR variants, resulting in 365 kb of assembled alpha satellite DNA (**Fig. 2a** and **Supplementary Data 1**). The centromeric locus contains a 301-kb array that is composed of the DYZ3 HOR, with a 5.8-kb consensus sequence, repeated in a head-to-tail orientation without repeat inversions or transposable element interruptions^{6,11,12}. The assembled length of the RP11 DYZ3 array is consistent with estimates for 96 individuals from the same Y haplogroup (R1b) (**Supplementary Fig. 3**; mean: 315 kb; median: 350 kb)^{13,14}. This finding is in agreement with pulsed-field gel electrophoresis (PFGE) DYZ3 size estimates from previous physical maps, and from a Y-haplogroup matched cell line (**Supplementary Fig. 4**).

Pairwise comparisons among the 52 HORs in the assembled DYZ3 array revealed limited sequence divergence between copies (mean 99.7% pairwise identity). In agreement with a previous assessment of sequence variation within the DYZ3 array⁶, we detected instances of a 6.0-kb HOR structural variant and provide evidence for seven copies within the RP11 DYZ3 array that were present in two clusters separated by 110 kb, as roughly predicted by previous restriction map estimates⁸. Sequence characterization of the DYZ3 array revealed nine HOR haplotypes, defined by linkage between variant bases that are

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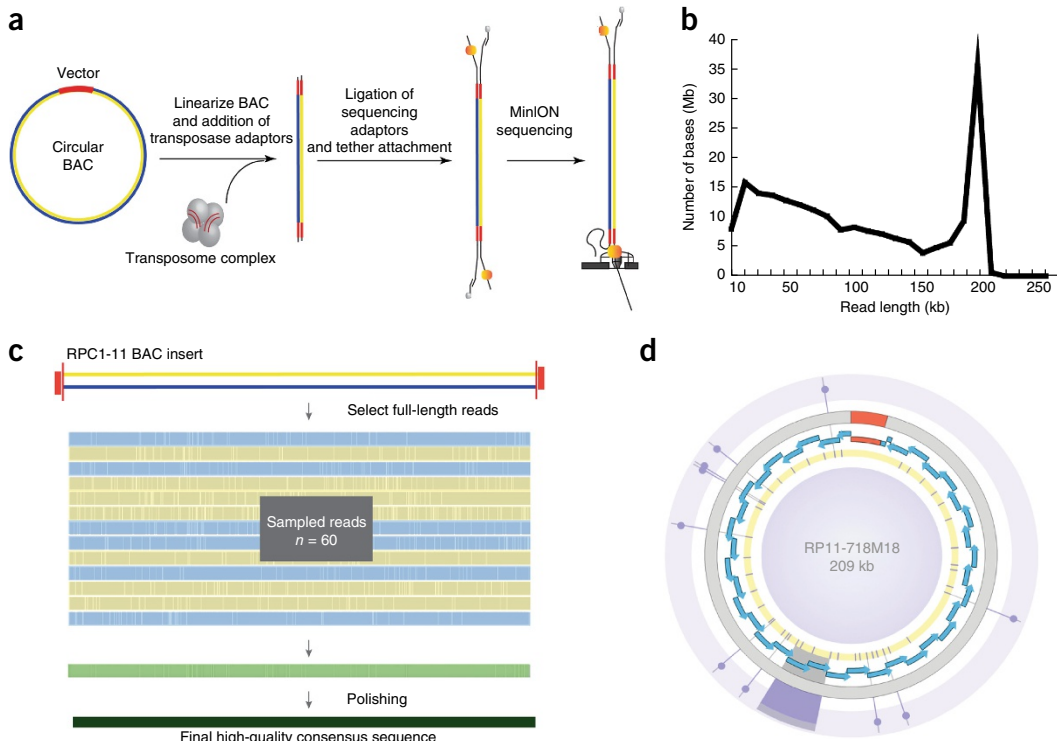


Figure 1 BAC-based longboard nanopore sequencing strategy on the MinION. (a) Optimized strategy to cut each circular BAC once with transposase results in a linear and complete DNA fragment of the BAC for nanopore sequencing. (b) Yield plot of BAC DNA (RP11-648J18). (c) High-quality BAC consensus sequences were generated by multiple alignment of 60 full-length 1D reads (shown as blue and yellow for both orientations), sampled at random with ten iterations, followed by polishing steps (green) with the entire nanopore long-read data and Illumina data. (d) Circos representation²⁰ of the polished RP11-718M18 BAC consensus sequence. Blue arrowheads indicate the position and orientation of HORs. Purple tiles in yellow background mark the position of the Illumina-validated variants. Additional purple highlight extending from select Illumina-validated variants are used to identify single-nucleotide-sequence variants and mark the site of the DYZ3 repeat structural variants (6 kb) in tandem.

frequent in the array (**Supplementary Fig. 5**). These HOR haplotypes were organized into three local blocks that were enriched for distinct haplotype groups, consistent with previous demonstrations of short-range homogenization of satellite-DNA-sequence variants^{6,15,16}.

Functional centromeres are defined by the presence of inner centromere proteins that epigenetically mark the site of kinetochore assembly^{17–19}. To define the genomic position of the functional centromere on the Y chromosome, we examined the enrichment profiles of inner

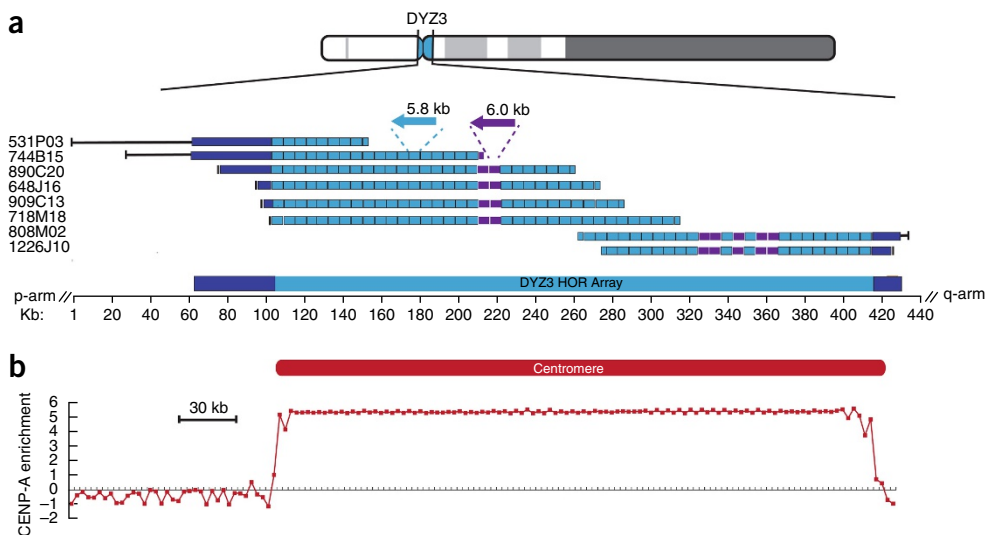


Figure 2 Linear assembly of the RP11 Y centromere. (a) Ordering of nine DYZ3-containing BACs spanning from proximal p-arm to proximal q-arm. The majority of the centromeric locus is defined by the DYZ3 conical 5.8-kb HOR (light blue). Highly divergent monomeric alpha satellite is indicated in dark blue. HOR variants (6.0 kb) indicated in purple. (b) The genomic location of the functional Y centromere is defined by the enrichment of centromere protein A (CENP-A), where enrichment (~5–6x) is attributed predominantly to the DYZ3 HOR array.

kinetochore centromere protein A (CENP-A), a histone H3 variant that replaces histone H3 in centromeric nucleosomes, using a Y-haplogroup-matched cell line that offers a similar DYZ3 array sequence (Fig. 2b and Supplementary Data 2)^{5,14,19}. We found that CENP-A enrichment was predominantly restricted to the canonical DYZ3 HOR array, although we did identify reduced centromere protein enrichment extending up to 20 kb into flanking divergent alpha satellite on both the p-arm and q-arm side. Thus, we provide a complete genomic definition of a human centromere, which may help to advance sequence-based studies of centromere identity and function.

We applied a long-read strategy to map, sequence, and assemble tandemly repeated satellite DNAs and resolve, for the first time to our knowledge, the array repeat organization and structure in a human centromere. Previous modeled satellite arrays¹⁴ are based on incomplete and gapped maps, and do not present complete assembly data across the full array. Our complete assembly enables the precise number of repeats in an array to be robustly measured and resolves the order, orientation, and density of both repeat-length variants across the full extent of the array. This work could potentially advance studies of centromere evolution and function and may aid ongoing efforts to complete the human genome.

METHODS

Methods, including statements of data availability and any associated accession codes and references, are available in the [online version of the paper](#).

Note: Any Supplementary Information and Source Data files are available in the [online version of the paper](#).

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AUTHOR CONTRIBUTIONS

K.H.M. and H.E.W. conceived the project. K.H.M., M.J., D.J.T., D.S., H.E.O., and M.A. designed the experiments; M.J. and H.E.O. were involved with BAC sample preparation; M.J. and H.E.O. performed MinION sequencing and base-calling; M.J. and K.H.M. analyzed the BAC sequencing data and validation analyses;

K.H.M. performed the pulsed-field gel electrophoresis array length estimates; K.V.B. contributed FISH analysis; K.H.M., M.J., and H.E.O. contributed to analysis and figure generation; M.A., D.J.T., D.S., H.E.W., B.P., and D.H. provided technical advice; all authors contributed to the writing, editing, and completion of the manuscript.

COMPETING INTERESTS

M.A. and M.J. are consultants to Oxford Nanopore Technologies. D.T. and D.S. are employed by Oxford Nanopore Technologies.

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ONLINE METHODS

BAC DNA preparation and validation. Clones of bacterial artificial chromosomes (BACs) used in this study were obtained from BACPAC RPC1-11 library, Children's Hospital Oakland Research Institute in Oakland, California, USA (<https://bacpacresources.org/>). BACs that span the human Y centromere, RP11-1081I4, RP11-1226J10, RP11-808M02, RP11-531P03, RP11-909C13, RP11-890C20, RP11-744B15, RP11-648J16, RP11-718M18, and RP11-482A22, were determined based on previous hybridization with DYZ3-specific probes, and confirmed by PCR with STSs sY715 and sY78 (ref. 9). Notably, DYZ3 sequences, unlike shorter satellite DNAs, have been observed to be stable and cloned without bias^{5,21}. The RP11-482A22 BAC was selected as our control since it had previously been characterized by nanopore long-read sequencing²², and presented ~134 kb of assembled, unique sequence present in the GRCh38 reference assembly to evaluate our alignment and polishing strategy. BAC DNA was prepared using the QIAGEN Large-Construct Kit (Cat No./ID: 12462). To ensure removal of the *Escherichia coli* genome, it was important to include an exonuclease incubation step at 37 °C for 1 h, as provided within the QIAGEN Large-Construct Kit. BAC DNAs were hydrated in TE buffer. BAC Insert length estimates were determined by pulsed-field gel electrophoresis (PFGE) (data not shown).

Longboard MinION protocol. MinIONs can process long fragments, as has been previously documented²². While these long reads demonstrate the processivity of nanopore sequencing, they offer insufficient coverage to resolve complex, repeat-rich regions. To systematically enrich for the number of long reads per MinION sequencing run, we developed a strategy that uses the Oxford Nanopore Technologies (ONT) Rapid Sequencing Kit (RAD002). We performed a titration between the transposase from this kit (RAD002) and circular BAC DNA. This was done to achieve conditions that would optimize the probability of individual circular BAC fragments being cut by the transposase only once. To this end, we diluted the 'live' transposase from the RAD002 kit with the 'dead' transposase provided by ONT. For PFGE-based tests, we used 1 µl of 'live' transposase and 1.5 µl of 'dead' transposase per 200 ng of DNA in a 10-µl reaction volume. This reaction mix was then incubated at 30 °C for 1 min and 75 °C for 1 min, followed by PFGE. Our PFGE tests used 1% high-melting agarose gels and were run with standard 180° field inversion gel electrophoresis (FIGE) conditions for 3.5 h. An example PFGE gel is shown in **Supplementary Figure 6**.

For MinION sequencing library preparation, we used 1.5 µl of 'live' transposase and 1 µl of 'dead' transposase (supplied by ONT) per 1 µg of DNA in a 10-µl reaction volume. Briefly, this reaction mix was then incubated at 30 °C for 1 min and 75 °C for 1 min. We then added 1 µl of the sequencing adaptor and 1 µl of Blunt/TA Ligase Master Mix (New England BioLabs) and incubated the reaction for 5 min. This was the adapted BAC DNA library for the MinION. R9.4 SpotON flow cells were primed using the protocol recommended by ONT. We prepared 1 ml of priming buffer with 500 µl running buffer (RBF) and 500 µl water. Flow cells were primed with 800 µl priming buffer via the side loading port. We waited for 5 min to ensure initial buffering before loading the remaining 200 µl of priming buffer via the side loading port but with the SpotON open. We next added 35 µl RBF and 28 µl water to the 12 µl library for a total volume of 75 µl. We loaded this library on the flow cell via the SpotON port and proceeded to start a 48 h MinION run.

When a nanopore run is underway, the amplifiers controlling individual pores can alter voltage to get rid of unadapted molecules that can otherwise block the pore. With R9.4 chemistry, ONT introduced global flicking that reversed the potential every 10 min by default to clear all nanopores of all molecules. At 450 b.p.s., a 200 kb BAC would take around 7.5 min to be processed. To ensure sufficient time for capturing BAC molecules on the MinION, we changed the global flicking time period to 30 min. This is no longer the case with an update to ONT's MinKNOW software, and on the later BAC sequencing runs we did not change any parameters. We acknowledge that generating long (>100 kb) reads presents challenges, given the dynamics of high-molecular-weight (HMW) DNA for ligation, chemistry updates, and delivery of free ends to the pore, reducing the effective yield. We found that high-quality and a large quantity of starting material (i.e., our strategy is designed for 1 µg of starting material that does not show signs of DNA shearing and/or degradation when evaluated by PFGE) and reduction of smaller DNA fragments were necessary for the longboard strategy.

Protocol to improve long-read sequence by consensus and polishing.

BAC-based assembly across the DYZ3 locus requires overlap among a few informative sequence variants, thus placing great importance on the accuracy of base-calls. Therefore, we employed the following strategy to improve overall base quality. First, we derived a consensus from multiple alignments of 1D reads that span the full insert length for each BAC. Further, polishing steps were performed using realignment of all full-length nanopore reads for each BAC. As a result, each BAC sequencing project resulted in a single, polished BAC consensus sequence. To validate single-copy variants, useful in an overlap-layout-assembly strategy, we included Illumina data sets for each BAC. Illumina data were not used to correct or validate variants observed multiple times within a given BAC sequence due to the reduced mapping quality.

MinION base-calling. All of the BAC runs were initially base-called using Metrichor, ONT's cloud basecaller. Metrichor classified reads as pass or fail using a Q-value threshold. We selected the full-length BAC reads from the pass reads. We later base-called all of the BAC runs again using Albacore 1.1.1, which included significant improvements on homopolymer calls. This version of Albacore did not contain a pass/fail cutoff. We re-performed the informatics using Albacore base-calls for full-length reads selected from the pass Metrichor base-calls. We selected BAC full-length reads as determined by observed enrichment in our yield plots (shown in **Supplementary Fig. 7** the read versus read length plots converted to yield plots to identify BAC length min-max selection thresholds).

Full-length reads used in this study were determined to contain at least 3 kb of vector sequence, as determined by BLASR²³ (*-sdp TupleSize 8 -bestn 1 -nproc 8 -m 0*) alignment with the pBACe3.6 vector (GenBank Accession: U80929.2). Reads were converted to the forward strand. Reads were re-oriented relative to a fixed 3-kb vector sequence, aligning the transition from vector to insert.

Derive BAC consensus sequence. Reoriented reads were sampled at random (blasr_output.py). Multiple sequence alignment (MSA) was performed using kalign²⁴. We determined empirically that sampling greater than 60 reads provided limited benefit to consensus base quality (**Supplementary Fig. 8**). We computed the consensus from the MSA whereas the most prevalent base at each position was called. Gaps were only considered in the consensus if the second most frequent nucleotide at that position was present in less than ten reads. We performed random sampling followed by MSA iteratively 10×, resulting in a panel of ten consensus sequences, observed to provide a ~1% boost in consensus sequence identity (**Supplementary Fig. 8**). To improve the final consensus sequence, we next performed a final MSA on the collection of ten consensus sequences derived from sampling.

Polish BAC consensus sequence. Consensus sequence polishing was performed by aligning full-length 1D nanopore reads for each BAC to the consensus (BLASR²⁵, *-sdp TupleSize 8 -bestn 1 -nproc 8 -m 0*). We used pysamstats (<https://github.com/alimanfoo/pysamstats>) to identify read support for each base call. We determined the average base coverage for each back, and filtered those bases that had low-coverage support (defined as having less than half of the average base coverage). Bases were lower-case masked if they were supported by sufficient sequence coverage, yet had <50% support for a given base call in the reads aligned.

Variation validation. We performed Illumina resequencing (MiSeq V3 600bp; 2 × 300 bp) for all nine DYZ3-containing BACs to validate single-copy DYZ3 HOR variants in the nanopore consensus sequence. Inherent sequence bias is expected in nanopore sequencing²², therefore we first used the Illumina matched data sets to evaluate the extent and type of sequence bias in our initial read sets, and our final polished consensus sequence. Changes in ionic current, as individual DNA strands are read through the nanopore, are each associated with a unique 5-nucleotide k-mer. Therefore in an effort to detect inherent sequence errors due to nanopore sensing, we compared counts of 5-mers. Alignment of full-length HORs within each polished BAC sequence to the canonical DYZ3 repeat demonstrated that these sequences are nearly identical, where in RP11-718M18 we detected 1,449 variant positions (42%

mismatches, 27% deletions, and 31% insertions) across 202,582 bp of repeats (99.5% identity). Although the 5-mer frequency profiles between the two data sets were largely concordant (**Supplementary Fig. 9**), we found that poly(dA) and poly(dT) homopolymers were overrepresented in our initial nanopore read data sets, a finding that is consistent with genome-wide observations. These poly(dA) and poly(dT) over-representations were reduced in our quality-corrected consensus sequences especially for 6-mers and 7-mers.

K-mer method. Using a k-mer strategy (where $k = 21$ bp), we identified exact matches between the Illumina and each BAC consensus sequence. Illumina read data and the BAC-polished consensus sequences were reformatted into respective k-mer library (where $k = 21$ bp, with 1 bp slide using Jellyfish v2 software²⁵), in forward and reverse orientation. K-mers that matched the pBACe3.6 sequence exactly were labeled as 'vector'. K-mers that matched the DYZ3 consensus sequence exactly¹⁴ were labeled as 'ceny'. We first demonstrated that the labeled k-mers were useful in predicting copy number. Initially, we showed how the ceny k-mer frequency in the BACs predicted the DYZ3 copy number, relative to the number observed in our nanopore consensus (**Supplementary Fig. 10a**). DYZ3 copy number in each consensus sequence derived from nanopore reads was determined using HMMER3 (ref. 26) (v3.1b2) with a profile constructed from the DYZ3 reference repeat. By plotting the distribution of vector k-mer counts (**Supplementary Fig. 10b** for RP11-718M18), we observed a range of expected k-mer counts for single-copy sites. DYZ3 repeat variants (single-copy satVARs) were determined as k-mers that (1) did not have an exact match with either the vector or DYZ3 reference repeat, (2) spanned a single DYZ3-assigned variant in reference-polished consensus sequence (i.e., that particular k-mer was observed only once in the reference), (3) and had a k-mer depth profile in the range of the corresponding BAC vector k-mer distribution. As a final conservative measure, satVARs used in overlap-layout-consensus assembly were supported by two or more overlapping Illumina k-mers (**Supplementary Fig. 10**). To test if it was possible to predict a single-copy DYZ3 repeat variant by chance, or by error introduced in the Illumina read sequences, we ran 1,000 simulated trials using our RP11-718M18 Illumina data. Here, we randomly introduced a single variant into the polished RP11-718M18 DYZ3 array (false positive). We generated 1,000 simulated sequences, each containing a single randomly introduced single-copy variant. Next, we queried if the 21-mer spanning the introduced variant was (a) found in the corresponding Illumina data set and (b) if so, we monitored the coverage. Ultimately, none of the simulated false-positive variants (21-mer) met our criteria of a true variant. That is, although the simulated variants were identified in our Illumina data, they had insufficient sequence coverage to be included in our study. Greater than 95% of the introduced false variants had $\leq 100\times$ coverage, with only one variant observed to have the maximum value of $300\times$. True variants were determined using this data set with values from 1,100–1600 \times , as observed in our vector distribution.

Alignment method. We employed a short-read alignment strategy to validate single-copy variants in our polished consensus sequence. Illumina-merged reads (PEAR, standard parameters²⁷) were mapped to the RP11 Y-assembled sequence using BWA-MEM²⁸. BWA-MEM is a component of the BWA package and was chosen because of its speed and ubiquitous use in sequence mapping and analysis pipelines. Aside from the difficulties of mapping the ultra-long reads unique to this work, any other mapper could be used instead. This involves mapping Illumina data to each BAC consensus sequence. After filtering those alignments with mapping quality less than 20, single-nucleotide DYZ3 variants (i.e., a variant that is observed uniquely, or once in a DYZ3 HOR in a given BAC) were considered "validated" if they had support of at least 80% of the reads and had sequence coverage within the read depth distribution observed in the single-copy vector sequence for each BAC data set.

To explore Illumina sequence coverage necessary for our consensus polishing strategy we initially investigated a range (20–100 \times) of simulated sequence coverage relative to a 73-kb control region (hg38 chrY:10137141–10210167) within the RP11-531P03 BAC data. Simulated paired read data using the ART Illumina simulator software²⁹ was specified for the MiSeq sequencing system (MiSeq v3 (250 bp), or 'MSv3'), with a mean size of 400 bp DNA fragments

for paired-end simulations. Using our polishing protocol, where reads are filtered by mapping quality score (i.e., at least a score of 20: that the probability of correctly mapping is \log_{10} of 0.01×10 , or 0.99), base frequency was next determined for each position using pysamstats, and a final, polished consensus was determined by taking the base call at any given position that is represented by sufficient coverage (at least half of the determined average across the entire BAC) and is supported by a percentage of Illumina reads mapped to that location (in our study, we required at least 80%). If we require at least 80% of mapped reads to support a given base call, we determine that $30\times$ coverage is sufficient to reach 99% sequence identity (or the same as our observed identity using our entire Illumina read data set, indicated as a gray dotted line in **Supplementary Fig. 11**). If we require at least 90% of mapped reads to support a particular variant it is necessary to increase coverage to $70\times$ to reach an equivalent polished percent identity.

To evaluate our mapping strategy, we performed a basic simulation using an artificially generated array of ten identical DYZ3 (5.7 kb) repeats. We then randomly introduced a single base change resulting in a new sequence with nine identical DYZ3 repeats and one repeat distinguished by a single-nucleotide change (**Supplementary Fig. 12**). We first demonstrate that we are able to confidently detect the single variant by simulating reads from the reference sequence containing the introduced variant of varying coverage and Illumina substitution error rate. Additionally, we investigated whether we would detect the variant as an artifact due to Illumina read errors. To test this, we next simulated Illumina reads from a DYZ3 reference array that did not contain the introduced variant (i.e., ten exact copies of the DYZ3 repeat). We performed this simulation $100\times$, thus creating $100\times$ reference arrays each with a randomly placed single variant. Within each evaluation we mapped in parallel simulated Illumina reads from (a) the array containing introduced variant sequence and (b) the array that lacked the variant. In experiments where reads containing the introduced variant were mapped to the reference containing the variant, we observed the introduced base across variations of sequence coverage and increased error rates. To validate a variant as "true," we next evaluated the supporting sequence coverage. For example in $100\times$ coverage, using the default Illumina error rate we observed 96 "true" calls out of 100 simulations, where in each case we set a threshold such that at least 80% of reads that spanned the introduced variant supported the base call. We found that Illumina quality did influence our ability to confidently validate array variants by reducing the coverage. When the substitution error was increased by 1/10th we observed a decrease to only 75 "true" variant calls out of $100\times$ simulations. Therefore, we suspect that Illumina sequencing errors may challenge our ability to completely detect true-positive variants.

In our alternate experiments, although simulated Illumina reads from ten identical copies of the DYZ3 repeat were mapped to a reference containing an introduced variant, we did not observe a single simulation and/or condition with sufficient coverage for "true" validation. We do report an increase in the percentage of reads that support the introduced variant as we increase the Illumina substitution error rate, however, the range of read depth observed across all experiments was far below our coverage threshold. We obtained similar results when we repeated this simulation using sequences from the RP11-718M18 DYZ3 array.

Finally, standard quality Illumina-based polishing with pilon 21 was applied strictly to unique (non-satellite DNA) sequences on the proximal p and q arms to improve final quality. Alignment of polished consensus sequences from our control BAC from Xq24 (RP11-482A22) and non-satellite DNA in the p-arm adjacent to the centromere (Yp11.2, RP11-531P03) revealed base-quality improvement to $>99\%$ identity.

Prediction and validation of DYZ3 array. BAC ordering was determined using overlapping informative single-nucleotide variants (including the nine DYZ3 6.0 kb structural variants) in addition to alignments directly to either assembled sequence on the p-arm or q-arm of the human reference assembly (GRCh38). Notably, physical mapping data were not needed in advance to guide our assembly. Rather these data were provided to evaluate our final array length predictions. Full-length DYZ3 HORs (ordered 1–52) were evaluated by MSA (using kalign²⁴) between overlapping BACs, with emphasis on repeats 28–35 that define the overlap between BACs anchored to the p-arm or q-arm (**Supplementary Fig. 13**). RPC1-11 BAC library has been previously

referenced as derived from a known carrier of haplogroup R1b^{30,31}. We compared our predicted DYZ3 array length with 93 R1b Y-haplogroup-matched individuals by intersecting previously published DYZ3 array length estimates for 1000 Genome phase 1 data^{13,14} with donor-matched Y-haplogroup information³². To investigate the concordance of our array prediction with previous physical maps of the Y-centromere we identified the positions of referenced restriction sites that directly flank the DYZ3 array in the human chromosome Y assembly (GRCh38)^{6,7,33}. It is unknown if previously published individuals are from the same population cohort as the RPC1-11 donor genome, therefore we performed similar PFGE DYZ3 array PFGE length estimates using the HuRef B-lymphoblast cell line (available from Coriell Institute as GM25430), previously characterized to be in the R1-b Y-haplogroup³⁴.

PFGE alpha satellite Southern. High-molecular-weight HuRef genomic DNA was resuspended in agarose plugs using 5×10^6 cells per 100 μ L 0.75% CleanCut Agarose (CHEF Genomic DNA Plug Kits Cat #: 170-3591 BIORAD). A female lymphoblastoid cell line (GM12708) was included as a negative control. Agarose plug digests were performed overnight (8–12 h) with 30–50 U of each enzyme with matched NEB buffer. PFGE Southern experiments used 1/4–1/2 agarose plug per lane (~5–10 μ g) in a 1% SeaKem LE Agarose gel and 0.5 \times TBE. CHEF Mapper conditions were optimized to resolve 0.1–2.0 Mb DNAs: voltage 6V/cm, runtime: 26:40 h, in angle: 120, initial switch time: 6.75 s, final switch time: 1 m 33.69 s, with a linear ramping factor. We used the Lambda (NEB; N0340S) and *Saccharomyces cerevisiae* (NEB; N0345S) as markers. Methods of transfer to nylon filters, prehybridization, and chromosome-specific hybridization with 32P-labeled satellite probes have been described³⁵. Briefly, DNA was transferred to nylon membrane (Zeta Probe GT nylon membrane; CAT# 162-0196) for ~24 h. DYZ3 probe (50 ng DNA labeled ~2 c.p.m./mL; amplicon product using previously published STS DYZ3 Y-A and Y-B primers³⁶) was hybridized for 16 h at 42 °C. In addition to standard wash conditions³⁵, we performed two additional stringent wash (buffer: 0.1% SDS and 0.1 \times SSC) steps for 10 min at 72 °C to remove non-specific binding. Image was recovered after 20 h exposure.

Sequence characterization of Y centromeric region. The DYZ3 HOR sequence and chromosomal location of the active centromere on the human chromosome Y is not shared among closely related great apes³⁷. However, previous evolutionary dating of specific transposable element subfamilies (notably, L1PA3 9.2–15.8 MYA³⁸) within the divergent satellite DNAs, as well as shared synteny of 11.9 kb of alpha satellite DNA in the chimpanzee genome Yq assembly indicate that the locus was present in the last common ancestor with chimpanzee (Supplementary Fig. 14).

Comparative genomic analysis between human and chimpanzee were performed using UCSC Genome Browser liftOver³⁹ between human (GRCh38, or hg38 chrY:10,203,170–10,214,883) and the chimpanzee genome (panTro5 chrY:15,306,523–15,356,698, with 100% span at 97.3% sequence identity). Alpha satellite and adjacent repeat in the chimpanzee genome that share limited sequence homology with human were determined using UCSC repeat table browser annotation⁴⁰.

The location of the centromere across primate Y-chromosomes was determined by fluorescence *in situ* hybridization (FISH) (Supplementary Fig. 14). Preparation of mitotic chromosomes and BAC-based probes were carried according to standard procedures⁴¹. Primate cell lines were obtained from Coriell: *Pan paniscus* (Bonobo) AG05253; *Pan troglodytes* (Common Chimpanzee) S006006E. Male gorilla fibroblast cells were provided by Stephen O'Brien (National Cancer Institute, Frederick, MD) as previously discussed⁴². The HuRef cell line³⁴ (GM25430) was provided through collaboration with Samuel Levy. BAC DNAs were isolated from bacteria stabs obtained from CHORI BACPAC. Metaphase spreads were obtained after a 1 h 15 min colcemid/karyomax (Gibco) treatment followed by incubation in a hypotonic solution. Cells were counterstained with 4',6-diamidino-2-phenylindole (DAPI) (Vector). BAC DNA probes were labeled using Alexa flour dyes (488, green and 594, red) (ThermoFisher). The BAC probes were labeled with biotin 14-dATP by nick translation (Gibco). And the chromosomes were counterstained with DAPI. Microscopy, image acquisition, and processing were performed using standard procedures.

Epigenetic mapping of centromere proteins. To evaluate similarity between the HuRef DYZ3 reference model (GenBank: GJ212193) and our RP11 BAC-assembly we determined the relative frequency of each k-mer in the array (where $k = 21$, with a 1-bp slide taking into account both forward and reverse sequence orientation using Jellyfish software) normalized by the total number of observed k-mers (Supplementary Fig. 15), with the Pearson correlation coefficient. Enrichment across the RP11 Y assembly was determined using the log-transformed relative enrichment of each 50-mer frequency relative to the frequency of that 50-mer in background control (GEO Accession: GSE45497 ID: 200045497), as previously described⁵. If a 50-mer is not observed in the ChIP background the relative frequency was determined relative to the HuRef Sanger WGS read data (AADD00000000 WGS)³⁴. Average enrichment values were calculated for windows size 6 kb (Fig. 2). Additionally, CENP-A and C paired read data sets (GEO Accession: GSE60951 ID: 200060951)⁴³ were merged (PEAR²⁷, standard parameters) and mapped to all alpha satellite reference models in GRCh38. Reads that mapped specifically to the DYZ3 reference model were selected to study enrichment to the HOR array. The total number of bases mapped from CENP-A and CENP-C data versus the input controls was used to determine relative enrichment. Second, reads that mapped specifically to the DYZ3 reference model were aligned to the DYZ3 5.7 kb in consensus (indexed in tandem to avoid edge-effects), and read depth profiles were determined. To characterize enrichment outside of the DYZ3 array CENP-A, CENP-C and Input data were mapped directly to the RP11 Y-assembly. Reads mapping to the DYZ3 array were ignored. Read alignments were only considered outside of the DYZ3 array if no mismatches, insertions, or deletions were observed to the reference and if the read could be aligned to a single location (removing any reads with mapping score of 0). Sequence depth profiles were calculated by counting the number of bases at any position and normalizing by the total number of bases in each respective data set. Relative enrichment was obtained by taking the log-transformed normalized ratio of centromere protein (A or C) to Input.

Statistics. The Pearson correlation coefficient was used to determine a positive linear relationship in our data sets (as shown in Supplementary Figs. 10a and 15a). Simulation experiments using Illumina short read data were performed using 100 replicates. Representative gel image shown (Supplementary Fig. 6) was repeated ten times, or once for each BAC in our study, with consistent results. Representative Southern Blot (shown in Supplementary Fig. 4a), was repeated twice with different restriction enzymes with the same results. Centromere Y position analysis using FISH on a panel of primates were repeated at least two times, and results were invariable between experiments and between hybridization patterns within multiple metaphase spreads within a given experiment.

Code availability. This study used previously published software: alignments were performed using BLASR²³ (version 1.3.1.124201) and BWA MEM²⁸ (0.7.12-r1044). Consensus alignments were obtained using kalign²⁴ (version 2.04). Global alignments of HORs used needle⁴⁴ (EMBOSS:6.5.7.0). Repeat characterization was performed using RepeatMasker (Smit, AFA, Hubley, R & Green, P. *RepeatMasker Open-4.0*. 2013-2015; <http://www.repeatmasker.org>). Satellite monomers were determined using profile hidden Markov model (HMMER3)²⁶. Jellyfish (version 2.0.0)²⁵ was used to characterize k-mers. Illumina read simulations was performed using ART (version 2.5.8)²⁹. PEAR²⁷ (version 0.9.0) was used to merge paired read data. Comparative genomic analysis between human and chimpanzee were performed using UCSC Genome Browser liftOver³⁹. Additional scripts used in preparing sequences before consensus generation are deposited in GitHub: <https://github.com/khmiga/CENY>.

Life Sciences Reporting Summary. Further information on experimental design is available in the Life Sciences Reporting Summary.

Data availability. Sequence data that support the findings of this study have been deposited in GenBank with reference to BioProject ID PRJNA397218, and SRA accession codes SRR5902337 and SRR5902355. BAC consensus sequences and RP11-CENY array assembly are deposited under GenBank accession numbers MF741337–MF741347.

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▶ Experimental design

1. Sample size

Describe how sample size was determined.

A single sample (RP11) was sequenced and so this is not applicable.

2. Data exclusions

Describe any data exclusions.

We excluded short reads from our analysis - defined as reads that are not represent the full length of the BAC insert and/or offered less than 4 kb of vector sequence.

3. Replication

Describe whether the experimental findings were reliably reproduced.

We reliably reproduced the MinION base statistics and consensus polishing results using a control RP11-482A22 BAC from the X chromosome

4. Randomization

Describe how samples/organisms/participants were allocated into experimental groups.

Read ids were randomly selected from a scrambled index of all full length reads. We optimized our study by obtaining random sampling of 10, 30, 60, 90 reads. Improvements were 98-99% identity for 60x, with only slight improvements with greater read depth.

5. Blinding

Describe whether the investigators were blinded to group allocation during data collection and/or analysis.

No blinding was required for this study.

Note: all studies involving animals and/or human research participants must disclose whether blinding and randomization were used.

6. Statistical parameters

For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or in the Methods section if additional space is needed).

n/a Confirmed

- The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement (animals, litters, cultures, etc.)
- A description of how samples were collected, noting whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
- A statement indicating how many times each experiment was replicated
- The statistical test(s) used and whether they are one- or two-sided (note: only common tests should be described solely by name; more complex techniques should be described in the Methods section)
- A description of any assumptions or corrections, such as an adjustment for multiple comparisons
- The test results (e.g. P values) given as exact values whenever possible and with confidence intervals noted
- A clear description of statistics including central tendency (e.g. median, mean) and variation (e.g. standard deviation, interquartile range)
- Clearly defined error bars

See the web collection on [statistics for biologists](#) for further resources and guidance.

► Software

Policy information about [availability of computer code](#)

7. Software

Describe the software used to analyze the data in this study.

This study used previously published software: alignments were performed using BLASR (version 1.3.1.124201) and BWA MEM (0.7.12-r1044). Consensus alignments were obtained using kalign (version 2.04). Global alignments of HORs used needle (EMBOSS:6.5.7.0). Repeat characterization was performed using RepeatMasker (Smit, AFA, Hubley, R & Green, P. RepeatMasker Open-4.0. 2013-2015; <http://www.repeatmasker.org>). Satellite monomers were determined using profile hidden Markov model (HMMER3). Jellyfish (version 2.0.0) was used to characterize k-mers. Illumina read simulations was performed using ART (version 2.5.8). PEAR (version 0.9.0) was used to merge paired read data. Comparative genomic analysis between human and chimpanzee were performed using UCSC Genome Browser liftOver. Additional scripts used in preparing sequences before consensus generation are deposited in GitHub: <https://github.com/khmiga/CENY>.

For manuscripts utilizing custom algorithms or software that are central to the paper but not yet described in the published literature, software must be made available to editors and reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). *Nature Methods* [guidance for providing algorithms and software for publication](#) provides further information on this topic.

► Materials and reagents

Policy information about [availability of materials](#)

8. Materials availability

Indicate whether there are restrictions on availability of unique materials or if these materials are only available for distribution by a for-profit company.

No restrictions are associated with this work

9. Antibodies

Describe the antibodies used and how they were validated for use in the system under study (i.e. assay and species).

No antibodies were used in this study. ChIP Seq datasets (using CENPA and CENPC antibodies) were obtained from two previously published studies.

10. Eukaryotic cell lines

a. State the source of each eukaryotic cell line used.

HuRef (Coriell GM25430); A female lymphoblastoid cell line (Coriell GM12708); Pan paniscus (Bonobo) Coriell: AG05253; Pan troglodytes (Common Chimpanzee) Coriell: S006006E. Male gorilla fibroblast cells were provided to Dr. Willard's lab previously by Dr Stephen O'Brien (National Cancer Institute, Frederick, MD).

b. Describe the method of cell line authentication used.

Cell line authentication was determined based information and quality assurance from Coriell biorepository. Source validates cells as described here: https://www.coriell.org/0/pdf/CC_Process_Flow.pdf

c. Report whether the cell lines were tested for mycoplasma contamination.

Cells were tested for mycoplasma contamination by biorepository

d. If any of the cell lines used are listed in the database of commonly misidentified cell lines maintained by [ICLAC](#), provide a scientific rationale for their use.

No commonly mis-identified cell lines were used.

► Animals and human research participants

Policy information about [studies involving animals](#); when reporting animal research, follow the [ARRIVE guidelines](#)

11. Description of research animals

Provide details on animals and/or animal-derived materials used in the study.

No animals were used in this study.

Policy information about [studies involving human research participants](#)

12. Description of human research participants

Describe the covariate-relevant population characteristics of the human research participants.

The study did not involve human research participants.

ChIP-seq Reporting Summary

Form fields will expand as needed. Please do not leave fields blank.

▶ Data deposition

1. For all ChIP-seq data:

- a. Confirm that both raw and final processed data have been deposited in a public database such as [GEO](#).
- b. Confirm that you have deposited or provided access to graph files (e.g. BED files) for the called peaks.

2. Provide all necessary reviewer access links.
The entry may remain private before publication.

The Centromere protein A (CENP-A) ChIP-seq data used in this study have been previously published, GEO Accession: GSE45497 and GSE60951. Enrichment files (k-mer track and bed file are provided as NBT_SupplementaryDara2.txt and NBT_SupplementaryData3.bed

3. Provide a list of all files available in the database submission.

NBT_SupplementaryDara2.txt
 NBT_SupplementaryData3.bed

4. If available, provide a link to an anonymized genome browser session (e.g. [UCSC](#)).

Genome browser session is not available

▶ Methodological details

5. Describe the experimental replicates.

This is not applicable to this study. ChIP seq data used is previously published and described (PMID: 23230266 & 25927077)

6. Describe the sequencing depth for each experiment.

This is not applicable to this study. ChIP seq data used is previously published and described (PMID: 23230266 & 25927077)

7. Describe the antibodies used for the ChIP-seq experiments.

Anti-CENP-A (Abcam cat #Ab13939)
 anti-CENP-C (Abcam cat # 33034); as described in PMID: 23230266 & 25927077.

8. Describe the peak calling parameters.

Enrichment was determined as the ratio of the normalized frequency of ChIP-seq data (Anti-CENP-A (Abcam cat #Ab1393 or CENP-C) relative to input control

9. Describe the methods used to ensure data quality.

This is not applicable to this study. ChIP-seq data used is previously published and described (PMID: 23230266 & 25927077)

10. Describe the software used to collect and analyze the ChIP-seq data.

Jellyfish (version 2.0.0; Marçais & Kingsford(2011)) to determine normalized enrichment values