

Diversity and evolution of centromere repeats in the maize genome

Paul Bilinski · Kevin Distor · Jose Gutierrez-Lopez · Gabriela Mendoza Mendoza · Jinghua Shi · Kelly Dawe · Jeffrey Ross-Ibarra

Received: date / Accepted: date

Abstract Centromere repeats are found in most eukaryotes and play a critical role in kinetochore formation. Though they exhibit considerable diversity both within and among species, little is understood about the mechanisms that drive centromere repeat evolution. Here, we use maize as a model to investigate how a complex history involving polyploidy, fractionation, and recent domestication has impacted the diversity of the maize CentC repeat. We first validate the existence of long tandem arrays of repeats in maize and other taxa in *Zea*. We show that genetic similarity among CentC copies is highest within these arrays, suggesting that tandem duplications are the primary mechanism for the generation of new copies. In spite of this, we see little evidence that CentC variants form distinct genetic groups, instead finding that homoplasious mutations have likely homogenized CentC diversity. We identify Cent4, a repeat initially thought to be specific to the chromosome 4 centromere, as an LTR retrotransposon that has increased drastically in frequency through domestication. Although the two ancestral subgenomes of maize have contributed nearly equal numbers of centromeres, our analysis shows that a vast

Paul Bilinski : Kevin Distor : Jose Gutierrez-Lopez : Gabriela Mendoza Mendoza : Jeffrey Ross-Ibarra
Department of Plant Sciences, UC Davis
Davis, California, 95616

Jeffrey Ross-Ibarra
The Genome Center and Center for Population Biology, University of California
Davis, California, 95616
E-mail: rossibarra@ucdavis.edu

Jinghua Shi : R. Kelly Dawe
Department of Plant Biology, University of Georgia
Athens, Georgia, 30602

R. Kelly Dawe
Department of Genetics, University of Georgia
Athens, Georgia, 30602

majority of all CentC repeats derive from only one of the parental genomes. Finally, by comparing maize with its wild progenitor teosinte, we find that the abundance of CentC has decreased through domestication.

Keywords Centromere · Evolution · Subgenome · sweetscience

Introduction

In spite of the rapid growth of sequenced plant genomes, plant centromeres remain poorly understood and relatively cryptic, due largely to their highly repetitive content. Centromere repeats are highly diverse across taxa and their turnover appears to be very rapid (Melters et al, 2013). However, little is known about the genetic mechanisms that produce centromere repeat diversity. Domesticated maize (*Zea mays* ssp. *mays*) has a high quality genome assembly (Schnable et al, 2009) including complete sequence of two centromeres (Wolfgruber et al, 2009), and the breadth of research into maize centromeres makes it one of the best systems in which to investigate the processes governing centromere repeat evolution.

Maize centromeres are comprised primarily of the 156bp satellite repeat CentC and a family of retrotransposons (CRM), both of which interact with kinetochore proteins such as CENH3 (Wolfgruber et al, 2009; Zhong et al, 2002), and both repeats show considerable variation in local abundance across taxa (Albert et al, 2010). But while there is considerable effort in investigating the molecular function of maize centromere repeats (Ananiev et al, 1998; Nagaki et al, 2003; Wolfgruber et al, 2009), we know comparatively little about the evolution of the sequences themselves. CRM elements are better understood, including the age and insertion preferences of different CRM families (Wolfgruber et al, 2009; Sharma et al, 2008). In contrast, studies (Wang and Bennetzen, 2012) have only examined the flanking sequences to CentC islands despite their known association with functional centromeres. To date, there is no in-depth characterization of the genetic diversity of centromere repeats in the maize genome.

In this paper, we describe the patterns of diversity of centromere repeats across the maize genome. We investigate the genetic diversity of these repeats, including whether the differential ancestry of maize centromeres (Wang and Bennetzen, 2012) has led to chromosome-specific variation as found in other species (Kawabe and Nasuda, 2005; Pontes et al, 2004) and how genetic relatedness among individual repeats varies spatially along chromosomes and across the genome. We then compare centromere abundance across a number of maize lines, including to its wild relatives, the teosintes. We find that CentC copies do not form genetic groups consistent with ancient whole genome duplications or chromosome specificity. Instead, we show higher genetic similarity within clusters, potentially indicating the predominance of tandem duplications in the formation of new CentC copies. Lastly, we use low coverage sequencing and cytological analyses data to show that domesticated maize has less CentC than its wild relatives.

Methods

CentC Repeat Identification and Diversity

We downloaded 218 previously annotated CentC sequences (Ananiev et al, 1998; Nagaki et al, 2003) from Genbank. We then searched the maize genome (5b60, www.maizesequence.org) with megaBLAST (McGinnis and Madden, 2004) using the 218 annotated CentCs as a reference. We kept hits with a length of over 140bp and a minimum bit score of 100. After meeting the bit score threshold, the longest hit was retained. We defined CentC's as being in tandem if the CentC's start location was within 1000bp of the start location of another CentC.

All 12,162 CentC sequences were aligned using 7 iterations of Muscle (Edgar, 2004) with default parameters. A Jukes-Cantor distance matrix of all sequences was calculated with PHYLIP ((Felsenstein, 1989) <http://evolution.genetics.washington.edu/phylip.html>), and an unrooted neighbor joining tree was built based on the distance matrix.

We used principle coordinate analysis (PCoA) to cluster CENTC variants based on their genetic distances. Eigenvalues from the PCoA were used to determine the number of statistically significant clusters using the Tracy-Widom distribution (Patterson et al, 2006).

We employed the software SpaGeDi ((Hardy and Vekemans, 2002) <http://ebe.ulb.ac.be/ebe/Software.html>) to estimate the spatial autocorrelation of sequence similarity of CENTC repeats in the completely sequenced centromeres 2 and 5. We calculated Moran's I statistic using Jukes-Cantor genetic distance and measures of physical distance between CENTC repeats in base pairs. Confidence intervals for the values of I were estimated by 20,000 random permutations of the physical distances.

Statistical analyses were performed in R with the packages ape (Paradis et al, 2004) and RMTstat (Perry et al, 2009). We compared clusters to chromosome of origin and syntenic maps of maize ancient tetraploidy (Schnable et al, 2011) to determine if the genetic history of maize left a footprint on CentC similarity.

Read Mapping and Genome Size Correction

We mapped Illumina reads from a broad panel of *Zea* species (Chia et al, 2012; Tenaillon et al, 2011) to a reference consisting of the full complement of 12,162 CentC variants identified in the B73 genome. We also used previously published whole genome chromatin immunoprecipitation (ChIP) (Wolfruber et al, 2009; Wang et al, 2013) with CenH3. Reads were mapped using Mosaik v1.0 (<https://code.google.com/p/mosaik-aligner/>). We first optimized mapping parameters by relaxing mapping stringency and evaluating the number of successfully mapped reads with each combination. Consistent with parameters from previous studies mapping reads to repetitive elements

(Tenaillon et al, 2011), we required homology to remain at a minimum of 80%. For other non-default parameters, we permuted over many values of hash size, alignment candidate threshold, percent of read aligning, and maximum number of hash positions per seed to find a combination that produced believable alignments. We selected an optimum combination of parameters just below the parameters where we observed a large increase in the total number of reads aligning (Supplementary Figure 1). The parameter boundary at which percentage of reads mapping increased disproportionately was identified, and parameters just below this threshold were selected. Our final set of parameters for tandem repeats used an initial hash size of 8, an alignment candidate threshold of 15 bases, 20% percent of mismatching bases, a minimum of 30% overlap to the reference, and stored the top 100 hits for alignment. After reads were mapped, we calculated the percentage of total reads hitting the given reference and multiplied this value by the relative genome size of each accession as reported in Chia et al (2012) and Tenaillon et al (2011). The total number of reads mapping did not change drastically when using one random copy of CentC versus the full AGPv2 reference, suggesting that our parameters are sufficiently broad to capture genome-wide CentC. Because library preparation has an effect on estimates of repeat abundance (see results), we only used individuals from maize HapMap v2 (Chia et al, 2012) with libraries prepared using identical methods.

We used a different set of mapping parameters for long repeats such as transposable elements. Previous studies (Schnable et al, 2009) estimated that approximately 85% of maize genome derives from transposable elements. Using the short read libraries from Tenaillon et al (2011), we selected parameters so that approximately 85% of the library mapped to the maize transposable element database (maizetdb.org) with a minimum homology of 80%. The final parameters for TEs were a hash size of 10, alignment candidate threshold of 11, 80% homology excluding non-aligned portions of the read, and a 30% minimum overlap.

Furthermore, we wanted to know the accuracy of our pipeline in measuring CentC content and therefore designed a simulation to vary CentC content in an artificial genome (code available at: <https://github.com/kddistor/dnasims>). In short, our simulations altered the copy number of CentC repeats over a given genome size, therefore changing the percentage of the genome deriving from the repeat. Illumina reads were simulated from each of the DNA strings and mapped using our pipeline. Relative differences in abundance were captured well, though our pipeline underestimated overall quantity of CentC. Altering the increments of CentC change, we found that our pipeline could accurately capture differences of 0.05% change in CentC abundance, suggesting that differences greater than 0.05% are likely to be biologically real (Supplementary Figure 4).

Simulation of homoplasious mutations

We wanted to better understand the diversity present in all CentC copies across the genome. Looking at our alignment, we found that the sequences had a pairwise percent identity of 65.8 with only 2.2% of sites identical, suggesting large sequence diversity. However, these statistics do not inform us whether the amount of diversity observed would be expected due to cumulative mutation over time. Therefore, we used simulations in python to count the number of sites with shared diversity across copies of CentC (code available on [GITHUB](#)). Our simulation assumed that CentC has been evolving for the 1 million years since the divergence of maize and *Tripsacum* (Ross-Ibarra et al, 2009), a closely related genus whose centromere repeat shares a large amount of homology (Melters et al, 2013). We assumed a constant copy number, a mutation rate of 3×10^{-8} , and one generation per year.

Sequencing

Library preparation and sequencing was performed according to the methods cited in Melters et al 2012. Using those protocols, we sequenced one individual from *mays*, *mexicana*, *parviglumis*, and *Z. luxurians* with Pacific Biosciences (Pacific Biosciences, Menlo Park, CA) technology. Approximately 200Mbp of reads were produced from each cell, and reads with length greater than 600bp were retained for analysis of tandem CentC content using BLAST (Supplementary Table 2). CentC copies were considered in tandem if the read had 4 CentC copies within 300bp of each other.

FISH

Fluorescent in situ hybridization (FISH) was used to compare abundance of CentC repeats in chromosomes of a maize x *parviglumis* F1 hybrid and a maize x *Z. luxurians* F1 hybrid. FISH protocols closely followed those of X and Y. **Kelly, would you mind helping fill this in?**

Results

Centromere repeats in the maize genome

We found a total of 12,162 CentC variants in the maize reference genome and the unassembled BACs. Of these 12,162 copies, 8,259 were unique over their full length. No CentC occurred more than 10 times in the genome, and the vast majority (>75%, Supplemental Table 1) of non-unique CentC variants occurred only twice. Of the 2,266 non-unique CentC sequences, only 3 were tandem, identical duplicates. Genome-wide CentC locations also show that

nearly all of the 10,639 CentC copies are found in one of the 248 clusters identified on chromosomes 1-10; only 14 occurred as solo copies. Cluster size varied from single CENTC copies to 84KB, with a mean of 7KB (approximately 45 CentC copies). Chromosomes varied greatly in copies of CentC, though we know that centromere assemblies for all of the chromosomes are not complete. For example, in mapping reads from ChIP with CENH3 to an oat plant with a single maize chromosome (OMA) ([whats the ref here? which paper did jiming send us reads from? Gaby?](#)), we find reads from many chromosomes have hits to the unassembled BACs (Supplemental table ??). In particular, chromosome 6 had many more reads aligning to the BACs than it did to its own centromere, suggesting a particularly lacking assembly. Examining total repeat number, Chromosome 7 had the most CentC, with 3,200 copies, while chromosome 6 had the fewest with 32 copies.

We used long-read Pacific Biosciences sequencing to verify that most CentC is in tandem arrays. We sequenced whole genome ($\approx 0.1X$) libraries from 4 *Zea* species. In spite of the low coverage, we recovered reads containing CentC sequence from all four taxa (Supplementary Table 2). In one 6.7KB read from the maize reference line B73, for example, we identified approximately 40 independent CentC copies in tandem, and similar arrays were seen in all four *Zea* species analyzed. These results show that overall structure of the repeats has been maintained for the approximately 140,000 years since the *luxurians-mays* divergence (Hanson et al, 1996; Ross-Ibarra et al, 2009) and that a majority of CentC is found in tandem arrays (Supplementary Table 2).

We then identified how many large clusters of CentC were retained from each of the two parental genomes that comprise the extant maize genome, referred to here as subgenome 1 and subgenome 2 (Figure 1). Previous work identified the parental genome for individual chromosomal segments (Schnable et al, 2011) and centromeres (Wang and Bennetzen, 2012). Because large clusters are less likely to be misassembled, we focused our analyses on the 52 clusters >10KB in length (Supplementary Fig

In addition to CentC, previous studies have described a second high copy number centromere repeat specific to maize chromosome 4 (Page et al, 2001). BLAST analyses of Cent4 sequences, however, revealed that all 15 had high homology to the poorly characterized LTR retrotransposon RLX_sela, which was previously shown to be associated with heterochromatic knobs (Tenaillon et al, 2011; Chia et al, 2012), suggesting that Cent4 is actually a pericentromeric retrotransposon. Whole-genome cenH3 chromatin immuno-precipitation data from Wolfgruber et al (2009) shows no significant over-representation of cent4 compared to five known non-centromeric TE's, suggesting the cent4 repeat is not involved in kinetochore formation on chromosome 4.

Relatedness of CentC in the maize genome.

CentC copies in the maize genome exhibit tremendous diversity: the overall pairwise identity in our alignment was only 65%, and 98% of sites in the

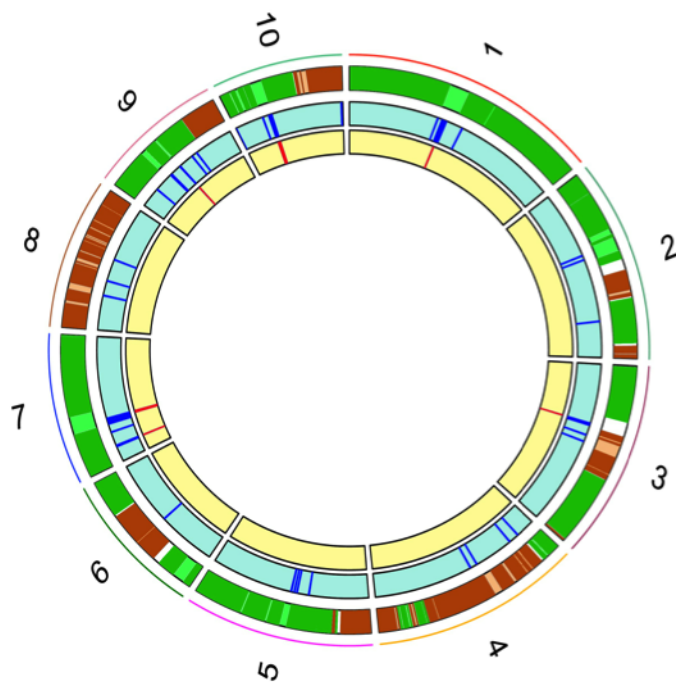


Fig. 1 CentC repeat location in relation to the maize subgenomes. High confidence regions are colored with darker colors while low confidence regions are colored with lighter colors. Breakpoints between the subgenomes remain uncolored to indicate uncertainty. The middle ring, shaded in blue, displays the locations of all CentCs across the genome. The inner ring, shaded in yellow, displays the locations of all CentC clusters greater than 20KB in length.

alignment had at least 2 variants. Such diversity led us to ask whether genetic groups of CentC variants could be distinguished. We performed principle coordinate analyses from a genetic distance matrix estimated from our alignment and assigned individual repeats to genetic clusters following the approach of Patterson et al (2006). We found 58 significant clusters, but observed no pattern of groupings that revealed chromosome specificity of CentC's or the impact of historical tetraploidy (Supplemental Table 4).

The tandem nature of CentC suggests it increases in copy number through local duplications that produce initially identical copies. This predicts that local clusters of CentC should be more closely related than CentC from different clusters. Analysis of genetic and physical distance among CentC repeats on chromosomes 2 and 5 indeed shows this pattern (Figure 2), revealing significant spatial autocorrelation of CentC variants over distances up to 10-50KB (Supplementary Figure 2 and 3).

The decreased genetic distance among CentCs in local clusters on chromosome 2 and 5 suggest that many of the genetic groupings discovered in our genome-wide analysis should correspond to local clusters of repeats. We see no evidence of this, however, as repeats within individual clusters are frequently

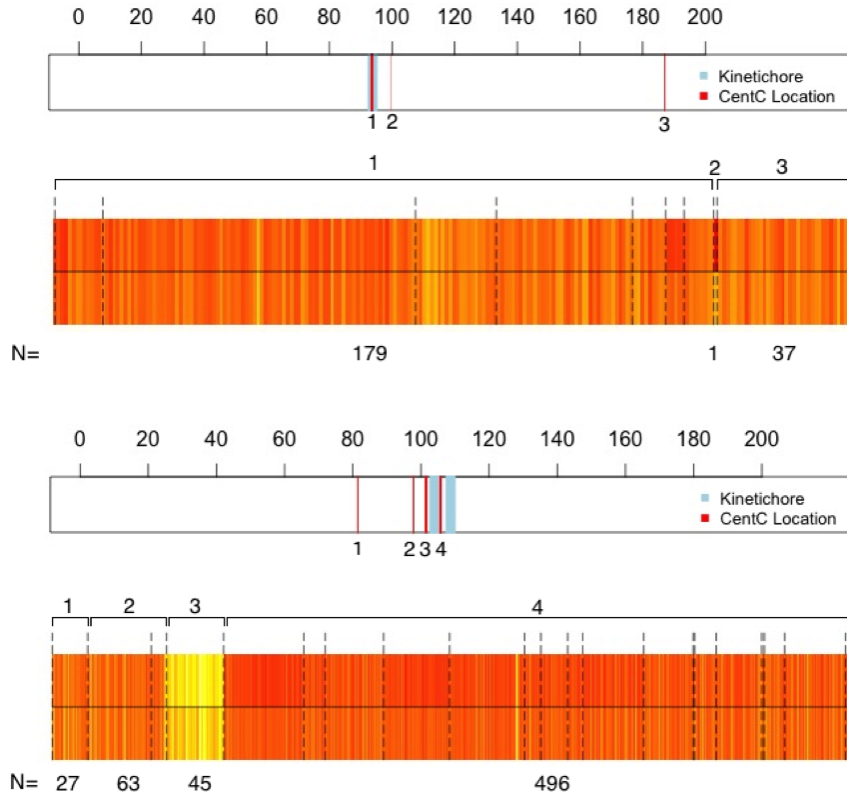


Fig. 2 CentC physical location and genetic relatedness for (a) chromosome 2 and (b) chromosome 5. On the physical map, red lines show locations of numbered CentC clusters and blue blocks show the location of the active kinetochores. Scale bar is in MB. Below each physical map is shown a heatmap of genetic relatedness of each CentC to (top) other copies within its island of tandem repeats delineated by dotted lines and (bottom) all other copies on the chromosome. Darker colors indicate higher relatedness. The total number of CentC in each cluster is shown below the map.

found in different genetic groups as defined by PCoA (Figure 3). A comparison of all pairs of CentC reveals a likely explanation: of the ≈ 74 million possible pairs, approximately 6 million share ≥ 2 mutations different from the genome-wide consensus, likely grouping in genetic clusters despite their physical distance. Looking at several triplets at random from our alignment confirms that two sequences in one PCoA assignment share greater pairwise identity than two sequences adjacent to one another in different PCoA groups. A simple forward simulation (see Methods) suggests this pattern could be due entirely to homoplasy rather than long-distance movement of CentC repeats. By stochastically applying mutations to an initially homogeneous group of repeat sequences, we find that plausible parameter values produce ≈ 10 million pairs of repeats sharing ≥ 2 mutations.

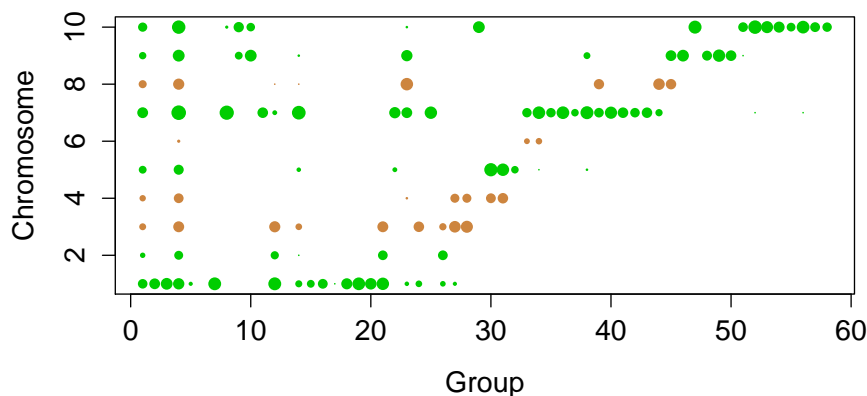


Fig. 3 Presence of CentC in each of the hierarchical groups. The 58 clusters found to be statistically significant in forming genetic groups are represented on the x-axis and chromosome of origin for CentC's on the y-axis. The intersection shows the number of CentC's in the group found on that chromosome, scaled logarithmically. CentC counts from chromosomes whose centromeres were derived from subgenome 1 are colored green and those from subgenome 2 are colored brown.

Variation in CentC abundance in *Zea*

Shotgun sequence data from the maize HapMap v2 (Chia et al, 2012), reveals a significantly greater abundance of CentC in teosinte than in domesticated maize ($p < 0.01$; Figure 4). Teosintes have more CentC than inbred maize. Further support for differences between maize and its wild relatives comes from shotgun sequence data from *Z. luxurians* (Tenaillon et al, 2011). Analysis of these data find nearly twice as much CentC in *Z. luxurians* as the maize inbred B73. To corroborate these results, we performed fluorescent in-situ hybridization of F1 crosses between inbred maize and teosinte to determine if cytological observations agreed with our sequencing findings. Cytology needed to be performed on an F1 cross so that we could compare relative probe fluorescence of the chromosomes within a single individual. FISH data supports our observation that the teosintes *parviglumis* and *Z. luxurians* have more CentC than inbred maize (Figure 4). Using whole genome shotgun PacBio long reads, we further investigated the overall structure of repeats across the different *Zea* species. Percentages of the libraries showing tandem repeats did not differ greatly across the taxa (Supplementary Figure 2).

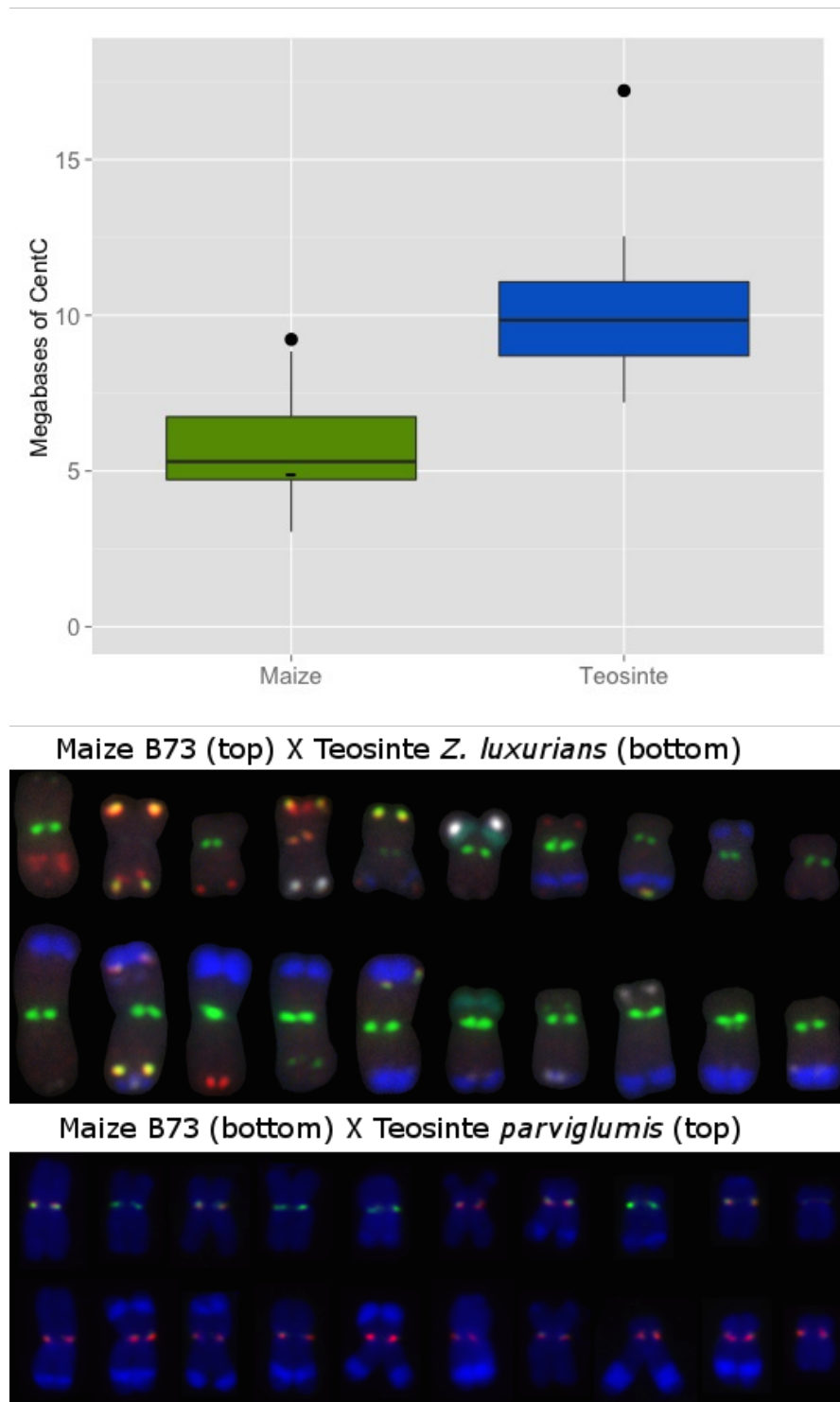


Fig. 4 (a) Mb of CentC in genomic libraries of maize and teosinte. Box plots show data from Chia et al (2012). Points show data for maize inbred B73 and the teosinte *Z. luxurians* from Tenaillon et al (2011). For comparison, the data point of maize inbred B73 in Chia et al (2012) is shown with a tick mark on the box plot. (b) FISH images from an F1 hybrids between maize and the teosinte *parviglumis*, and maize and the teosinte *Z. luxurians*. Upper row shows the karyotype for the teosinte and the lower row shows the maize karyotype. Red=CentC, Green=CRM.

Discussion

Our study traces the changes in centromere repeat genetic relatedness across maize chromosomes to show how ancient tetraploidy and subsequent evolution has impacted centromeres. Most interestingly, we show that most large arrays of CentC in the maize reference genome derive from maize1 subgenome, which is known to have lower gene loss and higher average expression of its genes than maize subgenome 2 Schnable et al (2011). Regulatory microRNA's are known to correspond to centromeric repeats (Reinhart and Bartel, 2002). Therefore, it may be worthwhile to further investigate the potential correlation between centromere retention and gene expression as the phenomenon of higher expression of genes from one ancestor may be very common across plants (*A. thaliana* Cheng et al (2012) and *Gossypium* Kapp and Wendel unpublished). **Two comments about this: first, its a terrible sentence so help make this better, because i think the point is really nice. second, the best citations i found for cotton was an unpublished ref that is still not published from what i can gather. advice? e.g. <http://www.plosone.org/article/info%3Adoi%2F10.1371%2Fjournal.pone.0036442> also check out papers that cite Schabs: http://scholar.google.com/scholar?cites=5313714612108479532&as_sdt=2005&sciodt=0,5&hl=en** There are several possible explanations for the differential contribution of subgenome 1 to the diversity of CentC in the extant maize genome. We put forth several hypotheses that may explain the strong bias for CentC to be located within a block from subgenome 1. Subgenome 2 may have had severely reduced quantities of CentC and therefore never contributed equally. The CentC from subgenome 2 may have decayed rapidly and thus less of it remains. Lastly, the CentC from subgenome 1 may have expanded in copy number since the allopolyploidy event. Though we are unable to conclusively identify the reason, the lack of a large number of identical tandem duplicates and lack of close proximity highest genetic relatedness pairs suggests that any large scale expansion of CentC has not been recent. Also, CentC's in subgenome 2 cluster do not appear to have accumulated excess mutation, suggesting that they are not decaying more rapidly than their subgenome 1 counterparts. We therefore believe that it is most likely that the subgenomes had an unequal contribution of CentC at the formation of the allopolyploid, though more research about the ancient parents is required.

We also explored how genetic relatedness between CentC's correlated with location along the physical map. According to our PCoA analyses, CentC copies across the genome do not form distinct genetic groups correlating with past origin (Figure 3). The lack of subgenome or chromosome grouping in modern CentC suggests a little differentiation between the repeats since the ancestral divergence. Given the lack of a genome wide pattern, we also wanted to investigate genetic relatedness on individual chromosomes. We hypothesized that most copies of CentC arose from local duplication rather than transposition, and therefore the genetic distance between CentC's would be lowest across CentC's within a large tandem array. **This part is ugly, and needs help** From the reference genome, we chose to investigate chromosomes 2 and 5,

since they have been sequenced from end to end, allowing for inferences about the relationship between diversity and physical proximity (Wolfgruber et al, 2009). Furthermore, the role of the repeat arrays on chromosomes 2 and 5 appear very different, as the largest array on chromosome 2 interacts with the kinetochore, while no array on chromosome 5 does (Wolfgruber et al, 2009). We find that repeats on chromosomes 2 and 5 are most highly genetically related to neighboring copies (Figure 2). This relationship was recapitulated in analyses using SpaGeDi, where CentC's within approximately 10-50KB of one another were more genetically similar (Supplemental Data 2 and 3). Though we did not investigate the relationship between CentC location and genetic similarity on the other chromosomes due to incomplete sequencing of centromeres, we observe that many CentC's within an array fall into the same significant grouping in our Tracy-Widom clustering analysis, suggesting that high local similarity of CentC's is a genome-wide phenomenon. This local relatedness suggests that most CentC is evolving through tandem duplication and not a long distance mechanism such as transposition that had been previously suggested (Shi et al, 2010). Concerning chromosomes 2 and 5, the higher local relatedness of all CentC clusters regardless of presence within the kinetochore suggests that CentC interaction with kinetochore proteins does not have a detectable change on their rates of evolution.

When investigating why a pair of chromosomes on two different chromosomes are each other closest genetic relative, we revealed that homoplasmy in mutations is common across CentC variants. We suggest that copies of CentC are sufficiently old within the genome that homoplasious mutations cause physically distant CentC's to be highly genetically related. Importantly, we also do not see PCoA group capturing full clusters across chromosomes in a way that would be consistent with retrotransposition. We speculate that a vast majority of the CentC's exist as a result of very old duplications meaning that mutations have had a long time to accumulate. Roughly 80% of the CentC repeats have their closest genetic relative on the same chromosome, an observation we would expect if CentC's on a chromosome share ancestry. However, only 14% of closest genetic pairs are found within 10KB of each other, suggesting that their tandem duplication is old and that most CentC's have persisted within the genome for a long time. **This is an important sentence, and I can't seem to get it to communicate my point properly. perhaps because im making too many points**

When studying CentC changes through domestication, our findings show that the repeat has experienced a decrease in copy number over time without a noticeable change in the structure of the repeat arrays. Using PacBio long read sequencing, we confirm genome-wide the observation we see on chromosomes 2 and 5, that most copies of CentC exist in large tandem arrays in all *Zea* taxa. From short read sequencing data, we show that modern maize has reduced genomic abundance of CentC when compared to the teosintes, a finding confirmed through FISH (Figure 4). Both teosintes within the species *mays* and its sister species *luxurians* have elevated levels of CentC. Lower levels of centromere repeats in inbred maize contrasts previous studies that

characterized the changing abundance of other common repeats. For example, the abundance of most transposable element families increased after domestication (Chia et al, 2012). Knowing that TE abundance increases in domesticated maize, we might have expected CentC content to increase as well if centromere size had to expand alongside most other repetitive content. Alternatively, due to their structural role in kinetichore formation, we might have also expected centromere repeats to be largely excluded from the genome wide fluctuations in repetitive content assuming that selection exists to maintain centromere size. Instead, the decrease in CentC content may indicate an active process of removing CentC. We hypothesize that the removal of CentC content may correlate with genome size, as the sharp decrease in knob content through domestication actually led to an overall smaller genome size in inbred maize. The correlation between functional centromere size and genome size has been observed in grass species (Zhang and Dawe, 2012). Further investigation would be required to discover whether only those copies of CentC outside of the active kinetochore are being deleted.

We also sought to further characterize the chromosome specific repeat cent4 (Page et al, 2001). Previous work showed that Cent4 probes lagged behind CentC probes in cell division (Jiang et al, 2002), suggesting that the repeat is not involved in the active kinetochore and rather located in the pericentromere. A lack of enrichment of cent4 in ChIP data agrees with a pericentromeric location. Using the published cent4 sequences, we identify the cent4 repeat as the LTR retrotransposon RLX_sela in the transposable element database (Baumcom et al, 2009). Characterization of the TE consensus sequence shows that it contains repetitive motifs with homology to both knob and telomere repeats. It lacks any of the protein sequences necessary for autonomous transposition, such as GAG and POL complexes. Previous work in rice has documented the presence of nonautonomous LTR retrotransposons in or near the centromere (Jiang et al, 2002). However, the RLX_sela sequence also appears to be missing some of the necessary primer binding sites that would distinguish it as a nonautonomous TE, suggesting that it may be a tandem repeat, and further sequencing of the region will be required to uncover the role of cent4. Given that the centromere and pericentromere on chromosome 4 show signs of strong selection during domestication (Hufford et al, 2012), it is plausible that the increase in abundance of Cent4 seen in domesticated maize is due to the effects of linked selection on a locus important for domestication rather than novel insertions or duplications.

In conclusion, our study pairs bioinformatics and cytology to validate observations of decreasing centromere repeat content through maize domestication. We show that patterns of CentC similarity are consistent with tandem duplications and maize's allopolyploid history has left no significant differentiation between CentC's from the different maize subgenomes. We also identify the chromosome 4 specific repeat, cent4, as the TE RLX_sela and show that it has increased through domestication, unlike a majority of the other TEs (Chia et al, 2012).

Acknowledgements We wish to thank Pacific Biosciences for sequencing resources. We thank Vince Buffalo, Lauren Sagara, Anne Lorant, Michelle Stitzer, Gernot Presting, NSF summer exchange program interns Cesar Alvarez Mejia, Aurelio Hernandez Bautista, and Siddharth Bhadra-Lobo for helpful discussion.

US-NSF grant IOS-0922703

References

- Albert P, Gao Z, Danilova T, Birchler J (2010) Diversity of chromosomal karyotypes in maize and its relatives. *Cytogenet Genome Res* 129(1-3):6–16
- Ananiev EV, Phillips RL, Rines HW (1998) Chromosome-specific molecular organization of maize (*zea mays* l.) centromeric regions. *Proceedings of the National Academy of Sciences* 95(22):13,073–13,078
- Baucom RS, Estill JC, Chaparro C, Upshaw N, Jogi A, Deragon JM, Westerman RP, SanMiguel PJ, Bennetzen JL (2009) Exceptional diversity, non-random distribution, and rapid evolution of retroelements in the b73 maize genome. *PLoS Genet* 5(11):e1000732
- Cheng F, Wu J, Fang L, Sun S, Liu B, Lin K, Bonnema G, Wang X (2012) Biased gene fractionation and dominant gene expression among the subgenomes of *Brassica rapa*. *PLoS ONE* 7(5):e36442
- Chia JM, Song C, Bradbury PJ, Costich D, de N Leon, Doebley J, Elshire RJ, Gaut B, Geller L, Glaubitz JC (2012) Maize hapmap2 identifies extant variation from a genome in flux. *Nature genetics* 44(7):803–807
- Edgar RC (2004) Muscle: multiple sequence alignment with high accuracy and high throughput. *Nucleic Acids Research* 32(5):1792–1797
- Felsenstein J (1989) Phylip - phylogeny inference package (version 3.2). *Cladistics* 5(2):164–166
- Hanson MA, Gaut BS, Stec AO, Fuerstenberg SI, Goodman MM, Coe EH, Doebley JF (1996) Evolution of anthocyanin biosynthesis in maize kernels: The role of regulatory and enzymatic loci. *Genetics* 143(3):1395–1407
- Hardy OJ, Vekemans X (2002) spagedi: a versatile computer program to analyse spatial genetic structure at the individual or population levels. *Molecular Ecology Notes* 2(4):618–620
- Hufford M, Xu X, van Heerwaarden J, Pyhajarvi T, Chia J, Cartwright R, Elshire R, Glaubitz J, Guill K, Kaeppler S, Lai J, Morrell P, Shannon L, Song C, Springer N, Swanson-Wagner R, Tiffin P, Wang J, Zhang G, Doebley J, McMullen M, Ware D, Buckler E, Yang S, Ross-Ibarra J (2012) Comparative population genomics of maize domestication and improvement. *Nat Genet* 44:808–811
- Jiang N, Bao Z, Temnykh S, Cheng Z, Jiang J, Wing RA, McCouch SR, Wessler SR (2002) Dasheng: a recently amplified nonautonomous long terminal repeat element that is a major component of pericentromeric regions in rice. *Genetics* 161(3):1293–1305

- Kawabe A, Nasuda S (2005) Structure and genomic organization of centromeric repeats in arabidopsis species. *Mol Genet Genomics Molecular Genetics and Genomics* 272(6):593–602
- McGinnis S, Madden TL (2004) Blast: at the core of a powerful and diverse set of sequence analysis tools. *Nucleic Acids Research* 32(suppl 2):W20–W25
- Melters D, Bradnam K, Young H, Telis N, May M, Ruby J, Sebra R, Peluso P, Eid J, Rank D, Garcia J, DeRisi J, Smith T, Tobias C, Ross-Ibarra J, Korf I, Chan S (2013) Comparative analysis of tandem repeats from hundreds of species reveals unique insights into centromere evolution. *Genome Biology* 14(1):R10
- Nagaki K, Song J, Stupar RM, Parokonny AS, Yuan Q, Ouyang S, Liu J, Hsiao J, Jones KM, Dawe RK, Buell CR, Jiang J (2003) Molecular and cytological analyses of large tracks of centromeric dna reveal the structure and evolutionary dynamics of maize centromeres. *Genetics* 163(2):759–770
- Page BT, Wanous MK, Birchler JA (2001) Characterization of a maize chromosome 4 centromeric sequence: evidence for an evolutionary relationship with the b chromosome centromere. *Genetics* 159(1):291–302
- Paradis E, Claude J, Strimmer K (2004) Ape: Analyses of phylogenetics and evolution in r language. *Bioinformatics* 20(2):289–290
- Patterson N, Price AL, Reich D (2006) Population structure and eigenanalysis. *PLoS Genet* PLoS Genet 2(12):e190
- Perry P, Johnstone I, Ma Z, Shahram M (2009) Rmtstat: Distributions and statistics from random matrix theory. 2009. R software package version 01
- Pontes O, Neves N, Silva M, Lewis MS, Madlung A, Comai L, Viegas W, Pikaard CS (2004) Chromosomal locus rearrangements are a rapid response to formation of the allotetraploid arabidopsis suecica genome. *Proceedings of the National Academy of Sciences* 101(52):18,240–18,245
- Reinhart BJ, Bartel DP (2002) Small rnas correspond to centromere heterochromatic repeats. *Science* 297(5588):1831–1831
- Ross-Ibarra J, Tenaillon M, Gaut BS (2009) Historical divergence and gene flow in the genus *zea*. *Genetics* 181(4):1399–1413
- Schnable JC, Springer NM, Freeling M (2011) Differentiation of the maize subgenomes by genome dominance and both ancient and ongoing gene loss. *Proceedings of the National Academy of Sciences* 108(10):4069–4074
- Schnable PS, Ware D, Fulton RS, Stein JC, Wei F, Pasternak S, Liang C, Zhang J, Fulton L, Graves TA (2009) The b73 maize genome: complexity, diversity, and dynamics. *science* 326(5956):1112–1115
- Sharma A, Schneider KL, Presting GG (2008) Sustained retrotransposition is mediated by nucleotide deletions and interelement recombinations. *Proceedings of the National Academy of Sciences* 105(40):15,470–15,474
- Shi J, Wolf SE, Burke JM, Presting GG, Ross-Ibarra J, Dawe RK (2010) Widespread gene conversion in centromere cores. *PLoS biology* 8(3):e1000327
- Tenaillon MI, Hufford MB, Gaut BS, Ross-Ibarra J (2011) Genome size and transposable element content as determined by high-throughput sequencing in maize and *zea luxurians*. *Genome biology and evolution* 3:219

- Wang H, Bennetzen JL (2012) Centromere retention and loss during the descent of maize from a tetraploid ancestor. *Proceedings of the National Academy of Sciences* 109(51):21,004–21,009
- Wang K, Wu Y, Zhang W, Dawe RK, Jiang J (2013) Maize centromeres expand and adopt a uniform size in the genetic background of oat. *Genome research* pp gr-160,887
- Wolfgruber TK, Sharma A, Schneider KL, Albert PS, Koo DH, Shi J, Gao Z, Han F, Lee H, Xu R (2009) Maize centromere structure and evolution: sequence analysis of centromeres 2 and 5 reveals dynamic loci shaped primarily by retrotransposons. *PLoS genetics* 5(11):e1000,743
- Zhang H, Dawe RK (2012) Total centromere size and genome size are strongly correlated in ten grass species. *Chromosome Research* 20(4):403–412
- 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. *The Plant Cell Online* 14(11):2825–2836

Supplemental Material

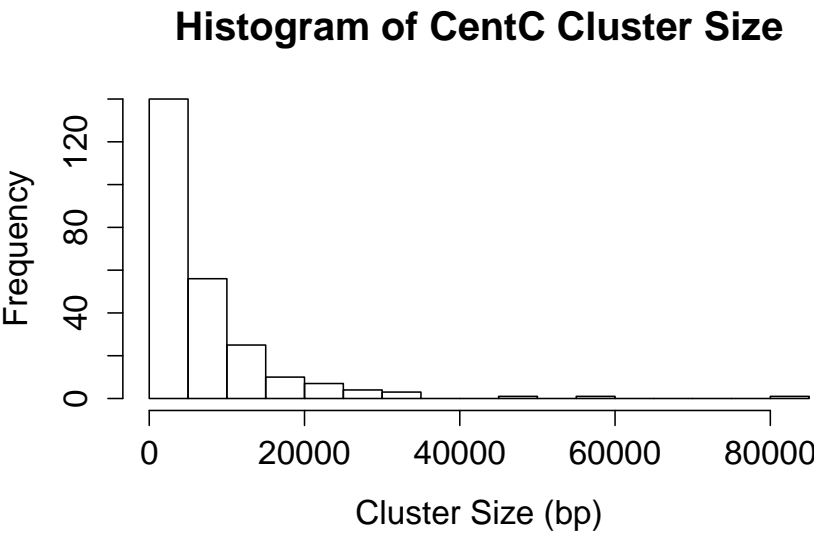


Fig. 1 CentC cluster size across all chromosomes.

Table 1 CentC Occurence Count In Maize RefGenv2

Occurrences	Number of CentC's
1	8259
2	1233
3	263
4	89
5	31
6	13
7	7
8	0
9	0
10	1

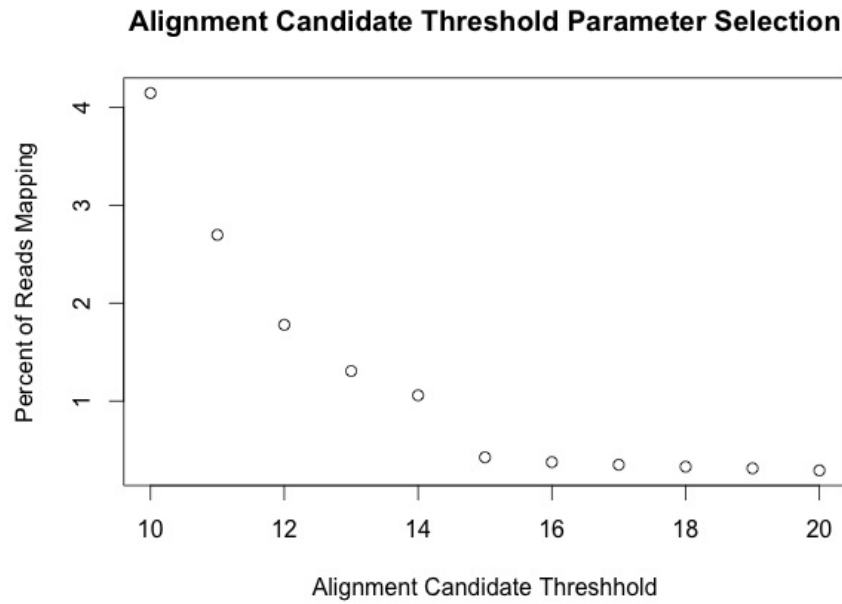


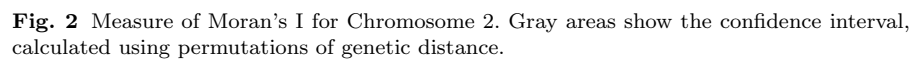
Fig. 1 Parameter selection for Alignment Candidate Threshold (ACT) for Mosaik. All other parameters were kept constant while ACT was changed. ACT was the parameter for which a non-linear pattern was observed. We selected to use an ACT of 15, the value for which we observed the greatest relative decrease between total percent mapping values. The sharp change suggests that, at a lower ACT, we may be mapping a non-CentC element to our reference.

Table 2 PacBio Read Counts and Tandem CentC

Maize Line	Reads over 600bp	Reads with ≥ 4 CentC	% Reads Showing Tandem CentC
B73	237995	30	0.030252736
<i>luxurians</i>	156964	79	0.050330012
<i>mexicana</i>	141939	150	0.1056792
<i>parviglumis</i>	227050	89	0.039198414

Table 3 ChIP Reads mapping to Unassembled from different Oat-Maize Addition (OMA) Lines

File Key	Maize Chr	Percent Reads Aligning to Unassembled BACs
JJ1BU (OMA 6.34)	6	21.35
JJ1BR (OMA 1.36)	1	15.56
JJ1CF (OMA 9.41)	9	6.16
JJ1CH (OMA 8.05)	8	6.69
JJ1CG (OMA 10.26)	10	9.21
JJ1CI (OMA 8.05)	8	8.5

[illegible]

[illegible]

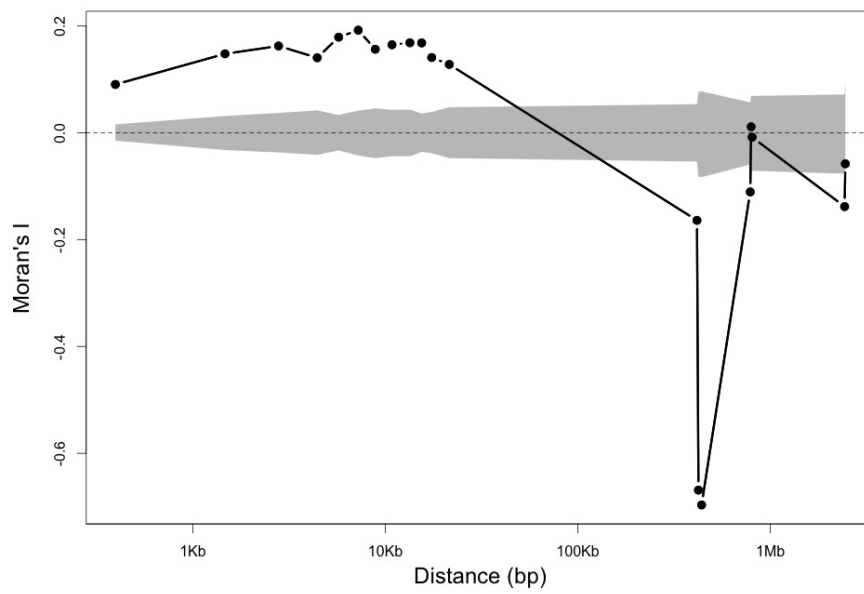


Fig. 3 Measure of Moran's I for Chromosome 5. Gray areas show the confidence interval, calculated using permutations of genetic distance.

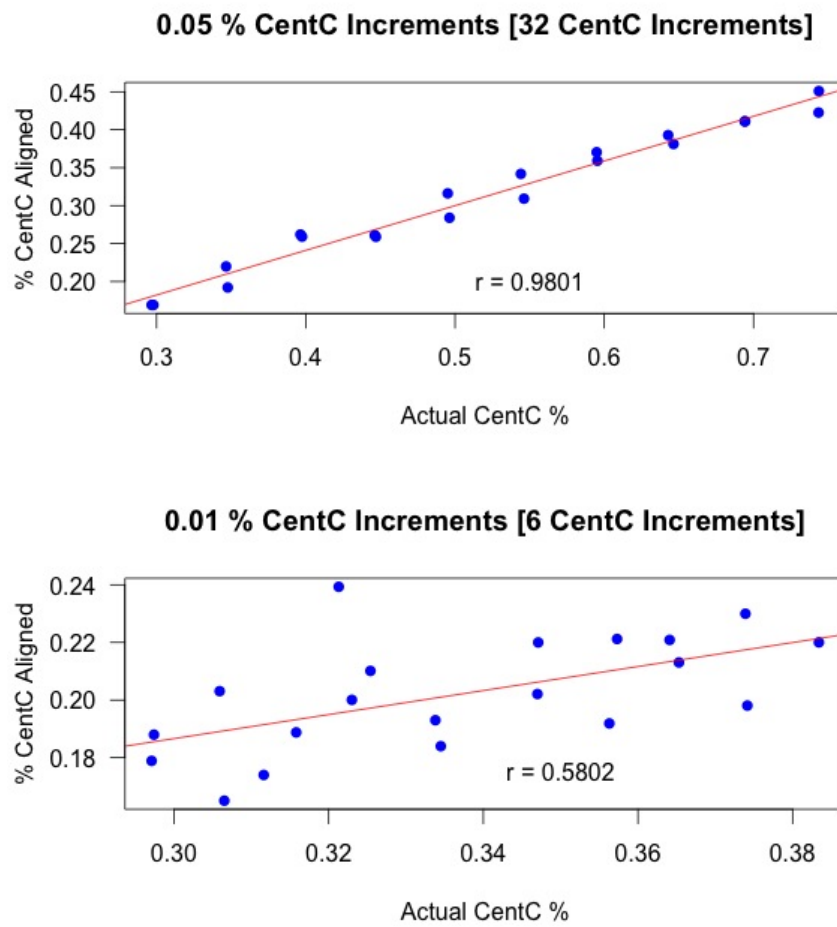


Fig. 4 Graphs showing our ability to capture changes in CentC repeat abundance under constant genome size. We simulated 10MB of DNA with varying CentC content and simulated Illumina reads from the DNA. Reads were mapped with our Mosaik pipeline, and several simulations at each percentage of genomic content were performed.