

## Research

# Chickens possess centromeres with both extended tandem repeats and short non-tandem-repetitive sequences

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The centromere is essential for faithful chromosome segregation by providing the site for kinetochore assembly. Although the role of the centromere is conserved throughout evolution, the DNA sequences associated with centromere regions are highly divergent among species and it remains to be determined how centromere DNA directs kinetochore formation. Despite the active use of chicken DT40 cells in studies of chromosome segregation, the sequence of the chicken centromere was unclear. Here, we performed a comprehensive analysis of chicken centromere DNA which revealed unique features of chicken centromeres compared with previously studied vertebrates. Centromere DNA sequences from the chicken macrochromosomes, with the exception of chromosome 5, contain chromosome-specific homogenous tandem repetitive arrays that span several hundred kilobases. In contrast, the centromeres of chromosomes 5, 27, and Z do not contain tandem repetitive sequences and span non-tandem-repetitive sequences of only ~30 kb. To test the function of these centromere sequences, we conditionally removed the centromere from the Z chromosome using genetic engineering and have shown that the non-tandem-repeat sequence of chromosome Z is a functional centromere.

[Supplemental material is available online at <http://www.genome.org>. The sequence data from this study have been submitted to GenBank (<http://www.ncbi.nlm.nih.gov/Genbank>) under accession nos. AB556430–AB556736.]

The contribution of DNA sequences to centromere formation remains controversial (Marshall et al. 2008). While there exist large arrays of repetitive sequences in human centromere regions (alphoid sequences; alpha satellite DNA) that contribute to the efficient construction of human artificial chromosomes (Harrington et al. 1997; Ikeno et al. 1998; Ohzeki et al. 2002), analysis of human neocentromeres suggests that these repetitive sequences are not absolutely essential for centromere formation (Marshall et al. 2008). To understand the molecular basis of centromere formation, it is necessary to define both the protein complexes that associate with these regions and their underlying centromere DNA sequences. We have previously isolated and analyzed multiple centromere-localized proteins from chicken DT40 cells (Okada et al. 2006; Fukagawa 2004, 2008; Hori et al. 2008; Amano et al. 2009). Due to high rates of homologous recombination, DT40 cells provide the unique ability to conduct genetic modifications to analyze the consequences of specific changes to centromere DNA (CenDNA) sequences to kinetochore formation. In addition, while mammalian centromere sequences have been defined, the analysis of centromeres in a distant vertebrate would provide an important evolutionary view of CenDNA. While previous studies have analyzed the chicken whole genome, sequence information from the chicken centromere

was not defined (International Chicken Genome Sequencing Consortium 2004). Here, we report the comprehensive analysis of centromere DNA sequences in chicken DT40 cells.

## Results

### Identification of chicken centromeric DNA

To identify and characterize CenDNA from chickens, we used the centromere-specific histone H3 variant CENPA (Palmer and Margolis 1987). CENPA is a defined marker for active centromeres including neocentromeres (Marshall et al. 2008), and thus DNA associated with CENPA represents bona fide functional CenDNA. To characterize chicken CenDNAs, we created a DT40 cell line stably expressing CENPA-Flag. We next performed chromatin immunoprecipitation (ChIP) experiments in the CENPA-Flag DT40 cell line with anti-Flag antibodies and isolated and cloned the DNAs from the immunoprecipitates. In total, we sequenced 292 clones (CAIP clones, Supplemental Table S1). To confirm that these DNA clones represent centromere sequences, we performed FISH analysis for ~100 of these clones to examine the centromere localization of these sequences (Fig. 1; Supplemental Table S1).

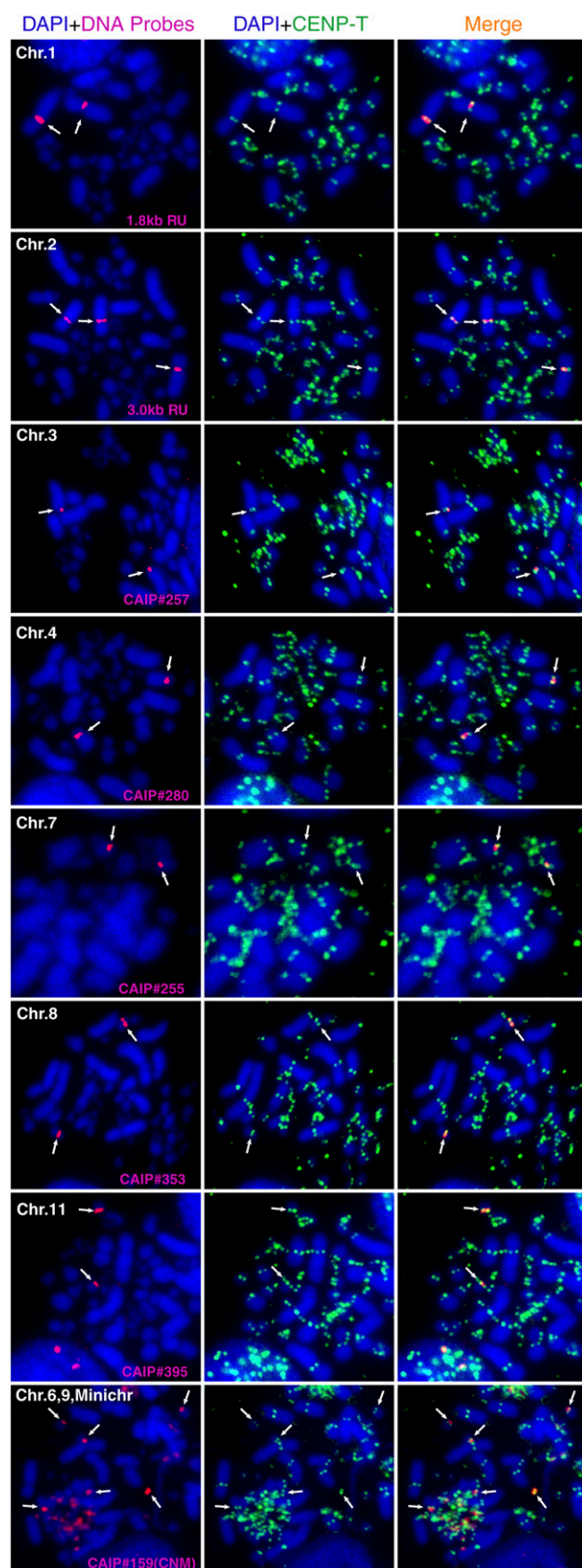
Chicken cells contain 10 pairs of macrochromosomes (chromosomes 1–10), 28 pairs of small chromosomes called microchromosomes, and Z/W sex chromosomes (Masabanda et al. 2004). Previous reports identified a 42-bp tandem repeat (CNM sequence) that localizes to centromeres from some of microchromosomes and on chromosomes 6 and 9 (Matzke et al. 1990;

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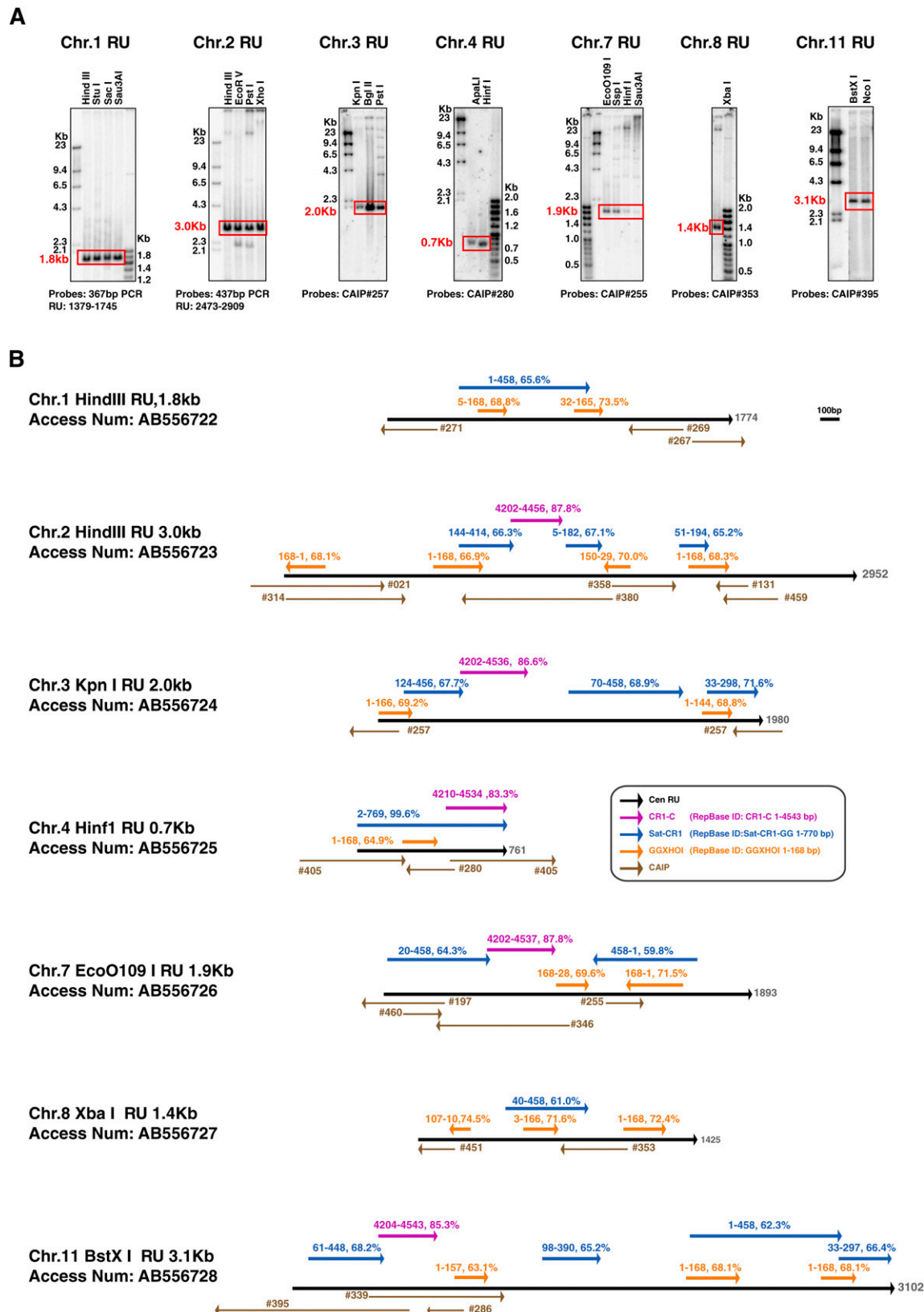


Wang et al. 2002; Krasikova et al. 2006). Consistent with these reports, we found CNM sequences in 15 of our CENPA-associated DNA clones (Supplemental Table S1). FISH analysis with a clone containing the CNM sequence confirmed its localization to a subset of microchromosomes and to chromosomes 6 and 9 (Fig. 1). Although the CenDNA sequences of the other macrochromosomes were previously unclear, our FISH analysis revealed distinct centromere sequences for chromosomes 1, 2, 3, 4, 7, 8, and 11 (Fig. 1). Centromere sequence of chromosome 5 will be described later. Unfortunately, we have not identified a specific sequence for chromosome 10 among the 292 clones. Interestingly, each CenDNA from chromosomes 1, 2, 3, 4, 7, 8, and 11 has a specific sequence and we did not detect cross hybridization with other centromeres (see Methods for hybridization condition). For each of these macrochromosomes, we isolated approximately three independent clones. As chicken have 39 chromosome pairs, we calculate that  $\sim 117$  ( $39 \times 3$ ) of the isolated clones were from centromere, suggesting that centromere DNA was strongly enriched in the CENPA ChIP-based cloning experiment (see also Supplemental Fig. S1C).

We next analyzed the detailed sequence of each chicken CenDNA. First, genomic DNA from DT40 cells was digested with several restriction enzymes and Southern hybridization analysis was performed with each CenDNA as a probe to determine whether it contained repetitive elements. As shown in Figure 2A, centromeres from several chromosomes have distinct repeated units with chromosomes 1, 2, 3, 4, 7, 8, and 11 having 1.8-, 3.0-, 2.0-, 0.7-, 1.9-, 1.4-, and 3.2-kb repeats, respectively. The sequences from each repeat-unit have been deposited in the DDBJ/EMBL/GenBank DNA database, and the corresponding accession numbers are shown in Figure 2. While global homology between repeat-unit sequences is  $\sim 50\%$ , we found common features in repeat-unit sequences. An important feature of these repeat sequences is that all these sequences contain a homology region ( $>60\%$  homology) with Satellite CR-1 sequences (Fig. 2B). In addition, a limited region of the CR1-C retrotransposon (4200–4500 bp) shows  $>80\%$  homology with the repeat-unit sequences of Cen2, Cen3, Cen4, Cen7, and Cen11 (Fig. 2B), suggesting that Cen2, Cen3, Cen4, Cen7, and Cen11 are at least partially derived from a common ancestral sequence.

The repeat-unit from chromosome 8 is reported as a PIR repeat (Wang et al. 2002; Li et al. 2007) which contains a 21-bp consensus sequence in which clusters of (A)3-5 and (T)3-5 are separated by 5–7 bp (for a consensus sequence of gaaataccncttttctcc). As the CNM repeat also contains a sequence related to this 21-bp PIR repeat, we examined whether CenDNAs identified in macrochromosomes show homology with this 21-bp sequence (RepbaseID: GGXHOI). We found that all repeat sequences have regions with homology with this GGXHOI 21-bp sequence ( $>65\%$  homology) (Fig. 2B). We also found the other 21-bp satellite sequence PO41 (Wicker et al. 2005) in CAIP clones (CAIP #217, #258, #381; Supplemental Table S1), but these clones did not localize to centromeres of macrochromosomes by FISH analysis, while signals were detected in mini chromosomes (Supplemental Table S1). However, it is very hard to conclude whether PO41 really colocalizes to

**Figure 1.** Identification of centromere DNA in chicken. Using DNAs precipitated with CENPA, FISH analysis was performed. Probes are indicated in each panel. FISH signals are shown in red. Centromeres are stained with anti-CENPT antibodies (green). CenDNAs of chromosomes 1, 2, 3, 4, 7, 8, and 11 are identified. These sequences are chromosome-specific. The clone containing CNM repeats hybridized with microchromosomes and chromosomes 6 and 9.



**Figure 2.** DNA sequence of centromere of each macro-chromosome is specific. (A) Using DNA clones, which hybridized with centromere region by FISH, DNA sequence of each repeat-unit was characterized. Each DNA sequence was deposited in DDBJ/EMBL/GenBank DNA database. Accession numbers of Cen1, Cen2, Cen3, Cen4, Cen7, Cen8, and Cen11 are AB556722, AB556723, AB556724, AB556725, AB556726, AB556727, and AB556728, respectively. Length of each repeat-unit was determined by Southern hybridization. Probes are indicated in each panel. (B) Homology region in each repeat-unit sequence with known interspersed repeats is shown. Percentage of identity and position of repeats are also shown. Sequences of interspersed repeats are from RepBase. RepBase IDs of CR1-C, Satellite CR1, and GGXHOI sequence are CR1-C, Sat-CR1-GG, and GGXHOI, respectively. Li and Leung (2006) have previously shown that Cen4 contains a CR-1 repeat.



centromere on mini-chromosomes due to limit of resolution of the FISH experiments.

In total, the Satellite CR-1/CR1-C/GGXHOI motifs mentioned above cover 41% (Cen1), 47% (Cen2), 82% (Cen3), 100% (Cen4), 81% (Cen7), 50% (Cen8), and 66% (Cen11) of the repeat-units (Fig. 2B). These data suggest that all the centromere repeat-unit sequences may originate from one or two progenitor sequences through cycles of interchromosomal transfer, divergence, rearrangement, and amplification, as has been suggested for human alpha satellite DNA (Shepelev et al. 2009).

### Long-range organization of centromeres of chicken chromosomes 1 and 2

Our results demonstrate that centromeres from most chicken macrochromosomes contain tandem repetitive DNA sequences. To examine the long-range organization of these centromere regions, we focused on the centromeres from chromosomes 1 and 2 (Cen1 and Cen2) and isolated bacteria artificial chromosome (BAC) clones (Hori et al. 2000) that cover these regions (Fig. 3). We characterized BAC clones that contain both repetitive and flanking unique sequences (Fig. 3A,D) and defined the exact boundary between these sequences (Fig. 3; accession nos. AB556732–AB556736). Each human centromere contains a different homogeneous higher-order repeated unit (HOR) with a length ranging from 0.5 to 6.5 kb (Alexandrov et al. 2001; Shepelev et al. 2009). The kinetochore is formed at the HOR region in the long alphoid array (Schueler et al. 2001; Fukagawa 2004; Shepelev et al. 2009). In the case of chicken Cen1 and Cen2, we found that repeat-unit sequences isolated from BAC clones show high similarity (~99%) with each other, suggesting regions containing high homology repeat-units from Cen1 and Cen2 correspond to the HOR region of human alphoid sequences and form the active centromere. In human centromere, there is a region of heterochromatin that contains divergent alphoid sequences. These adjacent divergent regions in human centromere are inactive centromeres of primate ancestors and have a clearly distinct origin from HOR sequences of alphoid DNA (Kazakov et al. 2003; Schueler and Sullivan 2006; Shepelev et al. 2009). In contrast, we did not find any evidence for such “old” centromere sequence near repeat-unit sequences on chicken Cen1 and Cen2.

To determine the exact length of the entire repetitive arrays, we performed pulsed-field gel electrophoresis (PFGE) analysis with DT40 genomic DNA. We identified a length polymorphism for Cen1 with one homolog containing a repetitive array of 350 kb and the other 450 kb. These correspond to ~190 and ~250 copies of the Cen1 repeat sequence (Fig. 3B,C). The length of the repetitive array for Cen2 is 490 kb corresponding to ~160 copies of the Cen2 repeat sequence (Fig. 3E,F). The long-range organization around the centromere region of chicken chromosomes 1 and 2 is summarized in Figure 3C,F, respectively. Both Cen1 and Cen2 contain several hundred kilobases of repetitive arrays. However, based on our studies of the other chicken centromeres described below, we do not believe that the kinetochore is formed along the entire region of these long repetitive arrays.

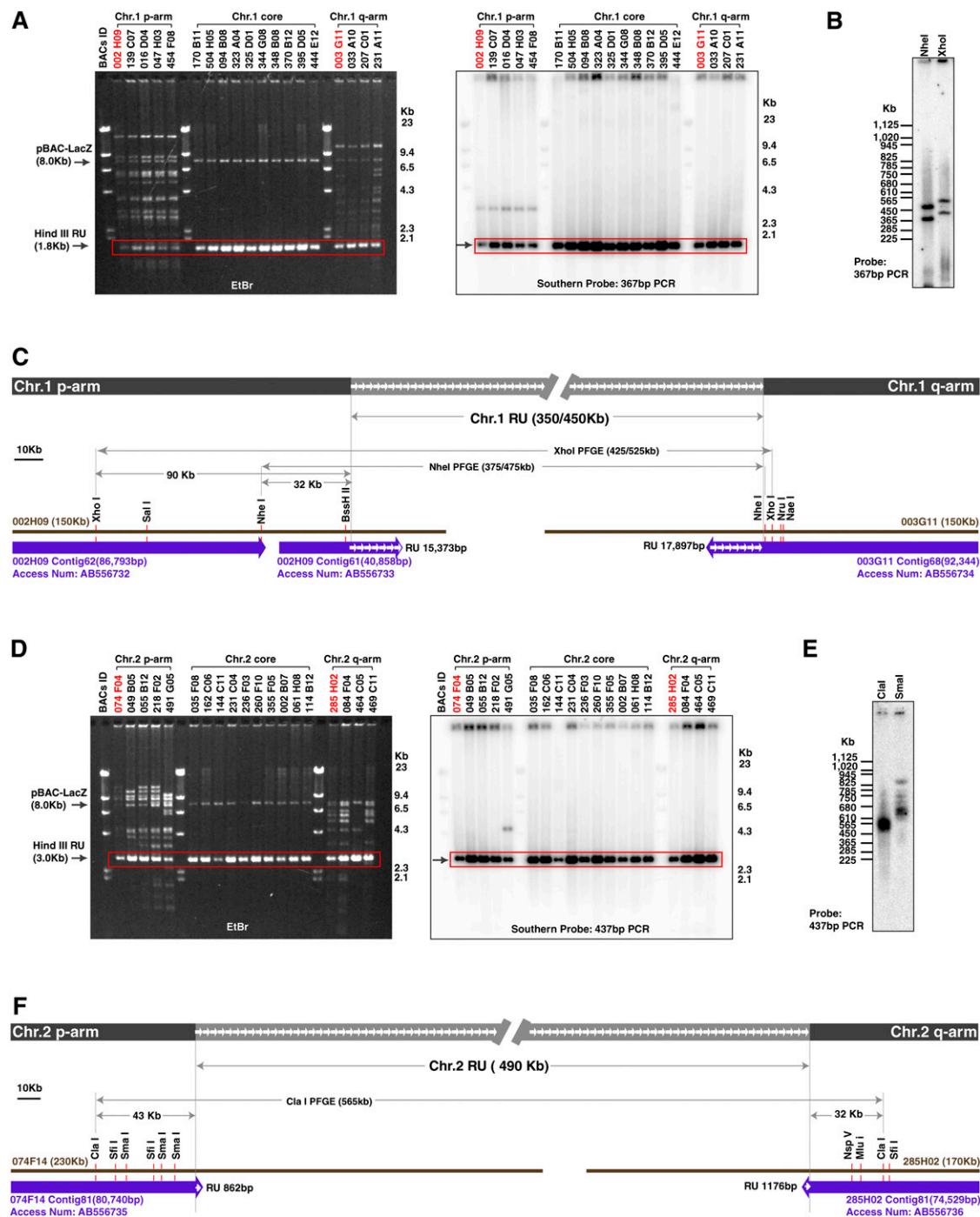
### Centromeres have non-tandem-repetitive sequences in several chicken chromosomes

Although we identified CenDNAs for most of the chicken chromosomes, sequence information for chromosomes 5 and Z was

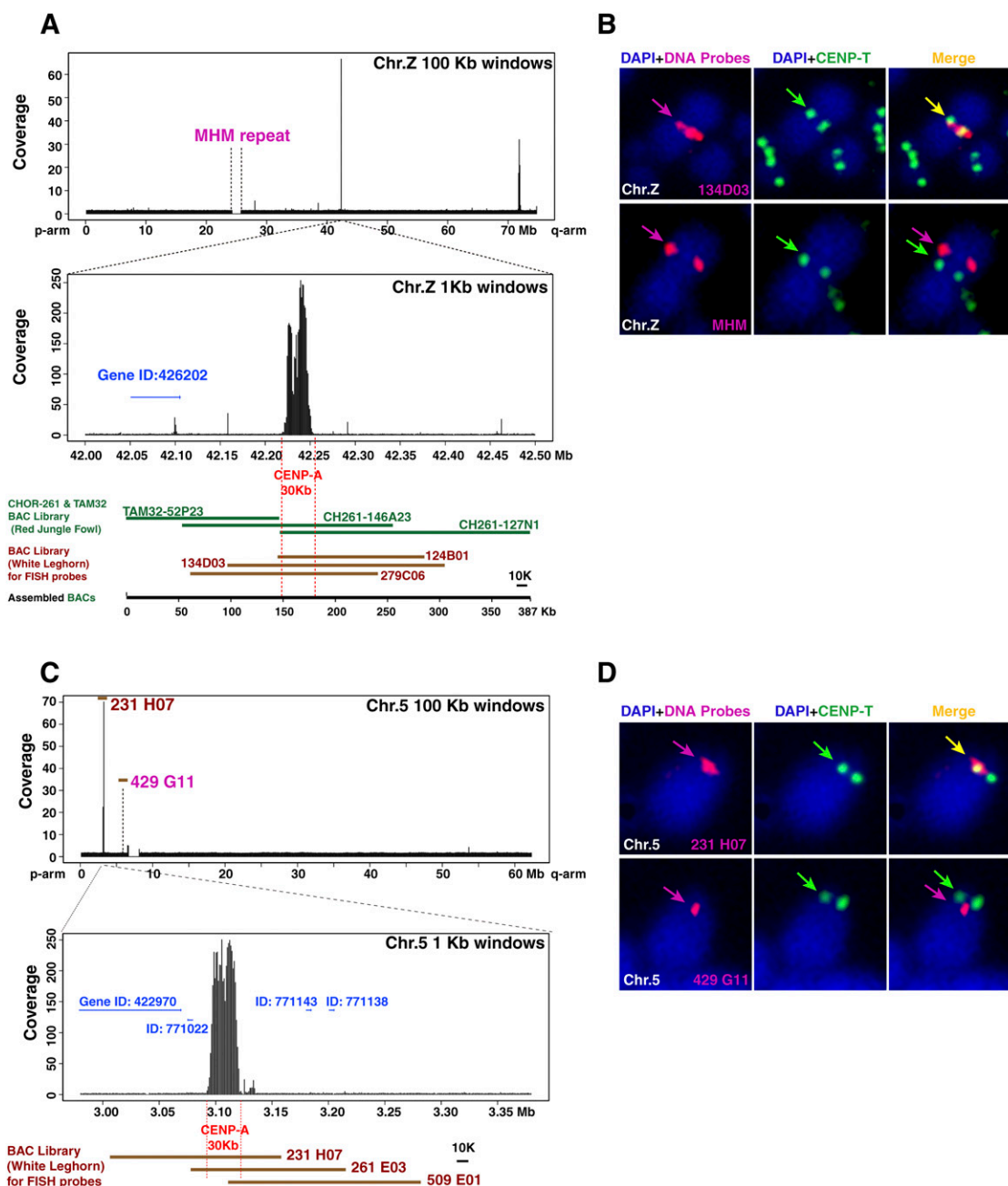
lacking. To further define the chicken centromeres, we used next generation deep sequencing to analyze CENPA-associated DNAs (ChIP with massively parallel sequencing [ChIP-seq] analysis). As the chicken genome information deposited in the NCBI database does not contain most of the centromeric repetitive sequences (International Chicken Genome Sequencing Consortium 2004) that we identified above, we could not map CENPA-associated DNAs to the majority of chromosomes. However, we identified a strong enrichment of sequences that mapped to the 42-Mb position of the Z chromosome and the 3.1-Mb position of chromosome 5 (Fig. 4). Both regions lack tandem repetitive sequences. Chromosome Z contains a long-range repetitive array, termed MHM repeats, and spans a ~2-Mb region (Fig. 4A; Teranishi et al. 2001). Chromosome 5 also may contain a long-range repeat sequence. To confirm which region colocalizes with centromere proteins, we isolated BAC clones containing both of these repeat sequences and the non-tandem-repetitive CENPA-associated DNAs and performed FISH analysis. As shown in Figure 4B,D, a BAC clone containing the corresponding CENPA-associated DNA, but not the long-range repeat sequences, colocalized with each centromere, indicating that the long-range repetitive arrays in chromosomes Z and 5 do not function as centromere DNA. Although we observed a second peak in Figure 4A (72-Mb region of Z chromosome), we did not observe colocalization of the region with CENPT, a second marker for a functional centromere (Hori et al. 2008). High-resolution DNA mapping indicates that the major CENPA-associated region spans ~30 kb in both chromosomes 5 and Z (Fig. 4A,C). In addition, we also found a CENPA-associated sequence on one micro-chromosome, which is devoid of tandem repeat (chromosome 27; Supplemental Fig. S2). This sequence also spans ~30 kb. These 30-kb centromere sequences were deposited in the DDBJ/EMBL/GenBank database with accession numbers AB556729–AB556731. We examined sequences of the 292 CENPA associated clones analyzed in Supplemental Table S1 and found that several clones contain these sequences (Supplemental Table S1). We compared the sequence homology between these three non-tandem-repetitive centromeres and found that there is an enrichment of retro-transposon sequences in CenZ and Cen27: Regions with homology with CR-1 and LTR elements (>70% homology) cover 55% of CenZ and 37% of Cen27. However, we did not detect further common features between these non-tandem-repetitive centromere sequences (Supplemental Fig. S3).

### Conditional removal of centromere sequences by genetic engineering in DT40 cells

Chicken DT40 cells provide a powerful system for genetic manipulations creating the possibility to modify CenDNA efficiently. To analyze the function of the centromere sequences we identified, we sought to create a cell line in which CenDNA is conditionally removed. For these studies, we chose to focus on the non-tandem-repetitive centromere sequence from chromosome Z. To remove CenDNA, we inserted *loxP* sequences at both flanking sites of the Z chromosome centromere region using homologous recombination (Fig. 5A). Each target construct containing *loxP* and a distinct drug resistance gene was transfected sequentially into DT40 cells. Homologous recombination was confirmed by Southern hybridization (Fig. 5A,C). We then introduced a Mer-Cre-Mer construct (Verrou et al. 1999), which expresses an active form of Cre recombinase in the presence of hydroxytamoxifen (OHT), into cells containing *loxP* sequences at both centromere flanking sites. In this resulting cell line, addition of OHT should conditionally remove



**Figure 3.** Genome organization of centromere region of chromosome 1 and chromosome 2. (A) DNAs from BAC clones hybridized with Cen1 were digested with HindIII and characterized by Southern analysis. Various BAC clones containing unique sequences and the 1.8-kb repeat-unit sequence were obtained. (B) A pulsed-field gel electrophoresis (PFGE) analysis was performed to determine the length consisting of the repeat-unit. Information of restriction enzyme sites is shown in C. (C) Genome organization around Cen1 region. Sequences of contigs from BAC002H09 and BAC003G11 are deposited in the DDBJ/EMBL/GenBank database with accession numbers AB556732–AB556734. Sequencing of BAC clones identified the exact boundary between unique region and repeat region. The length consisting of the repeat-unit was determined by a PFGE analysis. (D) DNAs from BAC clones hybridized with Cen2 were digested with HindIII and characterized by Southern analysis. Various BAC clones containing unique sequences and the 3.0-kb repeat-unit sequence were obtained. (E) A PFGE analysis was performed to determine the length consisting of the repeat-unit. Information of restriction enzyme sites is shown in F. (F) Genome organization around Cen2 region. Sequences of contigs from BAC074F14 and BAC285H02 are deposited in the DDBJ/EMBL/GenBank database with accession numbers AB556735 and AB556736, respectively. Sequencing of BAC clones identified the exact boundary between unique region and repeat region. The length consisting of the repeat-unit was determined by a PFGE analysis.



**Figure 4.** Centromere region is devoid of tandem-repeated sequences in chromosomes 5 and Z. (A) A distribution of DNA precipitated with CENPA in entire chromosome Z. Major peak is around position of the 42 Mb. High-resolution profile in 1000-bp windows indicates that the binding region of CENPA is restricted in the 30-kb region. Detail sequence information is available in several BAC contigs. As part of sequence information around the 42-Mb position was absent in the NCBI database, we remapped the DNA against BAC contig (387 kb) covering these genomic region and the CENPA associated-DNA was mapped in the 30-kb region, which does not contain tandem-repetitive sequence. Gene ID around CENPA-associated DNA is shown. (B) FISH analysis using a BAC clone (134D03) containing CENPA binding region of chromosome Z (red in *top* panel). Centromeres are immunostained with anti-CENPT antibodies (green). FISH signals of the 134D03BAC colocalized with centromeres, while FISH signals of the MHM repeat sequence (BamHI 2.2-kb unit) (red in *bottom* panel) did not localize with CENPT. (C) A distribution of DNA precipitated with CENPA in entire chromosome 5. Major peak is around position of the 3 Mb. High-resolution profile in 1000-bp windows indicates that the binding region of CENPA is restricted in the 30-kb region. Gene IDs around CENPA associated DNA are shown. Clone numbers of BACs are indicated. (D) FISH analysis using a BAC clone (231H07) containing CENPA binding region of chromosome 5 (red in *top* panel). Centromeres are immunostained with anti-CENPT antibodies (green). FISH signals of the 231H07 BAC colocalized with CENPT, while FISH signals of a BAC clone (429G11) around chromosome 5 satellite repeat sequence (red of *bottom* panel) did not localize with CENPT.

the complete centromere region (Fig. 5B). To test this, we performed PFGE analysis after addition of OHT in cells containing Mer-Cre-Mer and confirmed deletion of the centromere region

(Fig. 5D). The Z sex chromosome is present in only a single copy in DT40 cells. To determine the consequences of removing the centromere region, we examined cell growth upon the addition of





OHT. Cell growth ceased 2 d after addition of OHT and almost all cells had died 4 d after addition (Fig. 5E). We also examined chromosome behavior during mitosis after addition of OHT using FISH analysis with a probe for chromosome Z. As shown in Figure 5F, we frequently observed metaphase cells in which almost all chromosomes were aligned at the metaphase plate, but the chromosome Z had not aligned. During anaphase, sister Z chromosomes were not segregated into daughter cells and were left at the cell midzone (Fig. 5G). These observations indicate that cells with inserted *loxP* sequences completely lack centromere function after activation of Cre recombinase. We also investigated the stability of chromosome Z after deletion of the centromere region. We observed an increase in cells that lost chromosome Z or had two Z chromosomes, indicating that centromere deletion results in chromosome instability (Fig. 5H,I).

## Discussion

In this paper, we performed a comprehensive analysis of chicken CenDNA that demonstrates that (1) many chicken centromeres contain homogeneous repetitive arrays and the repeat-units of each centromere are chromosome-specific, (2) centromeres from three chromosomes (chromosomes 5, 27, and Z) span non-tandem-repetitive sequences of ~30 kb, and (3) cells in which the identified centromere sequence is conditionally removed fail to segregate that chromosome.

Human alphoid DNA is chromosome-specific (Shepelev et al. 2009). Each active centromere contains a different higher-order repeated unit (HOR) that is related to each other, but unique for each chromosome. Sequence similarity between HORs is 70%–88% (Alexandrov et al. 2001; Shepelev et al. 2009). The copy number of HORs ranges from a hundred to thousands. This feature is similar to what we described here for the chicken centromere. The main difference between human and chicken centromeres is that the composition of chicken centromere repeat-units is much more complex and includes not only one satellite repeat, but a satellite (21 bp, GGXHOI) and pieces of CR1 (Fig. 2). The 21-bp sequence (GGXHOI) is also observed in W heterochromatin regions in chicken genome and shows unusual electrophoretic behavior, which is attributed to the occurrence of DNA curvatures (Kodama et al. 1987). It is possible that a specific DNA-associated protein recognizes these DNA curvatures and localize to centromeres and heterochromatin similar to the mammalian protein CENPB (Masumoto et al. 1989).

In many species, chromosomes contain satellite sequences (Fukagawa 2004; Rudd et al. 2006; Yan and Jiang 2007). Human and mouse active centromeres are composed exclusively of satellite DNA. There are no retrotransposons in active centromeres, although there are transposons in dead centromeres surrounding the active one (Schueler et al. 2001; Kazakov et al. 2003). On the

other hand, in rice there are centromere-specific transposons in addition to the CentO repeat (Yan and Jiang 2007). As we also found retrotransposon-like sequences in chicken centromeres, chicken centromere is quite similar to that of rice. However, it appears in rice that there is no homogeneous core of defined repeat-units. In this respect, the chicken centromere is perhaps more similar to the human centromere. In *Drosophila* and *Xenopus*, chromosomes do not contain common centromere sequences (Sun et al. 1997; Edwards and Murray 2005), although the detailed sequence information from each centromere remains unclear. Extensive comparison with centromere sequences from various species will be important to understand centromere evolution.

We also identified non-tandem-repetitive sequences of ~30 kb for three chromosomes. The complete Z chromosome (~75 Mb) is 15 times bigger than that of chromosome 27 (~5 Mb), suggesting that the size of the kinetochore is independent of chromosome size. Considering that similar levels of constitutive centromere proteins are present at each chicken chromosome (Jonston et al. 2010), we propose that the kinetochore region of each chromosome should occupy only ~30 kb, even for chromosomes with long-range repetitive arrays. Therefore, most long-range repetitive sequences may serve as a mark for the centromere region within a chromosome rather than a specific functional element. In addition, for chicken chromosomes with repetitive centromere sequences, each of these repetitive sequences is quite divergent. We propose that centromeres without tandem-repetitive sequences represent a primitive form of these regions and that tandem-repetitive sequence were subsequently added during evolution. This idea is consistent with previous models regarding the formation of evolutionary new centromeres (ENC) (Ventura et al. 2007). Many ENCs have been identified in vertebrate and plant cells (Kasai et al. 2003; Carbone et al. 2006; Yan and Jiang 2007; Han et al. 2009). However, these ENCs have already acquired large arrays of satellite repeats. Very recently, an ENC without tandem-repetitive sequences was identified on chromosome 11 of domestic horse (Wade et al. 2009). Chickens contain three putative ENCs without tandem-repetitive array sequence that provide important insights into centromere evolution and function. Allshire and Karpen (2008) proposed that the centromere is formed by a sequence-independent epigenetic mechanism. Human neocentromere and chicken putative ENCs would be important materials to understand centromere formation. We also found an enrichment of retrotransposons in putative chicken ENCs like human neocentromeres (Chueh et al. 2009). Retrotransposons may ultimately contribute to the continued evolution of the centromere (Carone et al. 2009).

Finally, we have established an experimental system by which the centromere can be conditionally removed. It is possible to perform such an experiment using yeast cells (Ishii et al. 2008). However, there is a strong advantage to using cytogenetics in vertebrate DT40 cells compared with yeast cells. Therefore, our

**Figure 5.** Deletion of centromere region of chromosome Z results in chromosome instability. (A) To delete entire centromere region of chromosome Z, *loxP* sequences were inserted into both flanking sites of the centromere region using homologous recombination. CENPA binding region is also shown. (B) Expected genotype after recombination between both *loxP* sites upon activation of Cre recombinase. The 127-kb region should be removed after addition of OHT. (C) Confirmation of *loxP* insertions at both flanking sites of the centromere region. Information of restriction map and probes (#1 and #4) for Southern hybridization are shown in A. (D) Confirmation of recombination between both *loxP* sites. PFGE analysis was performed. Information for the restriction map and the probe (Bleo) for Southern hybridization is shown in B. After recombination a novel 97-kb band is observed. (E) Growth curve of DT40 cells after removal of centromere DNA. (F) Representative images of abnormal metaphase cells in which Z chromosome is not aligned at metaphase plate. Z chromosome was painted with a Z-specific macrosatellite DNA probe (red). (G) Representative images of abnormal anaphase cells in which sister Z chromosomes are left at midzone. Z chromosome was painted with a Z-specific macrosatellite DNA probe (red). (H) Representative images of FISH analysis with a Z-specific macrosatellite DNA probe (red). Cells with two copies of chromosome Z or lacking chromosome Z are observed. (I) Numbers of cells containing chromosome Z after removal of centromere sequence. As a control, numbers of cells containing chromosome 1 were scored. Total cell numbers (*n*) for the measurements are also shown.



conditional centromere knockout cell line represents a powerful system to investigate the specific requirements for centromere function in vertebrate cells.

## Methods

### Immunoprecipitation, cloning, and sequencing of centromere DNA

A CENPA-Flag expression construct under control of the CMV promoter was created. Flag was fused to the C-terminal end of CENPA. DT40 cells expressing CENPA-Flag grow similarly to wild-type DT40 cells. For immunoprecipitation of these cells with an anti-Flag antibody, a nuclear fraction of cells expressing CENPA-Flag was collected and digested with excess MNase (3 U/mL) (Takara) at 37°C for 2 h. The samples were solubilized in 0.5 M NaCl. Anti-Flag M2-beads (Sigma) were incubated with the sample for 2–4 h at 4°C and washed four times with 1 mL of buffer B (20 mM Tris-HCl at pH 8.0, 5 mM EDTA, 500 mM NaCl, and 0.2% Tween 20) and eluted with buffer B in the presence of 3× Flag peptide (Sigma). DNA was extracted from immunoprecipitates and cloned into pBluescript. Alternatively, the DNA was subjected to Illumina DNA sequencing. Sequenced DNAs were mapped into a Chicken Genome database with the MAQ program (Li et al. 2008).

### FISH and immunofluorescence

Two-hundred-ninety-two clones containing CENPA associated DNA was sequenced (Supplemental Table S1) and ~100 clones were randomly selected to use for FISH analysis. DT40 metaphase spreads were prepared by the cytospin method and fixed in 4% paraformaldehyde for 15 min at room temperature. After a brief treatment with 0.5% NP-40 in PBS, glass slides were dehydrated in ethanol. Chromosome DNAs were heat-denatured for 5 min at 95°C. DNA probes were labeled with Biotin-16-dUTP (Roche) by nick translation procedure. Two hundred nanograms of probes were dissolved in a hybridization buffer (0.04 M phosphate buffer at pH 7.0, 2× standard saline citrate [SSC], 50% formamide, 10% dextran sulphate, 1× Denhart's solution, 100 µg/mL of salmon sperm DNA, and 0.5 mM EDTA) denatured for 10 min at 75°C. Hybridization was carried out at 37°C in a humidity chamber overnight. Slides were washed in 2× SSC, 50% formamide in 1× SSC, 0.1× SSC for 10 min at 42°C. Percentage base pair mismatches allowed in this hybridization were estimated from the difference in the melting temperature ( $T_m$ ) for the hybridization or washing. The  $T_m$  was calculated for a perfectly matched hybrid having the average base pair composition of probes ( $G + C = 47.8\%$ ) and an average length of biotin-labeled 200-bp probes in the 2× SSC (0.39 M Na<sup>+</sup>), 50% formamide for the hybridization or 0.1× SSC (0.0195 M Na<sup>+</sup>), 50% formamide for the washing system. Using these parameters,  $T_m$ s for hybridization and washing are 86°C and 62°C. It was assumed that the  $T_m$  of a DNA–DNA hybrid decreased by 1°C for every 1% base pair mismatch present in the hybrid (Bonner et al. 1973), ~20% mismatch was allowed in our FISH experiment. After washing out probes, Cy3-anti-avidin were added to slides and incubated at 37°C for 30 min. Rabbit-anti CENPT antibodies (Hori et al. 2008) were used as a centromere marker. All immunofluorescence and FISH images were collected with a cooled EM CCD camera (QuantEM; Roper Scientific Japan) mounted on an Olympus IX71 inverted microscope with a 100× objective lens together with a filter wheel and a DSU confocal unit. Z-sections ( $n = 15\sim 25$ ) were acquired at 0.3-µm steps. All subsequent analysis and processing of images were performed using Metamorph software (Molecular Devices Japan). For detection of Z chromosome, Z-specific macrosatellite DNA (Hori et al. 1996) was used as a probe

### Southern blot analysis

For Southern hybridization, 5–10 µg of genomic DNAs of DT40 cells were digested with various restriction enzymes as indicated in the figures and subjected to 0.7% agarose gel electrophoresis. Probes indicated in the figures were prepared. After blotting of gels to nylon membranes, hybridizations with probes were carried out in phosphate-SDS buffer (0.5 M Na-phosphate at pH 7.0, 1 mM EDTA, 7% SDS, 1% bovine serum albumin) and incubated overnight at 62°C. Membrane was sequentially washed in 2× SSC, 1× SSC, and 0.5× SSC buffer, all containing 0.5% SDS for 20 min at 65°C. Autoradiogram was scanned by a Storm 860 system (Molecular Dynamics).

### Deletion of centromere DNA

Target constructs containing *loxP* sequence and several resistance cassettes under control of the beta-actin promoter were created by a standard method. Target constructs were transfected with a Gene Pulser II electroporator (Bio-Rad) into DT40 cells. All DT40 cells were cultured at 38°C in Dulbecco's modified medium supplemented with 10% fetal calf serum, 1% chicken serum, 2-mercaptoethanol, penicillin, and streptomycin. To activate the Cre recombinase, 4-hydroxytamoxifen (OHT; Sigma) was added to the culture medium to a final concentration of 100 nM.

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## Chickens possess centromeres with both extended tandem repeats and short non-tandem-repetitive sequences

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