New insights into mammalian sex chromosome structure and evolution using high-quality sequences from bovine X and Y chromosomes

Ruijie Liu1, Wai Yee Low1, Rick Tearle1, Sergey Koren2, Jay Ghurye3, Arang Rhie2, Adam M. Phillippy2, Benjamin D. Rosen4, Derek M. Bickhart3, Timothy P. L. Smith6, Stefan Hiendleder1 and John L. Williams1*

Abstract

Background: Mammalian X chromosomes are mainly euchromatic with a similar size and structure among species whereas Y chromosomes are smaller, have undergone substantial evolutionary changes and accumulated male specific genes and genes involved in sex determination. The pseudoautosomal region (PAR) is conserved on the X and Y and pair during meiosis. The structure, evolution and function of mammalian sex chromosomes, particularly the Y chromosome, is still poorly understood because few species have high quality sex chromosome assemblies.

Results: Here we report the first bovine sex chromosome assemblies that include the complete PAR spanning 6.84 Mb and three Y chromosome X-degenerate (X-d) regions. The PAR comprises 31 genes, including genes that are missing from the X chromosome in current cattle, sheep and goat reference genomes. Twenty-nine PAR genes are single-copy genes and two are multi-copy gene families, OBP, which has 3 copies and BDA20, which has 4 copies. The Y chromosome X-d1, 2a and 2b regions contain 11, 2 and 2 gametologs, respectively.

Conclusions: The ruminant PAR comprises 31 genes and is similar to the PAR of pig and dog but extends further than those of human and horse. Differences in the pseudoautosomal boundaries are consistent with evolutionary divergence times. A bovidae-specific expansion of members of the lipocalin gene family in the PAR reported here, may affect immune-modulation and anti-inflammatory responses in ruminants. Comparison of the X-d regions of Y chromosomes across species revealed that five of the X-Y gametologs, which are known to be global regulators of gene activity and candidate sexual dimorphism genes, are conserved.

Keywords: Genomes, Livestock, Bovine, Sex chromosomes, Pseudoautosomal region

Background

The sex chromosomes evolved from ancestral autosomes in dioecious lineages and have become extensively differentiated in structure and gene content [1, 2]. Mammalian X chromosomes are mainly euchromatic with a similar size and structure among species, and have retained most of the ancestral X genes [3, 4]. In contrast, Y chromosomes have undergone substantial evolutionary changes, accumulated male specific genes and genes involved in sex determination, and have lost 95% of the ancestral genes [5]. As a consequence, the Y chromosome is much smaller than the X chromosome and comprises mainly the pseudoautosomal (PAR), X-degenerate (X-d) and ampliconic regions [6].

The PAR is conserved on the X and Y, pairing and recombining at meiosis [7]. Most mammals have a single PAR region but the human sex chromosomes are an exception with a second PAR at the distal ends of the X and Y chromosomes [8]. The PAR plays an essential role in normal sexual development and loss of the PAR is associated with male sterility in humans [9]. Despite its critical role in fertility and disease, the PAR is one of the least well-characterised parts of most mammalian
genomes. Previous studies have described the genes at the bovine pseudoautosomal boundary and PAR gene content [10–12], but currently there is neither a complete gene map nor a precise size available for the bovine PAR.

The X-d regions of the Y chromosome contain single-copy genes, pseudogenes, which appear to be surviving relics of the autosomes from which the Y chromosome evolved, and ampliconic regions, which consist of large heterochromatic blocks rich in repetitive sequences [9, 13, 14]. Both the X-d and ampliconic regions are male-specific. Unlike the highly conserved PAR, the structure and gene content of the X-d regions differ among mammalian species. The human X-d regions are interrupted by several large blocks of ampliconic sequences, while the X-d regions of chimpanzee include a single ampliconic block [9, 15]. A study of bovine Y chromosome gene expression has contributed information on genes in X-d regions [16], but interpretation of the data is limited by the relatively poor quality of the available Y chromosome assembly.

The complex and highly repetitive Y ampliconic regions are difficult to assemble, particularly from short sequence reads. Many mammalian genome sequencing projects have used a female subject to avoid having to resolve X and Y haplotypes, and therefore do not include the Y chromosome. Only a few species, including human [6], chimpanzee [15], rhesus macaque [17], mouse [18], pig [19] and horse [20], have well characterised and assembled Y chromosomes.

The Bos taurus taurus reference genome assembly Btau_5.0.1 [21] (NCBI Project ID:20275) was assembled from short and long sequence reads of BAC clones and contains a Y chromosome sequence. A Bos taurus indicus Y chromosome was created by alignment of short read sequences to Btau_4.0 [22], and therefore will be missing any larger indicine-specific features. These sex chromosome assemblies are incomplete and inconsistent, hindering studies on sex chromosome evolution and the dissection of the molecular architecture of sexually dimorphic phenotypic traits.

In the present article, we report high quality assemblies of bovine X and Y chromosomes, created from long read sequences and optical mapping data, using a trio binning approach [23] that exploited the high level of DNA sequence divergence between the two subspecies of domestic cattle [24–26], Bos taurus taurus and Bos taurus indicus. We present a detailed gene map of the complete bovine PAR and X-d regions from these assemblies and discuss the evolutionary changes and functional aspects in these regions in comparison with other mammals.

**Results**

**Assembly and annotation of the cattle sex chromosomes**

The bovine X and Y chromosomes were assembled from whole genome sequence of a hybrid male with a Bos taurus taurus (Angus) sire and a Bos taurus indicus (Brahman) dam [23] (see URLs). The assembled chromosomes presented here are the Brahman X chromosome which comprises 146 Mb in 106 contigs with 983 genes, and the Angus Y chromosome which comprises 16 Mb in 67 contigs with 51 unique genes (Table 1 and Additional file 1: Table S1). These sequence assemblies have been deposited at NCBI (X: CM0011833.1; Y: CM0011803.1). The full length of the cattle Y chromosome has been estimated as ~ 50 Mb, at least half of which is in the highly repetitive region [27]. As in other species [19, 20, 28], even with long read sequencing, we could not assemble the ampliconic highly repetitive region [27] or the heterochromatic regions. Full annotation of the Brahman X and Angus Y chromosomes are available from Ensembl release v97 (UOA_Brhaman_1 and UOA_Angus_1). Analysis of the PAR and X-degenerate regions are presented below.

**Identification of the cattle PAR**

Alignment of assembled Brahman X and Angus Y chromosomes to each other identified a 6.8 Mb region with 99% sequence identity that extends from the start of the assembled X chromosome sequence (CM0011833.1) to 2933 bp distal to GPR143, after which sequence identity decreases to 86% for 348 bp and then drops abruptly to ~15% for the next 1 Mb (Fig. 1). The X chromosome PAR is assembled in one contig while the Y chromosome PAR only has two contig gaps. This enabled us to precisely define the PAR boundary and size. The PAR on the Brahman X and Angus Y chromosomes contained 31 genes in the same order. Of these, 29 are single-copy genes and two are multi-copy gene families, OBP, which has 3 copies and BDA20, which has 4 copies (Additional file 1: Table S4). The Brahman X chromosome PAR contains 12 genes that are missing from the proximal end of the X chromosome.

| Table 1 Length and number of gaps for Mammalian Sex chromosomes |
|---------------------------------------------------------------|
| **X Length (bp)** | **X Gaps** | **Y Length (bp)** | **Y Gaps** |
|-------------------|-----------|-------------------|-----------|
| Cattle-Brahman/Angus | 146,049,346 | 91 | 15,624,455 | 69 |
| Cattle-Nelore | 82,205,613 | 10,873 | 14,991,264 | 35,040 |
| Cattle-Hereford | 139,002,886 | 55 | 38,719,986 | 18 |
| Water buffalo | 143,477,029 | 65 | – | – |
| Goat | 115,943,529 | 319 | – | – |
| Sheep | 132,936,813 | 2968 | – | – |
| Pig | 125,778,992 | 10 | 15,567,420 | 12 |
| Dog | 123,180,702 | 1032 | – | – |
| Horse | 127,806,490 | 300 | 8,967,074 | 560 |
| Human | 154,893,106 | 28 | 26,415,094 | 55 |

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Liu et al. BMC Genomics (2019) 20:1000
chromosome in the current Hereford reference genome ARS-UCD1.2 (Additional file 1: Table S3).

**Identification of cattle X-degenerate regions**

Additional genes outside the PAR showed between 60 and 96% sequence identity between the X and Y chromosomes and are located in X-degenerate regions of the Y chromosome. The first of these regions, X-d1, is located distal to the PAR and spans 1.48 Mb, between 6.84 Mb and 8.32 Mb. X-d1 contains 11 single-copy protein coding genes. The corresponding region on the X chromosome spans 35 Mb and contains 10 X-d1 homologues in a different order but misses RPL23AY, which is located on chromosome 19 (Fig. 2). A 3 Mb ampliconic region immediately distal to X-d1 contains the male-specific Y (MSY) gene families PRAMEY, TSPY, and HSFY. At the distal end of the ampliconic region, the second X-degenerate region, X-d2a, spans 1.63 Mb and contains two single copy genes, UBE1Y and TXLNGY. The X chromosome homologs of these two genes are separated by a 44 Mb interval that contains 285 X chromosome-specific genes. Distal to X-d2a lies a 4.5 Mb ampliconic segment containing the bovine specific MSY genes ZNF280AY and ZNF280BY, which are
equivalent to TSPY and HSFY found in other species. The copy numbers of multi-copy MSY gene families are listed in Additional file 1: Table S2 and the complex ar-
rangement of multi-copy genes is presented in Add-
tional file 1: Figure S3. The distal end of chromosome Y contains the third X-degenerate region, X-d2b, which extends over 1.3 Mb and includes SRY and two copies of RBMY. The X chromosome homologs of these, SOX3 and RBMX, are located in a 5 Mb segment at the distal end of the X chromosome.

**Comparison of sex chromosome structure in mammals**

Alignment of the Brahman X chromosome with the water buffalo X chromosome [29] revealed a high level of co-linearity, with one large inversion and five small inversions at the distal end of the chromosome (Additional file 1: Figure S1b). The Brahman and water buffalo X chromosomes are 30 and 25 Mb longer, respectively, than the goat X chromosome, which consists of two scaffolds with a combined length of 116 Mb [30]. The goat X chromosome shows excellent co-linearity overall with the sheep X chromosome (Additional file 1: Figure S1c-d) but both showed numerous break points.
and several inversions, particularly on the short arm, in comparison with the Brahman and water buffalo X chromosomes. Non-ruminant mammalian X chromosomes, i.e. human, pig, dog and horse, revealed a striking similarity in the pattern of rearrangements in comparison to the *Bos taurus indicus* (Brahman) X chromosome (Additional file 1: Figure S1e-h). These consisted predominantly of 5 large inversions.

Alignment of the Angus Y chromosome assembly with pig, horse and human Y chromosomes showed limited co-linearity which was confined to the PAR and X-degenerate regions (Additional file 1: Figure S2a-c).

**Gene content and order of the mammalian PAR**

There is a very high level of conservation of synteny among mammalian PARs (Fig. 3). PLCXD1 is the most proximal PAR gene in human, horse, Brahman cattle and water buffalo. At their proximal ends, the PAR regions in the Hereford cattle reference genome, and sheep, goat and pig assemblies are truncated distal to *DHRSX*, *CLRF2*, *CD99* and *GYG2* respectively. In comparison to all the other species, the goat PAR has an inversion of three genes (*TBL1X*, *GPR143*, *SHROOM2*) close to the ruminant PAR boundary. This region is contained in one contig of the goat assembly and may thus be a contig orientation error, rather than a goat-specific rearrangement. The human sex chromosomes are an exception amongst mammals and have PARs at the proximal and distal ends [8]. The PAR1 in human is equivalent to the single PAR of other mammalian species, but is much shorter, with a distal boundary proximal to *XG*. The PAR of horse is the shortest with the distal boundary at *PRKX* (Fig. 3).

**Fig. 3** Comparison of the X chromosome PAR genes in cattle and other mammals. PAR genes (dark blue) and X specific genes (light blue) are indicated for each species. Red dashed lines represent genes with different order relative to the Brahman cattle. The pseudoautosomal boundary gene is the same in all ruminants except goat which has an inversion. Copy number of *OBP* and *BDA20* varies among species. The copy number of *OBP* and *BDA20* are listed in Additional file 1: Table S3. At their distal end, the pig and dog PAR extend beyond *GPR143* with a boundary distal to *SHROOM2*. In comparison to all the other species, the goat PAR has an inversion of three genes (*TBL1X*, *GPR143*, *SHROOM2*) close to the ruminant PAR boundary. This region is contained in one contig of the goat assembly and may thus be a contig orientation error, rather than a goat-specific rearrangement. The human sex chromosomes are an exception amongst mammals and have PARs at the proximal and distal ends [8]. The PAR1 in human is equivalent to the single PAR of other mammalian species, but is much shorter, with a distal boundary proximal to *XG*. The PAR of horse is the shortest with the distal boundary at *PRKX* (Fig. 3).
PAR gene family expansions in different lineages

The OBP gene family, which is distal to PRKX in all species, is within the PAR of all ruminants, pig and dog, but is outside the PAR of horse, and is missing from the human X chromosome. This gene family is expanded in ruminants (Fig. 3). The BDA20 gene family is immediately distal to the OBP family and present in all ruminants for which data are available, including Yak [31], Deer [32] and Chiru [33], but is not found in other mammals (Fig. 3). The BDA20 family shows differential expansion in the different ruminant species, with two or more copies with 74–91% nucleotide sequence identity at mRNA level in cattle [21], sheep [34], goat [30] and buffalo [29] (Fig. 3, Additional file 1: Table S4). In contrast, ARSF, a member of the ARS family, has been reported as a PAR gene in other mammalian species, but is not found in any of the ruminant PARs [29, 30, 34].

Comparison of X-degenerate Y chromosome regions

Most of the X-Y paired genes of cattle, pig and horse that are outside the PAR are found in the X degenerate region, X-d1, located adjacent to the PAR (Fig. 4). Of the 11 genes in the cattle X-d1 region 8 are in common with horse and pig X-d1 regions, but the gene order differs between the three species. RPL23AY is only found in the cow X-d1, while TMS4BAY is found in the horse and pig X-d1 regions and the human X-d3 region which is missing from cow X-d regions. Five additional bovine gametologs are found in two X-d2 regions, X-d2a and X-d2b, which correspond to the single X-d2 in horse and pig. Cattle X-d2a is distal to X-d1 and contains 2 genes, UBE1Y and TXLNGY. Both genes are found in the pig X-d2 region but UBE1Y is in an ampliconic region of the horse Y chromosome. The cattle X-d2b region contains SRY and is in a telomeric position similar to the X-d2 region of pig. The cattle X-d2b region contains two copies of RBMY, which is also duplicated in the horse X-d2 [20].

Discussion

The trio-binning approach facilitated the construction of the most complete bovine X and Y chromosome sequence assemblies available to date. Alignment of the Brahman cattle X with Hereford X revealed a major inverted region. In both assemblies this inverted region ends at contig breakpoints and could be an assembly artefact rather than a true biological difference. We could not resolve this discrepancy using RH and linkage maps [35–37] because the marker density was insufficient. Alignments of the Brahman cattle X chromosome

![Fig. 4 Comparison of Y chromosome X-degenerate regions in cattle and other mammals. The human Y chromosome is different from other mammals with 7 X-d regions and one unique X-transposed region (pink). Eight X-d1 genes are shared among cattle, pig and horse but the gene order and orientation are not the same. The cattle X-d2 is split into and X-d2a X-d2b regions with 2 and 3 genes respectively. The cattle Y chromosome diagram does not include the unknown heterochromatic regions.](image-url)
with those of the water buffalo, goat and sheep, revealed a high level of co-linearity, but with numerous break points and several inversions. Many of these structural differences are consistent with gross karyotypic rearrangements that differentiate goat and sheep X chromosomes from those of cattle and water buffalo [38]. The evolutionary separation of goat and sheep from cattle and buffalo occurred 20–25 MYBP, which pre-dates the separation of goat and sheep from each other, 10–15 MYBP, and of buffalo from cattle, about 12 MYBP [39]. This evolutionary history is consistent with the differences in X chromosome structure we see among these species [24, 40].

The X chromosomes of human, pig, dog and horse have a strikingly similar pattern of differences relative to the Brahman cattle X chromosome. Ruminants are even-toed ungulates which separated from odd-toed ungulates (including the pig and horse) more than 60 MYBP, suggesting that the rearrangements occurred after this split but prior to the separation of the ruminants, about 25 MYBP [24].

Comparison of the Brahman cattle X chromosome with other mammalian reference genomes suggest the X chromosome assemblies for the Hereford cattle, sheep, goat and pig are incomplete at the proximal end of the PAR. The proximal PAR genes of Brahman cattle, water buffalo, human and horse are the same. Genes missing from the proximal end of the Hereford cattle X chromosome have been misplaced on various autosomes, whereas the genes missing from sheep, goat and pig, are found on unplaced contigs. Given that the common ancestor of cattle, water buffalo, human and horse existed about 96 MYBP, it is more parsimonious to suggest the gene order and structure of the proximal PAR are conserved and not assembly artefacts. The assembly of the Brahman cattle and water buffalo [29] X chromosomes may have benefited from the use of much longer PacBio reads and improved assembly algorithms.

The distal PAR boundary in ancient species is thought to lie within the AMEL locus [41]. However, there was no substantial identity of the Brahman X and Angus Y chromosomes for the region between 2933 bp distal to GPR143 and AMELX. The PAR boundary in cattle is therefore just distal to GPR143. The distal ends of the PARs of water buffalo and sheep also lie close to GPR143.

The expansion of OBP and BDA20 gene families in ruminants suggests they have a specific role in these species. These genes are members of the lipocalin family which are involved in immune-modulation and anti-inflammatory responses [42]. For example, their expression changes after exposure of cattle to ticks [43]. The ARS gene family in the ruminant PAR is missing ARSF that is found in non-ruminant species, suggesting that it has been lost during ruminant evolution.

Recombination of mammalian sex chromosomes only occurs in the PAR. This may explain why the MSY regions are rich in repetitive sequences [2, 44]. The co-linearity between Angus and Hereford Y chromosomes is limited to X-d and PRAMEY regions, while the numerous repetitive sequences in ampliconic regions appear expanded in the Hereford Y. This may be due to the use of various BAC clones from several individuals to assemble the Hereford Y chromosome [45]. The Angus Y chromosome assembly shows some alignment in some isolated areas with pig, horse and human Y chromosomes, which is mainly in X-degenerate and ampliconic regions. This is consistent with rapid evolution of non-recombining Y chromosome sequences [3].

We identified 16 X-Y paired genes in 3 X-degenerate regions on the Angus Y chromosome, which were interspersed by ampliconic regions that contain multiple copies of bovine-specific PRAMEY, ZNF280AY and ZNF280BY genes. PRAMEY genes are exclusively expressed in testis and are involved in spermatogenesis during testicular maturation [46]. ZNF280BY and ZNF280AY are multi-copy Y-genes transposed from an autosome. The temporal and spatial expression patterns of these genes also suggests that they play a role in spermatogenesis [47]. While horse, pig and cattle X-d1 are similar in gene content, gene order is very different in the three species. RPL23AY is cattle-specific and a second copy of this gene, RPL23A, is found on chromosome 19, located in a conserved block in mammals with intron-exon structure and six identical intron-less pseudogenes with 91% sequence identity to this gene are found on cattle chromosomes 3, 9, 10, 14, 22 and 29, suggesting that these copies have arisen by retrotransposition.

Cattle have lost several ancestral Y genes from X-d regions including KDSMD, TMSB4Y and TXLNGY. Ten of the 16 gametologs are conserved in the X-d regions of horse, 8 in pig and 5 in human. The genes in the X-d regions, do not recombine, and hence diverge over time, allowing for the possibility of sex-specific selection. They may therefore be involved in sexual dimorphism [48]. Five of these (DDX3Y, EIF1AY, USP9Y, UTY, ZFY) are global regulators of gene activity expressed across a broad range of adult tissues and could have profound effects on sexual development [48].

Conclusions
The quality of assemblies achieved for the X chromosome, and for the PAR, X-d and ampliconic regions of the Y chromosome, using the trio-binning approach, enabled us to examine and compare major structures of the bovine sex chromosomes, both between the sex
chromosomes and among species. Alignment of the Brahman X and Angus Y chromosomes precisely identified boundaries and gene content of the PAR region and indicated that the proximal end of the PAR is truncated in the sheep, goat and current bovine reference genome assemblies. The sequence data revealed expansions of gene families in the ruminant PAR region that have previously been associated with immune function, and conservation of gametologs that are known dosage sensitive regulators of gene expression. The sex chromosome assemblies and the annotation presented are valuable resources for the molecular characterization of sex-specific phenotype in livestock and other species.

Methods

Sample collection
All animal work was approved by the University of Adelaide Animal Ethics Committee (No. S-094-2005). Briefly, a Brahman cow was bought by the University of Adelaide from a farm (Kiowa, Kingstown, New South Wales) and was transported to SARDI experimental farm at Struan South Australia where it was inseminated with semen of an Angus bull bought by the University from American Breeder Services, Australia. At day 153 post-insemination, the cow was humanely killed by stunning and exsanguination at a commercial abattoir (Dalriada Abattoir, Keith, South Australia) as per standard operating procedures. The uterus was recovered and the male fetus removed and immediately humanely killed by stunning and exsanguination. Lung tissue was collected and snap frozen in liquid nitrogen. Details of the contig creation for this assembly using the trio binning method have been previously described [23]. Briefly, DNA was extracted from fetal lung, paternal semen, and maternal uterine tissue. Long-read libraries of the fetus were prepared for sequencing on the Sequel platform as suggested by the manufacturer (Pacific Biosciences, Menlo Park, CA). Short-read libraries of the sire and dam were prepared for sequencing on the NextSeq500 platform as recommended by the manufacturer (Illumina, Inc. San Diego, CA). Approximately 60x short-read coverage of the dam and sire were produced, and 134x long-read coverage for the fetus. Parent-specific kmers were identified, long reads sorted into bins by parental origin, and independent haploid assemblies constructed using triocanu (implemented in Canu v1.7).

Sex chromosomes assembly and validation
Haplotype resolved paternal and maternal contigs constructed using Canu [23] were scaffolded independently using Hi-C reads [49] and an optical map (Bionano tools v1.3.0), which were then consolidated into chromosome specific groups of scaffolds and the sex chromosome scaffolds selected for the male and female assembly. The high density of repetitive elements on both the X and Y chromosomes made assembly difficult, breaking sequence contiguity even with long sequence reads, so additional markers were used to validate the order and orientation of scaffolds. For the X chromosome, the USDA-MARC Bovine linkage Map [35], and two RH maps, the BovGen RH map [37] and SUNbRH7000-rad map [36] were used to place, order and orientate scaffolds. For the Y chromosome, known genes in cattle [16, 22], pig (Sscrofa11.1) and human (GRCh38.p12) assemblies were used to identify Y-specific scaffolds. Cattle Y chromosome RH map markers [50] were used to guide ordering and orientation of the Y-specific scaffolds. To exclude scaffolds incorrectly identified as Y sequence, 26 scaffolds which were shorter than 50kb and contained fewer than three known Y genes, were manually inspected. These shorter scaffolds were partitioned into 50-kb bins and aligned with the CHORI-240 Bovine BAC library Y specific clones (see URLs) using BLASTN. Six sequences with less 90% alignment were removed. Further details on X and Y chromosomes scaffolds identification and orientation is given in Additional file 1: Note 1 and 2, respectively.

Conflict resolution
The RH X chromosome marker order and orientation was generally in agreement with the X chromosome assembly. However, 15 out of 84 markers from the BovGen RH map and 18 out of 93 markers from the SUNbRH7000-rad map suggested a possible assembly error. Comparison of the RH maps showed that 5 markers from BovGen RH map and 4 markers from SUNbRH7000-rad map had a consistent arrangement that differed from the Brahman X assembly. Both RH maps showed an inversion of a 1,286,607 bp contig in a scaffold that is made up of 4 contigs. With the exception of this scaffold, other RH marker inconsistencies would have required contigs to be broken and rearranged. The Brahman X and ARS-UCD1.2 X agreed in all these regions so the order and orientation of scaffolds that agreed best with Hi-C and optical map was retained.

Outside of the PAR, the Y chromosome assembly is made up of small scaffolds with average length of 98,509 bp. The ordering of scaffolds was completely dependent on the 52 RH markers and no contig breaks were required.

X and Y chromosome alignments
In order to compare intra- and inter-species sex chromosomes differences, pairwise alignments were carried out for cattle (Brahman, Angus, Hereford, Nelore), water buffalo, goat, sheep, human, pig, dog and horse using the aligner Lastz v1.04 [51] with default settings.
Brahman X is reverse complement to Hereford X. Prior to alignments, the Hereford X and Nelore X are reverse orientated for comparison. Repeats in the X and Y chromosomes were masked by Repeatmasker v4–0-7 using cow RepBase23.08 [52]. The number of gaps was calculated using a custom python script. The number of inversions were calculated by Smash with a 1 Mb block size [53]. Further details on the alignment parameters is given in Additional file 1: Note 3.

**X and Y chromosome gene annotations**

All X chromosome genes from ARS-UCD1.2 were re-mapped to the Brahman X chromosome using the Exonerate v2.4 software [54]. For the PAR, detailed manual annotation was carried out. A total of 42 protein-coding PAR genes were used to annotate the X chromosome. These genes came from the following sources: 1) 15 human PAR1 genes [8]; 2) 22 candidate cattle PAR genes [11]; 3) horse [55], dog [56] and pig [57] PAR genes. A cattle PAR gene, arylsulfatase E (ARSE), was reported in a FISH mapping study [11] but the sequence is not available from NCBI. However, three other arylsulfatases paralogs on the X chromosome were available and were used to search for other similar members on the PAR.

The current cattle reference genome assembly ARS-UCD1.2 does not have a Y chromosome, therefore 18 Y-specific genes from the X-degenerate regions were selected from a study of Y chromosome transcribed sequences [16], in addition, MSY genes from pig and human Y chromosomes were used for the Angus Y chromosome annotation. Further details on gene annotation are given in Additional file 1: Note 4 and Additional file 1: Table S1.

**Conservation of synteny in mammalian PARs**

Each of the X and Y chromosomes PARs were contained in single scaffold. To estimate the boundary of PAR, X and Y scaffolds containing PARs were partitioned into 1 Mb windows and aligned against each other using BLASTN. From the alignments, the first bin from the proximal end that had coverage below 80% and percentage identity below 80% was identified as the pseudoautosomal boundary. The gene order of PAR genes between closely related species was analysed using Bioconductor Gviz plots [58].

**Supplementary information**

Supplementary information accompanies this paper at https://doi.org/10.1186/s12864-019-6364-z.

Additional file 1: Figure S1. Alignment of the Brahman X with other mammalian X chromosomes. Figure S2. Alignment of the Angus cattle Y against other mammalian Y chromosomes. Figure S3. Multi-copy genes in Angus Y ampliconic region. Table S1. Summary of X and Y chromosome protein-coding genes. Table S2. Copy numbers of the protein-coding genes in the bovine MSY. Table S3. List of PAR genes missing in the current Hereford, sheep, goat and pig assemblies.

**Table S4.** Copy numbers of OBP and BDA20 genes in each species.

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**Authors’ contributions**

JLW and TPLS conceived and managed the project. SH designed breeding experiments, provided dam mtDNA typing, tissue and DNA samples and clarified the single PAR in cattle using BAC information. TPLS created, sequenced, and curated sequencing data. SK, AR and AMP assembled the contigs and provided guidance on scaffolding. JG and SK provided the Hi-C scaffolds. DMB and BDR compared scaffolding using various programs. WYL consolidated all data on assemblies to produce final chromosome-level haplotype-resolved genomes. RL and BDR validated the sex chromosome assemblies using various bovine markers. RL annotated the sex chromosomes and compared them to other mammals. RL, WYL, RT, SH and JH LW drafted the manuscript and all authors read, edited and approved the final manuscript.

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**Availability of data and materials**

Sequence data is available from the BioProject Accession PRJNA432857. Analysis of Angus and Brahman genome assemblies, https://github.com/llow/llow/llow/BrahmanAngusAssemblyScripts; the CHORI-240 Bovine BAC Library is available at https://bpicoursesources.org/fitlersAval.php; ArrowGrid, https://github.com/skoren/ArrowGrid; RepeatMasker, http://www.repeatmasker.org.

**Ethics approval**

All animal work was approved by the University of Adelaide Animal Ethics Committee, approval No. S-094-2005.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

**Author details**

1The Davies Research Centre, School of Animal and Veterinary Sciences, University of Adelaide, Roseworthy, South Australia, Australia. 2Genome Informatics Section, Computational and Statistical Genomics Branch, National Human Genome Research Institute, Bethesda, MD, USA. 3Center for Bioinformatics and Computational Biology, Lab 3104A, Biomolecular Science Building, University of Maryland, College Park, MD, USA. 4Animal Genomics and Improvement Laboratory, ARS USDA, Beltsville, MD, USA. 5Cell Wall Biology and Utilization Laboratory, ARS USDA, Madison, WI, USA. 6US Meat Animal Research Center, ARS USDA, Clay Centre, NE, USA.

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