A high-resolution comparative map of porcine chromosome 4 (SSC4)


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Summary

We used the IMNpRH12 000-rad RH and IMpRH7 000-rad panels to integrate 2019 transcriptome (RNA-seq)-generated contigs with markers from the porcine genetic and radiation hybrid (RH) maps and bacterial artificial chromosome finger-printed contigs, into 1) parallel framework maps (LOD≥10) on both panels for swine chromosome (SSC) 4, and 2) a high-resolution comparative map of SSC4, thus and human chromosomes (HSA) 1 and 8. A total of 573 loci were anchored and ordered on SSC4 closing gaps identified in the porcine sequence assembly Sscrofa9. Alignment of the SSC4 RH with the genetic map identified five microsatellites incorrectly mapped around the centromeric region in the genetic map. Further alignment of the RH and comparative maps with the genome sequence identified four additional regions of discrepancy that are also suggestive of errors in assembly, three of which were resolved through conserved synteny with blocks on HSA1 and HSA8.

Keywords comparative map, radiation hybrid map, RNA-seq, sequence assembly, SSC4, swine.

High-resolution radiation hybrid (RH) maps provide a ‘blueprint’ for genome sequence assembly and comparative mapping across genomes (Lewin et al. 2009). A RH-based bacterial artificial chromosome (BAC) contig map serves as the template for sequencing and assembly of the swine genome, but does not provide a robust comparative syntenic map that identifies breakpoints within and between minimally sequenced regions of the genome. Because closure of contig gaps would improve overall sequence assembly, we selected SSC4, one of the most deeply sequenced chromosomes in the draft sequence assembly, to test whether we could significantly and rapidly improve the pathway to final assembly by integrating RNA sequence (RNA-seq) data into a high-resolution comparative RH map of SSC4 and human HSA1 and HSA8.

Mapping vectors from the IMNpRH12 000-rad panel and IMpRH7 000-rad panel were first merged using Carthage (Schiex & Gaspin 1997), and the merged data set (Table S1) was analysed using a maximum multipoint likelihood linkage strategy. Three linkage groups with 89, 63 and 421 markers, respectively (Table S2), were initially mapped to SSC4 at a 2pt LOD≥10, and an FW map for each linkage group was built simultaneously at a likelihood ratio of 1000:1 (Fig. S1c, d). A total of 328 FW markers were ordered on the SSC4 IMNpRH12 000-rad FW map, over an accumulated map distance of 11 207.3 cR12 000 and 5507.8 cR7 000 (Tables S1 & S2). This yielded a Kb/cR12 000 ratio of ~13.0, a twofold increase in resolution over the IMpRH7 000-rad FW map (Table S2) and slightly lower than the 2.2- to 3.0-fold increase reported for several other porcine chromosomes (Yerle et al. 2002; Liu et al. 2005, 2008; Ma et al. 2009).

A total of 245 non-FW markers (Table S2) were then added to the RH12 000-rad FW map of SSC4 using Carthage (Schiex & Gaspin 1997), which improved the resolution of the SSC4...
map to \(\sim 254.8\) Kb/marker (146 Mb/573), a >3-fold increase in marker density on SSC4 over the 0.8–1.0 Mb/marker interval on the current IMpRH7,000-rad maps (Meyers et al. 2005; Rink et al. 2006; Humphray et al. 2007).

Seven FPC contigs (ctg) (4001–4005, 4007 and 4009), (http://www.sanger.ac.uk/Projects/S_scrofa/WebFPC/porcine/small.shtml) were identified, where 134 BACs/BEs were shared between the RH and FPC contig maps (Fig. S1c–e) with identical order of BAC end sequence (BES) in both the RH map and BACs, except for a small region of inconsistency that was observed for four BESs (362B20G08, 306B10F11, 274A10B02 and 310B20A04) bridging FPC ctg4004 and ctg4005 (Fig. S1c–e). The first two BESs, 362B20G08 and 306B10F11, mapped at the end of ctg4004, while BES 274A10B02-310B20A04 were at the beginning of ctg4005. In contrast, 274A10B02_306B10F11_362B20G08_310B20A04 were ordered within the same linkage group in the RH map (Fig. S1c–e), suggesting how contigs ctg4004 and ctg4005 became separated in the FPC maps.

Microsatellites (MSs) binned in the genetic map of SSC4 (http://www.marc.usda.gov/genome/swine/swine.htm) were all ordered on the RH12,000 map, providing a significant increase in map resolution. A discrepancy of the centromeric region (Fig. S1b–d) on the genetic map resulted from an inversion of five MSs (SWR362, SW1998, SW1003, SW1513 and SW1520) (Fig. S1b, c). SWR362 and SW1998 mapped to SSC4p, while SW1003, SW1513 and SW1520 mapped to SSC4q in the RH map (Fig. S1c, d). We validated MS order by blasting the sequences of the five MSs against the NCBI pig high-throughput genomic sequence database (http://blast.ncbi.nlm.nih.gov/Blast.cgi) to identify BACs containing the MS sequence and to check the map positions of the BACs in the FPC maps. BAC CH242-417I24 contains SWR362, while three overlapping BACs, CH242-512E4, CH242-18C1 and CH242-55N4, all contain SW1998, indicating that their order in FPC ctg4003 is 422B20A10_CH242-417I24_418A20F02_CH242-512E4_408A10F07 (http://www.sanger.ac.uk/Projects/S_scrofa/WebFPC/porcine/large.shtml), identical to their order in the RH map (Fig. S1c–e). Similar analyses identified that BACs CH242-321A10, CH242-38705 and CH242-62L8 contain SW1520, SW1513 and SW1003, respectively. The order of these BACs in FPC ctg4004 is 285A10D09_CH242-62L8_286A10B03_CH242-321A10_422B20A12, again identical to their order in the RH map (Fig. S1c–e) (http://www.sanger.ac.uk/Projects/S_scrofa/WebFPC/porcine/large.shtml). The results suggest that the limited number of meioses (104 animals) in the two-generation backcross population used to generate the swine genetic map (Rohrer et al. 1994) and low rate of recombination in this chromosomal region in the reference population (Yu et al. 2001; DeWan et al. 2002) have confounded the linkage ordering in the centromeric region of SSC4 from SW1520 to SW362 (Fig. S1b) (http://www.marc.usda.gov/genome/swine/swine.htm).

The comparative map of SSC4 and HSA1 and HSA8 (Fig. 1) revealed that a large block of sequence (0–93 Mb) from the distal region of SSC4p to the proximal region of SSC4q1.3 (Fig. S1a) is highly conserved within the long arm (48.4–146 Mb) of HSA8. The remaining of SSC4q (q1.3–2.5, Fig. S1a) from 93 to 136 Mb is conserved with HSA1 segments 1p22.3–12 (86–120 Mb) and 1q21.1–1q24.3 (145–171 Mb) (Fig. 1). However, this conservation in synteny was not complete. Firstly, the centromeric region (120–145 Mb) of HSA1 is not conserved with SSC4q (Fig. 1), indicating that a centromere origination and/or a genome-rearrangement event occurred between the two species during evolution (Liu et al. 2005). Second, two genes (PTK2 and TADA1L) in SSC4-HSA8 and two genes (TBP and PLA2G4A) in SSC4-HSA1 are not syntenic (Fig. 1), suggesting micro-rearrangements between the two species. The length ratio of the corresponding sequence blocks for SSC4-HSA8 is \(\sim 1:1\) (93/97 Mb) and SSC4-HSA1 \(\sim 1:1.4\) (43/60 Mb), which may explain the higher gene density (4X) observed in the SSC4q1.3–2.5 region (Figs 1 and S1).

Approximately 42 million Illumina RNA-seq reads (46-bp pair-end) generated from porcine macrophage and lymph node RNA were assembled de novo into contigs using a hybrid protocol incorporating ABySS-P (Simpson et al. 2009) and SSAKE (Warren et al. 2007). The resulting contigs were extended using PCAP (Huang et al. 2003). Reads were aligned to the pig genome sequence assembly (Sscrofa9) using GSNAI (Wu & Watanabe 2005). All high-throughput sequencing data were managed using the Alpheus pipeline and database resource (Miller et al. 2008). De novo assembled contigs, porcine sequences currently assembled in the NCBI unigene database, and previously assembled ESTs were aligned to the porcine genome using GMAP (Wu & Nacu 2010). Porcine transcripts derived from this combined resource were aligned to the human RefSeq protein data set and the strongest hit was taken as the value for a specific gene in that sample. All data from Alpheus alignments were loaded into a modified version of the Comparative Map and Trait Viewer (CMTV) (Sawkins et al. 2004). Approximately 22.5 million reads aligned to the NCBI Unigene: 15.7 million of the reads aligned uniquely; 17.5 million of the reads aligned to Sscrofa9; and 12.8 million reads aligned uniquely. Aligned vs. uniquely aligned reads in RNA-seq data provide a measure of the uniqueness of genes or of their components (e.g. a motif may be repeated in different genes). Hybrid assembly (methods) yielded 44 356 contigs longer than 100 bp. The contig N50 was 257, and the B2 000 was 257 785 bp, or 2.4% of the total assembly. Overall, we identified 17 328 de novo assembled contigs that aligned to the pig genome: 4863 contigs aligning only to the human genome, with 221 contigs aligning better to the human genome than porcine, suggesting that the sequence in those regions of the porcine genome is incomplete; and 15 737 contigs aligned to both species. The length ratio of the corresponding sequence blocks for SSC4-HSA8 is \(\sim 1:1\) (93/97 Mb) and SSC4-HSA1 \(\sim 1:1.4\) (43/60 Mb), which may explain the higher gene density (4X) observed in the SSC4q1.3–2.5 region (Figs 1 and S1).
genomes. Contigs were then aligned to the SSC4 genome sequence (Sscrofa9) and the RH map (Fig. 2). A total of 2019 contigs mapped to SSC4. Two hundred and forty-eight of these contigs (Table S3) overlapped with sequences of genes/ESTs mapped on the IMNpRH212000-rad RH panel, and the remaining 1171 contigs provided additional coding sequence for the RH map (Fig. 2). Several large blocks of SSC4 genomic sequence, i.e. 3–5 Mb, 22–27 Mb and 88–91 Mb (Fig. 2), did not match contig sequences, suggesting that a select group of genes are expressed in porcine macrophages and lymph nodes.

Finally, the high-resolution integrated map created through pairwise alignment of marker and sequence positions on the RH and SSC4 sequence maps, respectively, (Fig. S1c, d) identified four small regions (Fig. 2, regions a–d) of potential inconsistency. They were reanalysed using the conserved synteny between HSA8/HSA1 and SSC4 to determine whether the SSC4 RH or sequence map was correct based on agreement with the human sequence. Marker identity and order in the regions b, c and d on the SSC4 RH map (Fig. 2) are in agreement with the corresponding sequences on HSA8 and 1, eliminating the potential discrepancies identified in regions b-d (Fig. 2) and suggesting that a re-examination of the SSC4 sequence assembly in these three regions is necessary. The potential discrepancy identified as region a (Fig. 2) was not resolved because of limitations in the current level of resolution of the SSC4 RH map, sequence assembly and the HSA8 assembly.

We also analysed two genes (PTK2 and TADA1L) that were not in synteny between SSC4 and HSA8 (Fig. 1) and found that the human PTK2 gene has two copies; PTK2 at 141.7 Mb and PTK2B at 27.3 Mb on HSA8 (http://www.ncbi.nlm.nih.gov/mapview/map_search.cgi?chr=hum_chr_in&query=PTK2). BLAT alignment (http://genome.ucsc.edu/) of the pig PTK2 sequence (accession no. BW999672) against the human genome (Build 37) identified an orthologous copy at 27.3 Mb, but not the copy at 141.7 Mb on HSA8, suggesting that SSC4 may have only one copy of PTK2, similar to the human PTK2B gene, which is conserved between SSC4 and HSA8 (Fig. 1). A BLAT alignment also showed that the pig TADA1L gene (accession no. BI336804) has an orthologous sequence on HSA1 (score 360; similarity 94.2%) and a potentially orthologous sequence on HSA8 (score 305; similarity 86.1%). The human genome does, in fact, have two copies of TADA1L, one on HSA1 at 166.8 Mb and the other on HSA8 at 5.7 Mb (Fig. 1), suggesting that the assignment of the pig TADA1L to the SSC4 region in synteny with HSA8 is correct on the RH map, although we do not know if the pig has more than one copy of the TADA1L gene. Two additional genes, TAF12 and PLA2G4A, are not in synteny between SSC4 and HSA1 (Fig. 1), but the multiple copies of the human TAF12 may indicate a species-specific gene duplication and account for the lack of conservation in the position of this amplified gene family between human and pig.
We then used the comparative RH map to estimate gap distance in the FPC (and sequence) map. Analysis of the genes and BACs adjacent to the gap between ctg4004 and ctg4005 (Fig. S1) identified two porcine genes, $\text{FCGR2B}$ and $\text{DUSP12}$, and one MS, UMNp591, (Fig. S1) that mapped to BAC CH242-447O4, located at the terminus of ctg4004 in the FPC database (http://www.sanger.ac.uk/cgi-bin/Projects/S_scrofa/WebFPCdirect.cgi?contig=4004&clone=CH242-447O4). In contrast, the pig $\text{NDUFS2}$ and $\text{FCER1G}$ genes were located in BAC CH242-28G15, which maps to ctg4005 and overlaps with 310B20A04 (http://www.sanger.ac.uk/cgi-bin/Projects/S_scrofa/WebFPCdirect.cgi?contig=4005&clone=CH242-28G15). The ctg4004/ctg4005 boundary corresponds to a region of conserved synteny on HSA8. Human $\text{FCGR2B}$ starts at 161647639 bp, while $\text{FCER1G}$ ends at 161188748 bp, suggesting that the gap between ctg4004 and ctg4005 in the pig FPC map is $\approx 328$ Kb [(161647639–161188748)/1.4].

In conclusion, a comparative RH map to estimate gap distance in the FPC (and sequence) map. Analysis of the genes and BACs adjacent to the gap between ctg4004 and ctg4005 (Fig. S1) identified two porcine genes, $\text{FCGR2B}$ and $\text{DUSP12}$, and one MS, UMNp591, (Fig. S1) that mapped to BAC CH242-447O4, located at the terminus of ctg4004 in the FPC database (http://www.sanger.ac.uk/cgi-bin/Projects/S_scrofa/WebFPCdirect.cgi?contig=4004&clone=CH242-447O4). In contrast, the pig $\text{NDUFS2}$ and $\text{FCER1G}$ genes were located in BAC CH242-28G15, which maps to ctg4005 and overlaps with 310B20A04 (http://www.sanger.ac.uk/cgi-bin/Projects/S_scrofa/WebFPCdirect.cgi?contig=4005&clone=CH242-28G15). The ctg4004/ctg4005 boundary corresponds to a region of conserved synteny on HSA8. Human $\text{FCGR2B}$ starts at 161647639 bp, while $\text{FCER1G}$ ends at 161188748 bp, suggesting that the gap between ctg4004 and ctg4005 in the pig FPC map is $\approx 328$ Kb [(161647639–161188748)/1.4].

In conclusion, a comparative mapping approach allowed us to integrate results from the IMNpRH2 12 000-rad and IMpRH7 000-rad panels, RNA-seq data, genetic and BAC FPC maps, to estimate the size of the remaining contigs on SSC4, to identify the gaps between porcine BAC contigs along the tiling path of SSC4, and to aid sequence assembly.

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References


integrated RH map of porcine chromosome 10. BMC Genomics 10, 211.


Supporting information

Additional supporting information may be found in the online version of this article.

Figure S1 High-resolution RH comprehensive and comparative maps of porcine chromosome (SSC) 4.

Table S1 Mapping vectors used in this study.

Table S2. Markers in the SSC4 12 000- and 7 000-rad RH maps.

Table S3. SSC4 marker information.

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