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Single nucleotide polymorphism discovery in bovine liver using RNA-seq technology

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Abstract

Background

RNA-seq is a useful next-generation sequencing (NGS) technology that has been widely used to understand mammalian transcriptome architecture and function. In this study, a breed-specific RNA-seq experiment was utilized to detect putative single nucleotide polymorphisms (SNPs) in liver tissue of young bulls of the Polish Red, Polish Holstein-Friesian (HF) and Hereford breeds, and to understand the genomic variation in the three cattle breeds that may reflect differences in production traits.

Results

The RNA-seq experiment on bovine liver produced 107,114,407 raw paired-end reads, with an average of approximately 60 million paired-end reads per library. Breed-wise, a total of 345.06, 290.04 and 436.03 million paired-end reads were obtained from the Polish Red, Polish HF, and Hereford breeds, respectively. Burrows-Wheeler Aligner (BWA) read alignments showed that 81.35%, 82.81% and 84.21% of the mapped sequencing reads were properly paired to the Polish Red, Polish HF, and Hereford breeds, respectively. This study identified 5,641,401 SNPs and insertion and deletion (indel) positions expressed in the bovine liver with an average of 313,411 SNPs and indel per young bull. Following the removal of the indel mutations, a total of 195,380, 152,712 and 205,318 raw SNPs expressed in bovine liver were identified for the Polish Red, Polish HF, and Hereford breeds, respectively. Breed-wise, three highly reliable breed-specific SNP-databases (SNP-dbs) with 31,562, 24,945 and 28,194 SNP records were constructed for the Polish Red, Polish HF, and Hereford breeds, respectively. Using a combination of stringent parameters of a minimum depth of ≥10 mapping reads that support the polymorphic nucleotide base and 100% SNP ratio, 4,368, 3,780 and 3,800 SNP records were detected in the Polish Red, Polish HF, and Hereford breeds, respectively. The SNP detections using RNA-seq data were...
Abbreviations: SNP, Single nucleotide polymorphism; RNA-seq, RNA sequencing; HF, Holstein-Friesian; KASP™, competitive allele-specific PCR; BWA, Burrows-Wheeler Aligner; MAS, marker assisted selection; GAS, gene assisted selection; GS, genomics selection; QTL, Quantitative trait locus; OTNs, quantitative trait nucleotides; DEGs, differentially expressed genes; CEGs, co-expressed genes; GBS, Genotyping-by-Sequencing; BTA, Bos Taurus autosome; indel, insertion and deletion; SNP-db, SNP database; HWE, Hardy Weinberg equilibrium; HT, high-throughput; NGS, next generation sequencing; RIN, RNA Integrity Number; QC, quality control; WGS, whole genome sequence; GWAS, genome-wide association; eQTL, expression quantitative trait loci; TAGs, trait associated genes; CTNS, cystinosin, lysosomal cystine transporter; PAH2A, prolyl 4-hydroxylase subunit alpha 2; IQGAP, IQ motif containing GTPase activating protein homologue; CGs, candidate genes; IGFBP2, insulin-like growth factor 2 binding protein 2; GAD1, Glutamate decarboxylase 1; IGFBP2, insulin-like growth factor binding protein 2; IGFBP5, insulin-like growth factor binding protein 5; MSTN, myostatin; LEP, leptin receptor; APOA2, apolipoprotein A2; SLC44A5, solute carrier family 44, member 5; NTNG1, neurotrophin 3 receptor; IGFBP1, insulin-like growth factor binding protein 1; IgFBP3, insulin-like growth factor binding protein 3; LEP, leptin; NPY, neuropeptide Y; NSIG1, insulin induced gene 1; DPP6, dipeptidyl aminopeptidase-like protein 6; HGF, hepatocyte growth factor; IGFBP6, insulin-like growth factor binding protein 6; IGF-1, insulin-like growth factor 1; WNT10B, wingless-type MMTV integration site family member 10B; MYF5, myogenic factor 5; MYF6, myogenic factor 6; IGFBP7, insulin-like growth factor binding protein 7; CCSER1, coiled-coil serine-rich protein 1; PKD2, polycystic kidney disease 2; NCAPG, non-SMC condensin I complex subunit G; KONIP4, k channel interacting protein 4; CAST, calpastatin; NPM1, nucleophosphin; PROP1, PROP paired-like homeobox 1; IGFBPL1, insulin-like growth factor binding protein-like 1; ATP6V1B2, ATPase, H+ transporting, lysosomal 9QA, V1 subunit B2; IGFR2, insulin-like growth factor 2 receptor; RXF6, regulatory factor X 6; MAP3K5, mitogen-activated protein kinase 5; GTF3C5, general transcription factor IIIc, polypeptide 5; RALGDS, ras association; SDC1, syndecan 1; DMNT3B, DNA (Cytosine-5-) methyltransferase 3 beta; KSRH, growth hormone releasing hormone; FOXA2, forkhead box A2; HNF4A, hepatocyte nuclear factor 4, alpha; successfully validated by kompetitive allele-specific PCR (KASP™) SNP genotyping assay. The comprehensive QTL/CG analysis of 110 QTL/CG with RNA-seq data identified 20 monomorphic SNP hit loci (CARTPT, GAD1, GDF5, GHRH, GHR, GRB10, IGFBPL1, IGFL1, LEP, LHX4, MC4R, MSTN, NKA1, PLAG1, POU1F1, SDR16C5, SH2B2, TOX, UCP3 and WNT10B) in all three cattle breeds. However, six SNP loci (CCSER1, GHR, KCNNP4, MTSS1, EGFR and NSMCE2) were identified as highly polymorphic among the cattle breeds.

Conclusions

This study identified breed-specific SNPs with greater SNP ratio and excellent mapping coverage, as well as monomorphic and highly polymorphic putative SNP loci within QTL/CGs of bovine liver tissue. A breed-specific SNP-db constructed for bovine liver yielded nearly six million SNPs. In addition, a KASP™ SNP genotyping assay, as a reliable cost-effective method, successfully validated the breed-specific putative SNPs originating from the RNA-seq experiments.

Background

With the advancement of high-throughput (HT) NGS technology, transcriptome complexity and its dynamics can now be revealed and explored at different levels. Over the past few years, several sequencing-based technologies have been developed to analyse the transcriptomes in an unprecedented manner, and have revolutionized human and animal genome research [1]. Currently, the most widely used HT-RNA sequencing (RNA-seq) technology utilizes the NGS reads of the entire transcriptome including all transcripts produced in a tissue sample, which was not previously characterized as transcribed sequences and novel isoforms. Moreover, the most important potential applications of RNA-seq technology include the identification of differentially expressed genes (DEGs), co-expressed genes (CEGs) and differences in single nucleotide polymorphism (SNP) variation between experimental groups, such as samples from different (i) tissues (tissue-specific RNA-seq experiments) [2–4], (ii) treatment groups (trait-specific or trait-associated RNA-seq experiments) [5, 6], and (iii) populations (population-based breed-specific RNA-seq experiments) [7–9]. In this study, we utilized an experimental design similar to that described in our recent RNA-seq study on bovine pituitary tissue [9].

Most of the recent studies detecting SNPs using RNA-seq in domestic animals have mainly been focused on the identification of a large number of polymorphisms, with the aim of discovering causative variants involved in phenotypes affecting economic traits of interest in different domestic animal species, e.g., sheep [10], goat [11], pig [12], horse [13], chicken [14] and cattle [15–20]. Furthermore, SNP markers have increasingly been used in cattle breeding improvement programmes, e.g., marker-assisted selection (MAS), gene-assisted selection (GAS) and genomic selection (GS) [21–23], as a means of conventionally improving phenotypic selection. It is noteworthy that SNP detection for economic traits has great potential in the genetic improvement of cattle through the implementation of MAS, GAS and GS programmes, which have been highly recommended to the global cattle breeding programme.

In this study, we have chosen to perform RNA-seq on bovine liver tissue because of its highly robust metabolic activity [24], and because it is one of the most common target organ sites for body growth, feed utilization or feed efficiency and developmental trait assessments.
Results

mRNA sequencing and read alignment

mRNA sequencing of bovine liver at single-nucleotide resolution was carried out using two biological replicates of poly(A)-enriched mRNA of young bulls aged 6, 9, and 12 months from three cattle breeds. These mRNA samples were first converted into barcoded strand-specific dUTP RNA-seq libraries, followed by HT sequencing on the Illumina NextSeq 500 sequencer. The HT sequencing produced a total of 107,114,407 raw paired-end reads with a length of 156 bases. The reads were de-multiplexed to assign reads to each sequenced sample according to its index. The FASTQ sequence dataset of each library (Table 1) was submitted to the NCBI-SRA database with NCBI-SRA experiment number SRS1296732 (https://www.ncbi.nlm.nih.gov/sra?linkname=bioproject_sra_all&from_uid=312148).

Using the Burrows-Wheeler Aligner (BWA) program under default conditions, the breed-specific liver transcripts were mapped to the bovine reference genome (UMD3.1 assembly plus Y chromosome). Our read alignment results showed that 98.55% sequencing reads (105,567,9630) were successfully aligned to the UMD3.1 bovine reference genome. Furthermore, the acquired BWA mapping profile results showed that 81.35%, 82.81% and 84.21% of the mapped sequencing reads were properly paired for the Polish Red, Polish HF, and Hereford breeds, respectively (Tables 2–4).

SNP discoveries in cattle breeds

Breed-specific raw SNP-db records. Using the SAMtool package, a total of 5,641,401 (~5.6 million) breed-specific SNPs and indel positions expressed in the bovine liver were detected with the RNA-seq reads, with an average of 313,411 (~0.31 million) SNPs and indels per young bull (Table 5). Breed-wise, this raw SNP-db comprised 1,995,571 (35.4%), 1,556,048 (27.6%), and 2,089,782 (37%) SNPs and indels for the Polish Red, Polish HF, and Hereford breeds, respectively. Following the removal of the indel mutations, a total of 1,953,804 (35.3%), 1,527,120 (27.6%), and 2,053,184 (37.1%) raw SNPs expressed in bovine liver were recovered from the Polish Red, Polish HF, and Hereford breeds, respectively. Using the SAMtool package, single-base substitutions (SNPs) and small indels were also identified. In this study, a total of 41,767, 36,604 and 28,934 indel mutations were identified in the Polish Red, Polish HF, and Hereford breeds, respectively.

SNP distribution in Venn plot. In our initial SNP analyses, a stringent filtering parameter of read count with a minimum depth of 5 SNP reads that support the polymorphic nucleotide base and existed in both replicates was utilized to allow the identification of approximately 0.8 million SNPs among the three cattle breeds as shown in Venn diagram (Fig 1).

Breed-specific SNP-db records. For the detection of breed-specific putative SNPs expressed in bovine liver, only the records in the raw SNP-db (S1–S18 Tables) were combined into one file to construct a highly reliable SNP-db with 84,701 SNP hit records. Three highly reliable breed-specific SNP-dbs comprising 31,562 (37.27%), 24,945 (29.45%) and 28,194
SNP records were constructed for the Polish Red, Polish HF, and Hereford breeds, respectively (S19–S21 Tables).

**Error removals:** During SNP detection using SAMtool, some records were observed as more than one SNP mutation. Such records were considered as error records and were excluded from the SNP-db. In our study, a total of 4381, 1164 and 1202 such error records in the Polish Red, Polish HF and Hereford breeds were observed and excluded. After removal of the error records, a total of 27,182, 23,781 and 26,992 SNP-db records were recovered from the Polish Red, Polish HF and Hereford breeds, respectively (S22–S24 Tables).

**SNP filtering.** We utilized the stringent parameter of a minimum depth of 10 SNP reads that support the polymorphic nucleotide base with a SNP ratio of 100%, because the SNP filtering criteria of 10 SNP reads with a SNP ratio of 100% could cover and explain all the HT SNP variations compared with the SNP filtering criteria of ≥10 SNP reads with a SNP ratio of 90% [9].

Initially, for the SNP filtering analysis, we utilized stringent parameters with a minimum depth of ≥10 SNP reads that support the polymorphic nucleotide base and identified 15,197, 11,346 and 12,455 SNP records for the Polish Red, Polish HF, and Hereford breeds, respectively (S25–S27 Tables). Similarly, by utilizing stringent parameters of 100% SNP ratios, we identified 10,206, 9,684 and 9,778 SNP records for the Polish Red, Polish HF, and Hereford breeds, respectively (S28–S30 Tables). Finally, the combination of both stringent filtering parameters of ≥10 reads, and a 100% SNP ratio, yielded a total of 4,368, 3,780 and 3,800 SNP records for the Polish Red, Polish HF, and Hereford breeds, respectively (S31–S33 Tables).

**Breed comparison.** Comparison of the breed-specific SNP records resulted in the identification of common and unique SNPs. For each breed, three comparisons were made to find common and unique SNPs within the investigated breeds. For example, i) a comparison between breed-1 (Polish Red) with breed-2 (Polish HF); ii) a comparison between breed-1 and breed-3 (Hereford); and iii) a comparison of breed-1 to both breed-2 and breed-3.

In this study, a total of 50 and 81 unique SNP loci were identified in the Polish Red breed that were not present in either the Polish HF or Hereford breeds, respectively (S34 and S35 Tables). In addition, only six unique SNP records were identified in the Polish Red breed that was not detected in either Polish HF or Hereford breeds (S36 Table).

Similarly, the SNP data comparison of the Polish HF breed to other breeds resulted in the identification of 7 and 22 unique SNPs that were not present in the Polish Red or Hereford breeds, respectively (S37 and S38 Tables, respectively). However, no single unique SNP record was identified in the Polish HF breed that was not present in either the Polish Red or Hereford breeds, respectively (S39 Table). Finally, the SNP data comparison of the Hereford breed to other breeds identified a total of 80 and 41 unique SNPs that were not detected in Polish Red or Polish HF breeds, respectively (S40 and S41 Tables). Furthermore, a total of nine unique SNP records of the Hereford breed were identified, which were not present in either the Polish Red or Polish HF breeds (S42 Table).

**De novo SNPs.** Deep and extensive SNP analysis provided evidence that certain regions of the *Bos taurus* genome were still unknown (base = N, according to recent mapping to UMD3.1). Using the SNP filtering criteria (≥10 SNP reads with SNP ratio of 100%), a total of 217, 193 and 265 best candidates of de novo SNP reads were identified for the Polish Red, Polish HF, and Hereford breeds, respectively (S43–S45 Tables).

**Breed-specific SNP discovery and QTL/CG analysis**

For the QTL/CG analysis on identified breed-specific SNP data, a set of 110 QTL/CG loci (http://www.animalgenome.org/cgi-bin/QTLdb/index) (S46 Table) was comprehensively
investigated for each breed and ages of the young bulls (S47–S52 Tables). We summarized, for each breed, the chromosomal locations and SNP locations of identified putative SNP loci hits of RNA-seq data on 110 potential CGs from the bovine QTL-db (S47–S49 Tables).

Table 1. Description of submitted FASTQ sequences of bovine liver of all 18 young bulls from three cattle breeds using RNA-seq.

<table>
<thead>
<tr>
<th>Breed</th>
<th>Age</th>
<th>Animal ID</th>
<th>Library name</th>
<th>SRA Run</th>
<th>MBases</th>
<th>Mbytes</th>
<th>SRA Experiment</th>
<th>SRA accession no.</th>
<th>FastQ file size</th>
</tr>
</thead>
<tbody>
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<td>6m</td>
<td>6938</td>
<td>CP19</td>
<td>SRR3176171</td>
<td>8488</td>
<td>4041</td>
<td>SRRX1590547</td>
<td>SRX1590547</td>
<td>1 ULLUMINA (NextSeq 500) run: 28.8M spots, 8.9G bases, 3.9Gb downloads</td>
</tr>
<tr>
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<td>CP27</td>
<td>SRR3176230</td>
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<td>1834</td>
<td>SRRX1590600</td>
<td>SRX1590600</td>
<td>1 ULLUMINA (NextSeq 500) run: 13.4M spots, 4.1G bases, 1.8Gb downloads</td>
</tr>
<tr>
<td>Herford</td>
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<td>CP35</td>
<td>SRR3176234</td>
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<td>SRX1590604</td>
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<td>8603</td>
<td>CP29</td>
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</tr>
<tr>
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</tr>
</tbody>
</table>

Table 2. Transcriptome mapping profile of liver tissue from the Polish Red cattle breed aligned to bovine reference UMD3.1 genome assembly.

<table>
<thead>
<tr>
<th>Age</th>
<th>Total</th>
<th>Mapped</th>
<th>Paired in sequencing</th>
<th>read1</th>
<th>read2</th>
<th>Properly paired</th>
<th>With itself and mate mapped</th>
<th>Singletons</th>
<th>With mate mapped to a different chr (map Q &gt; 5)</th>
<th>With mate mapped to a different chr (map Q = 5)</th>
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<td>9m</td>
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<td>3987897</td>
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</tr>
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<td>345069302</td>
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</table>

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doi:10.1371/journal.pone.0172687.t002
Tables, we further summarized the numbers of putative SNP hits of RNA-seq data, identified within breed (ages) and between breeds. Overall, we have identified i) monomorphic breed-specific SNPs, and ii) highly polymorphic breed-specific SNPs within the investigated 110 QTL/CGs loci (S50–S52 Tables).

**Identification of breed-specific monomorphic SNP loci in bovine QTL/CG db.** Breed-wise, a total of 32, 28, and 29 monomorphic SNP hits on 110 QTL/CG genes were identified in the Polish Red, Polish HF and Hereford breeds, respectively. Among the identified monomorphic SNP hit loci, 20 CGs (CARTPT, GAD1, GDF5, GHRH, GRB10, IGFBPL1, IGFL1, LEP, LHXA, MC4R, MHC, NKAIN1, PLAG1, POU1F1, SDR16C5, SH2B2, TOX, UCP3 and WNT10B) were monomorphic to all investigated breeds. Furthermore, SNP hits in the BMP8B, GHSR, and RFX6 CGs were monomorphic only to the Polish Red and Polish HF breeds, while a SNP hit in the GDF5 CG was monomorphic only to the Polish HF breed, and SNP hits in the GHRHR and IGFL1 CGs were monomorphic only to the Hereford breed.

**Identification of highly polymorphic breed-specific SNP loci in the bovine QTL/CG db.** Highly polymorphic breed-specific SNP loci were summarized as top-10 and top-20 SNP hits (S50–S52 Tables). Among the top 10 SNP hits, five CGs (CCSER1, GHR, KCNIP4, MTSS1, EGFR and NSMCE2) were highly polymorphic to all three cattle breeds. Furthermore, SNP hits in the IGF-I CG were highly polymorphic to the Polish Red and Polish HF breeds, SNP hits in the CAST CG were highly polymorphic to the Polish HF and Hereford breeds, and SNP hits in the LEPR CG were highly polymorphic to the Polish Red and Hereford breeds. In addition, SNP hits in the BMP8B, GHSR, and RFX6 CGs were monomorphic only to the Polish Red and Polish HF breeds, whereas SNP hits in the DLK2, MYF5, and PROP1 CGs were monomorphic to both the Polish HF and Hereford breeds. Monomorphic SNP loci in the BTG4, NPM1, NPy and SIX3 CGs were detected in the Polish Red and Hereford breeds. In addition, SNP hits in the AMPD1, CAPN3, MYF6 and SDR16C6 CGs were monomorphic only to the Polish Red breed, while a SNP hit in the GDP10 CG was monomorphic only to the Polish HF breed, and SNP hits in the GHRHR and IGFL1 CGs were monomorphic only to the Hereford breed.

### Table 3. Transcriptome mapping profile of liver tissue from the Polish HF cattle breed aligned to bovine reference UMD3.1 genome assembly.

<table>
<thead>
<tr>
<th>Age</th>
<th>Total</th>
<th>Mapped</th>
<th>Paired in sequencing</th>
<th>read1</th>
<th>read2</th>
<th>Properly paired</th>
<th>With itself and mate mapped</th>
<th>Singletons</th>
<th>With mate mapped to a different chr (map Q &gt; 5)</th>
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</table>

doi:10.1371/journal.pone.0172687.t003

### Table 4. Transcriptome mapping profile of liver tissue from the Hereford cattle breed aligned to bovine reference UMD3.1 genome assembly.

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<th>Age</th>
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<th>read2</th>
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doi:10.1371/journal.pone.0172687.t004
Phylogenetic analysis of the breed-specific SNP-db

Additionally, a phylogenetic tree was constructed using a JTT matrix-based model with superior log maximum likelihood values [28] and molecular evolutionary genetics analysis version 7 (MEGA7) software [29] to examine the close relationship of the investigated bulls within and between breeds. Results showed that all three cattle breeds were separated from each other, but clustered together for each breed. The percentage of trees in which the associated taxa clustered together is shown next to the branches (Fig 2).

Breed-specific SNP validation

Selection of nine breed-specific putative SNPs. A breed-specific SNP validation experiment was carried out using a subset of nine putative SNPs derived from the Polish Red (n = 2), Polish HF (n = 4) and Hereford (n = 3) breeds, on single-plex KASP™ genotyping assays (LGC Genomics) based on fluorescently labelled allele-specific PCR primers (Table 6). The selection of nine putative breed-specific SNPs was performed according to the RNA-seq experimental results (S53 Table). We selected three SNPs that were uniquely expressed in only one cattle breed (CTNS and P4HA2 SNP gene loci specific to the Polish Red breed and IQGAP2 SNP gene locus specific to the Hereford breed), and the remaining six SNP gene loci that were specific to all the breeds (GHR, IGF2R, IGF2BP3, IGFBP4 SNP gene loci specific to the Polish HF breed, and GHR and IGF2 SNP gene loci specific to the Hereford breed, respectively). Based on the RNA-seq experimental results, detailed information of the primer design of the nine selected breed-specific SNP loci, such as the candidate genes and their symbols, genome
locations, SNP positions, UMD 3.1 chromosome and gene map positions, and both 5’ and 3’ flanking regions of 200 bp sequences at the SNP mutation site, is illustrated in S54 Table.

The selected subset of the nine breed-specific putative SNP markers from the Polish Red, Polish HF and Hereford breeds worked well in KASP™ SNP genotyping assay and did not reveal either non-amplification or ambiguous clustering, except for a few samples due to poor DNA quality (Figs 3–11).

**KASP™ SNP assay analysis.** Initially, an additional analysis for the estimation of a direct relationship between breed and the detected SNP polymorphisms was performed using the PROC MIXED SAS 9.2 package with age as a random effect. Two SNP markers, *IGFBP4* (Polish HF) and *IGF2* (Hereford) genes were identified as homozygous at the SNP loci, and therefore were excluded for further statistical SNP validation analysis (S55 Table). The results of the remaining seven SNP loci did not show any significant association between age and SNPs selected for the validation. However, there were significant and highly significant associations.
Fig 2. The phylogenetic relationship among RNA-seq samples of bovine liver using Maximum likelihood method based on the JTT matrix-based model. An unrooted phylogeny tree of 18 bulls’ samples representing developmental ages 6 months (6A, 6B), 9 months (9A, 9B), and 12 months (12A, 12B) of Polish Red, Polish-HF and Hereford breeds. The percentage of trees in which the associated taxa clustered together is shown next to the branches. All nodes were robust at 100% bootstrap support. The scale bar denotes substitutions per site.

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between breed and the selected SNPs (S56 Table). Furthermore, a chi-square analysis showed significant differences in genotypes and allele frequencies for the **CTNS** gene SNP locus specific to the Polish Red breed, **IGF2R** gene SNP locus specific to the Polish HF breed, and the **GHR** and **IQGAP2** genes SNP loci specific to the Hereford breed (S57 Table).

### Table 6. The distribution (n) of investigating young bulls in a breed-specific experimental design representing bovine liver transcriptome.

<table>
<thead>
<tr>
<th>Breeds</th>
<th>RNA-seq experiment</th>
<th></th>
<th></th>
<th></th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Breeds</td>
<td>6 months</td>
<td>9 months</td>
<td>12 months</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hereford</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Polish HF</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Polish Red</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>6</td>
<td>6</td>
<td>6</td>
<td>18</td>
<td></td>
</tr>
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<td>SNP validation</td>
<td>RNA-seq experiment</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Breeds</td>
<td>6 months</td>
<td>9 months</td>
<td>12 months</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hereford</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td>Polish HF</td>
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<td>5</td>
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</tr>
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<tr>
<td>Total</td>
<td>15</td>
<td>14</td>
<td>15</td>
<td>44</td>
<td></td>
</tr>
</tbody>
</table>

doi:10.1371/journal.pone.0172687.t006

**Fig 3.** KASP™ SNP genotyping assay of BTA19_24970466 locus of Polish Red **CTNS** gene showing the data for single KASP™ assays on a single cluster plot.

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**Statistical analysis using Genepop software.** Using the Fisher’s Exact Probability test, the genetic differentiation of SNP alleles and SNP genotype results revealed significant differences in SNP allele frequencies for the *CTNS, IGF2R, GHR* (Polish HF breed), *IGF2BP3, GHR* (Hereford breed) and *IQGAP2* SNP loci (S58 Table). Regarding the genotype frequencies, significant differences were observed in the *CTNS, P4HA2, IGF2R, GHR* (Polish HF breed), *IGF2BP3, GHR* (Hereford breed), and *IQGAP2* SNP loci (S59 Table) in all investigated cattle breeds. Moreover, the genetic differentiation comparison of the SNP alleles (S60 Table) and genotypes (S61 Table) in all SNP loci and all investigated cattle breeds were also performed, using Fisher’s Exact Probability test and Fisher’s Exact G test. Results showed highly significant differences in the SNP alleles (S60 Table) and SNP genotypes (S61 Table) among breeds to validate the breed-specific SNP markers. For SNP validation statistics using a Markov chain method, the selected SNP markers were further examined by testing the deviation from Hardy Weinberg equilibrium (HWE) for each SNP locus (S62 Table), and for each breed population (S63 Table). The results based on genetic differentiation of the investigated SNPs across all loci showed significant differences in the *CTNS, IGF2R, IGF2BP3, GHR* (Polish HF), and *IQGAP2* SNP loci (S62 Table). Furthermore, the results based on genetic differentiation of the investigated SNPs across all breeds revealed significant differences among them (S63 Table).
Discussion

mRNA sequencing and read alignment

RNA-seq technology has great potential in identifying genetic variation at many loci, with respect to SNP polymorphisms and gene expression patterns across different organ tissues. In the present study, we identified approximately 107 million raw paired-end reads with an average length of 156 bases in bovine liver tissue compared to 113 million raw paired-end reads detected in the bovine pituitary gland [9]. This variation in the RNA-seq yield from different organ tissues might be due to technical variation, such as differences in the quality and quantity of the RNA recovered during tissue-specific sample preparations, batch effects in library preparation [30, 31], flow cell and lane effects caused by the Illumina sequencing platform, or adapter bias [32, 33]. However, both the liver and pituitary gland tissues revealed higher percentages (99.4% and 98.5%, respectively) of read alignment to the UMD3.1 bovine reference genome. Furthermore, using the BWA program under default conditions, showed that 94.39%, 93.04% and 83.46% of the mapped sequencing reads of the pituitary gland tissues were properly paired for the Polish Red, Polish HF, and Hereford breeds, respectively [9].
The FASTQ sequences of both tissues were submitted with the NCBI-SRA experiment number SRS1296732 (http://www.ncbi.nlm.nih.gov/sra?linkname=bioproject_sra_all&from_uid=312148). Currently, there are 24 SRA records (FASTQ sequence dataset) for bovine liver tissue at the SRA NCBI database [34] (https://trace.ncbi.nlm.nih.gov/Traces/sra/sra.cgi?view=studies&term=(bos%20taurus%20liver)%20NOT%20cluster_dbgap%5BPROP%5D).

**SNP discovery in cattle breeds**

One of the potential applications of the RNA-seq technology is to identify several thousands of SNPs and as well as to construct the tissue-specific SNP datasets. In our study, we constructed breed-specific SNP-db records for bovine liver tissue, with more than 5.6 million SNP records compared to 13.7 million SNP-db records for the bovine pituitary gland tissue [9]. The differences in the SNP data yield in tissue-specific RNA-seq experiments in the bovine liver and pituitary gland might be due to pre- and post-quality assessment of read alignments, *i.e.*, quality control (QC) metrics before alignment [35]. In most SNP discoveries based on bovine RNA-seq studies [7–9, 16–20], selection of stringent parameters in the SNP filtering process is undoubtedly a critical task for identifying the most reliable novel putative SNPs. Using the stringent parameter of a minimum depth of 10 SNP reads that support the polymorphic nucleotide base with a SNP ratio of 100%, we identified 20,573, 31,978 and 30,052 breed-specific

![Image](image.png)
SNP records for bovine pituitary gland in the Polish Red, Polish HF and Hereford breeds, respectively [9]. However, in the current study, we identified 31,562, 24,945 and 28,194 breed-specific SNP records for bovine liver tissue in the Polish Red, Polish HF and Hereford cattle breeds, respectively.

It should be emphasized that, in cattle breeding practices, identification of breed-specific gene-associated SNPs can serve as suitable markers for trait-associated studies and can be effectively utilized in genomic selection (GS) programmes [36]. In our study, we investigated 110 QTL/CG loci to identify novel putative gene-associated SNPs within a breed, with respect to the animal’s age, and between breeds. Results based on within-breed QTL/CG analysis revealed that approximately one third of the selected QTL/CG loci were monomorphic in all investigated breeds. These findings indicate that the dairy and beef cattle breeds might have certain genetic selection signatures due to fixation of certain genotypes (monomorphic SNPs). It is noteworthy that, between breeds, the QTL/CG analysis identified CCSER1, GHR, KCNIP4, MTSS1, EGFR and NSMCE2 gene loci as highly polymorphic. Moreover, analysis of the 76 QTL/CGs in bovine pituitary gland RNA-seq data demonstrated that KCNIP4, CCSER1, DPP6, MAP3K5 and GHR genes loci were highly polymorphic in all the investigated cattle breeds [9]. These results indicate that there is still an abundant genetic variation in the Polish dairy and
beef cattle breeds due to the high degrees of polymorphism in the CGs of economically important traits, which could be exploited in GS and other breeding programmes.

In cattle, many studies have utilized QTL/CG analysis with respect to whole genome sequence (WGS) data, genome-wide association study (GWAS) data and expression quantitative trait loci (eQTL) data, to investigate the causal relationships of variant-trait and variant-gene expression, respectively [37–40]. Using human and bovine data, Seo et al. [5] have successfully demonstrated the application of the association test approach based on RNA-seq analysis to identify trait associated genes (TAGs).

In this study, we have validated and revealed that breeds of the investigated bulls were separated from each other using an ITT matrix based maximum likelihood model [28] and MEGA 7 [29]. With regards to bovine pituitary gland tissue, we successfully utilized another phylogenetic analysis method using the SNPphylo model [41], which revealed that the three cattle breeds were clustered together for each breed and were separated from each other [9].

Finally, the utilization of KASP™ genotyping assay technology in our SNP validation experiment has confirmed the presence of seven breed-specific SNP markers, which are expressed in bovine liver tissue. In addition, a similar SNP experiment has been successfully performed to validate the expression of the seven SNP markers in bovine pituitary tissue [9].
Thus, the SNP experiment using KASP<sup>TM</sup> genotyping assay technology is an effective and reliable method to validate RNA-seq in various bovine organ tissues.

**Conclusions**

Breed-specific SNP discovery using NGS-based RNA-seq in bovine liver tissue has been reported to provide a global view of the complexity of the bovine liver transcriptome. Our study has developed three breed-specific SNP-dbs based on expressed genes in the bovine liver, which might provide valuable resources for trait-associated genomic and genome-wide association studies. Our study has demonstrated the utility of QTL/CG analysis on RNA-seq SNP data to identify putative trait-associated SNPs from the bovine QTL-db. Our phylogenetic analyses have shown that all three cattle breeds were separated from each other long ago and that each breed is represented as unique genetic clusters. The transcriptome sequencing (RNA-seq) technique employed in the current study is similar to the Genotyping-by-Sequencing (GBS) and it has good potential for improving the accuracy of genomic selection because it involves several millions of SNPs covering the entire transcriptome, which increases the chance of identifying QTLs or quantitative trait nucleotides (QTNs) in linkage disequilibrium with SNPs. Furthermore, these results suggest that there are wide differences in the liver...
transcriptomes between the breeds, which could be useful to study the mechanisms underlying genetic variability in meat quality and other production traits in cattle.

Materials and methods

Experimental design

The bovine liver tissue samples were collected from 18 young bulls of three cattle breeds (Table 6) stationed at the Institute of Genetics and Animal Breeding (IGAB), Polish Academy of Science (PAS), Jastrzębiec, Poland. The investigated animals were selected randomly, and after the purchase at birth, they were reared in the experimental farm of IGAB, PAS, Jastrzębiec, Poland, with uniform environmental and feeding conditions. After slaughtering, the collected liver tissues were immediately kept in liquid nitrogen, and stored at -80°C. All procedures involving animals were conducted in accordance with the guiding principles for the care and use of research animals. The investigating research materials were approved by the local ethics commission of IGAB, PAS, Jastrzębiec, Poland (permission No. 3/2005). The experimental designs, with detailed laboratory procedures including isolation of total RNA, library preparation and NGS sequencing using the Illumina NextSeq 500 High Output/300
cycle platform and kits (Illumina), were performed as previously described in a recent study [9].

Bioinformatics analysis

Complete bioinformatics analysis, including read alignment of RNA-seq data to the reference genome, breed-specific SNP detections, QTL/CG analysis, phylogenetic analysis, SNP validation by KASP™ genotyping assay, was performed as previously described in a recent study [9]. However, i) the SNP filtering criterion, ii) the number of QTL/CG loci to investigate the RNA-seq SNP-db, and iii) SNP markers selected for SNP validation by KASP™ genotyping assay were slightly modified from our recent bovine pituitary gland RNA-seq study [9].

**SNP filtering criterion.** To construct the bovine liver SNP-db, stringent parameters of a minimum depth of 10 SNP reads that support the polymorphic nucleotide base with a SNP ratio of 100%, were carried out using Microsoft office excel program in the following manner:

- Stringent parameter of SNP filtering with minimum depth of 10 SNP reads that support the polymorphic nucleotide base,
- Stringent parameter of SNP filtering with a SNP ratio of 100%,
Combining the stringent parameters of a minimum depth of 10 SNP reads that support the polymorphic nucleotide base with a SNP ratio of 100%.

Using Microsoft office excel program, the bovine liver SNP-dbs of the 18 young bulls were combined according to breed and were further trimmed to one SNP-db that was highly reliable and specific to the three breeds.

**Breed-specific SNP discovery and QTL/CG analysis**

Based on a publicly available animal QTL database (http://www.animalgenome.org/cgi-bin/QTLdb/index), a total of 110 potential QTL/CGs (S46 Table) for bovine body growth and developmental trait were included to investigate the RNA-seq SNP-db of bovine liver using Microsoft Office Excel.

**Phylogenetic analysis.** The phylogenetic analysis of the breed-specific SNP-dbs of bovine liver transcriptome was inferred by using the Maximum Likelihood method based on the JTT matrix-based model [28]. The phylogenetic tree(s) of the 18 bull samples representing the Polish Red, Polish HF and Hereford breeds were constructed, firstly with heuristic search by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using a JTT model, and then by selecting the topology with superior log likelihood value. Finally, the evolutionary analyses were conducted in MEGA7 [29].

**SNP validation by KASP™ genotyping assay**

The primer sequences of nine breed-specific SNPs from the Polish Red (n = 2), Polish HF (n = 4), Hereford (n = 3) breeds selected for KASP™ SNP genotyping assay to test the SNP validation experiment are presented in Table 7. The complete methodological procedure of the KASP™ genotyping assay, including statistical analysis using Genepop software (http://genepop.curtin.edu.au) was performed as previously described in a recent study [9].

**Supporting information**

S1 Table. RNA-seq SNP-db of bovine liver tissue of young bull-1 of Polish Red cattle aged 6 months.

(XLSX)

S2 Table. RNA-seq SNP-db of bovine liver tissue of young bull-2 of Polish Red cattle aged 6 months.

(XLSX)
S3 Table. RNA-seq SNP-db of bovine liver tissue of young bull-3 of Polish Red cattle aged 9 months.
(XLSX)

S4 Table. RNA-seq SNP-db of bovine liver tissue of young bull-4 of Polish Red cattle aged 9 months.
(XLSX)

S5 Table. RNA-seq SNP-db of bovine liver tissue of young bull-5 of Polish Red cattle aged 12 months.
(XLSX)

S6 Table. RNA-seq SNP-db of bovine liver tissue of young bull-6 of Polish Red cattle aged 12 months.
(XLSX)

S7 Table. RNA-seq SNP-db of bovine liver tissue of young bull-7 of Polish HF cattle aged 6 months.
(XLSX)

S8 Table. RNA-seq SNP-db of bovine liver tissue of young bull-8 of Polish HF cattle aged 6 months.
(XLSX)

S9 Table. RNA-seq SNP-db of bovine liver tissue of young bull-9 of Polish HF cattle aged 9 months.
(XLSX)

S10 Table. RNA-seq SNP-db of bovine liver tissue of young bull-10 of Polish HF cattle aged 9 months.
(XLSX)

S11 Table. RNA-seq SNP-db of bovine liver tissue of young bull-11 of Polish HF cattle aged 12 months.
(XLSX)

S12 Table. RNA-seq SNP-db of bovine liver tissue of young bull-12 of Polish HF cattle aged 12 months.
(XLSX)

S13 Table. RNA-seq SNP-db of bovine liver tissue of young bull-13 of Hereford cattle aged 6 months.
(XLSX)

S14 Table. RNA-seq SNP-db of bovine liver tissue of young bull-14 of Hereford cattle aged 6 months.
(XLSX)

S15 Table. RNA-seq SNP-db of bovine liver tissue of young bull-15 of Hereford cattle aged 9 months.
(XLSX)

S16 Table. RNA-seq SNP-db of bovine liver tissue of young bull-16 of Hereford cattle aged 9 months.
(XLSX)
S17 Table. RNA-seq SNP-db of bovine liver tissue of young bull-17 of Hereford cattle aged 12 months.
(XLSX)
S18 Table. RNA-seq SNP-db of bovine liver tissue of young bull-18 of Hereford cattle aged 12 months.
(XLSX)
S19 Table. SNPs filtering data set of Polish Red breed.
(XLSX)
S20 Table. SNPs filtering data set of Polish HF breed.
(XLSX)
S21 Table. SNPs filtering data set of Hereford breed.
(XLSX)
S22 Table. SNPs filtering data set with no errors of Polish Red breed.
(XLSX)
S23 Table. SNPs filtering data set with no errors of Polish HF breed.
(XLSX)
S24 Table. SNPs filtering data set with no errors of Hereford breed.
(XLSX)
S25 Table. SNPs filtering data set with a minimum depth of 10 SNP reads that support the polymorphic nucleotide base of Polish Red breed.
(XLSX)
S26 Table. SNPs filtering data set with a minimum depth of 10 SNP reads that support the polymorphic nucleotide base of Polish HF breed.
(XLSX)
S27 Table. SNPs filtering data set with a minimum depth of 10 SNP reads that support the polymorphic nucleotide base of Hereford breed.
(XLSX)
S28 Table. SNPs filtering data set with SNP ratio = 100% of Polish Red breed.
(XLSX)
S29 Table. SNPs filtering data set with SNP ratio = 100% of Polish HF breed.
(XLSX)
S30 Table. SNPs filtering data set with SNP ratio = 100% of Hereford breed.
(XLSX)
S31 Table. SNPs filtering data set with a minimum depth of 10 SNP reads that support the polymorphic nucleotide base and SNP ratio = 100% of Polish Red breed.
(XLSX)
S32 Table. SNPs filtering data set with a minimum depth of 10 SNP reads that support the polymorphic nucleotide base and SNP ratio = 100% of Polish HF breed.
(XLSX)
S33 Table. SNPs filtering data set with a minimum depth of 10 SNP reads that support the polymorphic nucleotide base and SNP ratio = 100% of Hereford breed.
(XLSX)

S34 Table. Breed comparison in SNPs filtering data set of Polish Red with Polish HF breed.
(XLSX)

S35 Table. Breed comparison in SNPs filtering data set of Polish Red with Hereford breed.
(XLSX)

S36 Table. Breed comparison in SNPs filtering data set of Polish Red with Polish HF and Hereford breeds.
(XLSX)

S37 Table. Breed comparison in SNPs filtering data set of Polish HF with Polish Red breed.
(XLSX)

S38 Table. Breed comparison in SNPs filtering data set of Polish HF with Hereford breed.
(XLSX)

S39 Table. Breed comparison in SNPs filtering data set of Polish HF with Polish Red and Hereford breeds.
(XLSX)

S40 Table. Breed comparison in SNPs filtering data set of Hereford with Polish Red breed.
(XLSX)

S41 Table. Breed comparison in SNPs filtering data set of Hereford with Polish HF breed.
(XLSX)

S42 Table. Breed comparison in SNPs filtering data set of Hereford with Polish Red and Polish HF breeds.
(XLSX)

S43 Table. Identification of De novo SNPs data set in Polish Red breed.
(XLSX)

S44 Table. Identification of De novo SNPs data set in Polish HF breed.
(XLSX)

S45 Table. Identification of De novo SNPs data set in Hereford breed.
(XLSX)

S46 Table. Lists of 110 SNP loci within the CGs bovine growth trait with representing full names, UMD3.1 genome locations, chromosomal locations and web links at bovine QTL-DB.
(XLSX)

S47 Table. Identification of putative SNPs hits of liver RNA-seq data on 110 potential candidate genes from bovine QTL-db in Polish Red cattle.
(XLS)

S48 Table. Identification of putative SNPs hits of liver RNA-seq data on 110 potential candidate genes from bovine QTL-db in Polish HF cattle.
(XLS)
S49 Table. Identification of putative SNPs hits of liver RNA-seq data on 110 potential candidate genes from bovine QTL-db in Hereford cattle.
(XLS)

S50 Table. Summary of identified putative SNPs hits of liver RNA-seq data on 110 potential candidate genes from bovine QTL-db in Polish Red cattle.
(XLS)

S51 Table. Summary of identified putative SNPs hits of liver RNA-seq data on 110 potential candidate genes from bovine QTL-db in Polish HF cattle.
(XLS)

S52 Table. Summary of identified putative SNPs hits of liver RNA-seq data on 110 potential candidate genes from bovine QTL-db in Hereford cattle.
(XLS)

S53 Table. RNA-seq experimental results of nine selected breed-specific SNPs loci utilized in SNP validation experiment.
(XLSX)

S54 Table. Primer design details of nine breed-specific SNPs generated from the bovine liver RNA-seq experiment.
(XLSX)

S55 Table. Distribution of KASP™ genotypes of nine selected breed-specific SNPs originating from the bovine liver RNA-seq experiment.
(XLSX)

S56 Table. Fixed effect of breeds and developmental ages on validated SNPs markers using REML mixed model procedure.
(XLSX)

S57 Table. SNPs genotypes and allele frequencies of selected breed-specific SNPs loci in all investigated cattle breeds.
(DOC)

S58 Table. Genetic differentiation of SNP alleles among investigated cattle breeds using the Fisher’s Exact Probability test.
(DOC)

S59 Table. Genetic differentiation of SNP genotypes among investigated cattle breeds using the Fisher’s Exact G test.
(DOC)

S60 Table. Genetic differentiation comparison of SNP alleles among cattle breeds using the Fisher’s Exact Probability test.
(DOC)

S61 Table. Genetic differentiation comparison of SNP genotypes among cattle breeds using the Fisher’s Exact G test.
(DOC)

S62 Table. Hardy-Weinberg test for genetic differentiation of investigated SNP loci using the Markov chain method.
(DOC)
S63 Table. Hardy-Weinberg test for genetic differentiation of investigated cattle breeds using the Markov chain method.

(DOC)

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Writing – review & editing: CSP HNK LF UC PS DK YF.

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