## 1 The Bovine Genome Variation Database (BGVD): Integrated Web-

## 2 database for Bovine Sequencing Variations and Selective Signatures

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- 16 **Running title:** Chen N et al / Bovine Variation and Selective Signature Database
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- 28
- 29 No. of words: 2048.
- 30 No. of figures: 4.
- 31 No. of tables: 1.

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### 33 Abstract

Next-generation sequencing has yielded a vast amount of cattle genomic data for the global 34 characterization of population genetic diversity and the identification of regions of the genome 35 under natural and artificial selection. However, efficient storage, querying and visualization of 36 such large datasets remain challenging. Here, we developed a comprehensive Bovine Genome 37 Variation Database (BGVD, http://animal.nwsuaf.edu.cn/BosVar) that provides six main 38 functionalities: Gene Search, Variation Search, Genomic Signature Search, Genome Browser, 39 Alignment Search Tools and the Genome Coordinate Conversion Tool. The BGVD contains 40 information on genomic variations comprising ~60.44 M SNPs, ~6.86 M indels, 76,634 CNV 41 regions and signatures of selective sweeps in 432 samples from modern cattle worldwide. Users 42 43 can quickly retrieve distribution patterns of these variations for 54 cattle breeds through an 44 interactive source of breed origin map using a given gene symbol or genomic region for any of the three versions of the bovine reference genomes (ARS-UCD1.2, UMD3.1.1, and Btau 5.0.1). 45 Signals of selection are displayed as Manhattan plots and Genome Browser tracks. To further 46 47 investigate and visualize the relationships between variants and signatures of selection, the Genome Browser integrates all variations, selection data and resources from NCBI, the UCSC 48 49 Genome Browser and AnimalQTLdb. Collectively, all these features make the BGVD a useful archive for in-depth data mining and analyses of cattle biology and cattle breeding on a global 50 51 scale.

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53 Keywords: Bovine; Sequence variation; Selective signatures; QTL; Web-database

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### 55 Introduction

Cattle are usually considered the most economically important livestock. The species numbers 56 more than 1.4 billion on a global scale, constituting some 800 extant cattle breeds (FAO, 2016, 57 http://www.fao.org/home/en/). Cattle are now kept on all inhabited continents, in contrasting 58 climatic zones and under very different conditions [1]. The different uses of cattle and the 59 selection for desired traits have resulted in diverse populations distributed across the world. To 60 meet projected global demands for food, initiatives such as the cattle genome project [2–5] are 61 generating resequencing data from breeds worldwide. The DNA-based selection tools built 62 from these data are further accelerating rates of genetic gain and improving animal health and 63 welfare [2]. However, the limited amount of variation data provided by dbSNP [6], restricted 64 access to the 1000 Bull Genomes Project [7], and the existence of only sporadic cattle databases 65

that are specialized in gene and quantitative trait locus (QTL) annotation [8–10] considerably hinder the utility of these data. Furthermore, accessing and integrating resequencing data in a highly interactive, user-friendly web interface, especially data for allele frequency resource and selection in natural populations, is a pre-requisite for identifying functional genes. Therefore, building a public data repository is vital for collecting a wide variety of cattle resequencing data and performing integrative, in-depth analyses within the research community.

Here, we develop the Bovine Genome Variation Database (BGVD), the first web-based 72 public database for accessing dense and broadly representative bovine whole-genome variation 73 74 data. The BGVD is a data repository that focuses on single nucleotide polymorphisms (SNPs), indels, copy number variations (CNVs), and selective signatures underlying domestication and 75 76 population bottleneck events. We have implemented a large number of summary statistics informative for the action of selection, such as nucleotide diversity (Pi) [11], heterozygosity 77 78  $(H_p)$  [12], integrated haplotype score (iHS) [13], Weir and Cockerham's  $F_{ST}$  [14], crosspopulation extended haplotype homozygosity (XP-EHH) [15], and the cross-population 79 80 composite likelihood ratio (XP-CLR) [16] (Table 1). Six early differentiated ancestral 81 populations were used for selection analysis: African taurine, European taurine, Eurasian taurine, East Asian taurine, Chinese indicine and Indian indicine. The current version of the 82 BGVD contains 60,439,391 SNPs, 6,859,056 indels, and 76,634 CNV regions derived from 83 432 cattle. With its functionalities for browsing for variations and their selection scores, the 84 BGVD provides an important publicly accessible resource to the research community to 85 facilitate breeding research and applications and provides information on dominant functional 86 loci and targets for genetic improvement through selection. 87

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### 89 Database structure and content

The BGVD includes SNPs, indels, CNVs, genomic selection, and other database resources
including NCBI, UCSC Genome Browser, AnimalQTLdb, KEGG, and AmiGO 2 for cattle. A
detailed description is provided in the following sections and documents on the homepage.

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### 94 Sample information

Our data set integrates genomes from previously published cattle genetic works [3–5,17–21], providing a total of 432 samples representing 54 breeds. All raw sequence data were obtained from the Sequence Read Archive (SRA) of NCBI. The set of samples is grouped by location of breed origin and contains the following number of individuals: 108 West European, 83 Central-

99 South European, 9 Middle East, 9 Tibetan, 28 Northeast Asian, 47 North-Central Chinese, 21

100 Northwest Chinese, 33 South Chinese, 24 Indo-Pakistan, and 70 African cattle. Geographic

information and other detailed information for each sample are provided on the homepage andthe corresponding 'Sample Table' page.

103

### 104 Variants information

Data were processed and loaded into the BGVD using the following pipeline according to 105 previously published protocols [5] (Figure 1A, see detailed description on the Documentation 106 107 page at of the website). First, short, 250 bp paired-end Illumina reads were aligned to the Btau 108 5.0.1 genome assembly (GCF 000003205.7) using BWA [22], resulting in an average of ~13X 109 coverage of the bovine genome among the cattle varieties. Duplicate reads were removed using Picard tools (http://broadinstitute.github.io/picard/). The Genome Analysis Toolkit (GATK) 110 111 was used to detect SNPs and indels [23]. A total of ~60.4 million autosomal SNPs and ~6.8 million autosomal indels were identified. Beagle was used to phase the identified SNPs [24]. 112 113 Annotation of SNPs and indels was carried out by using snpEff [25]. Minor allele frequency (MAF) for all cattle and allele frequencies for each breed and the "core" cattle group (see 114 Population structure section) were calculated with PLINK [26]. CNVcaller [27] was used to 115 discover CNVs, and 76,634 CNV regions (CNVR) were detected in 432 cattle genomes. Then, 116 the CNVs were annotated using Annovar [28]. Given that three versions of the bovine genome, 117 Btau 5.0.1, UMD3.1.1, and the newly released ARS-UCD1.2 (project accession: 118 NKLS0000000), are commonly used, produced liftOver chain 119 we files (Btau5.0.1ToUMD3.1.1.chain.gz and Btau5.0.1ToARS-UCD1.2.chain.gz) and converted 120 variation coordinates to those of the other two genomes using liftOver [29]. 121

122

### 123 **Population structure**

The population structure of all cattle was inferred using Eigensoft and ADMIXTURE [30,31], 124 based on the genome-wide unlinked SNP dataset, all according to previously published 125 126 protocols [5]. All 432 individuals were used for principal component analysis, and the results were consistent with our previous results [6], except that the African taurine cattle were split 127 form other taurine cattle (Figure 1B). To reduce the bias due to sample size, 10 individuals were 128 randomly selected for breeds that had more than 10 samples. A total of 317 cattle samples were 129 130 selected for estimating ancestral populations by setting K = 2 through K = 8 in ADMIXTURE (Figure 1C). Combining our previous results [5], in addition to five geographically distributed 131 132 ancestral groups (European taurine, Eurasian taurine, East Asian taurine, Chinese indicine, and

133 Indian indicine), African taurine was added in this study (Figure 1B).

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### 135 Selection evaluation

The BGVD provides signatures of selection for eight groups, six of which were the "core" cattle groups that we identified as ancestral groups and the other two of which were directly divided

- into two categories based on sub-species: *Bos indicus* and *Bos taurus*. Here, selective signals
- 139 were evaluated using six methods, namely, Pi,  $H_p$ , iHS,  $F_{ST}$ , XP-EHH, and XP-CLR (**Table 1**).
- 140

### 141 **Database implementation**

The web interface of the BGVD was built by combining an Apache web server, the PHP 142 language, HTML, JavaScript, and the relational database managements system MySQL. High-143 quality SNPs, indels, CNVs, selection scores and their corresponding annotations, classification 144 145 and threshold values, were processed with Perl scripts and stored in the MySQL database. The server application was written in PHP, and CodeIgniter was chosen as the model-view-146 147 controller (MVC) framework for the system. A client interface developed with HTML5 and JavaScript was used to implement search, data visualization and download. Moreover, we 148 introduced web-based software such as BLAST, BLAT, liftOver, and the UCSC Genome 149 Browser (hereafter referred to as 'Gbrowse') [29,32] into the BGVD. Information including 150 variations, selection scores, gene annotation, QTLs, and phastCons conserved elements of 20-151 way mammals and 100-way vertebrates was integrated into Gbrowse to facilitate global 152 presentation. 153

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### 155 Web interface and usage

The BGVD uses a series of user-friendly interfaces to display results. All the parts in our browser are dynamic and interactive. We provided six main functionalities: (i) Gene Quick Search, (ii) Variation Search, (iii) Genomic Selection Search, (iv) Genome Browser, (v) Alignment Search Tools (BLAT/BLAST), and (vi) Genome Coordinate Conversion Tool (liftOver).

For "Gene Quick Search", we integrated information from NCBI, AmiGO 2, and KEGG. Users can input a gene symbol to view all available information, including basic gene information (*e.g.*, genomic location, transcript and protein profile, relevant Gene Ontology (GO) ID, GO terms, and KEGG pathways), gene variations (*e.g.*, SNPs, indels, and CNVs), as well as selective signatures. We also provide links to Gbrowse and external databases (NCBI, AmiGO 2, and KEGG) to help the user obtain more information, such as gene/mRNA/protein
 sequence, KEGG Orthology (KO), and motif.

For "Variation Search", the BGVD allows users to obtain information on SNPs, indels, and CNVs by searching for a specific gene or a genomic region in three versions of the bovine genome (ARS-UCD1.2, UMD3.1.1, and Btau 5.0.1) (**Figure 2**A). Users can filter SNPs and indels further by "Advanced Search", in which certain parameters (Figure 2B), such as MAF and consequence type, can be set; this option enables users to narrow down the items of interest in an efficient and intuitive manner.

The results are presented in an interactive table and graph. For SNPs and indels, users can obtain related details including variant position, alleles, MAF, variant effect, rs ID and the allele frequency distribution pattern in 54 cattle breeds worldwide (Figure 2C) or in six "core" cattle groups (Figure 2D), which could help users dynamically visualize breed-specific (rs384881761, *KRT27*) [2] or ancestral group-specific (rs109815800, *PLAG1*) [33] variants and their global geographical distributions.

For CNVs, users can obtain information about CNVR, such as intersected genomic regions, CNV length, the closest gene, consequence type (**Figure 3**A), and copy number distribution in 432 individuals representing 49 cattle populations. We provide three types of display formats of copy number distributions in which the categories and haploid copy number of each individual can be viewed (Figure 3B–D), such as the "view" button, which produces a scatterplot (*MATN3*); "Gbrowse", which is linked to the "CNVR Bar" track (*KIT*); and the more detailed visualization "cnvBar" track, which generates a box-whisker plot (*CIITA*) [34].

In the genomic signature interface, users can select a specific gene symbol or genomic 187 region, one of the statistical methods (Pi,  $H_p$ , iHs,  $F_{ST}$ , XP-CLR, or XP-EHH), and a specific 188 "core" cattle group to view the selection scores (Table 1 and Figure 4A). In our database, the 189 190 selection scores are pre-processed by several algorithms (Z-transform and logarithm). The results are retrieved in a tabular format (Figure 4B). When users click the "show" button on the 191 table, selective signals are displayed in Manhattan plots or common graphics, where the target 192 193 region or gene is highlighted in a red/blue colour. In addition, the "Gbrowse" button can locate the position of the selection and differentiation profiles of specific groups (Figure 4C). To 194 demonstrate the function of our database, we extracted results for a number of putatively 195 selected genes detected by different methods: OR2T33 [35] (Figure 4B, C), STOM, EPB42 [3], 196 PLAG1 [33], MSRB3 [35], CDC42SE1 [36], R3HDM1 [37], and ASIP [5] (Figure 4C). 197

To further investigate the relationship between variations and signatures of selection, Gbrowse has been introduced to support our database. Currently, 57 tracks have been released

for the Btau 5.0.1 assembly. Users can search with a gene symbol or genomic region to see 200 SNPs, indels, CNVs, genomic signatures, QTLs, and conserved elements in the global view. 201 All search pages in the BGVD allow quick access to Gbrowse to deepen the functional inference 202 of the candidate gene or region by combining other tracks. Most noteworthy, the phased 203 haplotypes from six "core" cattle groups are displayed in "SNPs&Hap" track. The 'squish' or 204 'pack' view highlights local patterns of genetic linkage between variants. In the haplotype 205 sorting display, variants are presented as vertical bars with reference alleles in blue and alternate 206 alleles in red so that local patterns of linkage can be easily discerned when clustering is used to 207 visually group co-occurring allele sequences in haplotypes. We display different haplotypes of 208 the Bos taurus and Bos indicus groups in Figure 4C. We highlight that the tracks of selection 209 210 statistics from different populations are visualized in different colours (Figure 4D).

211 We also introduced two sequence alignment tools, webBlat, and NCBI wwwBLAST, as 212 well as a genome coordinate conversion tool (liftOver) [29] into the BGVD. The webBlat tool can be used to quickly search for homologous regions of a DNA or mRNA sequence, which can 213 214 then be displayed in Gbrowse. BLAST can find regions of local similarity between sequences, which can be used to infer functional and evolutionary relationships between sequences. The 215 liftOver tool is used to translate genomic coordinates from one assembly version into another. 216 Our database provides an online lift from Btau\_5.0.1 to UMD\_3.1.1 and from Btau\_5.0.1 to 217 ARS-UCD1.2. 218

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### 220 **Discussion**

By applying summary statistics to a relatively extensive data set from cattle genomes, we 221 provide a timely and expandable resource for the population genomics research community. An 222 associated user-friendly genome browser gives a representation of the genetic variation in a 223 genomic region of interest and offers functionality for an array of downstream analyses. We 224 expect that the database will prove useful for genome mining through the large number of test 225 statistics and the fine-grained character of resequencing data. We believe that this expandable 226 resource will facilitate the interpretation of signals of selection at different temporal, 227 228 geographical and genomic scales.

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### 230 Authors' contributions

NC, WF, and YJ conceived of the project and designed the research. NC and WF drafted the manuscript. TS, CL, YJ, HC, and ZZ revised the manuscript. NC, JS, and QC performed the

- data analyses. WF and JZ wrote the source code for the BGVD.
- 234

### 235 Competing interests

- 236 The authors declare that they have no competing interests.
- 237

### 238 Acknowledgments

- 239 The project was supported by the National Natural Science Foundation of China (31822052),
- 240 the National Thousand Youth Talents Plan (both to Yu Jiang), the National Beef Cattle and Yak
- 241 Industrial Technology System (Grant No. CARS-37) and the National Natural Science
- 242 Foundation of China (Grant No. 31872317) (both to Chuzhao Lei).
- 243

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### 329 Figure legends

# Figure 1 Analysis pipeline used to construct the database and population analysis of 432 cattle

- A. Analysis pipeline used to construct the database. B. Principal component analysis of 432
- 333 cattle; different numbers in B represent six "core" cattle groups. C. Model-based clustering of
- cattle breeds using the program ADMIXTURE with K = 2 to 8 (plotted in R).
- 335

# Figure 2 Screenshots of a single nucleotide polymorphism (SNP) data search and the results for two examples

- A. Search items involving rs ID, gene name and position of three bovine reference genomes. B.
  Advanced Search menu enabling filtering for minor allele frequency and consequence type. C.
  Detailed annotation of the rs384881761 locus of the *KRT27* gene and the allele frequency
  distribution pie-chart of 54 cattle breeds worldwide. D. Display format of the allele frequency
  for the rs109815800 locus of *PLAG1* among defined ancestral groups.
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# Figure 3 Screenshots of a copy number variation region (CNVR) data search and three types of display formats of the results

A. Search items involving the gene name and position of three bovine reference genomes. **B.** Results involving detailed annotation for the CNVR and copy number distribution patterns of 432 individuals representing 49 populations. An example of *MATN3*, which showed different copy numbers in the Holstein population. **C.** "CNVR Bar" track in the bar chart format in UCSC Genome Browser (Gbrowse). An example of the *KIT* gene, which is related to coat color in Herefords. **D.** The more detailed visualization "CNVR Bar" track in the format of a box-whisker plot, displaying copy number distribution in 49 cattle populations. An example of *CIITA*, which lies within a high-frequency gain CNVR identified in multiple breeds that showed nematoderesistance.

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# Figure 4 Screenshots of a search for genomic selection data and representation of the selection data

A. Search items involving gene name, position, and one of the statistical methods (nucleotide 358 diversity (Pi), heterozygosity (Hp), integrated haplotype score (iHS), Weir and Cockerham's 359  $F_{ST}$ , cross-population extended haplotype homozygosity (XP-EHH), and the cross-population 360 composite likelihood ratio (XP-CLR)), and specific "core" cattle groups. B. Detailed annotation 361 for the target gene or region in the variant grid and the corresponding selective signal at the 362 chromosome and whole-genome levels, respectively. An example of selective signal of the 363 OR2T33 gene in Eurasian taurine population. C.-E. The display of 57 tracks in UCSC Genome 364 Browser (Gbrowse) in the BGVD. Numbers 1-16 represent the corresponding tracks. (C) 365 Example of the OR2T33 gene in "SNPs&Hap" track. Different haplotypes of the Bos taurus 366 367 and Bos indicus groups are shown in blue and red, respectively. D. Examples of the six selection scores of the POFUT1 gene in the Chinese indicine (CN) group, and where each group is 368 represented by a different color. Here, we show  $F_{ST}$  scores of Indian indicine (IN) and East 369 Asian (EA) groups with orange and blue, respectively. E. Fifty-seven tracks in Gbrowse. 370 371

372 **Table** 

373 Table 1 Statistical terms for selection sweep in the Bovine Genome Variation Database374 (BGVD)

### Table 1 Statistical terms for selection sweep in the Bovine Genome Variation Database

### (BGVD)

Statistical term	Abbreviation	Population 1	Population 2	Windows
Nucleotide diversity	Pi	Indian indicine (IN)		30k
Heterozygosity	$H_{ m p}$	Chinese indicine (CN)		60k
Integrated haplotype score	his	East Asian taurine (EA) Eurasian taurine (EUA) European taurine (EUR) African taurine (AFR) <i>Bos indicus</i> (BIN) <i>Bos taurus</i> (BTA)		30k
Weir and Cockerham's Fst	$F_{\rm ST}$	Indian indicine (IN)	Other five groups	30k
Cross-population composite likelihood ratio	XP-CLR	Chinese indicine(CN)	Other five groups	30k
Cross-population extended haplotype homozygosity	ХР-ЕНН	East Asian taurine (EA) Eurasian taurine (EUA) European taurine (EUR) African taurine (AFR) <i>Bos indicus</i> (BIN)	Other five groups Other five groups Other five groups Other five groups <i>Bos taurus</i> (BTA)	30k



### Region

- Africa
- Central-South Europe
- Indo-Pakistan
- Middle East
- North-Central China
- Northeast Asia
- Northwest China
- South China
- Tibet, China
- West Europe

### Sub-species

- Bos indicus
- Bos taurus
- imes Hybrid

### Ancestry

- 1 European taurine
- 2 Eurasian taurine
- 3 East Asian taurine
- 4 Chinese indicine
- 5 Indian indicine
- 6 African taurine



### A SNPs (single nucleotide polymorphisms)

Please enter a dbSNP ID, or a gene symbol, or a chromosome location for one of the genome versions, such as Btau 5.0.1 (GCF\_000003205.7), UMD3.1.1 (GCF\_000003055.6) and ARS-UCD1.2 (GCF\_002263795.1), to obtain a SNP information and allele frequency distribution pattern in 54 world-wide cattle breeds or six "core" cattle groups.

Basic search				
	dbSNP ID:		e.g., rs384881761, rs109815800	
Or	Gene symbol:		e.g., PLAG1, KRT27, HOXD4	
Or	Chromosome location:		For Btau_5.0.1, e.g., 19:41811000-41811922, 19:41811922	
Or	Chromosome location:		For UMD_3.1.1, e.g., 19:41636098-41636961, 19:41636961	
Or	Chromosome location:		For ARS-UCD1.2, e.g., 19:40981387-40982250, 19:40982250	

#### B Advanced search

Minor allele frequency >= ~ (range: 0-1) Consequence type: 🗹 Transcript variant 🗸 Coding variant -Missense variant ☑ Initiator\_codon\_variant Start lost Stop lost Stop\_gained Stop\_retained\_variant Synonymous\_variant Non-coding variant -5\_prime\_UTR\_variant Start\_gained ☑ 3 prime UTR variant ☑ Intron variant ☑ Non\_coding\_transcript\_exon\_variant Splice variant -Splice acceptor variant Splice donor variant Splice region variant Intragenic variant -☑ Intragenic\_variant Intergenic variant -Upstream\_gene\_variant Downstream gene variant ☑ Intergenic variant Search Reset

#### C SNPs found **Details** Core\_group Position of Position of Gene Breed Chr Position Alleles MA MAF Consequence type Gene Variant ID UMD3.1.1 ARS\_UCD1.2 Visualization details frequency frequency 19 41811922 G/C C 0.006 missense variant KRT27 rs384881761 19:41636961 19:40982250 Show Show Ghrows Showing 1 to 1 of 1 entries Previous Next

Allele frequency distribution of world-wide cattle breeds Pie-chart on world map



 $\downarrow$ 

Ref

Alt 📃

Indian\_indicine(24):0.771 Chinese\_indicine(19):0.974 East\_Asian\_taurine(37):0.959 European\_taurine(38):0.039 Eurasian\_taurine(19):0.947 Africa\_taurine(10):1.000

### A CNVs (Copy number variations)

Showing 1 to 1 of 1 entries

0.0

Please enter a gene symbol or a chromosome location for one of the genome versions, such as Btau 5.0.1 (GCF\_000003025.7), UMD3.1.1 (GCF\_000003055.6) and ARS-UCD1.2 (GCF\_002263795.1), to obtain CNV region (CNVR) information of intersected genomic region, CNV length, the closest gene, consequence type and copy number distribution in 432 individuals representing 49 cattle populations.



Distribution of haploid copy number in different cattle breeds

Link to CNV details D Haploid Copy Number (CNV69269) Name of transcript: CNV69269 Name of gene: CNV69269 Total all median values: 53.87 Haploid\_Copy\_Number Maximum median value: 2.04 Haploid\_Copy\_Number in CentralChina\_Luxi Score: 999 Genomic position: Btau 5 0 1 25:9649602-9705600 Strand: + CNV69269 (CNV69269) 2.5 CIITA gene, which lies within a high frequency gain CNVR identified in multiple breeds 2.0 Haploid\_Copy\_Nun 1.0 1.5 0.5 0.0 N=25 16 21 CNV details, group category of each individual View all data points for CNV69269 (CNV69269)

Haploid Copy Number

# С

Visualization

1 Next

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Gbrowse

Previous

CNV21480

KIT gene in Hereford

### A Selective signatures

Please enter a gene symbol or a chromosome location (Btau 5.0.1, accession: GCF\_000003205.7), and select one of the statistical methods as well as specific "core" cattle populations to view the selection score.

#### Search by gene symbol or chromosome position Gene symbol: OR2T33 e.g., OR2T33 e.g., 7:43260001-43290000, 7:43280000 Or Chromosome location: Statistical methods: PI: Nucleotide diversity Hp: Heterozygosity iHS: Integrated haplotype score FST: Fixation index XP-CLR: Cross-population composite likelihood ratio XP-EHH: Cross-population extended haplotype homozygosity Population: Indian indicine Vs Other five groups Chinese indicine Vs Other five groups East Asian taurine Vs Other five groups Eurasian taurine Vs Other five groups European taurine Vs Other five groups African taurine Vs Other five groups Bos indicus Vs Bos taurus Search Reset Е B Selective region found FST: Eurasian taurine vs other five groups Selective region Gene symbol Window number\*size (bp) Visualization User query Signature figure OR2T33 7:43020001-43050000 OR2T33 1\*30000 Show Gbrowse Showing 1 to 1 of 1 entries Next Previous C Selective signatures at the Chr7 (target region or gene is highlighted in red, see Z(Fst) value > 0) 12.0 90-₽ 6.0 -JZ 3.0 ------ - - - - - - 2.576 0.0 Chr7 -3.0 C Genome-wide detection of selective signatures 14



#### С move $\langle \langle \langle \langle \rangle zoom in 1.5x 3x 10x base zoom out 1.5x 3x 10x 100x zoom out 1.5x 3x 10x zoom out 1.5x 3x 10x zoom out 1.5x 3x 10x 100x zoom out 100x zoom out 100x zoom out 100x zo$ 7:43,020,001-43,050,000 30,000 bp. 0R2T33 × go Btau\_5\_0\_1 Scale 7: 10 kb 1 43, 825, 8881 43,030,000 43,035,000| Gap Location 43.848.888 43.845.888 2 OC Percent in S-Base kindous Incol Open Predictions Gap 3 GC percent 4 5 6 100 vertebrates conservation by PhastCon Cons 100 Verts SNPs(432) 7 Haploid Copy Number Inde1s(432) 1 1 10 0 11 1 1 10 10 1 8 CNV24428 Q D 10 11 12 13 HS CN 14 YPEHH Value 15 D\_CN 16 Pi value - Chinese indicine Ch Mapping and Sequencing refresh Short Match **Base Position** Gap GC Percent dense $\lor$ dense dense 🗸 hide Genes and Gene Predictions refresh NCBI Genes QTL pack 🗸 hide $\sim$ 2 1 -**Comparative Genomics** refresh Cons 20 Mammals Cons 100 Vertebrates dense 🗸 dense V 4 3 -Variation and Repeats refresh SNPs & Haplotype SNPs(432) Indels(432) CNVR Bar $\times$ hide 5 hide pack 🗸 hide $\sim$ 8 -Selection Statistics based on Population Differentiation refresh 世 Fst IN Other Fst CN Other Fst EA Other Fst EUA Other Fst EUR Other Fst AFR Other hide $\vee$ 9 full hide $\vee$ hide < hide $\vee$ 11 10 hide $\sim$ XP-CLR CN Other XP-CLR EA Other XP-CLR EUA Other XP-CLR EUR Other Fst BIN BTA XP-CLR IN Other hide × 12 hide full hide hide hide $\vee$ XP-CLR AFR Other XP-CLR BIN BTA hide hide ✓ 13 -Selection Statistics based on Linkage Disequilibrium Structure refresh iHS IN iHS CN ihs ea ihs eua ihs eur iHS AFR hide $\vee$ full $\sim$ hide $\vee$ hide hide $\vee$ hide $\vee$ iHS BIN **iHS BTA** XPEHH IN Other XPEHH CN Other XPEHH EA Other **XPEHH EUA Other** hide V hide hide 🗸 full $\sim$ 14 hide hide XPEHH EUR Other XPEHH AFR Other **XPEHH BIN BTA** hide hide hide $\vee$ -Selection Statistics based on Allele Frequency Spectrum refresh Hp IN Hp CN Hp EA Hp EUA Hp EUR Hp AFR hide full 15 hide hide hide 🗸 $\sim$ hide Hp BIN Hp BTA hide hide -**Descriptive Statistics** refresh Pi IN Pi CN Pi EA Pi EUA Pi EUR Pi AFR 16 hide hide $\vee$ full 🗸 hide hide 🗸 hide 🗸 Pi BTA Pi BIN hide hide

UCSC Genome Browser on Btau 5 0 1 November 08th, 2017 Assembly (Btau 5 0 1)