

# BovineSNP50 Genotyping BeadChip

Featuring 54,609 evenly spaced SNP probes that span the bovine genome

#### **Highlights**

- Unrivaled Call Rates and Accuracy
   >99% average call rates and 99.9% reproducibility
- Comprehensive and Uniform Coverage Evenly distributed polymorphic SNPs with a mean gap of 49.4kb
- Simple Workflow PCR- and ligation-free protocol
- High-Throughput Format
   Up to 24 samples can be interrogated in parallel

#### Introduction

Illumina, in collaboration with the USDA ARS, University of Missouri, and the University of Alberta, has developed the second generation of a proven high-density, genome-wide genotyping array for the interrogation of genetic variation in cattle. The BovineSNP50 v2 BeadChip features 54,609 informative SNP probes that uniformly span the entire bovine genome, empowering applications such as genome-wide enabled selection, identification of quantitative trait loci, evaluation of genetic merit of individuals, and comparative genetic studies.

More than half of the probes on this BeadChip are designed to target novel SNPs discovered using Illumina's Genome Analyzer. Additional high-value content is derived from publicly available sources such as the bovine reference genome, Btau¹, and the Bovine HapMap Consortium data set². The BovineSNP50 BeadChip covers common SNPs validated in economically important beef and dairy cattle breed types and presents an average minor allele frequency (MAF) of 0.24 across all loci. Importantly, this BeadChip offers uniform coverage with an average probe spacing of 49.4kb to provide more than sufficient SNP density for robust genome-association studies in cattle.

The BovineSNP50 BeadChip is a multi-sample genotyping panel powered by Illumina's Infinium® HD Assay. This revolutionary assay provides the industry's highest call rates, allows for flexible content deployment, and enables the detection and measurement of copy number variation. In addition, the assay's PCR-free, single-tube sample preparation<sup>3,4</sup> significantly reduces labor and potential sample handling errors. Illumina's multi-sample format further reduces experimental variability and overall project cost by allowing researchers to interrogate up to 24 samples in parallel.

The combination of Illumina's proprietary assay technologies, unconstrained locus selection, and high-throughput format offers a comprehensive solution for whole-genome studies in beef and dairy cattle.

Figure 1: BovineSNP50 BeadChip

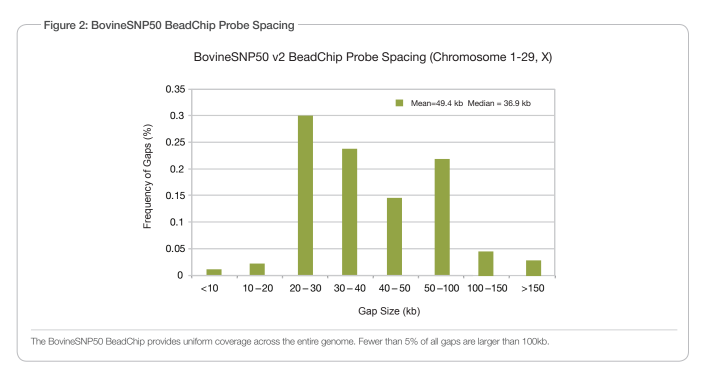


The BovineSNP50 BeadChip features more than 54,000 evenly spaced SNPs across the entire bovine genome.

# BovineSNP50 v2 BeadChip Content

Illumina scientists and collaborators set out to develop an informative and high-density SNP genotyping microarray that could be used to investigate genetic variation in any cattle breed. More than 12,000 probes were designed to target validated common SNPs (MAF ≥ 0.05) described by the Bovine HapMap Consortium. BeadChip developers also mined publicly available resources for common SNPs including the bovine reference genome¹ and whole-genome shotgun reads assembled by researchers at Baylor College of Medicine⁵. Additional content sources include parentage markers identified by researchers from the U.S. Meat Animal Research Center, Clay Center⁶, and SNPs identified by researchers at the USDA ARS through the comparison of Holstein bacterial artificial chromosomes (BAC) sequence dataⁿ to the bovine genome assembly (Table 1).

More than 24,000 SNP probes featured on the BovineSNP50 Bead-Chip target novel SNP loci that were discovered by sequencing three pooled populations of economically important beef and dairy cattle. Using Illumina's Genome Analyzer, researchers at the USDA ARS discovered more than 62,000 putative SNPs by deeply sequencing approximately 2% of the *Bos taurus taurus* genome<sup>8</sup>. More than 23,800 SNPs derived from this novel data source were chosen based upon their spacing, expected MAF, and Infinium HD Assay performance. In addition, BeadChip developers selected loci that target the largest unmapped contigs to ensure comprehensive coverage. All 54,609 SNP probes on the BovineSNP50 BeadChip have been validated in 19 common beef and dairy breeds (Table 3). This product targets evenly distributed SNPs that are polymorphic across the breeds tested and



provides an average probe spacing of 49.4kb and a median spacing of 36.9kb (Figure 2). Current research on linkage disequilibrium (LD) in multiple breeds of cattle suggests haplotype blocks of approximately 70kb on average<sup>9</sup>, indicating that the resolution offered by the BovineSNP50 chip is well within the resolution of LD in cattle. More than 54,060 SNP probes on the BovineSNP50 BeadChip map to UMD 3.0, the most current bovine reference genome assembly.

### **High-Quality Data**

The 54,609 SNP assays on the BovineSNP50 BeadChip were subjected to rigorous functional testing to ensure strong performance using Illumina's Infinium HD Assay. Whole-genome association studies are successful, in part, due to high call rates and accurately called genotypes. Since complex traits often have relatively small gene effects, potential associations may be missed if the assayed SNP in linkage disequilibrium with the SNP of interest has a low call rate or incorrect genotype call. Illumina ensures that every BovineSNP50 BeadChip offers > 99% call rate.

Table 2 shows the outstanding results from Illumina's internal validation of the BovineSNP50 BeadChip content using samples provided by the Bovine HapMap Consortium. Product developers identified and retained 4,290 loci that appear to have an adjacent or underlying deletion among the breeds sampled. Although these loci yield lower call rates when compared to the other loci on the panel, they were retained because they may provide biologically relevant information for traits of interest and for future improvements in the genome build.

The performance and content validation results clearly show the reliability and outstanding data quality the BovineSNP50 BeadChip delivers. With this BeadChip, researchers can predict the genetic merit for phenotypes of interest and investigate the genetic basis of variation among a multitude of cattle breed types.

# Solutions for Genotyping

As with all of Illumina's standard products, an optional Laboratory Information Management System (LIMS) and robotic automation are available to accurately and efficiently track samples throughout analysis. Researchers can genotype their own samples using the Illumina GenomeStudio® DNA analysis software or Illumina's FastTrack Genotyping Service.

Table 1: BeadChip Content Sources

Source	BovineSNP50 v1 Probes	BovineSNP50 v2 Probes	
Novel SNPs derived from sequencing common cattle breeds using Illumina's Genome Analyzer	23,840	24,181	
Bovine HapMap Data Set	12,298	12,342	
Btau assembly SNPs	9,361	9,404	
Whole-Genome Shotgun Reads*	5,808	6,038	
Holstein BAC Sequence Data	1,409	1,411	
Parentage**	116	120	
Other <sup>†</sup>	1,169	1,113	
Total	54,001	54,609	

<sup>\*</sup> Reads derived from six cows/breeds (Norwegian Red, Hostein, Brahman, Angus, Jersey and Limousin) compared against Btau2.0

<sup>\*\*</sup> Highly curated parentage markers that include 30 duplicates

<sup>†</sup> Includes common SNPs validated by the Institute for Food and Agricultural Sciences Alberta and INRA, the French International Institute of Agriculture

## **Product Summary**

Developed in collaboration with leading bovine researchers, the BovineSNP50 BeadChip features more than 54,000 evenly spaced SNP probes that span the bovine genome. This 24-sample BeadChip presents a high-throughput, cost-effective solution for whole-genome studies in beef and dairy cattle.

Table 2: Product Performance and Specifications

Parameter	Results	Product Specification
Average Call Rate*	99.7%	> 99%
Reproducibility	100%	> 99.9%
Mendelian Inconsistencies**	0.06%	< 0.1%

<sup>\*</sup> Based on 473 individuals from 19 major cattle breeds, 31 trios, and 1 replicate

Table 3: BeadChip Content Validation

Breed	Samples	Polymorphic Loci*	Mean MAF	Median MAF**
Angus	24	41,073	0.21	0.21
Beefmaster	23	43,114	0.22	0.22
Bos indicus Gir	21	23,567	0.11	0.02
Bos indicus Nelore	19	25,492	0.11	0.03
Brahman	22	29,444	0.13	0.07
Brown Swiss	21	35,971	0.19	0.17
Charolais	19	43,723	0.22	0.21
Guernsey	21	36,748	0.19	0.17
Hereford	24	42,132	0.22	0.23
Holstein	49	42,849	0.22	0.22
Jersey	23	35,346	0.18	0.15
Limousin	39	42,617	0.22	0.21
N'Dama	23	28,869	0.14	0.07
Norwegian Red	17	42,055	0.22	0.21
Piedmontese	21	41,912	0.22	0.21
Red Angus	10	42,388	0.21	0.20
Romagnola	21	38,524	0.20	0.19
Santa Gertrudis	21	41,783	0.22	0.21
Sheko	16	35,084	0.17	0.13
Overall	434	47,168	0.24	0.25

<sup>\*</sup> MAF > 0.05

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<sup>\*\*</sup> Based on 4 Holstein trios

<sup>\*\*</sup> Across all 54,609 loci

#### **Ordering Information**

Catalog No.	Product	Description
WG-450-2001	BovineSNP50 v2 Whole-Genome Genotyping Kit (for 48 samples)	Each package contains 2 BeadChips, along with reagents for amplifying, fragmenting, hybridizing, labeling, and detecting 48 DNA Samples
WG-450-2002	BovineSNP50 v2 Whole-Genome Genotyping Kit (for 288 samples)	Each package contains 12 BeadChips, along with reagents for amplifying, fragmenting hybridizing, labeling, and detecting 288 DNA Samples
WG-450-2003	BovineSNP50 v2 Whole-Genome Genotyping Kit (for 1152 samples)	Each package contains 48 BeadChips, along with reagents for amplifying, fragmenting hybridizing, labeling, and detecting 1152 DNA Samples

Each BovineSNP50 v2 DNA Analysis BeadChip can process 24 samples and analyze approximately 54,000 loci.

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