Infinium[™] Japanese Screening Array-24 v1.0 BeadChip

A powerful, high-quality, costeffective array for genetic studies in Japanese populations

- Features expert-selected content, including approximately 100K Japanese-specific markers and the option to add custom content
- Enables genotyping of clinical research variants for a broad range of applications, including complex disease studies, pharmacogenomics research, and more
- Maintains the same data quality of Illumina genotyping arrays with call rates > 99% and reproducibility > 99.9%

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Introduction

The Infinium Japanese Screening Array-24 v1.0 BeadChip provides a scalable solution for variant screening and precision medicine research in Japanese populations (Table 1). Built on the Infinium high-throughput screening (HTS) platform, it allows for rapid analysis of large sample sets, reducing the overall cost per sample. Designed in collaboration with a Japanese consortium, it harnesses content from the Infinium Asian Screening Array-24 v1.0 BeadChip and tag single nucleotide polymorphisms (SNPs) based on Japanese whole-genome sequencing (WGS) data. Combined with the iScan[™] System, integrated analysis software, and the Infinium HTS assay, this highdensity, 24-sample BeadChip (Figure 1) provides optimized content with the same high-quality, reproducible data that Illumina genotyping arrays have provided for over two decades. The Infinium Japanese Screening Array-24 v1.0 BeadChip will power growing biobank and translational research studies in the Japanese population.

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|------------|------------------|
| | |

Figure 1: Infinium Japanese Screening Array-24 v1.0 BeadChip— Built on the trusted 24-sample Infinium HTS platform.

Optimized Japanese content from WGS studies

The Infinium Japanese Screening Array-24 v1.0 BeadChip contains highly informative tag SNPs in Japanese populations (Table 2), including > 490K genome-wide backbone markers from the Infinium Asian Screening Array-24 v1.0 BeadChip and approximately 100K Japanese-specific markers contributed by expert human disease researchers and genomic service providers in Japan. Exclusive to this BeadChip, the clinical research content contains approximately 33K regional markers from various consortia members in Japan. The array combines curated clinical research variants and quality control (QC) markers for a broad range of clinical research and variant screening applications. These applications include disease association and risk profiling studies, pharmacogenomics research, disease characterization, lifestyle and wellness characterization, and marker discovery in complex disease research.¹⁻⁴

Table 1: Product information^a

| Feature | Description |
|--|--------------------------------|
| Species | Human |
| Total number of markers ^b | 736,847 |
| Capacity for custom bead types | 50,000 |
| Number of samples per BeadChip | 24 |
| Input requirement | 200 ng genomic DNA |
| Assay chemistry | Infinium HTS |
| Instrument | iScan System |
| Maximum sample throughput ^a | ~2304 samples/week |
| Scan time per sample | 2.5 minutes |
| Approximate values approximate and movimum | throughput pour yory doponding |

 Approximate values, scan times, and maximum throughput may vary depending on laboratory and system configurations.

b. Total number of markers calculated from the consortium manifest.

Table 2: Marker information

| Marker categories ^a No. of mark | |
|--|-----|
| Exonic markers ^b 73,737 | |
| Nonsense markers ^c 5608 | |
| Missense markers ^c 50,775 | |
| Synonymous markers ^c 11,636 | |
| Mltochondrial markers ^c 1262 | |
| Indels ^d 9286 | |
| X Y PAR/homolog | ous |
| Sex chromosomes ^c 27,445 6869 721 | |

a. Number of markers are calculated from the consortium manifest.

- b. RefSeq: NCBI Reference Sequence Database. Accessed August 30, 2020.5
- c. Compared against the UCSC Genome Browser. Accessed August 30, 2020.6

 NCBI Genome Reference Consortium, Version GRCh37. Accessed August 30, 2020.⁷

Abbreviations: indel, insertion/deletion; PAR, pseudoautosomal region.

Table 3: High-value content

| Content ^a | No. of markers ^b | Research application/note | Contentª | No. of markers ^b | Research application/note |
|--|--------------------------------|--|---|--------------------------------|--|
| ACMG ⁸ 59 2016 gene coverage | 15,156 | _ | GO ¹¹ CVS genes | 106,209 | Cardiovascular conditions |
| ACMG 59 all annotations | 13,115 | | Database of Genomic Variants ¹² | 573,087 | Genomic structural variation |
| ACMG 59 pathogenic | 5903 | _ Variants with known clinical significance identified from clinical WGS and WES samples | eQTLs ¹³ | 3982 | Genomic loci regulating mRNA expression levels |
| ACMG 59 likely pathogenic | 1953 | | Fingerprint SNPs ¹⁴ | 378 | Human identification |
| ACMG 59 benign | 580 | - | gnomAD ⁴ exome | 73,737 | WGS and WES results from unrelated individuals from various studies |
| ACMG 59 likely benign | 888 | _ | HLA genes ¹⁵ | 1103 | Disease defense, transplant rejection, and autoimmune disorders |
| ACMG 59 VUS | 2272 | - | Extended MHC ^{15,d} | 12,668 | Disease defense, transplant rejection, and autoimmune disorders |
| AIMs ^c | 2595 | Ancestry-informative markers | KIR genes ⁶ | 80 | Autoimmune disorders and disease defense |
| APOE ⁶ | 16 | Cardiovascular disease, Alzheimer's disease, and cognition | Neanderthal SNPs ¹⁶ | 1651 | Neanderthal ancestry and human population migration |
| ClinVar ¹ variants | 50,223 | | NHGRI-EBI GWAS catalog ¹⁷ | 22,103 | Markers from published GWAS |
| ClinVar pathogenic | 19,432 | - Relationships among variation, | Newborn/carrier screening gene coverage | 26,303 | Genes associated with childhood diseases included in the TruSight [™] Inherited Disease Sequencing Panel ⁷ |
| ClinVar likely pathogenic | 7684 | [–] phenotypes, and human health | NHGRI diseases | 19,492 | Markers related to various diseases from published studies |
| ClinVar benign | 13,134 | - | PharmGKB ^{2,17} | 4287 | Human genetic variation associated with drug responses |
| ClinVar likely benign | 6516 | - | RefSeq⁵ 3' UTRs | 16,350 | 3' untranslated regions |
| COSMIC ⁹ genes | 323,620 | Somatic mutations in cancer | RefSeq 5' UTRs | 7450 | 5' untranslated regions |
| CPIC ¹⁰ all | 250 | | RefSeq All UTRs | 23,073 | Untranslated regions |
| CPIC-A/B | 140 | | RefSeq | 362,588 | All known genes |
| CPIC-B | 17 | - | RefSeq +/- 10 kb | 427,037 | Regulatory regions |
| CPIC-C | 14 | Variants with potential guidelines to optimize drug therapy | RefSeq Promoters | 17,248 | 2 kb upstream to include promoter regions |
| CPIC-C/D | 109 | _ | RefSeq Splice Regions | 2696 | Variants at splice sites |
| CPIC-D | 76 | | | | |

a. Content are derived from the consortium manifest.

b. The number of markers for each category may be subject to change.

c. Based on internal calculations.

d. Extended MHC is a 8 Mb region.

Abbreviations: ACMG, American College of Medical Genetics; AIM, ancestry-informative marker; APOE, apolipoprotein E; COSMIC, catalog of somatic mutations in cancer; CPIC, Clinical Pharmacogenetics Implementation Consortium; EBI, European Bioinformatics Institute; eQTL, expression quantitative trait loci; gnomAD, Genome Aggregation Database; GO CVS, gene ontology annotation of the cardiovascular system; GWAS, genome-wide association study; HLA, human leukocyte antigen; KIR, killer cell immunoglobulin-like receptor; MHC, major histocompatibility complex; NHGRI, National Human Genome Research Institute; PharmGKB, Pharmacogenomics Knowledgebase; RefSeq, NCBI Reference Sequence Database; UTR: untranslated region, WES: whole-exome sequencing.

Broad clinical research applications

The clinical research content of the Infinium Japanese Screening Array-24 v1.0 BeadChip was designed through collaboration with medical genomics experts using multiple annotation databases¹⁻⁴ to create an informative, cost-effective panel for clinical research applications (Figure 2 and Table 3).

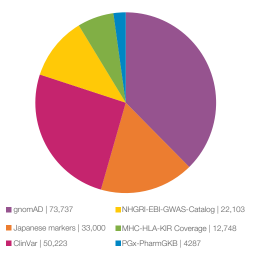


Figure 2: Clinical research content—Content was expertly selected from scientifically recognized databases to create a highly informative array for clinical research applications. Variant counts may be subject to change.

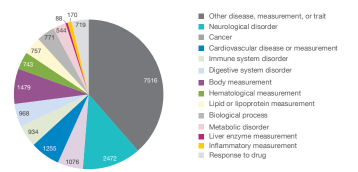


Figure 4: NHGRI disease categories—Clinical research content on the array features > 12,000 markers across a broad range of disease categories based on the NHGRI database. Variant counts are derived from consortium manifest and may be subject to change.

Extensive range of disease categories covered

Including over 18K variants with established clinical associations based on the ClinVar database,¹ clinical research content on the Infinium Japanese Screening Array-24 v1.0 BeadChip enables validation of disease associations, risk profiling, preemptive screening research, and pharmacogenomics studies. Variant selection includes a range of pathology classifications based on ClinVar American College of Medical Genetics and Genomics (ACMG) annotations (Figure 3A).⁸ There are over 7K disease and trait associations from the ClinVar database (Figure 3B) and over 12K variants selected from the NHGRI-EBI GWAS catalog³ (Figure 4), representing a broad range of phenotypes and disease classifications.

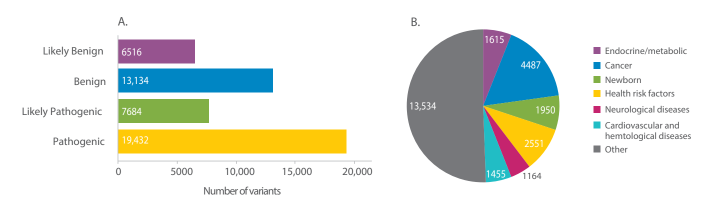


Figure 3: Broad coverage of disease categories—(A) Variants sorted by range of pathology classifications according to ClinVar ACMG annotations. (B) Infinium Japanese Screening Array-24 v1.0 clinical research content features > 7000 disease traits and associations based on categories within the ClinVar database. Variant counts may be subject to change.

QC markers for sample identification, tracking, and stratification

The Infinium Japanese Screening Array-24 v1.0 BeadChip includes QC and high-value markers for large-scale studies, enabling sample identification, tracking, ancestry determination, and stratification (Figure 5).

| | Ancestry informative (2337) |
|--------------|-------------------------------------|
| | Sex determination (2411) |
| | Blood phenotype (1337) |
| QC markers — | Mitochondrial (128) |
| | Human linkage (1050) |
| | Pseudo autosomal regions 1 & 2 (88) |
| | Fingerprinting (323) |

Figure 5: QC markers—QC variants on the array enable various capabilities for sample tracking such as sex determination, continental ancestry, human identification, and more. Data are derived from the consortium manifest.

Flexible content options

The Infinium Japanese Screening Array-24 v1.0 BeadChip is available as a predesigned content panel or can be customized to incorporate up to 50K custom bead types (Table 4). The DesignStudio" Microarray Assay Designer can be used to design targets such as SNPs, copy number variants (CNVs), and indels.

Table 4: Flexible content options

| Compatible content | No. of markers | Description |
|---------------------------------|-------------------|--|
| Custom content | ≤ 50,000 | Custom design virtually any target (eg, SNP, CNV, indel) using the DesignStudio Microarray Assay Designer |
| Multi-disease drop- in panel | ~50,000 | Fine-mapping content derived from exome sequencing and meta analysis of phenotype-specific consortia focused on the following traits: psychiatric, neurological, cancer, cardiometabolic, autoimmune, and anthropometric |

Summary

The Infinium Japanese Screening Array-24 v1.0 BeadChip provides a cost-effective solution for population-scale genetic studies, variant screening, and precision medicine research focusing on the Japanese population. The array builds on the success of the widely adopted Infinium Global Screening Array and Infinium Asian Screening Array. Using the iScan System, Infinium HTS Assay, and integrated analysis software, this high-density, 24-sample BeadChip provides optimized content for a broad range of clinical research applications.

Learn more

Illumina microarray solutions

For labs interested in higher throughput processing, contact your local account manager for information about Infinium HTS Extra high-throughput kit configurations.

Ordering information

| Infinium Japanese Screening Array-24 v1.0 BeadChip Kit | Catalog no. |
|---|-------------------------|
| 48 samples | 20040743 |
| 288 samples | 20040744 |
| 1152 samples | 20040745 |
| | |
| Infinium Japanese Screening Array-24+ v1.0 BeadChip Kitª | Catalog no. |
| 1 3 | Catalog no. 20040746 |
| Array-24+ v1.0 BeadChip Kita | |
| Array-24+ v1.0 BeadChip Kit ^a 48 samples | 20040746 |

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