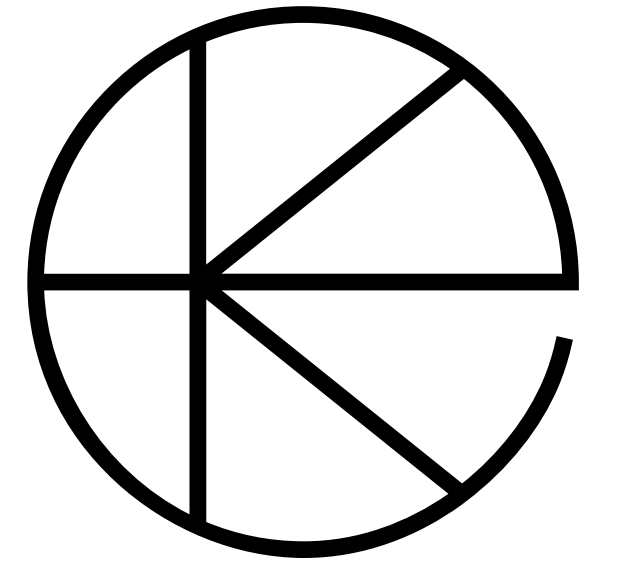


ekei epigenetics

Research Platform



Extremely low-cost, high-resolution epigenetics sequencing. Suited for fundamental research, large scale studies and AI algorithms training.

Introducing ekei epigenetics research platform, built on our innovative proprietary methods to revolutionize the adoption of epigenetics in various fields of research, pharmaceutical product development and clinical applications.

Epigenetics is a key element of life's adaption potential. This dynamic layer of biological information adapts to life cycles, development phases, and as a response to environmental factors and cellular stressors. In recent years, its study has revolutionized fields like longevity science, offering powerful ways to quantify biological ageing, and providing critical insights into complex diseases, including cancer progression.

Today's main barrier to entry is the high cost and limited data resolution of epigenetic analysis.

Our epigenetics platform is addressing these challenges by pushing the boundaries of the current next-generation sequencing solutions for high-throughput and high data coverage coupled with our proprietary wet-lab methods to deliver best-in-class epigenetic data from various type of DNA samples.

Our platform is optimized for flexibility and sensitivity. We support DNA analysis from various sources, including blood, saliva, buccal swabs, bulk tissue and cellular secretomes. Our methods excel with low DNA input, enabling minimally invasive collection protocols and preserving precious sample material.

Since each project is unique it may require information on various epigenetic factors in addition to the data from our core proprietary DNA methylation methods. Our platform allows for the combination of protocols to acquire the specific data needed, maximizing the potential to achieve target goals.

Engineered for the era of artificial intelligence, ekei generates the robust, high-dimensional datasets essential for modern machine learning. Empower your research teams working in life science to accelerate the discovery of new biomarkers with greater confidence, to create tailored biotech products and therapeutics, to validate clinical trial outcomes more effectively and to deepen the understanding of ageing and to potentially reverse it.

If you are working on solutions for anti-ageing healthcare, cosmetics, complex diseases therapeutics, cancer biomarkers, pharmaceutical drug discovery, biobanks, fundamental cell biology or even agricultural biotechnology such as optimized crop breeding programs, our ekei epigenetics research platform will provide invaluable information for your research.

Highlights

Whole Genome

Our methods are able to sequence epigenetic information genome-wide, providing a considerable advantage over existing methods such as methylationEPIC arrays or RRBS. We are able to target over **25M** unique CpG sites* in the human genome which includes up-to **9.6M CpGs** in annotated gene locations as well as promoter and enhancer regions.



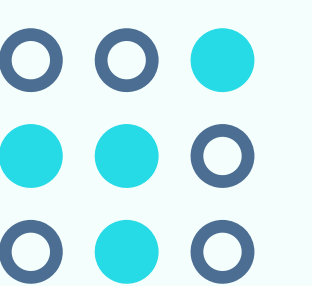
Ultra-low cost

The core strength of our services lies in the use of novel proprietary lab methods that when coupled with the most recent next-generation sequencing techniques allow for maximum throughput and best data value at ultra-low cost. Using multiplexing, it is now possible to capture a wide human methylome for less than **20,000 JPY** in large studies.



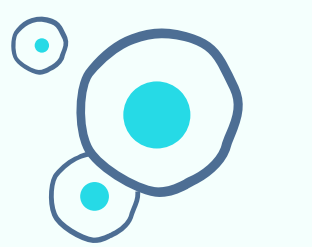
High resolution single channel

While traditional techniques output quantified methylation information per CpG site, our platform captures single channel information at each CpG site with increased sensitivity for highly or low methylated regions. With **only 50M reads** it is possible to capture high resolution information to highlight differentially methylated regions (DMRs) with strong confidence at genome-wide scale.



Best samples preservation

No chemically intense reaction such as bisulfite conversion is used on our platform, therefore we guarantee the **best-in-class DNA samples preservation**.



Innovative, customizable platform

In most cases, our partners are looking for an **end-to-end solution** to guide them in their epigenetics adoption journey. We bring our expertise to consult on which ideas can be transformative and together we design tailored protocols. We are also developing and running bioinformatics algorithms that will scavenge the valuable information from the data sequenced. Once the discovery goals are achieved, we propose to develop ultra-focused epigenetic kits at orders of magnitude lower costs tailored to look at discovered biomarkers to be implemented in our partners products pipeline or clinical services.



* Minimum total count of CpG locations recorded based on experimental observations (N=20) and using latest T2TCHM13v2 FASTA reference genome.
https://www.ncbi.nlm.nih.gov/datasets/genome/GCF_009914755.1/

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TARGET APPLICATIONS

ANTI-AGEING

PHARMA

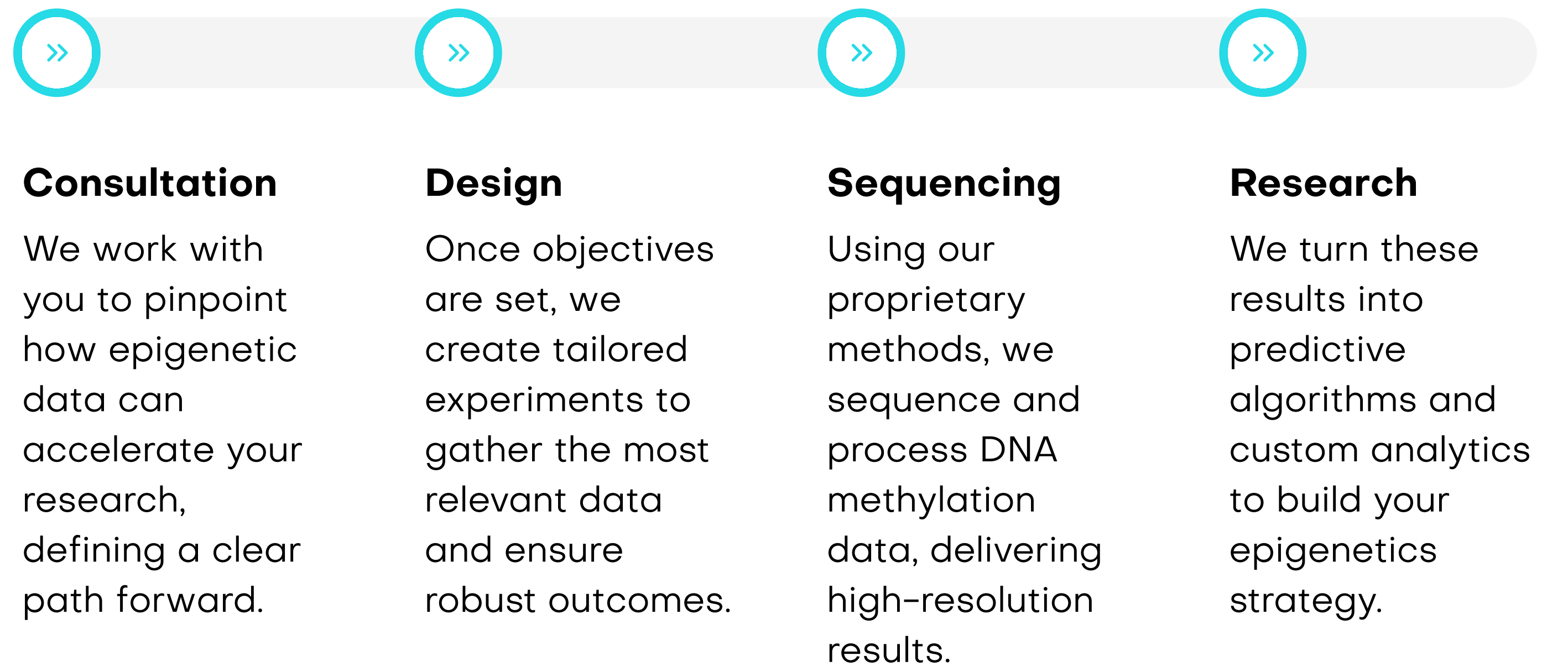
CELL BIOLOGY

AGRITECH

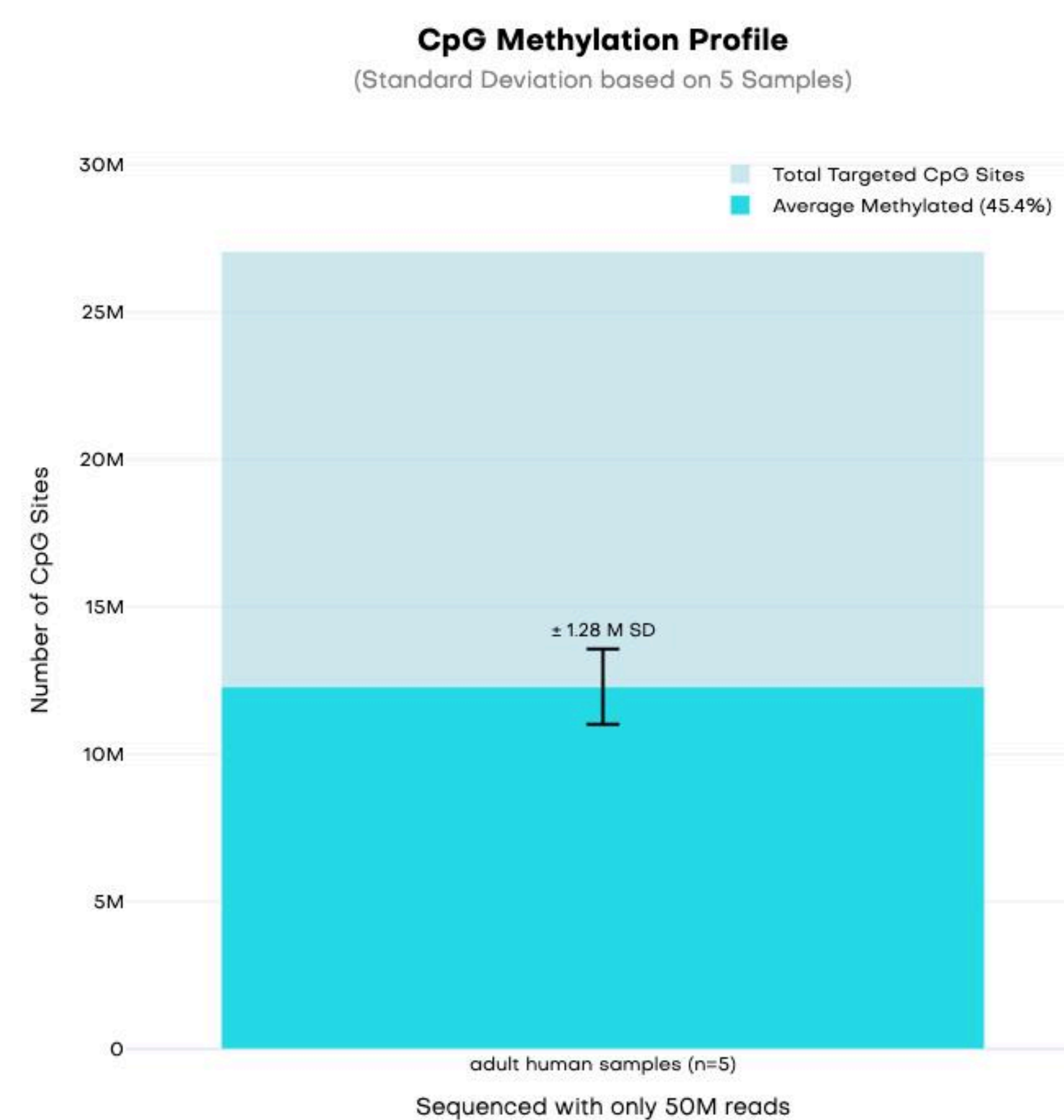
Features

- Supports high-throughput sequencing of highly multiplexed libraries, enabling cost-effective analysis of large sample cohorts.
- Provides extensive genome coverage for broad epigenetic insights by analyzing more than half of the entire methylome.
- Avoids harsh chemical treatments like bisulfite conversion, minimizing DNA degradation and preserving sample integrity.
- Enables analysis of methylation patterns in diverse genomic regions, including promoters, enhancers, gene bodies, and intergenic spaces.
- Detects methylation at CpH sites. Valuable for studying neurons, stem cells, plants, and other organisms with significant CpH methylation.
- Performs robust detection at only ~50 million reads for human sample (typically >10M mCpG sites identified), offering significant cost savings compared to deep WGBS. Sequencing depth is adjustable based on project goals.
- Accepts versatile sample input compatible with DNA extracted from various sources, including blood, bulk cells, tissues (e.g., skin), saliva, and buccal swabs. Please inquire regarding specific protocols for plasma/cfDNA.
- Allows customizable genomic annotation within specific genomic contexts by selecting annotations for promoters, enhancers, gene bodies, CpG islands, shores/shelves, and intergenic regions.
- Offers the option to add other epigenetic factors other than 5-mC such as 5-hmC, histone modifications with CHIP-seq or with ATAC-seq.
- Supports advanced bioinformatics with flexible analysis pipelines, including customizable options for biomarker discovery and data delivery formatted for machine learning applications.
- Utilizes secure data transfer methods tailored to client institutional requirements.
- Enables translation of discovery findings into cost-effective, targeted downstream assays suitable for product development or clinical service integration.

Collaboration Workflow



Specifications



	EPIC	RRBS	WGBS	
DNA coverage with CpGs targets	<1M	<5M	34M*	>25M**
DNA input quantity requirement	500ng	100ng	1000ng	20ng
Novel CpGs biomarkers discovery	✗	✗	✓	✓
Application for non-human DNA	✗	✗	✓	✓
High-throughput costs scaling	✗	✗	✗	✓
Low sequencing costs per CpG	✗	✗	✗	✓
No harsh chemical reaction	✗	✗	✗	✓

- Human DNA methylomes at base resolution show widespread epigenomic differences <https://doi.org/10.1038/nature08514>
- Reduced representation bisulfite sequencing for comparative high-resolution DNA methylation analysis <https://doi.org/10.1093/nar/gki901>
- A tiling resolution DNA microarray with complete coverage of the human genome <https://doi.org/10.1038/ng1307>

* 33,896,489 unique CpG sites were counted in the T2T-CHM13V2.0 human reference genome.
** Value based on experimental observations (N=20).

