

UNLEASH YOUR ULTIMATE SEQUENCING SPEED

Benchtop Genetic Sequencer **DNBSEQ-G99**

- **Rapid sequencing** Only 12 hrs for PE150 (from loading to FASTQ)
- Flexible throughput Independent loading and running of dual flow cells.
- **Bioinformatics integrated** Option to include built-in bioinformatics module to support sequencing and advanced analysis in a single machine.



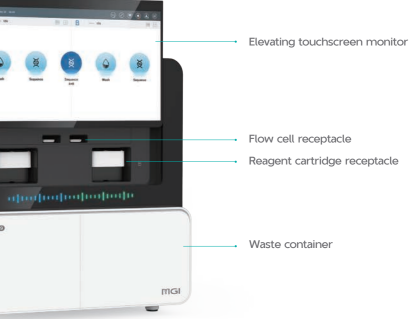
MGI'S PROPRIETARY **DNBSEQ**TM TECHNOLOGY

Genetic Sequencer **DNBSEQ-G99**



DNESEQ G99

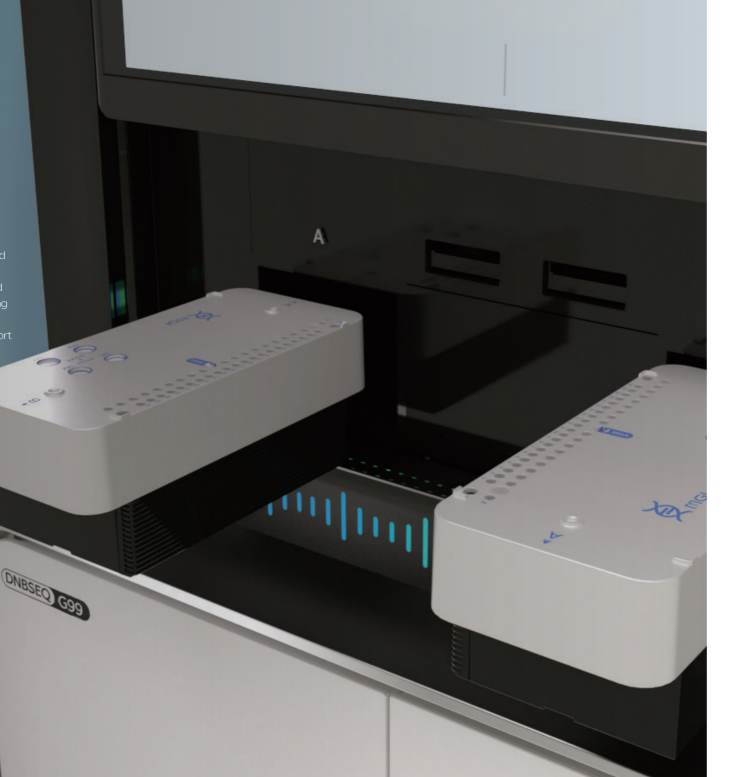
DNBSEQ-G99 is developed based on MGI's core DNBSEQ™ sequencing technology. Enabled by innovations in biochemistry, optics, fluidics, temperature control, and other core systems. DNBSEQ-G99 boasts the fastest speed amongst all medium-to-low throughput sequencers globally. DNBSEQ-G99 is especially applicable for targeted oncology panel sequencing, infectious disease sequencing. oncology methylation sequencing, small whole-genome sequencing, low-depth whole genome sequencing, individual identification, small panel sequencing of 24-28 samples, or whole-exome sequencing of 1-4 samples.



DNBSEQ-G99 also comes with an optional build-in bioinformatics module, which allows advanced analysis to begin automatically after the sequencing run. This facilitates a tremendously efficient and simple workflow, thus accelerating the application of omics technology to advance global life sciences and clinical research.

Designed for Simplicity

Newly designed flow cell, reagent cartridge, and user interface are introduced in DNBSEQ-G99. providing laboratory personnel with unparalleled ease and peace of mind in the entire sequencing workflow, A built-in bioinformatics module can also be included, achieving from sample to report all in one equipment.





Novel Reagent Cartridge Design

- Pre-loaded reagents within cartridge. one-step operation: simply press to load
- Sequencing and cleaning cartridges combined 2-in-1, cleaning initiates automatically after run.



Built-in Bioinformatics Module

- Advanced analysis begins automatically after run, and supports Bioanalysis by Sequencing (BBS) mode
- ZLIMS-compatible. achieve efficient workflow management and local data output





Intelligent Interaction

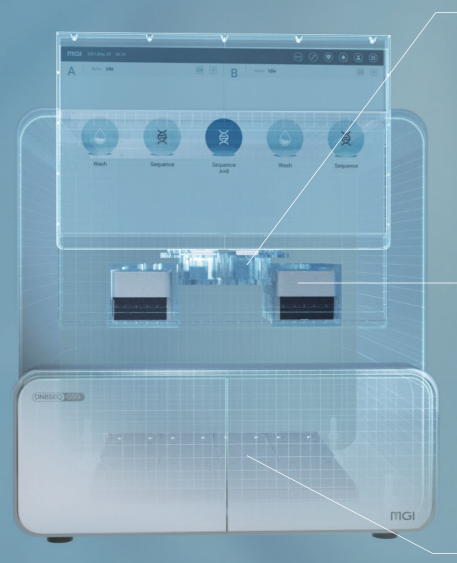
- Visualize the entire sequencing process in real-time
- Intuitive animations are included to guide flow cell loading, minimizing operational errors



Data Security

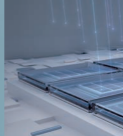
- Designed based on GDPR privacy protection requirement
- Secure storage to safeguard your sensitive data

Flexible Customize your run





Reagent cartridge



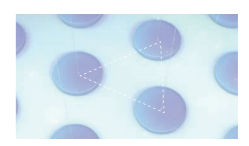
Analysis automatically begins afte



Bioinformatics module

Speed for your time-sensitive needs

To achieve the fastest sequencing speed in its range. DNBSEQ-G99 incorporates innovatively optimized flow cell. biochemistry process. fluidics. optics. temperature control and other core



High Density Flow Cell

- 600nm pitch high density patterning
- 68% more DNB loading per unit area
- Novel triangular configuration



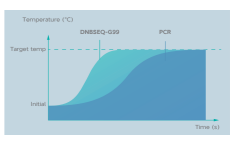
Surpassing the Optical **Diffraction Limit**

- In-house developed ultra-high quality objective lens
- Improvement of signal capture efficiency by decreasing scan area



Super Fast Biochemistry

- 10s rapid fluorescence reaction
- Biochemistry incubation reaction sped up from minutes-range to seconds-range



Rapid Temperature Control

- ~7 °C/s for heating and cooling
- Doubles the heating and cooling speed of conventional PCR instrument

Superior Performance

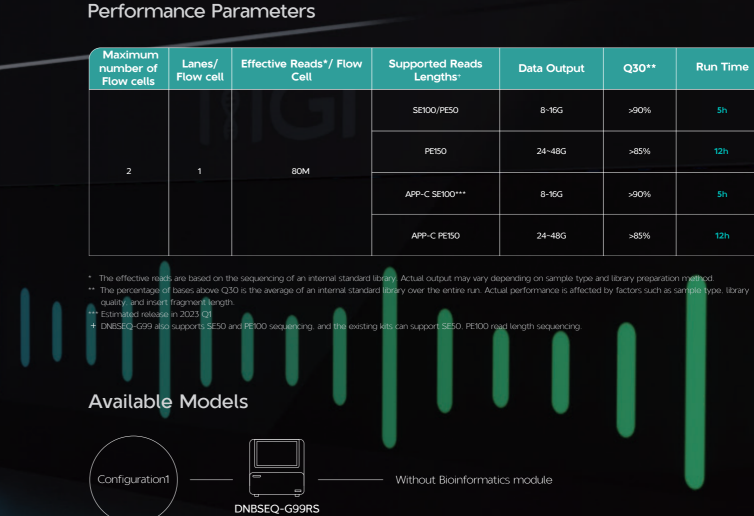
DNBSEQ-G99 delivers uncompromised high data quality. A multitude of applications can be executed on DNBSEQ-G99. such as targeted sequencing. small genome and Low pass WGS WGS sequencing, etc.

In addition. DNBSEQ-G99ARS supports the retrieval of data at intermediate time points under the Bioanalysis by Sequencing (BBS) mode. Users can obtain the first batch of summary report as quick as 2.5 hrs from the start of sequencing run (read length: SE40).

				Samples per Run		
Method	Application	Recommended read length	Data size per Sample	1 flow cell	2 flow cells	
				80M reads	160M reads	
	Oncology panel	PE150	Small panel: ~1Gb	24	48	
Targeted	Genetic disease diagnosis (small panel)	PE150	Thalassemia:~0.2Mb Deafness:~5Gb	4	8	
Capture/ Multiplex PCR	ATOPlex panel	PE100/PE150	Respiratory tract panel: 5M reads COVID-19 panel: 5M reads	16	32	
	WES	PE150 ~15Gb		1-2	2-4	
Methylation	Oncology targeted methylation panel	PE150	~5Gb	4	8	
Small Genome	Metagenomics for pathogen detection	SE50, SE100	20M reads	4	8	
Sequencing	Microbial WGS	PE100, PE150	Single bacterium: ~1Gb	16-24	32-48	
Low pass	NIPT	SE35 10M reads			16	
whole-genome sequencing	PGS	SE35		8	16	
Transcriptome sequencing	RNA-Seq	SE50/PE100/PE150	Expression profiling: ~1Gb Transcriptome: ~8Gb	2-4	4-8	

① Recommended data output and sample numbers are only for reference. actual application will require optimisation adjustments

Longer read lengths will be released in 2023.





ed Reads gths⁺	Data Output	Q30**	Run Time
/PE50	8~16G	>90%	5h
50	24~48G	>85%	12h
E100***	8-16G	>90%	5h
PE150	24~48G	>85%	12h



With Bioinformatics module

Len

SE100

APP-

Oncology Application Low Frequency Variants Detection

Experiment Scheme

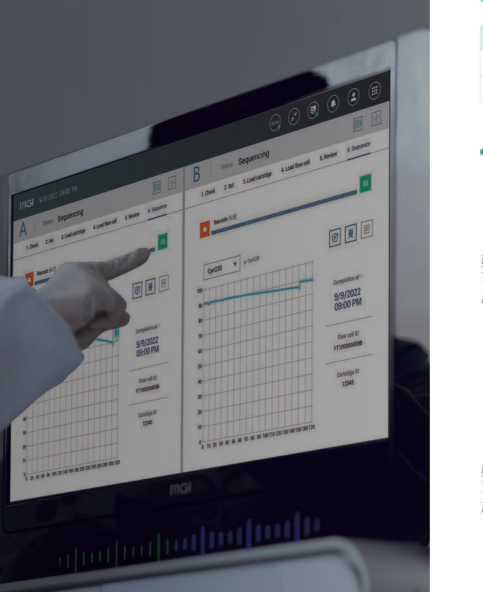
Sample: Lung cancer ctDNA standards, diluted to 1%. 0.5%, 0.2%, and 0.1% variant ctDNA samples Library prep: Targeted capture kit from third party Sequencing strategy: PE100 dual-barcode sequencing. 4 repeated runs were tested Objective: To test DNBSEQ-G99 variant detection capability

Sequencing Summary

The 4 runs generated 126M reads on average, Q30 >93%, with excellent uniformity observed.

Analysis Summary

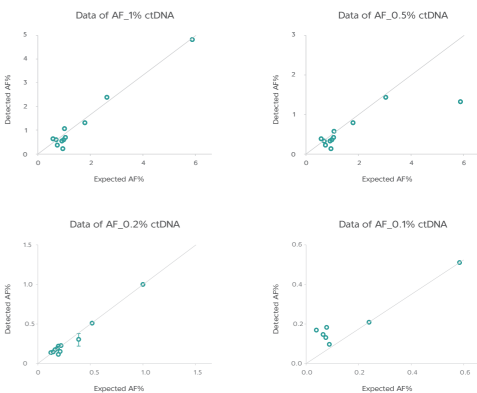
• 100% detection of SNV mutation sites in the samples (1%,0.5%,0.2%, 0.1%).

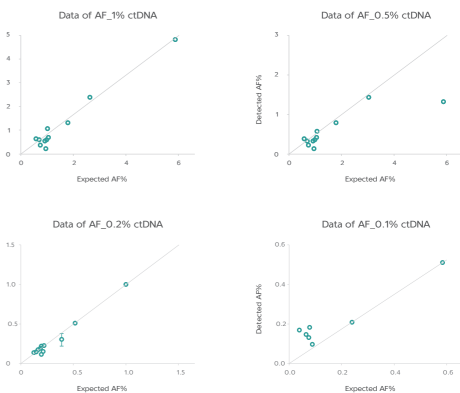


• Sequencing Result

	Total reads (M)	Q30	EstErr(%)	Time (h)
Mean value	126.22	94.00	0.22	9.15
Standard deviation	7.41	0.52	0.02	0.11

• Analysis Result





Oncology Application Methylation Sequencing (Targeted Capture)

9/9/2022 09:00 PM

12345

华大智道 MGI

Experiment Scheme

Sample: Fragmented DNA from 4 cell lines Library prep: Targeted capture double stranded library kit from third party Sequencing strategy: PE100 dual-barcode sequencing. 2 repeated runs were tested Objective: To evaluate DNBSEQ-G99's compatability with low-diversity libraries, and data uniformity for targeted methylation regions.

Sequencing Summary

Without addition of a spike-in balanced library, 2 runs had an averafe output of 101M reads, with Q30 >86%, showing good compatibility with the low diversity libraries.

Analysis Summary

- Highly concordant average methylation fractions (AMF) measured between both runs for all tested samples.
- For results of the same samples on different instruments: AMF of the samples is highly consistent.



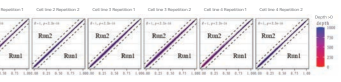
• Sequencing Results

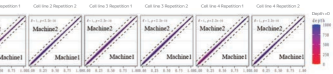
	Total reads (M)	Q30(%)	SplitRate (%)	Time (h)
Run 1	103.12	86.42	98.36	9.20
Run 2	99.08	90.57	98.02	9.15

• Analysis Results

1.00	-1, p+1 h-14	1 1-1.7-1.30		8-1,9+2.30
8,75	Run2	Run2	1	Run2
0,25	Runi	1	Run1	1
	0 0.25 0.50 0.75	1.000.00 0.23	5.51 4.75 1.90	1

Cell line 1 Repetition 1	Cell line 1 Repetition 2	Cell line 2 R
1.11 1.11 1.11 Machine2	Machine2	Machine2
		1
A.00 1.25 1.38 1.71 1.1	Machinel	1





Small Genome Sequencing Pathogen detection

Experiment Scheme

Sample: 4-pooled 1% reference microbial community standards Library prep: MGIEasy FS DNA Library Prep Set Sequencing strategy: PE100 single-barcode sequencing Objective: Assess DNBSEQ-G99's capability to identify unknown

Sequencing Summary

Output of 110M reads, Q30 >95%, exceeding data amount required for analysis.

Analysis Summary

- Pathogen fast identification (PFI) was used for analysis, and the pathogen identification results were consistent with reference microbial community in terms of detected species and abundance.
- Fluctuation in abundance CV was lower than 2%, indicating high accuracy.

• Sequencing Results

	Total reads (M)	Q30(%)	SplitRate(%)	Time(h)
Output	110.17M	95.22	96.95	9

• Analysis Results

Species	Sample 1	Sample 2	Sample 3	Sample 4	Standard abundance	mean	SD	CV
Salmonella enterica	15.87%	15.60%	15.72%	15.87%	12.00%	15.77%	0.13%	0.82%
Pseudomonas aeruginosa	14.12%	14.00%	13.83%	14.12%	12.00%	14.02%	0.14%	1.00%
Bacillus subtilis	13.30%	13.14%	13.46%	13.30%	12.00%	13.30%	0.13%	0.98%
Escherichia coli	11.75%	12.32%	11.90%	11.75%	12.00%	11.93%	0.27%	2.26%
Enterococcus faecalis	11.13%	11.30%	11.24%	11.13%	12.00%	11.20%	0.08%	0.71%
Listeria monocytogenes	11.12%	11.02%	11.12%	11.12%	12.00%	11.10%	0.05%	0.45%
Staphylococcus aureus	10.18%	10.23%	10.30%	10.18%	12.00%	10.22%	0.06%	0.59%
Limosilactobacillus fermentum	9.52%	9.48%	9.46%	9.52%	12.00%	9.50%	0.03%	0.32%
Cryptococcus neoformans	1.49%	1.49%	1.52%	1.49%	2.00%	1.50%	0.02%	1.33%
Saccharomyces cerevisiae	1.47%	1.42%	1.45%	1.47%	2.00%	1.45%	0.02%	1.38%

Small Genome Sequencing Phage Assembly

Experiment Scheme

Sample: 16 pure bacteriophage cultures Library prep: MGIEasy universal DNA library prep set Test Strategy: PE150 dual-barcode Test Purpose: Assess DNBSEQ-G99 capability for assembling whole bacteriophage genome

Sequencing Summary

Output of 106M reads, Q30 >92%, exceeding data amount required for analysis.

Analysis Summary

- Conserved protein genes for phage capsid, portal, and terminase were successfully detected in the G99 assemblies.
- G99 outperforms other platform for bacteriophage whole genome assembly, achieving higher assembly integrity.

• Sequencing Results

	Total reads
Output	106.99

• Analysis Results



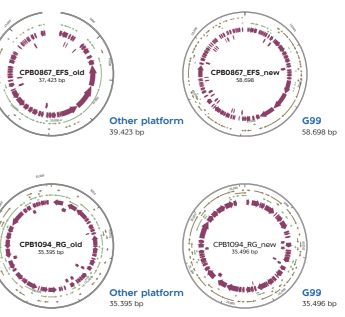








Q30(%) SplitRate(%) Time(h) s (M) M 92.32 97.97 12



Hardware Specifications

Model	DNBSEQ-G99RS	Outputs FASTQ files		
Model	DNBSEQ-G99ARS	Equipped with bioinformatics module for advanced analysis		
Dimensions	607*680*640 mm			
	Rated Voltage	100V-240V		
Power	Rated frequency	50/60Hz		
	Rated Power	1000 VA. [working current]: ≥10A		
	LCD touch screen			
Touch Screen	Touch screen size	1.5inch		
	Touch screen resolution	1920×1080		
Maximum Sound Pressure	75 dB(A)			
Shell Protection Grade	IPXO			
	Temperature	19-30 ℃		
Operating Environment	Relative Humidity	20-80 %RH		
Requirements	Atmospheric Pressure	70 kPa-106 kPa		
	Maximum Altitude (above sea level)	3000 m		
	CPU	Intel 19-10900e 2.80 GHz		
Computer	Internal Storage	64 GB		
Configurations	HDD	6 TB		
	Operating System	Windows 10		

Ordering Information

Cat. No	Product Name
900-000561-00	DNBSEQ-G99RS
900-000560-00	DNBSEQ-G99ARS
940-000409-00	High-throughput Sequencing Set (G99 SM FCL SE100/PE50)
940-000410-00	High-throughput Sequencing Set (G99 SM FCL PE150)
940-000413-00	High-throughput Sequencing Set (G99 SM FCL APP-C PE150)
940-000624-00	DNBSEQ-G99RS Cleaning Reagent Kit
Selected as needed	UPS

*Unless otherwise informed, StandardMPS and CoolMPS sequencing reagents, and sequencers for use with such reagents are not available in Germany, Spain. UK. Hong Kong, Sweden, Belgium, Italy, Finland, Czech Republic, Switzerland, Portugal, Austria and Romania. No purchase orders for StandardMPS products will be accepted in the USA until after January 1, 2023.

MGI Global Presence

Technical Support Available Globally

The MGI technical support team has a complete global coverage, including technical service centers in major international regions and multiple locations to maximize customer satisfaction.



Spare part centers in Shenzhen, Wuhan, Qingdao, Tianjin, Hong Kong (China), Brisbane (Australia). Singapore, and Riga (Latvia) to ensure sufficient supply of parts for machine maintenance.



Ane

Online technical support accessible worldwide. with a fully functioning call center (Toll-Free Hotline 4000-966-988) (9:00AM-12:00PM.13:00PM-18:00PM. Beijing time (GMT+8), workday). Multi-language online training courses coming soon.

Comprehensive Instrument Service and Warranty Plans Globally



Warehouses in Shenzhen. Wuhan, Qingdao. Tianjin. Hong Kong, Taipei. Singapore(Asia-Pacific), Brisbane (Australia Oceania), Riga (Latvia, Europe), and San Jose (the USA, Americas) are established to ensure sufficient supply of maintenance parts for major regions.



Free installation and system verification services (inclusive of necessary reagents and consumables) to quickly turn your investment into production.



MGL is responsible for any manufacturing defects or faults on the system within the warranty. Warranty covers labor parts and travel charges.



One free instrument preventive maintenance is provided with warranty, along with a variety of available extended warranty support plans.

d the world provide timely and effective technical support and

MGI Genetic Sequencers



DNBSEQ-E25 Reads per flow cell: 25 Number of flow cells: 1



DNBSEQ-G99 Reads per flow cell: 80M Number of flow cells: 2 Data output: 8-48GB



DNBSEQ-G50 Reads per flow cell: 100

Reads per flow cell: 100-500M Number of flow cells: 1 Data output: 10-150GB



DNBSEQ-G400 Reads per flow cell: 300-1800M Number of flow cells: 2 Data output: 55-14400GB



@About MGI Tech Co., Ltd

MGI Tech Co., Ltd. (referred to as MGI) is committed to building core tools and technology to lead life science through intelligent innovation. MGI focuses on R&D, production and sales of DNA sequencing instruments, reagents, and related products to support life science research, agriculture, precision medicine and healthcare. MGI is a leading producer of clinical high-throughput gene sequencers, and its multi-omics platforms include genetic sequencing, mass spectrometry, medical imaging, and laboratory automation. Founded in 2016, MGI has more than 1800 employees, nearly half of whom are R&D personnel. MGI operates in more than 80 countries and regions and has established multiple research and production bases around the world. Providing real-time, comprehensive, life-long solutions, its vision is to enable effective and affordable healthcare packages for all.



DNBSEQ-T7

Reads per flow cell: 5000M Number of flow cells: 4 Data output: 250-6000Gb



DNBSEQ-T10x4 Reads: 27.5-45Billion Number of flow cells: 8 Data output: 66-72TB



MGI Tech Co.,Ltd

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