

FASTASeq 300

High-throughput sequencing platform

—Flexible · Rapid · Easy-to-use



Company Introduction



GeneMind Biosciences

Explore Life's Mysteries for Better Healthcare

Established in 2012, GeneMind Biosciences Co., Ltd. is headquartered in Luohu, Shenzhen, with more than 10,000 square meters of working space including R&D lab and GMP production line.

Specializing in the independent R&D and manufacturing of molecular diagnosis technology platform centered on sequencing system, GeneMind is committed to working with genetic testing service providers and life science research institutions to build an industrial ecosystem that serves human life and health.

Milestones



Product Introduction

FASTASeq 300 High-throughput Sequencer

FASTASeq 300 is a desktop sequencer focused on targeted sequencing, whole-genome low-depth sequencing, with innovative breakthroughs in sequencing chemistry, high-density flowcell, fluid design and base identification algorithms, bringing users more flexibility, more consistent data quality and faster delivery.



Flexible

- Support two types of flow cells and multiple read lengths
- Multi-time output (MTO)
- Libraries can be loaded on each flow cell lane automatically or manually

Easy-to-use

- Pre-configured, plug-and-play cartridges
- Support RFID reader module
- Libraries can be directly used for sequencing after preparation
- Automatic post-run wash

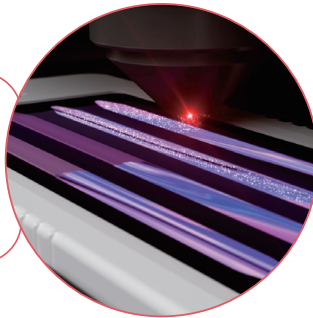
Rapid

- mNGS (SE50 4.5hrs)
- Targeted sequencing (PE75 11.5hrs)
- Targeted sequencing (PE150 19.5hrs)

Core Technologies

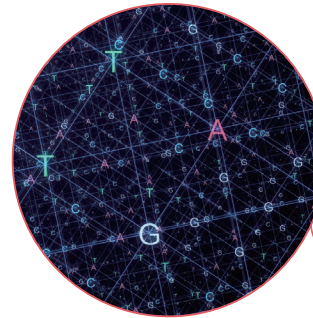
- **Constant temperature chemistry**
Sequencing speed increased by 1x

- Biochemical and signal acquisition processed in parallel, making sequencing more efficient



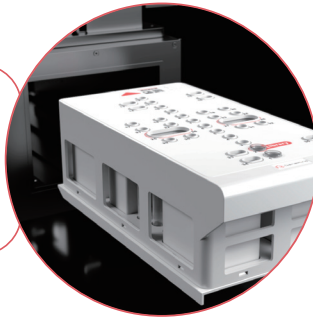
- **High-density flow cell**
Signal density increased by 1x

- New high-density flow cell combined with optimized Basecall algorithm
- Accurate identification of high density signals and efficient output of high quality data



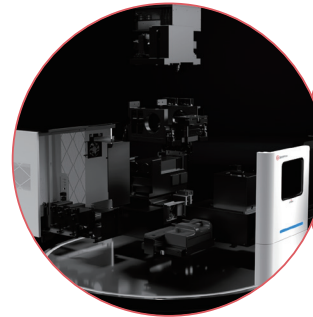
- **Bi-directional sequencing**
Reagent usage reduced by 50%

- Optimized reagent formulation and fluid configuration
- Promote a significant reduction in reagent consumption per unit of data output



- **Localisation**
90% of core components produced domestically

- Increased proportion of core components of sequencers made domestically
- More stable and controllable supply chain and production cycle



Configuration

Flow cell	Flowcell Type	Lane	Reads ¹	Read Length	Output	Q30 ¹	index (0) Sequencing Time (index 0)	index (8+8) Sequencing Time (index 8+8)
Single	FCM	4	100 M	SE50 ³	5 Gb	85%	4.5 hrs	5.5 hrs
				SE75	7.5 Gb	85%	6 hrs	7 hrs
				PE75	15 Gb	85%	11.5 hrs	13 hrs
				PE150	30 Gb	85%	19.5 hrs	21 hrs
	FCH	4	250 M	SE50 ³	12.5 Gb	85%	5 hrs	6.5 hrs
				SE75	18.5 Gb	85%	6.5 hrs	8 hrs
				PE75	37.5 Gb	85%	12.5 hrs	14 hrs
				PE150	75 Gb	85%	22.5 hrs	24 hrs

1.The number of reads and Q30 are based on sequencing using standard libraries; performance may vary depending on library type and quality, insert size, loading concentration, and other experimental factors.

2.Sequencing time includes the time from sample loading to base identification to generate the Basefile file.

3.SE50 sequencing mode is implemented based on the FASTASeq 300 sequencing kit V1.0 (FCM/FCH-D SE075-D).

Application

Application	Read length	Data/sample	FCM*1	FCH*1
			100M	250M
NIPT/PGT-A/CNV-Seq	SE75	5 M reads/sample	20	48
Small tumor panel (tissue)	PE75	0.2 Gb/sample	>48	>96
Small tumor panel (ctDNA)	PE150	5 Gb/sample	6	16
Large tumor panel (tissue)	PE150	5 Gb/sample	6	16
mNGS	SE75	20 M reads/sample	4	12
tpNGS	PE150	0.1-1 M reads/sample	>96	>192

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

Application summary



A total of 60 runs were counted (FASTASeq 300 FCH)

Parameter	Mean
Q30	90%
Output reads	310 M

Application

Application—PGT-A/CNV-Seq/NIPT

- Sequencing:FASTASeq 300 Chromosomal Abnormality Detection
- Sample type:Construct PGT-A and CNV-seq libraries based on PGT-A reference standard and CNV-seq reference standard
- Read length:SE75

Table 1. Consistency assessment of CNV-Seq results

CNV-Seq Reference Standard Sequencing								
No.	Sample	Effective data (reads)	Average GC	Average coverage	Average depth	Results	Reference standard	Consistency
1	GM12	4811803	38.15%	7.34%	0.159	+13	T13	Yes
2	GM15	5187339	38.46%	7.88%	0.173	+16	T16	Yes
3	GM21	5394891	38.26%	8.18%	0.178	-X	XO	Yes
4	GM27	5302386	38.18%	8.05%	0.177	-22 (q11.21, ~ 2.96Mb)	22q11.21 (del,2.5Mb)	Yes
5	GM31	4843680	38.50%	7.39%	0.162	-7 (q11.23, ~ 1.78Mb)	7q11.23 (del,1.8Mb)	Yes
6	GM40	5845360	38.06%	8.82%	0.192	+X—31%;-1 (p36.33-p36.22, ~ 10.65Mb)—31%	XXY—30%;1p36.33-p36.22 (del,11.0Mb)—30%	Yes
7	GM42	4906427	38.23%	7.48%	0.162	Normal	Regular Sample	Yes

Table 2. Consistency assessment of PGT-A results

PGT-A Reference Standard Sequencing								
No.	Sample	Effective data (reads)	Average GC	Average coverage	Average depth	Results	Reference standard	Consistency
1	GM02008	4641367	39.59%	5.92%	0.169	-11(q23.3-q25, ~ 14.7Mb)	-11(q23.3-q25, ~ 14.7Mb)	Yes
2	GM06473	5123946	39.33%	6.70%	0.183	-1(q43-q44, ~ 9.9Mb)	-1(q43-q44, ~ 9.9Mb)	Yes
3	GM06097	4964662	39.43%	6.72%	0.179	-17(p13.3-p13.2, ~ 4.2Mb)	-17(p13.3-p13.2, ~ 4.2Mb)	Yes
4	GM13325	5330814	39.49%	7.79%	0.197	-22(q11.21, ~ 2.3Mb)	-22(q11.21, ~ 2.3Mb)	Yes
5	GM25372	4115286	39.39%	5.42%	0.151	-17(p11.2, ~ 1.2Mb)	-17(p11.2, ~ 1.2Mb)	Yes
6	GM01359	3914767	40.17%	5.84%	0.151	+18	+18	Yes
7	GM02767	3872356	39.47%	5.08%	0.141	+21	+21	Yes

Conclusion:

FASTASeq 300 has excellent detection of chromosomal abnormalities.

Application

Application-Target tumor panel

- Sample type: HD832 (FFPE)
- Library prep: TruSight Oncology 500
- Sequencing platform: GenoLab M, FASTASeq 300, Competitor A (NS platform) and Competitor B (NV platform)
- Read length: PE150
- Analysis: Extract 100M reads/sample to do deep analysis

Platform	FASTASeq 300	GenoLab M	NS Platform	NV Platform	AF	Mut Number	FASTASeq 300	GenoLab M	NS Platform	NV Platform
Q30	91.33%	88.58%	83.45%	91.25%	AF ≤ 10%	9	88.89%	77.78%	77.78%	77.78%
Total SE reads (M)	100	100	100	100	10% < AF ≤ 20%	24	100.00%	100.00%	100.00%	100.00%
Fold80	1.41	1.43	1.41	1.56	20% < AF ≤ 30%	49	100.00%	100.00%	100.00%	100.00%
Unique mapping rate	99.34%	99.22%	97.45%	98.14%	AF > 30%	124	100.00%	100.00%	100.00%	100.00%
Target region rate	84.74%	84.37%	83.74%	83.05%	NGS no value	6	83.33%	83.33%	83.33%	83.33%
Target mean depth	1182	1551	1178	1165	Total	212	99.06%	98.58%	98.58%	98.58%

TAT: 24h (FASTASeq 300 FCH PE150 Dual index)

Conclusion:

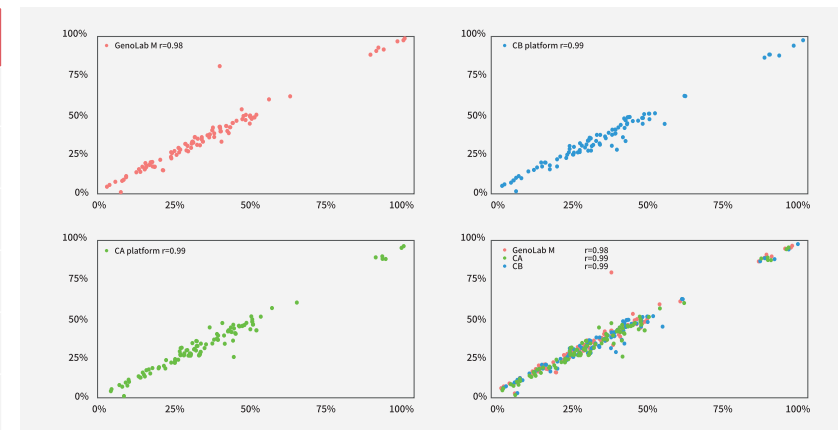
High data quality (Q30 > 91%, unique mapping rate > 99%) for FASTASeq 300, the sequencing time only takes 24h for PE150. And the results of FASTASeq 300 mutation detection are consistent with GenoLab M and other competitors.

Application

Targeted tumor panel

- Sample type: HD832 (FFPE)
- Library prep: TruSight Oncology 500
- Sequencing platform: GenoLab M, FASTASeq 300, Competitor A and Competitor B
- Read length: PE150
- Analysis: Extract 100M reads/sample to do deep analysis

AF	Mut Number	FASTASeq 300	GenoLab M	NS Platform	NV Platform
AF ≤ 10%	9	88.89%	77.78%	77.78%	77.78%
10% < AF ≤ 20%	24	100.00%	100.00%	100.00%	100.00%
20% < AF ≤ 30%	49	100.00%	100.00%	100.00%	100.00%
AF > 30%	124	100.00%	100.00%	100.00%	100.00%
NGS no value	6	83.33%	83.33%	83.33%	83.33%
Total	212	99.06%	98.58%	98.58%	98.58%



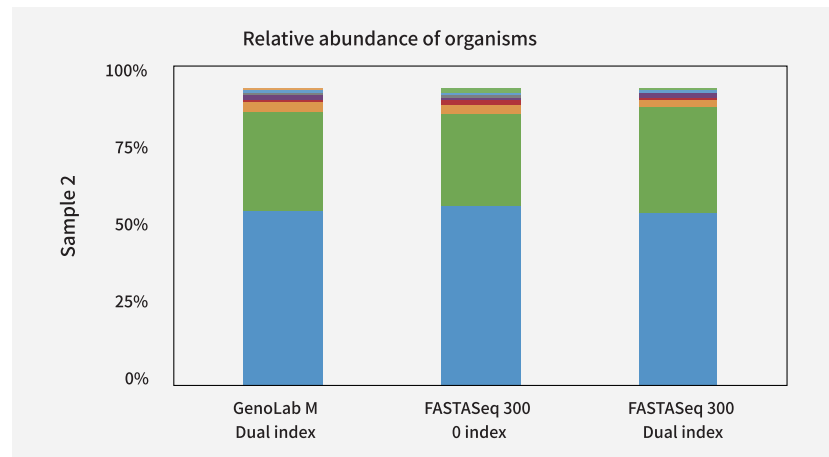
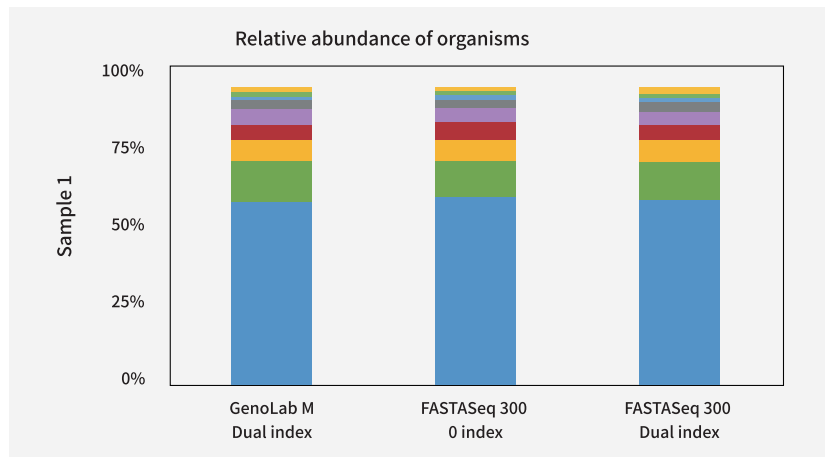
Conclusion:

The results of FASTASeq 300 mutation detection are consistent with GenoLab M and other competitors.

Application

Application-mNGS

- sample type: Clinical samples (Alveolar lavage fluid, sputum, urine, tissue, hydrothorax)
- Library: 6 mNGS libraries
- Sequencing platform: GenoLab M, FASTASeq 300
- Read length: SE50
- Data analysis: 50M reads/sample for deep analysis



Conclusion:

The microbial detection of FASTASeq 300 are consistent with GenoLab M, and the clinical judgment results of all test samples are 100% accurately detected.

Application

Application- De novo WGS (single bacteria assembling)

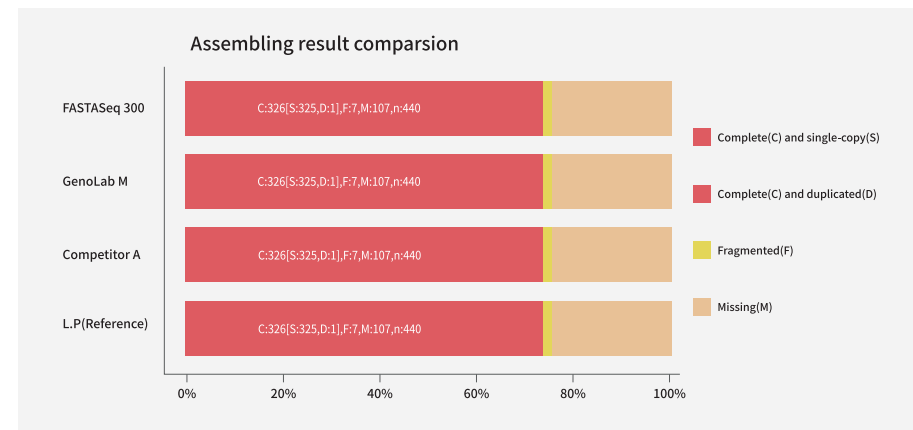
- Sample type: Using Legionella pneumophila subsp. Pneumophila construct WGS library
- Sequencing platform: FASTASeq 300, GenoLab M, Competitor A(CA)
- Read length: PE150
- Data analysis: 100× effective reads/sample for de nova analysis

Assembling results			
Platform	FASTASeq 300	GenoLab M	CA
Q30	92.10%	91.93%	90.60%
Total Length (bp)	3387746	3376652	3377221
Reference Length (bp)	3409194	3409194	3409194
Genome Fraction	99.02%	99.02%	99.04%
GC Content	38.22%	38.23%	38.23%
Reference GC Content	38.33%	38.33%	38.33%
Contigs	28	27	29
Largest Contig (bp)	989763	989763	989763
Contig N50 (bp)	363258	363258	363482

TAT: 19.5h (FASTASeq 300 FCM PE150 0 index)

Conclusion:

Data quality and assembling results are quite comparable with GenoLab M and CA, the sequencing time only takes 19.5h for PE150.



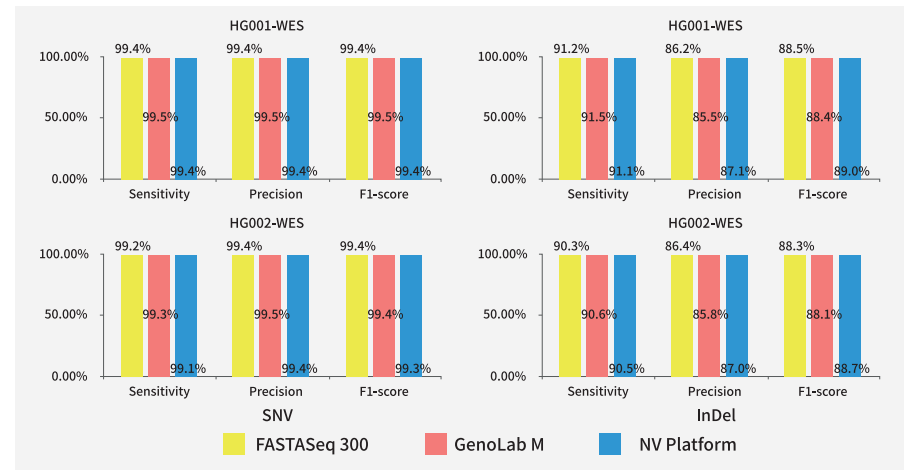
Application

Application-WES

- Sample type:HG001(NA12878), HG002(NA24835)
- Library prep:Agilent SureSelect Human All Exon V6
- Sequencing Platform:FASTASeq 300,GenoLab M, Competitor A(NV platform)
- Read length:PE150
- Analysis:12Gb/sample for deep analysis

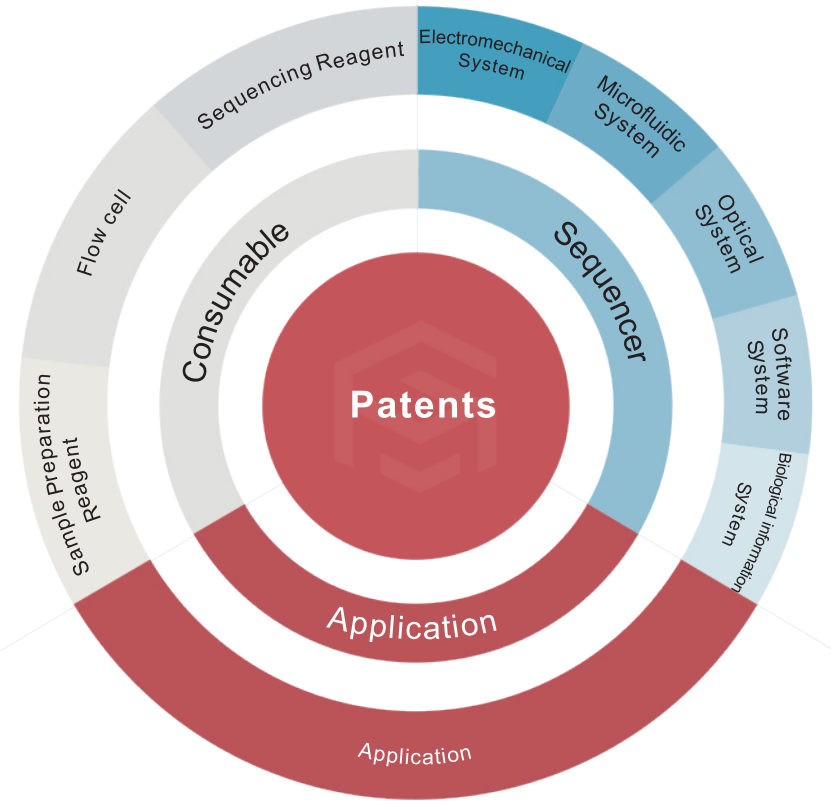
Platform	HG001-WES			HG002-WES		
	FASTASeq 300	GenoLab M	Competitor A	FASTASeq 300	GenoLab M	Competitor A
Raw data (Gb)	12	12	12	12	12	12
Clean data (Gb)	11.99	11.99	11.95	11.99	11.99	11.96
Properly paired reads	99.49%	99.62%	98.81%	99.60%	99.65%	98.92%
Capture efficiency	77.91%	77.93%	78.29%	78.78%	78.49%	78.91%
Mean sequencing depth	124.59	124.64	124.78	125.95	125.75	126.05
Duplication ratio	3.46%	5.48%	21.03%	5.79%	5.22%	19.60%
Coverage at least 0×	99.64%	99.64%	99.65%	99.87%	99.87%	99.85%
Coverage at least 4×	99.52%	99.52%	99.55%	99.65%	99.67%	99.56%
Coverage at least 10×	99.06%	99.05%	99.19%	98.57%	98.68%	98.16%
Coverage at least 30×	94.95%	95.16%	94.67%	89.56%	90.32%	87.33%

Conclusion: High data quality(Q30>91%, low duplication rate<5%)



Conclusion: FASTASeq 300 excels in sensitivity and precision for germline mutation (SNV and InDel) detection.

Intellectual Property & Qualifications



Sequencing systems are multidisciplinary specialities that combine optics, fluids, algorithms, chemistry and molecular biology. Since its establishment, GeneMind has been specializing in the independent R&D and manufacturing of molecular diagnosis technology platform centered on sequencing system. We have continuously innovated and accumulated a lot of intellectual property rights in the underlying technologies such as instrument hardware, reagent, flowcell and software algorithms.

As of October 2023, GeneMind has more than 300 granted domestic and foreign patents, and has successfully obtained ISO 13485 medical device quality management system certification. The sequencer, reagent kits and other products have also received CE certification, NMPA medical device approval.

300+
Patents granted

ISO 13485:2016
Quality Management System Certification

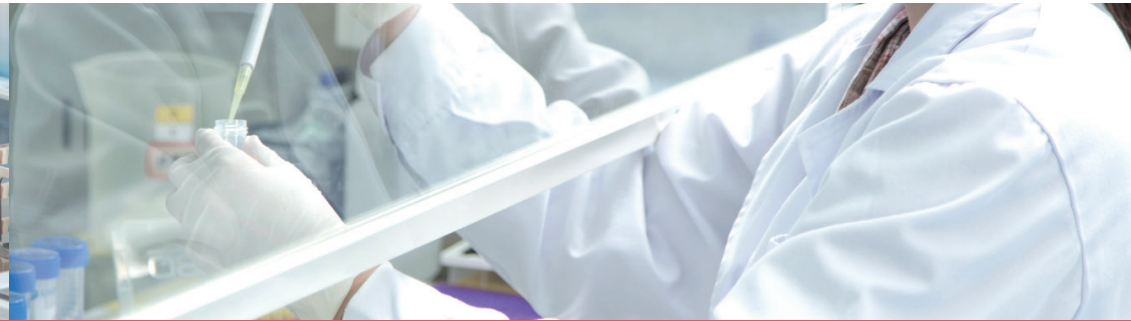
34
NMPA approval/ CE IVD registration

10+
Academic papers

Independent Research & Development

In addition to the innovative breakthroughs in sequencer development, GeneMind has also realized the independent R&D and production of core materials such as enzymes, nucleotide analogues, fluorochrome and flowcell. The company has the ability to develop and manufacture the total solution of "instrument-reagent-flow cell-software" independently, and are able to provide customers total solution with good quality, low cost and short delivery time.

The total area of R&D and production in use is nearly 10,000 square meters, with sequencer production factory, GMP reagent kit production line, flow cell laboratory, organic synthesis laboratory and enzyme engineering laboratory. The designed annual capacity of sequencer is 1000 units and the annual capacity of kits is 2.4 million tests.



● Sequencer Production Line
Designed with annual production capacity of 1000 sequencers

● Flow cell Lab
In-house production of sequencing flowcell

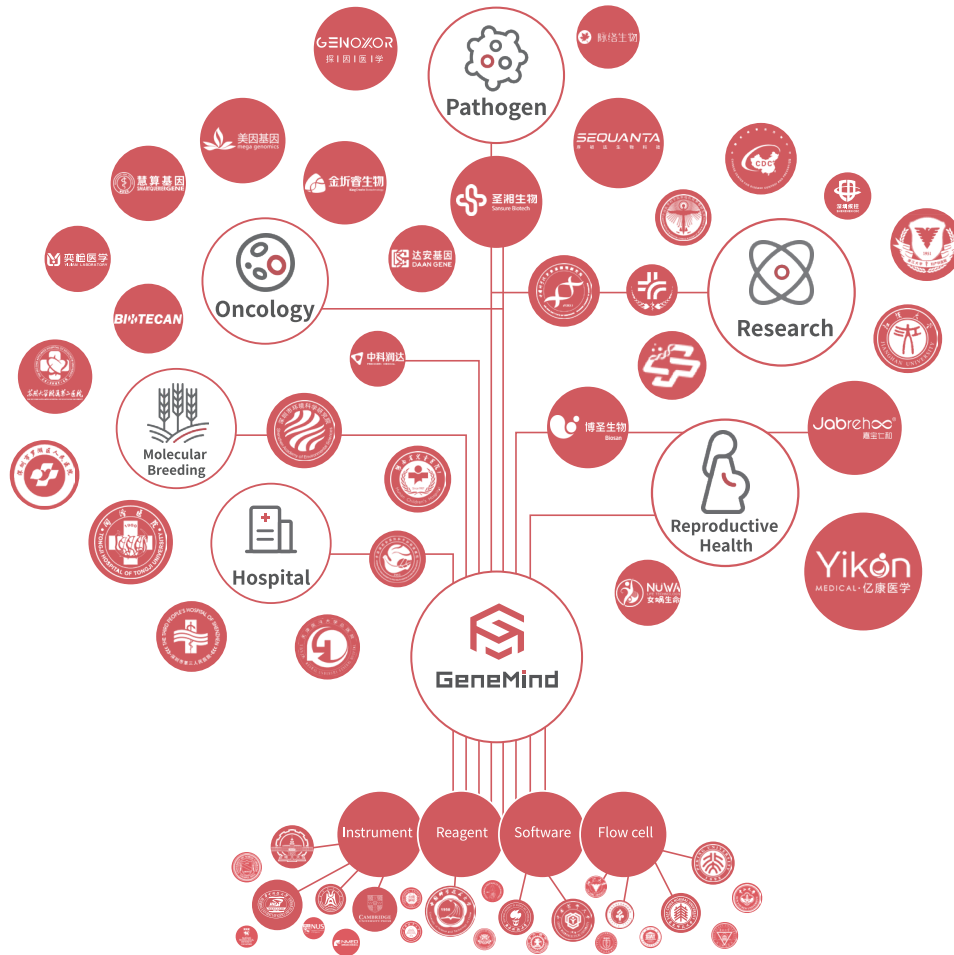
● Organic Synthesis Lab
Able to synthesize our own nucleic acid and fluorochrome



● Reagent Kits Production Line
Designed as a GMP workshop to produce kits for 2.4 million tests annually

● Enzyme Engineering Lab
Independent research and development of core enzyme

Partners



Product Information and Specification

Product Information

Sequencer	
SQ00019	FASTASeq 300 Sequencing Platform
Reagent Kit	
S000181	FASTASeq 300 Sequencing Set V1.0 (FCM-D SE075-D)
S000186	FASTASeq 300 Sequencing Set V1.0 (FCM 150cycles)
S000189	FASTASeq 300 Sequencing Set V1.0 (FCM 300cycles)
S000191	FASTASeq 300 Sequencing Set V1.0 (FCH-D SE075-D)
S000194	FASTASeq 300 Sequencing Set V1.0 (FCH 150cycles)
S000197	FASTASeq 300 Sequencing Set V1.0 (FCH 300cycles)

Product Specification

Specification	
Dimensions	684 mm × 644 mm × 615 mm
Weight	145 kg
Power Requirements	2000 VA
Power Requirements	100-240 V~, 50 /60 Hz
Operating Environment	Temperature:19°C-25°C Humidity: 20%-80% Relative Humidity, non-condensing Altitude:0-3000m
Instrument Control Computer	CPU: Intel Xeon Silver 4216 Memory: 32GB DDR4*6 Solid-state drive: 480GB Hard drive: 2TB Operating system: window10x64



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