



Company Introduction



Milestones



SURFSeq: Key Functionalities



Wide range application

Dual flow cell to support 5 different modes/run

Compatible with most common used applications



FCM PE150≤24h

Flexible FQ upload node, no need to wait for one flow cell finish the run



Good data quality Balanced mode: Q30≥90%, Q40≥85% Enhanced mode: Q30≥95%, Q40≥90%

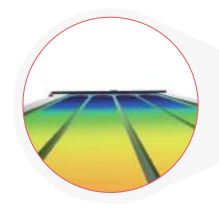
Lower start-up cost

Wide range selection for sample volume in one run

Most cost-effictive for each sample



Product Innovation



Patterned array flow cells enable 2X higher density of signals



HyEND advanced surface amplification technology makes over 80% utilization rate of each flow cell



Supports real-time FastQ generation to allow customers to obtain data from different applications of the same sequencing run

SURFSeq 5000 Specification

			Spec	ification											
Flow cell	Lane	Throughput (Reads/FC)	Туре	Read Length	Data output ¹ (FC×2)	Balance Mode	Enhanced Mode	TAT ²							
			100cycles	SE100	50 Gb × 2			12 hr							
FCM	4	500 M	Toocycles	PE50	50 Gb × 2			12 hr							
I CIVI	4	300 W	200cycles	PE100	100 Gb × 2			18 hr							
			300cycles	PE150	150 Gb × 2			24 hr							
			50cycles	SE50	100 Gb × 2	Q30≥90%	Q30≥95%	14 hr							
					2000 M	2000 M	2000 M	2000 M	2000 M	100cycles	SE100	200 Gb × 2	Q40≥85%	Q40≥90%	21 hr
FCH	4	2000 M	2000 M	2000 M						2000 M	2000 M	Toocycles	PE50	200 Gb × 2	
			200cycles	PE100	400 Gb × 2			34 hr							
			300cycles	PE150	600 Gb × 2			47 hr							
			Main	Features											
	Two flow cell with 5 differ	rent sequencing modes		Individually addre	essable flow cell lanes v	vith manual onboard	l independent lane lo	ading							
Support one or two flow cell in a time				User-friendly interface to easily switch read/index sequence											
	Obtain data in real-time a	at high-speed	Obtain data in batches based on different applications in one run												
	RFID recognize cartaidge	information		Automatical post-run wash											
	KFID recognize cartaidge	information		Automatical post-	-run wash										

^{1.}The percentage of bases above Q30 and run time is the average of an internal standard library over the entire run. The actual performance is affected by factors such as sample type, library quality, and insert fragment length.

^{2.}Run time was calculated based on dual flow cell mode, and includes sample loading, sequencing, base calling and data processing.

SURFSeq 5000 application

		Dete/essents	FCM*1	FCM*2	FCH*1	FCM*1+FCH*1	FCH*2
Application	Read length	Data/sample	500 M	1000 M	2000 M	2500 M	4000 M
WGS	PE150	120Gb/sample	1	2	4	5	8
Targeted panel (FFPE)	PE150	5Gb/sample	25	50	100	125	200
Targeted panel (ctDNA)	PE50	10Gb/sample	4	8	16	20	32
RNAseq	PE150	10Gb/sample	12	25	50	62	100
WES	PE150	10Gb/sample	12	25	50	62	100
Single cell	PE150	60Gb/sample	2	4	8	10	16

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

Application summary



Summay reports



30 runs data quality statistic analysis(FCH PE150)								
Indicator	Mean	SD	cv					
Output Reads	2300 M	150 M	10.2%					
Q30	92.3%	3.0%	3.2%					



Application-WGS (Human)

- > Sample type:Human standard library
- ➤ Library:5 WGS PCR-Free libraries
- > Sequencing platform:SURFSeq 5000(SF), Competitor A(CA)
- ➤ Read length:PE150
- ➤ Data analysis:100×effective reads/sample for deep analysis

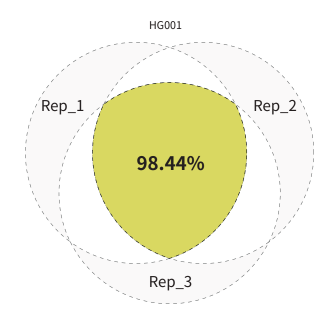
	Sample ID	Raw Q30%	Mapping Ratio	Error Ratio
UC001	SF_1	93.8%	99.9%	0.18%
HG001	CA_1	92.2%	99.6%	0.19%
110000	SF_2	93.9%	99.9%	0.19%
HG002	CA_2	92.8%	99.6%	0.19%
110003	SF_3	93.6%	99.8%	0.17%
HG003	CA_3	92.9%	99.5%	0.19%
115004	SF_4	94.1%	99.6%	0.18%
HG004	CA_4	92.6%	99.2%	0.19%
LICO05	SF_5	93.7%	99.8%	0.18%
HG005	CA_5	92.6%	99.5%	0.20%

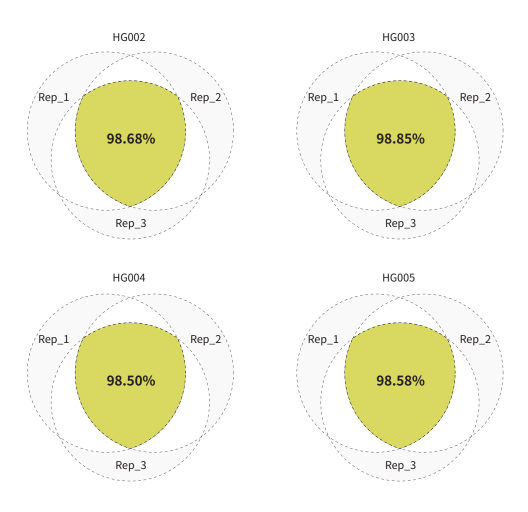
Conclusion:High data quality of WGS samples ensures accurate SNV (>99%) /InDel calling (>95%).



Application-WGS (Human)

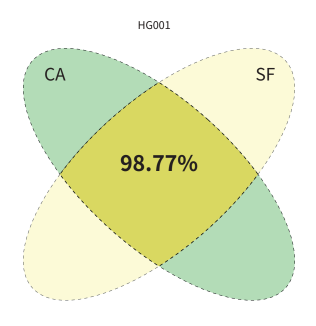
Conclusion:SNV/InDel detection in all samples showing a high consistency (>98%) in different batches.

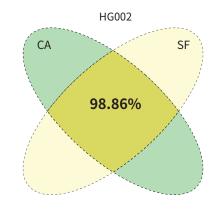


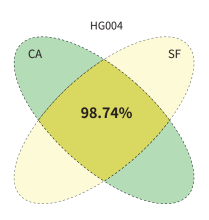


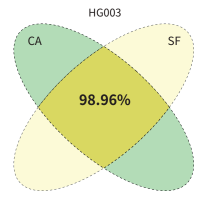
Application-WGS (Human)

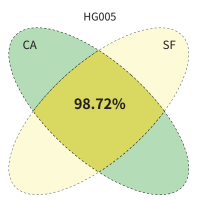
Conclusion: The SNV/InDel result is quite comparable between SF and CA platform (>98%).







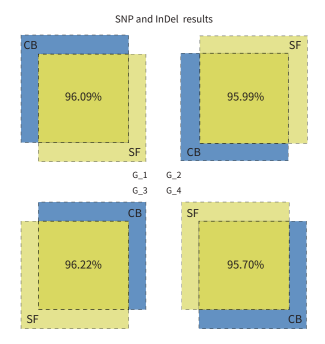




Application-WGS (Animal)

- > Sample type:Four chicken WGS standard libraries
- > Sequencing platform:SURFSeq 5000(SF), Competitor B(CB)
- > Read length:PE150
- ➤ Analysis:26Gb/sample for deep analysis

Sample ID	G_1		G_2		G_3		G_4	
Platform	SF	СВ	SF	СВ	SF	СВ	SF	СВ
Raw Q30	95.9%	91.9%	95.2%	91.0%	95.5%	90.3%	96.1%	90.8%
Mapping Ratio	99.7%	99.6%	99.7%	99.7%	99.7%	99.7%	99.7%	99.7%
Coverage ≥1×	98.9%	99.3%	99.2%	99.2%	99.3%	99.3%	98.9%	98.9%
Mean Coverage	25.6X	25.8X	25.4X	25.8X	25.3X	25.8X	25.6X	25.8X
SNP	6624k	6683k	6802k	6874k	6625k	6703k	6746k	6839k
InDel	871k	881k	890k	905k	878k	893k	892k	908k

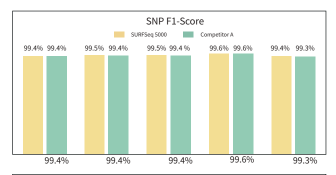


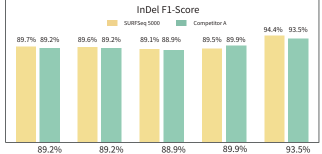
Conclusion:Q30 is higher than Competitor B.As for the mapping rate, coverage, SNP and Indel detection results are quite comparable with competitors.

Application-WES

- ➤ Sample type:5 Human WES libraries
- ➤ Library prep:Agilent SureSelect Human All Exon V6
- > Sequencing Platform:SURFSeq 5000(SF), Competitor A(CA)
- > Read length:PE150
- ➤ Analysis:100×effective reads/sample for deep analysis

Sample ID	HG	001	HG	002	HG	003	HG	004	HG	005
Platform	SF	CA								
Raw Q30	90.9%	94.9%	91.3%	94.9%	90.8%	94.6%	91.4%	94.8%	90.9%	94.7%
Mapping Ratio	99.9%	99.8%	99.9%	99.8%	99.9%	99.8%	99.9%	99.8%	99.9%	99.8%
Coverage ≥10×	99.0%	99.3%	99.2%	99.4%	98.6%	98.5%	99.0%	99.2%	98.2%	98.0%
Coverage ≥30×	93.3%	93.6%	93.1%	92.7%	89.1%	87.2%	93.3%	92.9%	87.0%	85.4%



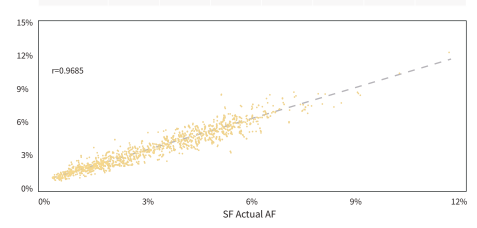


Conclusion:Q30 is a little bit lower than Competitor A.As for mapping rate, coverage, SNP and Indel detection results are quite comparable with competitors.

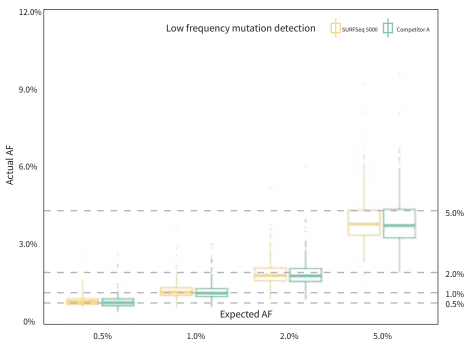
Application-Target commercial panel

- > Sample typr:Four Standard panels (0.5%,1.0%, 2.0%, 5.0%)
- Library prep:iGeneTech TargetSeq Pan-Cancer Panel and Twist cfDNA Pan-Cancer reference standard
- > Sequencing platform:SURFSeq 5000 (SF), Competitor A(CA)
- ➤ Read length:PE150
- ➤ Analysis:20Gb/sample to do deep analysis

Statictics result of variants calling									
Variant_ Recall		SNV_Rec	all (227)		InD	el_Recall (2	115)		
VAF	0.50%	1.00%	2.00%	5.00%	1.00%	2.00%	5.00%		
SF	99.0%	99.6%	99.6%	99.6%	68.6%	88.5%	89.8%		
CA	99.1%	99.6%	99.6%	99.6%	69.8%	86.1%	89.8%		



Conclusion: SURFSeq 5000 showing a quite comparable results with Competitor A in the low frequencing mutation detection.

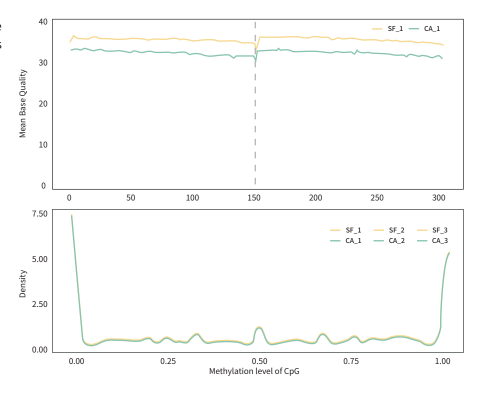


Application-WGBS

- > Samlp type:GIAB HG001
- Library prep:WGBS library
- > Sequencing platform:SURFSeq 5000 (SF), Competitor A(CA)
- > Read length:PE150
- ➤ Data analysis:120Gb/sample for deep analysis

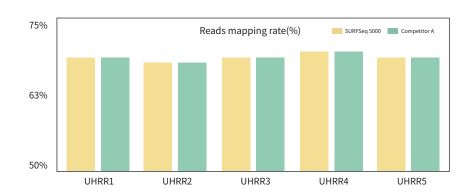
Conclusion: SURFSeq 5000 performs better than CA platform in Q30, mapping rate and duplication rate results. Methylation level analysis, SURFSeq 5000 showes excellent performance on intra-platform repeatability and consistency.

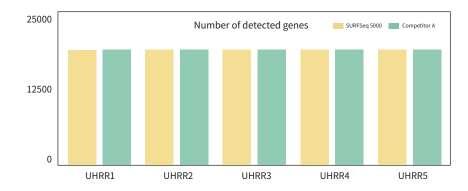
summary peport										
Platform	SF (0	0% Balance Lib	orary)	CA (I	Pooling with W	ES)				
Sample ID	SF_1	SF_2	SF_3	CA_1	CA_2	CA_3				
Raw_Q30	93.81%	94.17%	93.01%	88.23%	91.18%	90.22%				
Mapping Ratio	80.04%	81.53%	81.25%	78.10%	78.88%	75.68%				
Duplication Ratio	15.20%	19.40%	20.00%	21.50%	21.80%	19.60%				



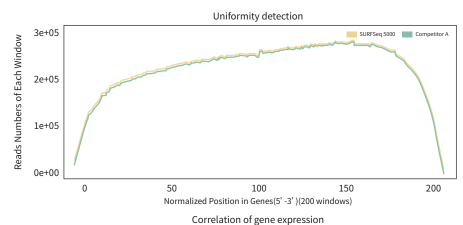
Application-RNAseq

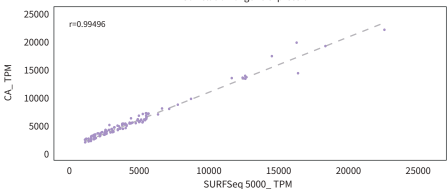
- > Sample type:5 UHRR (Universal Human Reference RNA) standard libraries
- > Sequencing platform:SURFSeq 5000 (SF), Competitor A(CA)
- > Read length:PE150
- > Data analysis:12Gb/sample for deep analysis





Conclusion: Mapping reads, detected gene numbers and the uniformity of transcription coverage are similar in two platforms. The correlation of gene expression reaches to 0.99, showing a high consistency.

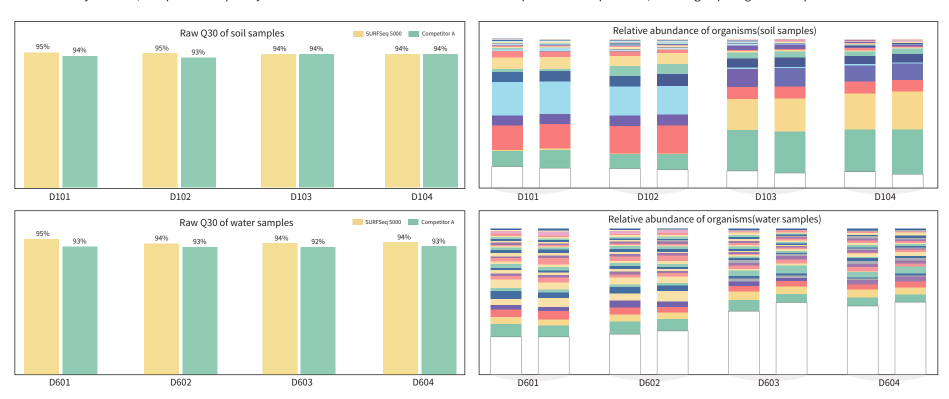




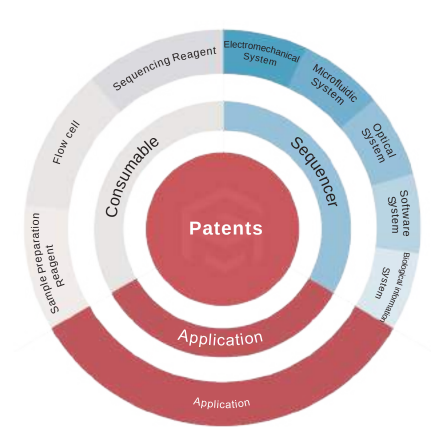
Application-mNGS

- > Sample type:Soil, water
- ➤ Library:8 mNGS libaries
- > Sequencing platform:SURFSeq 5000(SF), Competitor A(CA)
- > Read length: PE150
- > Data analysis:10Gb/sample for deep anaysis

Conclusion: SURFSeq 5000 Q30 and relative organism abdundance tested results are comparable to CA platform, showing a quite good data performance.



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.











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 fluorochron



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

Applications



Product Information

Product Type	Product Nme	Product Code
Sequencer	SURFSeq 5000 Sequencing System Set	SQ00023
	SURFSeq 5000 Sequencing Kit V1.0 (FCM 100cycles)	S000236
	SURFSeq 5000 Sequencing Kit V1.0 (FCM 200cycles)	S000237
	SURFSeq 5000 Sequencing Kit V1.0 (FCM 300cycles)	S000238
Reagent	SURFSeq 5000 Sequencing Kit V1.0 (FCH 50cycles)	S000239
	SURFSeq 5000 Sequencing Kit V1.0 (FCH 100cycles)	S000240
	SURFSeq 5000 Sequencing Kit V1.0 (FCH 200cycles)	S000241
	SURFSeq 5000 Sequencing Kit V1.0 (FCH 300cycles)	S000242

Specification						
Dimensions	1090 mm×690 mm×810 mm					
Weight	240 kg					
Power	2000 VA					
Power requirement	100 -240 V ~, 50/60 Hz					
Operating environment	Temperature: 19 °C-25 °C Humidity: 20%-80% relative humidity (no-condensing) Altitude: below 3000 meter					
Computer	CPU:Intel Xeon Gold 6342 Memory:256 GB Hard Drive 1: 2 TB SSD Hard Drive 2: 17 TB HDD System: Windows10 x64					



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