MGAP: Microbial Genome Analysis Pipeline



Can choose to assemble with or without parameters

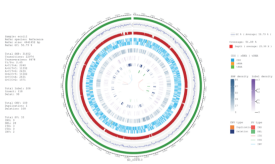
Assembly and Annotation

Assembly summary

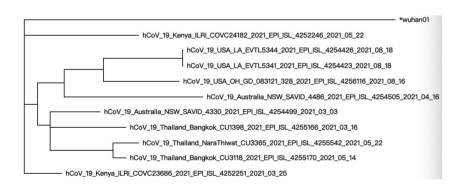
Genome Assemble is the process of putting nucleotide sequence into the correct order, including *De novo* and Reference guides.

ltems	Length(bp)/Count
N90	38197
N75	81471
N50	136226
Minimum	515
Maximum	358330
Average	52436.13
Count	89
Total	4666816

Comparison of assembly results with reference

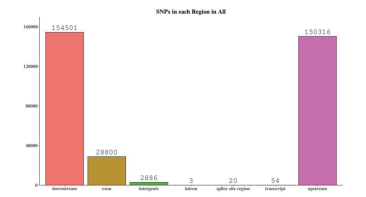


Evolutionary analysis



Species tree analysis based on genome-wide SNPs

Mutation detection



MGI metargetCOVID Software

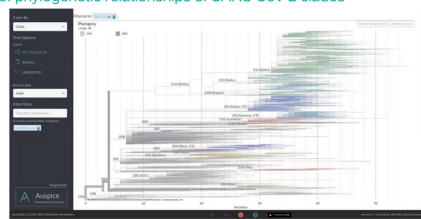
Histograms of Coverage Range



Summary of Result: assemble, variants calling, clade and lineage assign

Assemble Size(bp)	Num Ns	Num SNPs	Num INSs	Num DELs	Clade ID	Lineage ID
29890	75	43	0	3	21A (Delta)	AY.4

▼ Illustration of phylogenetic relationships of SARS CoV 2 clades







900-000399-00

■ Contact Us

Product

MGI Tech Co., Ltd.

+86-4000-688-114

Building 11, Beishan Industrial Zone, Yantian District, Shenzhen, CHINA

Platform of microorganisms Fast Identification and assembly evolution







The copyright of this brochure is solely owned by MGI Tech Co. Ltd., The information included in this brochure or part of, including but not limited to interior design, cover design and icons. is strictly forbidden to be reproduced or transmitted in any form, by any means (e.g. electronic, photocopying, recording, translating or otherwise) without the prior written permission by MGI Tech Co., Ltd.

Version: December 2022 | MGPE2022302



Platform of microorganisms Fast **Identification and Assembly Evolution**

PFI+MGAP Microbial bioinformatics software suite



PFI built-in microbial database



Load identification and assembly bioinformatics process



Equipped with laboratory information management system



External localized server

Platform of microorganisms Fast Identification and Assembly Evolution

PFI+MGAP Microorganisms analysis platform suite

MGI's Platform of microorganisms Fast Identification and assembly evolution is a one-stop platform developed specifically to simplify the analysis of Microbial MPS genetic data. The system includes three types of bioinformatics analysis software: Microorganisms Fast Identification Software (PFI), Microbial Genome Analysis Pipeline (MGAP) and metarget-COVID software. The system is also equipped with a professional laboratory information management system ZLIMS, providing a graphical operation interface, ready to use after booting.



Equipped with laboratory information management system: ZLIMS

Graphical interface, easy to use.

Full-process information management from sample entry, library preparation, sequencing, data analysis to the generation of bioinformatics reports.

External localization server

CPU	Intel Xeon Gold 6240*2
Memory	192GB DDR4
Solid state drive	2TB+250GB
Mechanical hard disk	30TB 7200 SATA 3.5 inch

PFI

Able to process metatranscriptomic data analysis, metagenomic data processing. Analyze and identify microbial nucleotide sequence in the original sample, and automatically generate identification results which provides a reference for accurate detection in the field of microorganisms.

MGAP

Contains multiplex analysis modules for pathogenic microorganisms sequencing data of bacteria, fungi, and virus, realizing from the sequencing data to genome assembly, system evolution analysis, and automatic calculation of mutation. The test results are for research reference, and customers can flexibly choose analysis modules according to actual needs.

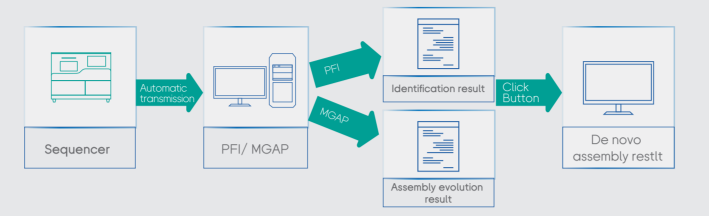
metargetCOVID

MGI metargetCOVID software can applicable for NGS data analysis of metagenomics sequencing or MGI ATOPlex platform, It's functions include variants detection, variants annotation and functional effect prediction, and assign sequence to the clade by Nextclade, implement the dynamic nomenclature SARS CoV 2 lineage by Pangolin.

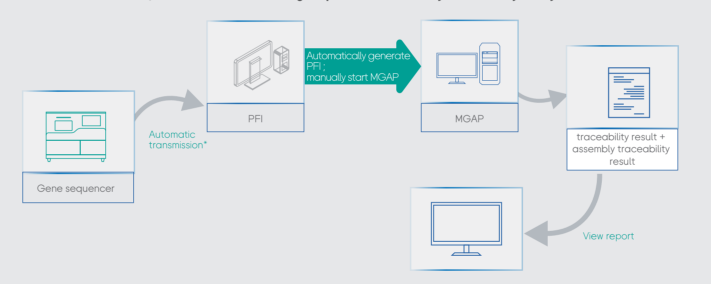
✓ PFI and MGAP operate separately

1.PFI: PFI is used for metagenomic species identification, and use the interest species FASTQ file to de novo assembly

2.MGAP: MGAP is used for assembly and traceability analysis of known pathogen species



☑ PFI for identification, and then select the target species for assembly traceability analysis



MGI metargetCOVID software is an bioinformatics workflow based on Linux operating system, which includes basic data QC, short reads alignment, variants detection, assign the clade and lineage.

One click to complete the virus sequence identification, assembly and Phylogeny analysis

Data QC	Sequence Alignment	Data Processing
Raw dataFiltered data	SPIKE-IN CONTROL sequence, Human GAPDH Gene as internal reference, and the target virus sequence (SARS-CoV-2 reference: MN908947.3)	 Virus identification Virus sequence assembly Mutation detection Variants detection Variants annotation Functional effect prediction Phylogeny analysis

PFI Built-in Species Database (

Host database

Includes more than 10 host species, such as humans, pigs, goats, sheep, mice, etc. In addition, the pipeline includes new host species automatically adding function ,customers can add more hosts as needed.

Host database		es more than species
2 2 2		(
01215151	8	More host species
@ \$ \$		can be customized

Pathogen database

Contains more than 20,000 species of microorganisms, including nCOV-19 sequences, and supports regular updates.

Pathogen database					
Classification of	€		\$	D.	
microorganisms	Viruses	Fungi	Bacteria	Archaea	Parasites
Species	8900+	9000+	8000+	300+	600+
Subspecies	3000+	300+	19000+	200+	50+

^{*}This function needs to be equipped with a localized high-throughput sequencer. It is recommended to use the DNBSEQ sequencing platform manufactured by MGI.



	•			
	Data QC	Remove poor quality reads included adapters and N bases.		
Data processing	Remove host	Remove host DNA and rRNA*		
		1. Database comparison, classification		
		2.Prediction of resistance gene		
	Species identification	3.Prediction of virulence gene		
		4.Activity analysis*		
		Tips: indicates that microorganism activity analysis is performed for an analyzed sample only when the sample contains both DNA and RNA data.		
	Denovo assembly	Manually start de novo assembly for target species		
	DNA/RNA classification, species			
Identification report	reads QC Chart	Starphiococcus aureus Entercoccus faculas Starbonius centrals Starbonius centrals Literalistandoslita ferrentum Bacha audela Piacutoroccus aureus Cryptococcus modernares Succharonyous centralista O 5 10 15 15 15 16 16 16 16 16 16 16 16 16 16 16 16 16		
Assembly report	3. Segments Contributing There is man account a manager from the first one of a manager from the manager f			
	De novo assembly			