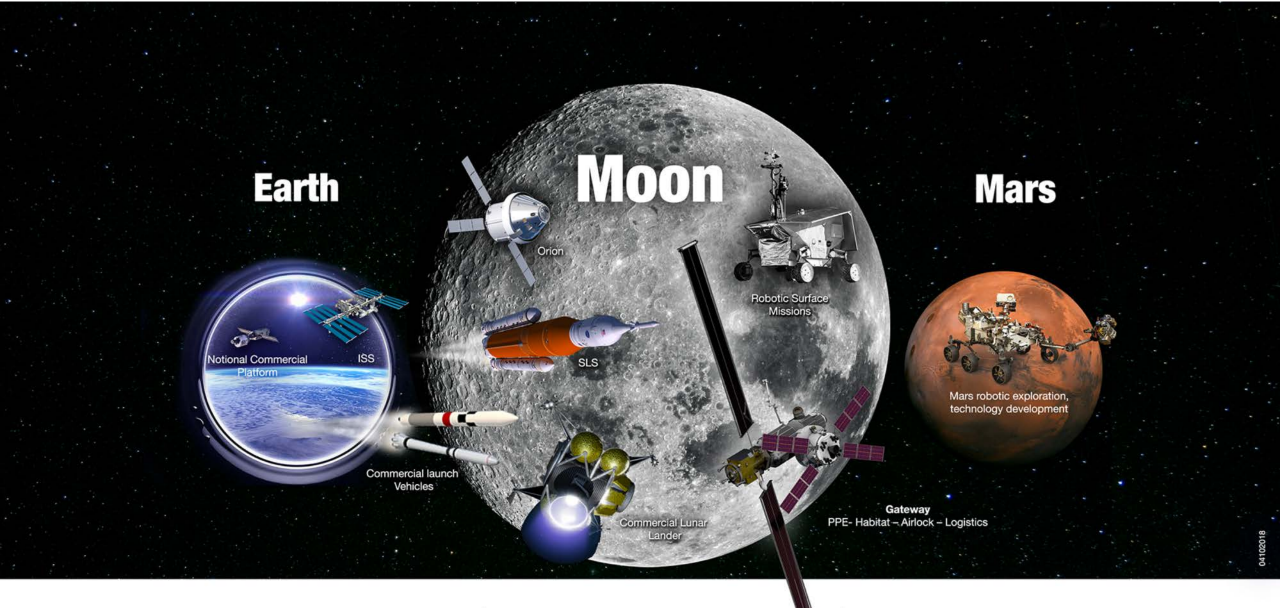
### Using the NASA GeneLab data system to study the metagenomes of spaceships and their occupants

genelab.nasa.gov

Jonathan Galazka, PhD GeneLab Project Scientist NASA Ames Research Center



In LEO Commercial & International partnerships In Cislunar Space A return to the moon for long-term exploration On Mars Research to inform future crewed missions

Image credits: NASA







GATEWAY

EXPLORE MOON to MARS



EXPLORE MOON to MARS

## MARS

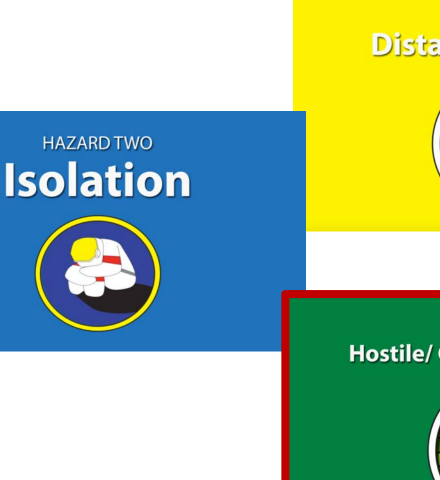
EXPLORE MOON to MARS

### Challenges of Spaceflight

# HAZARD ONE Space Radiation

### HAZARD FOUR Gravity Fields





### HAZARD THREE Distance from Earth



HAZARD FIVE Hostile/ Closed Environments



### Risk of Adverse Health Effects Due to Host-Microorganism Interactions

#### 😴 💈 Risk of Adverse Health Effects Due to Host-Microorganism Interactions

Last Published: 07/31/19 10:05:29 AM (Central)

Short Title: Microhost Element: Human Health Countermeasures (HHC) Evidence: Report Risk Master Logic Diagram: Diagram Point of Contact: Cherie Oubre

DRM Categories	Mission Duration	Operations			Long-Term Health		
		LxC	Risk Disposition *	LxC	Risk Disposition *		
Low Earth Orbit	6 months	1x3	Accepted with Monitoring	3x1	Accepted		
	1 year	1x3	Accepted with Monitoring	3x1	Accepted		
Deep Space Sortie	1 month	1x3	Accepted with Monitoring	3x1	Accepted		
Lunar Visit/ Habitation	1 year	1x3	Accepted with Monitoring	3x1	Accepted		
Deep Space Journey/Habitation	1 year	1x3	Accepted with Monitoring	3x1	Accepted		
Planetary	3 years	3x3	<b>Requires Mitigation</b>	3x3	<b>Requires Mitigation</b>		

#### Risk Ratings and Dispositions per Design Reference Mission (DRM) Category

Note: LxC is the likelihood and consequence rating.

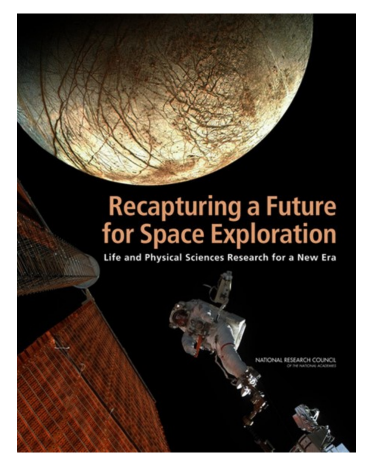
#### Risk Statement

Given that evidence collected during space flight indicates alterations in microbial virulence and astronaut immune function, there is a possibility that infectious disease will be enhanced during spaceflight missions.

#### Gaps (5)

- ➡ MICRO-01: We need to determine the efficacy of current countermeasures and the need for countermeasure development based on changes in microbial populations and characteristics. (Previous title: AEH 10)
- ➡ MICRO-02: We need to determine if spaceflight induces changes in diversity, concentration, and/or characteristics of medically significant microorganisms associated with the crew and environment aboard the International Space Station that could affect crew health. (Previous title: AEH 12)
- ➡ MICRO-03: We need to determine which medically significant microorganisms display changes in the dose-response profiles in response to the spaceflight environment that could affect crew health. (Previous title: AEH 13)
- ➡ MICRO-04: We need to determine how physical stimuli specific to the spaceflight environment, such as microgravity, induce unique changes in the dose-response profiles of expected medically significant microorganisms. (Previous title: AEH 14)
- ➡ MICRO-05: Current microbial standards identifying microbial risk limits need to be updated and microbial requirements need to be developed to include new technologies and future mission scenarios. (Previous title: AEH 15)

### 2011 NRC Decadal Survey: "Microbial observatory" in space



National Research Council. 2011. *Recapturing a Future for Space Exploration: Life and Physical Sciences Research for a New Era*. Washington, DC: The National Academies Press. https://doi.org/10.17226/13048.

"Species that are uncommon, or that have significantly increased or decreased in number, can be studied in a "microbial observatory" on the ISS, in groundbased facilities, or both."

"If these studies suggest that permanent changes have occurred within the species, approaches such as microarray analysis and whole-genome sequencing can be used to determine what modifications or mutations may have occurred to **shift the microbial population dynamics**."

"The continuing decline in the cost and increase in speed of genomic analysis should facilitate the comprehensive study of any changes in these microbial populations in space. Wide dissemination of this rich collection of raw data within the scientific community will allow a variety of scientific investigations."

### Microbial Tracking 1

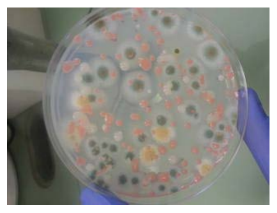
Microbial Tracking-1A (launched January 10, 2015) Microbial Tracking-1B (launched April 14, 2015) Microbial Tracking-1C (launched April 8, 2016)



Using adhesive tape as a sampling device, Venkateswaran and Karouia demonstrate one of the methods that station crew used to collect microorganisms from surfaces for the Microbial Tracking-1 investigation. Credit NASA.



One sampling location for the Microbial Observatory-1 investigation. Credit NASA.



A petri dish contains colonies of fungi grown from a sample collected aboard the International Space Station during the first of the three Microbial Tracking-1 flights. Credit NASA.

Checinska Sielaff A, Singh NK, Allen JE, Thissen J, Jaing C, Venkateswaran KJ. Draft genome sequences of biosafety level 2 opportunistic pathogens isolated from the environmental surfaces of the International Space Station. Genome Announcements. 2016 December 29; 4(6): e01263-16. DOI: 10.1128/genomeA.01263-16. PMID: 28034853. DOI: 10.1128/genomeA.01263-16

Knox BP, Blachowicz A, Palmer JM, Romsdahl J, Huttenlocher A, Wang CC, Keller NP, Venkateswaran KJ. Characterization of Aspergillus fumigatus isolates from air and surfaces of the International Space Station. mSphere. 2016 September-October; 1(5): e00227-16. DOI: 10.1128/mSphere.00227-16.PMID: 27830189. DOI: 10.1128/mSphere.00227-16

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Urbaniak C, van Dam P, Zaborin A, Zaborina O, Gilbert JA, Torok T, Wang CC, Venkateswaran KJ. Genomic Characterization and Virulence Potential of Two Fusarium oxysporum Isolates Cultured from the International Space Station. mSystems. 2019 March/April; 4(2): DOI: 10.1128/mSystems.00345-18. DOI: 10.1128/mSystems.00345-18

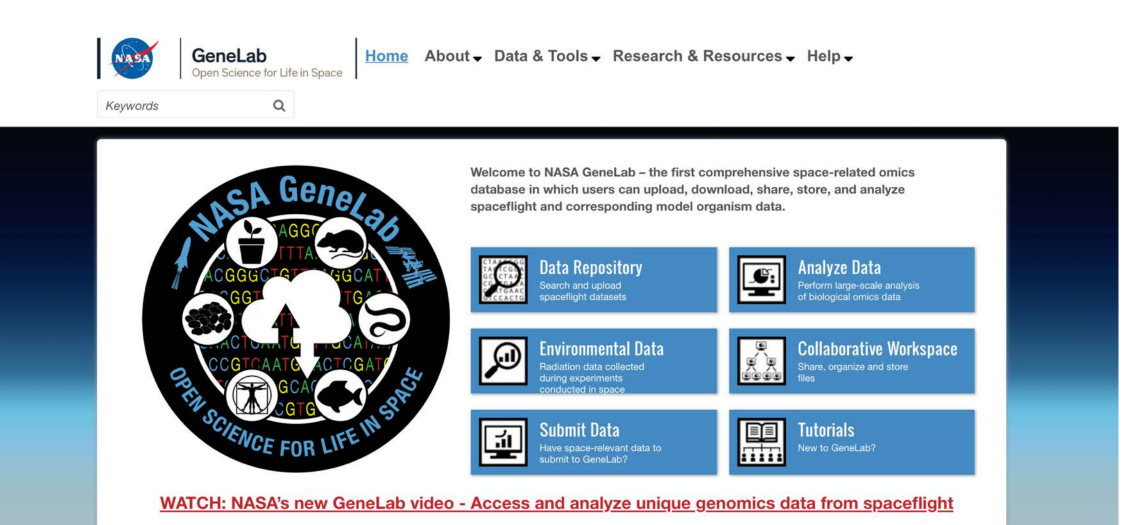
Xiao S, Venkateswaran KJ, Jiang SC. The risk of Staphylococcus skin infection during space travel and mitigation strategies. Microbial Risk Analysis. 2019 April; 1123-30. DOI: 10.1016/j.mran.2018.08.001. DOI: 10.1016/j.mran.2018.08.001

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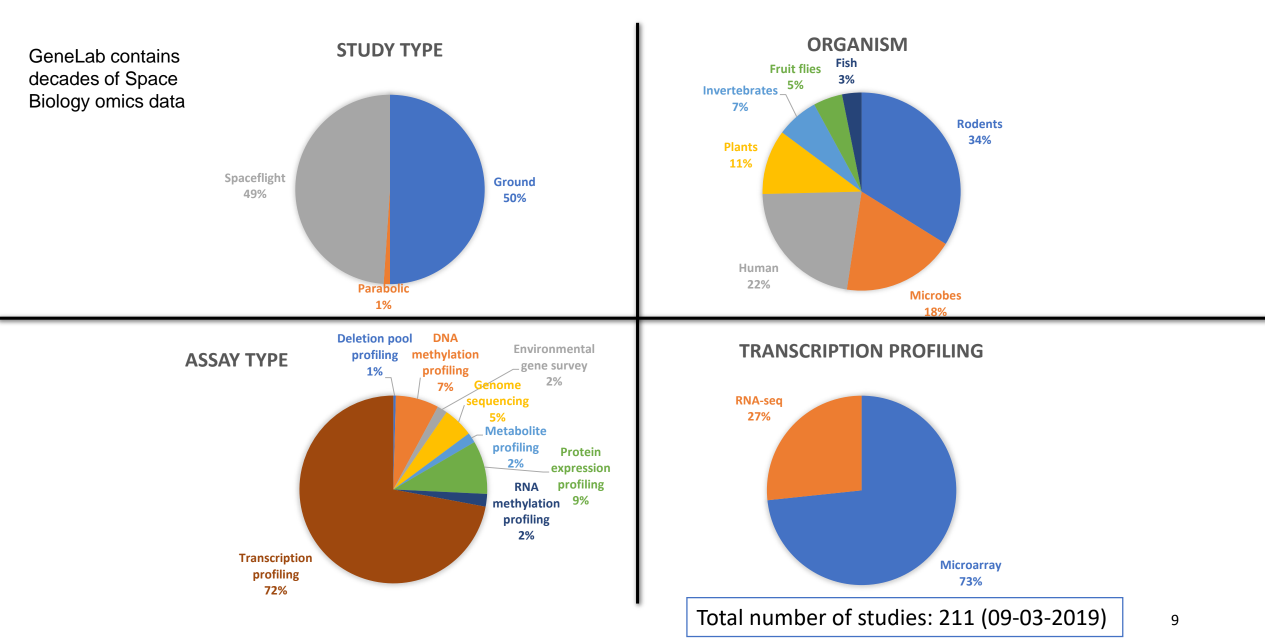
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### NASA GeneLab data system

- Mission: To enable scientific discovery and space exploration through multi-omics data-driven research.
- Currently funded by SLPSRA Space Biology program. Previously received funding from the ISS program.

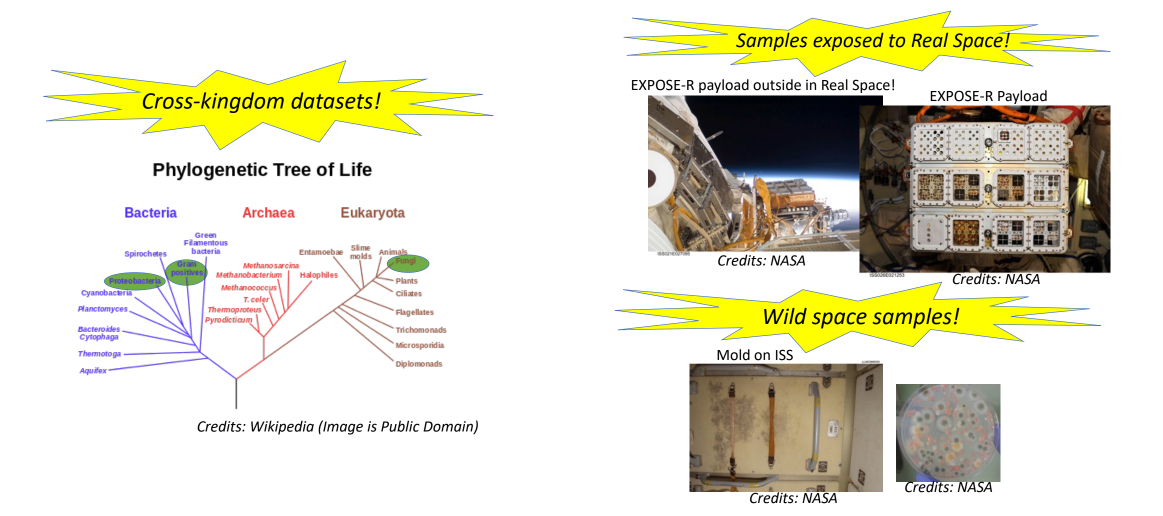


### GeneLab overview: Database content



### Microbe datasets on GeneLab

- Understanding how microbes respond the spaceflight is critical to: Astronaut health and safety, Space biotechnology, Planetary protection, Astrobiology
- 31 studies: 17 transcription profiling, 7 genome sequencing, 2 proteomic, 6 microbiome profiling



### Microbiome profiling datasets on GeneLab

GLDS	Title	Data type	Study type
GLDS-26	Microbiomes of the Dust Particles Collected from the International Space Station and Spacecraft Assembly Facilities	Amplicon	Spaceflight (ISS)
GLDS-65	Microbial Observatory (ISS-MO): Microbial diversity	Amplicon	Spaceflight (ISS)
GLDS-66	Microbial Observatory (ISS-MO): Antimicrobial resistance genes	Antibiotic resistance	Spaceflight (ISS)
GLDS-69	Microbial Observatory (ISS-MO): Microbial metagenomics	Whole Genome	Spaceflight (ISS)
GLDS-72	Space environmental factor impacts upon murine colon microbiota and mucosal homeostasis	Amplicon	Spaceflight (ISS)
GLDS-82	Microbial monitoring in the ISS-Kibo	Amplicon	Spaceflight (ISS)
GLDS-126	Analysis of dust samples from the Russian part of the ISS	Amplicon	Spaceflight (ISS dust)
GLDS-146	Mouse fecal microbiome after exposure to high LET radiation	Amplicon	Ground analog (Irradiated mice)
GLDS-191	Temporal dynamics of the gut microbiota in people sharing a confined environment, a 520-day ground-based space simulation.	Amplicon	Ground analog (Humans on space mission)
GLDS-170	NASA Aircraft Bioaerosol Collector (ABC)	Amplicon	Ground analog (stratosphere)
GLDS-200	Quantitative evaluation of bioaerosols in different particle size fractions collected on the International Space Station (ISS)	Amplicon	Spaceflight (ISS)
GLDS-212	Reproducible changes in gut microbiome reveal a shift in microbial and host metabolism during spaceflight (Peng Jiang)	Amplicon	Spaceflight (Mouse)

#### microbial observatory

🛛 All 🕑 GeneLab 🔄 NIH GEO 🔄 EBI PRIDE 🔄 ANL MG-RAST

#### Search Filters (GeneLab Only)

Project Type	$\sim$	Factors	$\sim$	Organisms	$\sim$	Assay Type	$\sim$	Clear
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Search results for: microbial observatory using filter(s):

Sort by Relevance \$ 25 \$

Total Search Results Found: 7

Getting the data

#### Microbial Observatory (ISS-MO): Study of BSL-2 bacterial isolates from the International Space Station https://genelab-data.ndc.nasa.gov/genelab/accession/GLDS-67

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In an on-going Microbial Observatory experimental investigation on the International Space Station (ISS) multiple bacterial isolates of Biosafety Level 2 (BSL-2) were isolated and identified. The antibiotic susceptibility pattern was tested in these BSL-2 isolates for the following antibiotics: cefazolin ciprofloxacin cefoxitin erythromycin gentamycin oxacillin penicillin rifampin tobramycin and many of the BSL-2 isolates showed multiple drug resistance. Among these isolates 21 strains were chos...

Organism: Klebsiella, Staphylococc... Factor: Microgravity Assay Type: genome sequencing Accession: GLDS-67 Pl/Contact: Kasthuri Venkateswaran Release/Publication Date: 07-Jul-2016

### Microbial Observatory (ISS-MO): Draft Genome Sequence of two Aspergillus fumigatus Strains Isolated from the International Space Station

#### https://genelab-data.ndc.nasa.gov/genelab/accession/GLDS-68



Aspergillus funigatus is a saprophytic filamentous fungus that is ubiquitous outdoors (soil decaying vegetation) and indoors (hospitals simulated closed habitats etc.). A. funigatus can adapt to various environmental conditions and form airborne conidia that are the inoculum for a variety of diseases (e.g. non- and invasive pulmonary infections allergic bronchopulmonary aspergillosis etc.) in immunocompromised hosts. In an on-going Microbial Observatory Experiments on the International Space Sta...

Organism: Aspergillus fumigatus Factor: spaceflight Assay Type: genome sequencing Accession: GLDS-68 Pl/Contact: Kasthuri Venkateswaran Release/Publication Date: 08-Jul-2016

#### Microbial Observatory (ISS-MO): Antimicrobial resistance genes

https://genelab-data.ndc.nasa.gov/genelab/accession/GLDS-66



The environmental samples were collected with the polyester wipes from eight different locations in the International Space Station (ISS) during two consecutive sampling sessions (three months apart) within the ISS Microbial Observatory Experiment. DNA extracted from each of the samples was used to create amplicon libraries based on customized panel of 500 antimicrobial resistance genes followed by next-generation sequencing. This is the first study of that shows the reservoir of antimicrobial g...

Organism: cellular organisms Factor: spaceflight Assay Type: environmental gene survey Accession: GLDS-66 Pi/Contact: Kasthuri Venkateswaran Release/Publication Date: 07-Jul-2016

#### Microbial Observatory (ISS-MO): Microbial diversity

https://genelab-data.ndc.nasa.gov/genelab/accession/GLDS-65



The environmental microbiome study was designed to decipher microbial diversity of the International Space Station surfaces in terms of spatial and temporal distributions using 16S and ITS iTag Illumina sequencing. We hypothesized that the microbial population of environmental surfaces changes in time due to astronauts xe2 x80 x99 activity and might be location specific. The environmental samples were collected with the polyester wipes from eight different locations in the ISS during two consecu...

Organism: cellular organisms Factor: spaceflight Assay Type: environmental gene survey Accession: GLDS-65 Pl/Contact: Kasthuri Venkateswaran Release/Publication Date: 07-Jul-2016

### Microbial Observatory (ISS-MO): Molecular characterization of Bacillus issensis sp. nov. isolated from various quarters of the International Space Station

https://genelab-data.ndc.nasa.gov/genelab/accession/GLDS-64



As part of an ongoing effort to catalogue microbial communities inhabiting the International Space Station (ISS) crew-associated environmental samples were collected from the Japanese Kibo Russian and US research modules. Initial analysis based on 16S rRNA gene sequencing identified 11 Bacillus isolates (two from Kibo Japanese Experiment Module (JEM) four from US Segment Harmony Node 2 and five from Russian Segment Zvezda module sites) all belonging to the Bacillus anthracis-B. cereus-B. thuring...

Organism: , Bacillus Factor: Spaceflight Assay Type: genome sequencing Accession: GLDS-64 PI/Contact: Kasthuri Venkateswaran Release/Publication Date: 08-Jul-2016

#### Microbial Observatory (ISS-MO): Indoor microbiome study of the International Space Station surfaces https://genelab-data.ndc.nasa.gov/genelab/accession/GLDS-69



Presented here is the environmental microbiome study of the International Space Station surfaces. The environmental samples were collected with the polyester wipes from eight different locations in the ISS during two consecutive sampling sessions (three months apart). The specific objective was to unveil the pool of genes for each location during two separate sessions to learn of functional and metabolic diversity of microorganisms in the ISS. The International Space Station (ISS) as a closed bu...

Organism: cellular organisms Factor: Spaceflight Assay Type: metagenome profiling Accession: GLDS-69 Pt/Contact: Kasthuri Venkateswaran Release/Publication Date: 06-Jul-2016

DESCRIPTION	PROTOCOLS	SAMPLES	ASSAYS/MEASUREMENTS	PUBLICATIONS	STUDY FILES	
	DESCRIPTION					
Study Description	locations in the ISS during two consecut of functional and metabolic diversity of n microgravity, radiation, and limited humu were used for sampling of eight various 1MM or Dragon capsule from SpaceX). A were processed to measure microbial to using Adenosine Triphosphate (ATP) as USE (The INA) and the International CFU/In2). The PMA-gPCR analysis sho polyester wipes cover much larger samp cultivable, total, and vable incrobial div	ive sampling sessions (three month- incorognainsm in the ISS. The Intern in presence. The mit bial diversity suffare locations on the ISS at differ difference of the ISS at difference of the order (R2A, Blood Agar, and Potato arguing and and attack and and and arguing and and and and and and say. and quantifiative thymerase ch sweat characteristic and and and sweat characteristic and and and say and and and and and and say and and and and and and and say and	pace Station surfaces. The environmental sample space Station (SS) as a closed bulk environmental sample sational Space Station (ISS) as a closed bulk environment estimation of the service service station of the growth of the service station of the service service station of the service service station of the growth Date seate A, and the service station of the growth Date seate A, and the service station of the growth Date seate A, and the service station of the service station of the service station of the service station Date seate A, and the service station of the service station Date seate A, and the service station of the service station Date seate and the power station of the service station Date setting of the service station of the service station of the at molecular techniques. The implemental health, the ISS maintenance and the general kore	ool of genes for each location during two investigated during this study. Polyester a lanalyzed inmodately upon the return to of bacteria and fungi were incubated at well as fungi. Subsequently, valide micr azade (PAA) treatment. The 15G-tag an optiestical and the study of the study of the optiestical and the study of the study optiestical and a sampled. Un of the ISS covering both cultivable and n on of the PAA assay before DAA extran	» separate sessions to obiome which is shap wipes and contact slid o the Earth (via Soyu 25C. The polyester wi obial burden was assist d metagenome analys contact slides. (102 te tilke contact slides, n-cultivable species. tion allowed distinguit	
Contacts	NAME	ROLE	ORGANIZATION	E-MAIL	E-MAIL	
	Kasthuri Venkateswaran	Princip. Investigate	or NASA	kjvenkat@jpl.nasa.go	rv .	
Study Design Factor(s)	FACTOR	ONTOL	DGY: CONCEPT			
	Spaceflight	Space F	light			
Organisms	collular organisms					
Assay(s)	ASSAY TYPE		TYPE	DEVICE PLATFORM		

GLDS-69: Microbial Observatory (ISS-MO): Indoor microbiome study of the International Space Station surfaces Version 1



915.39 MB



a Version: 1

Select a Version:

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DESCRIPTION PROTOCOL S ASSAYS/MEASUREMENTS PUBLICATIONS SAMPI ES STUDY FILES To view files, click on the folder of interest. 0 files selected All Files Study Metadata Files Download Selected Files Whole Genome Sequencing Data FILE SIZE 4 **RESOURCE CATEGORY RESOURCE DESCRIPTION** FILES GLDS-69\_wgs\_5816\_IIF1SW-P-M\_F2\_L1\_PMA\_FLT\_R2-Raw sequencing data and sequencing assembly files and reports 0 979.56 MB Whole Genome Sequencing Data clean.fastq.gz associated with this study. GLDS-69 wgs 5816 IIF1SW-P-M F2 L1 PMA FLT R1-Raw sequencing data and sequencing assembly files and reports 976 56 MB Whole Genome Sequencing Data clean.fastq.gz associated with this study. GLDS-69 wgs SRX3808529 IIIF7SW-P-M\_F3\_L7\_PMA\_FLT\_R2-Raw sequencing data and sequencing assembly files and reports 0 951.87 MB Whole Genome Sequencing Data clean.fastq.gz associated with this study. GLDS-69\_wgs\_5647\_IF8SW-P-M\_F1\_L8\_PMA\_FLT\_R1-Raw sequencing data and sequencing assembly files and reports 0 939.76 MB Whole Genome Sequencing Data clean.fastq.gz associated with this study. GLDS-69\_wgs\_SRX3808529\_IIIF7SW-P-M\_F3\_L7\_PMA\_FLT\_R1-Raw sequencing data and sequencing assembly files and reports 933.78 MB Whole Genome Sequencing Data clean.fasto.oz associated with this study.

Whole Genome Sequencing Data

Submitted Date: 03-May-2016

Raw sequencing data and sequencing assembly files and reports

associated with this study.

Release Date: 07-Jul-2016

amples were collected with the The specific objective was to unveil rganisms in the ISS. The GLDS-69\_wgs\_5629\_IFISW-M\_F1\_L1\_NORM\_FLT\_R2clean.fastq.gz

### How can YOU download and analyze this data?

Anyone:

- 1. Go to genelab.nasa.gov!
- 2. Download and begin analyzing!

Those with technical skills and an interest in collaborating with likeminded scientists:

1. Go to genelab.nasa.gov!

3. Join a NASA GeneLab Analysis Working Group (<u>https://genelab.nasa.gov/awg/join</u>)!

### Thank you for listening! Go to: genelab.nasa.gov



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Welcome to NASA GeneLab – the first comprehensive space-related omics database in which users can upload, download, share, store, and analyze spaceflight and corresponding model organism data.



WATCH: NASA's new GeneLab video - Access and analyze unique genomics data from spaceflight