





Draft Genome Sequences of Several Fungal Strains Selected for Exposure to Microgravity at the International Space Station

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ABSTRACT The whole-genome sequences of eight fungal strains that were selected for exposure to microgravity at the International Space Station are presented here. These baseline sequences will help to understand the observed production of novel bioactive compounds.

n a screening project of natural products, fungal strains isolated from environments associated with the Chernobyl nuclear power plant (ChNPP) accident (1) have been investigated. The radiation-tolerant microorganisms selected for exposure to microgravity at the International Space Station were known to produce valuable natural products; their genomic sequences coded for secondary metabolism pathways; or they displayed positive radiotropism.

Aspergillus niger, an industrially important filamentous fungus, contains a sequence resembling the fumonisin gene cluster, which suggests that the fungus has the genetic potential to produce carcinogenic fumonisins (2). A. niger also produces an abundance of naphtho-gamma-pyrone secondary metabolites, which have been shown to have antibacterial, antifungal (3), antitumor (4), and cytotoxic (3, 4) activity.

Aspergillus terreus is used to produce organic acids, such as itaconic acid (5), or enzymes, such as xylanases (6, 7). One of the most important secondary metabolites made by *A. terreus* is the cholesterol-lowering molecule lovastatin. Discovery of this potent compound revolutionized the treatment of hypercholesterolemia (8, 9).

Aureobasidium pullulans is an important producer of pullulan, a homopolysaccharide of glucose that is widely used in the food, pharmaceutical, and electronics industries (10, 11). The whole-genome sequence of *A. pullulans* revealed significant biotechnological potential but also the presence of virulence factors that cannot be overlooked (12)

Beauveria bassiana is an entomopathogenic fungus used to produce biodegradable, nonpoisonous, and cost-efficient bioinsecticides (13). Genomic analysis of *B. bassiana* exhibits its capacity to produce a plethora of secondary metabolites, such as oosporein, bassianin, beauvericin, or oxalic acid (14). Beauvericin possesses antimicrobial, antiviral, antifungal, and antitumor activity (15).

Cladosporium cladosporioides, a ubiquitous organism, produces cladosporin and isocladosporin—secondary metabolites known to have antifungal activities. Its genome has not been sequenced yet (16, 17).

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Singh et al.

TABLE 1 Statistical summary for the eight draft fungal genome sequences

	Strain			No. of	Genome		G+C content
Strain	designation	NCBI accession no.	Isolation location ^a	contigs	size (bp)	N ₅₀ (bp)	(%)
Aspergillus niger	JSC-093350089	MSJD00000000	ISS environmental surface	223	36,080,355	543,773	49.46
Aspergillus terreus	IMV 01167	MSJE00000000	Soil, Kirovograd region	331	31,580,414	482,632	52.24
Aureobasidium pullulans	IMV 00882	MSJF00000000	Wall surface, unit 4, ChNPP	879	40,984,331	98,085	51.01
Beauveria bassiana	IMV 00265	MSJG00000000	Wall surface, unit 4, ChNPP	735	35,190,057	138,299	51.72
Cladosporium cladosporioides	IMV 00236	MSJH00000000	Wall surface, unit 4, ChNPP	843	47,573,060	186,555	42.89
Cladosporium sphaerospermum	IMV 00045	MSJI00000000	Wall surface, unit 4, ChNPP	959	50,156,125	72,128	53.05
Fusarium solani	IMV 00293	MSJJ00000000	Wall surface, unit 4, ChNPP	876	51,318,644	812,484	47.58
Trichoderma virens	IMV 00454	MSJK00000000	Soil, 10-km ChEZ	197	42,025,033	1,319,489	48.44

International Space Station; ChNPP, Chernobyl nuclear power plant; ChEZ, Chernobyl exclusion zone.

Cladosporium sphaerospermum is a plant endophyte but also an allergen to immunocompromised populations. It has the capacity to produce a variety of allergens, such as enolase, mannitol, dehydrogenase, and aldehyde dehydrogenase (18).

Fusarium solani is a plant pathogen that produces multiple phytotoxins, such as marticin, isomarticin, anhydrofusarubin, and javanicin, that cause sudden death syndrome of soybean (19), for example.

Trichoderma virens is a common rhizosphere fungus beneficial to plants and reported to induce a defense response of cotton to *Rhizoctonia solani*–incited seedling disease (20, 21).

The whole-genome sequences of these eight fungal strains were obtained by shotgun sequencing performed on an Illumina HiSeq2500 platform with a paired-end module. The NGS QC toolkit version 2.3 (22) was used to filter the data for high-quality vector- and adaptor-free reads for genome assembly (cutoff read length for high quality: 80%; cutoff quality score: 20). High-quality vector-filtered reads were used for assembly with the MaSuRCA genome assembler (k-mer size = 70) (23). Data from the final assembly of the strains, including number of scaffolds, total size, N_{50} contig length, G+C content, and GenBank accession numbers, are given in Table 1.

Accession number(s). This whole-genome shotgun project has been deposited in DDBJ/ENA/GenBank under the accession numbers given in Table 1. The versions described in this paper are the second versions.

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Volume 5 Issue 15 e01602-16 genomea.asm.org **2**

Genome Announcement

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Volume 5 Issue 15 e01602-16 genomea.asm.org **3**