Genome sequence of *Stachybotrys chartarum* Strain 51-11

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Abstract: *Stachybotrys chartarum* strain 51-11 genome was sequenced by shotgun sequencing utilizing Illumina Hiseq 2000 and PacBio long read technology. Since *Stachybotrys chartarum* has been implicated in health impacts within water-damaged buildings, any information extracted from the genomic sequence data relating to toxin and allergen production or the fungi’s metabolism could be potentially useful.

*Stachybotrys chartarum* is a saprophytic fungus with world-wide distribution. In nature, it is isolated from soil, seeds and decaying organic matter. Indoors, *Stachybotrys chartarum* grows on wet cellulose-containing building material such as gypsum wallboard, cardboard and ceiling tiles [1].

*S. chartarum* is among the indoor fungal population frequently identified in water-damaged buildings and it has been linked to damp building-related illnesses (DBRIs) [2]. This fungus is capable of producing mycotoxins of which the macrocyclic trichotheccenes are among the most toxic [3, 4]. Likewise, several studies have shown the possible association of *S. chartarum* proteins, e.g. proteinases and hemolysins, with inflammation of the respiratory system [5, 6]. Recently, Semeiks et al reported the genomes of four *S. chartarum* strains in an effort to determine toxin chemotypes within the toxigenic strains [7].

*S. chartarum* toxigenic strain 51-11, an environmental isolate obtained from the cluster of idiopathic pulmonary hemorrhage cases (Cleveland, Ohio) [8], is part of the RTI International (Research Triangle Park, NC) microbial collection assigned with a unique identification number and stored at -80°C for long term preservation. Propagation and maintenance of this fungus was on potato dextrose agar (PDA) (Becton, Dickinson & Company, Sparks, MD) at 25°C. Colony morphology and microscopic observations were made to confirm the phenotype that is characteristic of this species. A single spore isolate was used for genomic sequencing. The fungal spores were grown in potato dextrose broth (PDB) (Becton, Dickinson & Company, Sparks, MD) and were shaken at 25°C. The resulting fungal mycelia were harvested using sterile mira cloth (EMD Millipore, Darmstadt, Germany) and washed with sterile distilled water. The mycelia were flash frozen in liquid nitrogen and stored at -20°C until the genomic DNA was isolated using a protocol described by Kohler et al [9].

Two genomic DNA libraries for Illumina sequencing: one paired-end library (insert size 350-390bp), and one mate-paired library (insert size 100-5000 bp) were produced. These two
libraries were sequenced on an Illumina Hiseq 2000 (Ilumina, San Diego, CA) at Argonne National Laboratory (Lemont, IL). The machine produced 460x coverage, and approximately 100x of that coverage was actually used for the assembly, though the whole dataset was exploited for error correction. Genome size was assessed using Kmer-counting tools, which pointed to a genome size of approximately 40 Mbases. The resulting 40 Mb error-corrected reads were assembled using MIRA [10] on a large-memory machine. The resulting contigs were hand-edited for quality. SSPACE (Leiden, The Netherlands) [11] basic 2.0 was run on the mate pair data to try to close gaps and produce better scaffolds. The resulting assembly contained 2843 scaffolds. Higher molecular weight DNA for PacBio long-read sequencing was also produced to close gaps in the Illumina assembly. Nine sequencing runs were performed, and after standard Quality Assurance/Quality Control cutoffs were applied, there were 284,625 reads, with ~736 million bases total (~2,600 bases per read), or ~18x coverage. Two hybrid assembly methods were compared, SSPACE-LongRead and PBJelly [12]. Multiple combinations using both programs were performed on a supercomputing cluster, and quality of assembly was assessed using standard metrics (N50, N90). Three iterations of SSPACE-LongRead alone (not in combination with PBJelly) showed the highest quality assembly, with the final number of scaffolds at 1,435, or approximately half of the Illumina assembly alone.

**Nucleotide sequence accession numbers:** submitted to GenBank. (Accession numbers pending)

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**References**


7. Semeiks J, Borek D, Otwinowski Z, Grishin NV: **Comparative genome sequencing reveals chemotype-specific gene clusters in the toxigenic black mold *Stachybotrys.*** *BMC Genomics* 2014, **15**:590-605


10. MIRA: Open source: [http://genome.cshlp.org/content/14/6/1147.full](http://genome.cshlp.org/content/14/6/1147.full)


12. PBJelly: Open source: [http://www.biomedcentral.com/1471-2105/15/180/abstract](http://www.biomedcentral.com/1471-2105/15/180/abstract)