Draft genome sequences of the oomycete

*Pythium insidiosum* strain CBS 573.85
from a horse with pythiosis and strain CR02
from the environment

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**Abstract**

*Pythium insidiosum* is an aquatic oomycete microorganism that causes the fatal infectious disease, pythiosis, in humans and animals. The organism has been successfully isolated from the environment worldwide. Diagnosis and treatment of pythiosis is difficult and challenging. Genome sequences of *P. insidiosum*, isolated from humans, are available and accessible in public databases. To further facilitate biology-, pathogenicity-, and evolution-related genomic and genetic studies of *P. insidiosum*, we report two additional draft genome sequences of the *P. insidiosum* strain CBS 573.85 (35.6 Mb in size; accession number, BCFO00000000.1) isolated from a horse with pythiosis, and strain CR02 (37.7 Mb in size; accession number, BCFR00000000.1) isolated from the environment.

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### Specifications Table

| Subject area | Biology |
|--------------|---------|
| More specific subject area | Microbiology, Genomics |
| Type of data | Genome sequence data |
| How data was acquired | IlluminaHiSeq 2000 and IlluminaHiSeq 2500 Next Generation Sequencing Platforms |
| Data format | Assembled genome sequences |
| Experimental factors | Genomic DNA was extracted from the *Pythium insidiosum* strains CBS 573.85 (an animal isolate) and CR02 (an environmental isolate). |
| Experimental features | Genome of the *P. insidiosum* strains CBS 573.85 and CR02 were sequenced and assembled. |
| Data source location | *Pythium insidiosum* strain CBS 573.85 was isolated from a horse in Costa Rica, and strain CR02 was isolated from the environment in Thailand. |
| Data accessibility | The draft genome sequences of *P. insidiosum* have been deposited in the DNA Data Bank of Japan (DDBJ) under the accession numbers: BCF000000000.1 (strain CBS573.85; https://www.ncbi.nlm.nih.gov/nuccore/BCFO000000000.1) and BCFR000000000.1 (strain CR02; https://www.ncbi.nlm.nih.gov/nuccore/BCFR000000000.1). |

### Value of the data

- Previously, only genome sequence data of *P. insidiosum* isolated from humans is available in the public databases.
- The first draft genome sequences of *P. insidiosum* isolated from a non-human animal with pythiosis and from the environment are now made available.
- The additional genome data will facilitate biology-, pathogenicity-, and evolution-related studies of *P. insidiosum*, through comparative genomic studies of *Pythium* species and related species.

### 1. Data

*Pythium insidiosum* is an aquatic oomycete microorganism that causes the lethal infectious condition, pythiosis, in humans and other animals [1,2]. The organism has been isolated from the environment in Australia, Thailand, Brazil and the United States [3–6]. Genome sequences of *P. insidiosum*, isolated from humans, are available and accessible in public databases [7,8]. We report two additional draft genome sequences of the organism isolated from a horse with pythiosis, as well as from the environment.

### 2. Experimental design, materials and methods

#### 2.1. Genome sequencing and assembly

Genomic DNA samples were extracted from *P. insidiosum* strain CBS573.85 (from an infected horse in Costa Rica) and strain CR02 (from an agricultural area in Thailand), using the conventional extraction method described by Lohnoo and co-workers [9]. rDNA sequence analysis was performed to confirm the identity of the organism [10–12]. The extracted genomic DNA of each of these two strains was subjected to preparation of a paired-end library for genome sequencing, using the IlluminaHiSeq 2500 (strain CBS573.85) or IlluminaHiSeq 2000 (strain CR02) platform (Yourgene Bioscience, Taiwan). Quality trims of the raw reads were executed by CLC Genomics Workbench.
(www.clcbio.com) to yield read lengths with at least 35 bases. The adaptor sequences were removed by Cutadapt 1.8.1 [13] to obtain a total of 34,651,034 reads with an average read length of 122 bases (4,238,414,330 total bases) for the strain CBS573.85, and a total of 27,436,541 reads with an average read length of 105 bases (2,888,290,738 total bases) for the strain CR02. All reads were assembled by Velvet 1.2.10 [14] to the total sequence length of 35,561,321 bases (number of contigs, 11,223; average contig length, 3169; \( N_{50}, 12,261 \); ‘N’ composition, 1.2%) for the strain CBS573.85, and 37,673,126 bases (number of contigs, 22,560; average contig length, 1670; \( N_{50}, 3553 \); ‘N’ composition, 2.7%) for the strain CR02. CEGMA analysis with 248 highly-conserved eukaryotic genes [15,16] reported 87% and 77% completeness of draft genomes of the strains CBS573.85 and CR02, respectively. MAKER2 [17] assigned 14,487 (strain CBS573.85) and 15,231 (strain CR02) open reading frames.

### 2.2. Data accessibility

The genome sequence data has been deposited in DDBJ under the accession numbers BCFO00000000.1 (strain CBS573.85) and BCFR00000000.1 (strain CR02).

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### Transparency document. Supplementary material

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