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Nathalie Diagne
Centre de Recherche de Bel Air

Erik Swanson
University of New Hampshire, Durham

Celine Pesce
University of New Hampshire, Durham, Celine.Pesce@unh.edu

Fatoumata Fall
Centre National de Recherches Agronomiques

Fatou Diouf
Centre National de Recherches Agronomiques

See next page for additional authors

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Authors
Nathalie Diagne, Erik Swanson, Celine Pesce, Fatoumata Fall, Fatou Diouf, Niokhor Bakhous, Dioumacor Fall, Mathieu Ndigue Faye, Rediet Oshone, Stephen Simpson, Krystalynne Morris, W. Kelley Thomas, Lionel Moulin, Diegane Diouf, and Louis S. Tisa

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Permanent Draft Genome Sequence of *Rhizobium* sp. Strain LCM 4573, a Salt-Tolerant, Nitrogen-Fixing Bacterium Isolated from Senegalese Soils

Nathalie Diagne,a,b,c Erik Swanson,c Céline Pesce,c Fatoumata Fall,a,b Fatou Diouf,a,b Niokhor Bakhoun,a,b Dioumacor Fall,a,b, d Mathieu Ndigue Faye,b,e Rediet Oshone,c Stephen Simpson,c Krystalyne Morris,c W. Kelley Thomas,c Lionel Moulin,f Diegane Diouf,a,b,g Louis S. Tisa c

Centre National de Recherches Agronomiques, Institut Sénégalais de Recherches Agricoles (CNRA/ISRA), Bambey, Senegal; Laboratoire Mixte International Adaptation des Plantes et Microorganismes Associés aux Environnements Stress Est (LAPSE), Centre de Recherche de Bel Air, Dakar-Bel Air, Senegal; University of New Hampshire, Durham, New Hampshire, USA; Laboratoire Commun de Microbiologie IRD/ISRA/UCAD, Centre de Recherche de Bel Air, Dakar, Senegal; Centre National de Recherches Forêtières, Institut Sénégalais de Recherches Agricoles (CNRA/ISRA), Hann Dakar, Senegal; Institut de Recherche Pour le Développement (IRD), UMR IPME 3439, Montpellier, France; Département de Biologie Végétale, Université Cheikh Anta Diop (UCAD), Dakar, Senegal

ABSTRACT The genus *Rhizobium* contains many species that are able to form nitrogen-fixing nodules on plants of the legume family. Here, we report the 5.5-Mb draft genome sequence of the salt-tolerant *Rhizobium* sp. strain LCM 4573, which has a G+C content of 61.2% and 5,356 candidate protein-encoding genes.

*Rhizobia* are a diverse group of alpha- and betaproteobacteria that form nitrogen-fixing symbiosis with legumes (1, 2). This group currently consists of more than 100 species dispatched in 13 genera (http://www.rhizobia.co.nz). During the past decade, the number of rhizobial species increased dramatically, especially in the genus *Rhizobium* (3). The symbiotic relationship with legumes results in the formation of a special structure on the root of the legume called the nodule (4, 5). Inside the nodule, the bacteria obtain their nutrients from the plant and in exchange produce a reduced form of nitrogen from atmospheric dinitrogen (process of biological nitrogen fixation, or BNF).

Rhizobial host plants belong to the *Leguminosae* family, which is the third largest family of angiosperms. The legume family includes roughly 730 genera and over 19,400 species (6) and is divided into three subfamilies: the *Caesalpinioideae*, *Mimosoideae*, and *Papilionoideae* (7). The *Rhizobium*-legume symbiosis provides several advantages for improving soil fertility and agricultural productivity (8). This symbiosis, by providing nitrogen to plants, limits the requirement of chemical fertilizers and thus groundwater pollution from nitrates (9). Legumes also serve as an alternative source of protein for human and animal consumption (10).

Many members of the genus *Rhizobium* are able to form nodules on a broad range of legumes, while others are very specific. *Rhizobium* sp. strain LCM 4573 was isolated from the rhizosphere of soil around *Prosopis juliflora* under saline conditions in the peanut basin in Senegal (11). The isolate also infects and forms root nodules on *Acacia seyal* plants. Under in vitro culture conditions, this strain is able to tolerate up to 700 mM NaCl. Because of these properties, this strain could potentially be used in association with leguminous plants for the reforestation of saline lands. The *Rhizobium* sp. strain LCM 4573 genome was sequenced to provide information on its physiology.
and to identify molecular markers that are involved in its tolerance to salinity.

Sequencing of the draft genome of *Rhizobium* sp. strain LCM 4573 was performed at the Hubbard Center for Genome Studies (University of New Hampshire, Durham, NH, USA) using Illumina technology techniques (12). A standard Illumina shotgun library was constructed and sequenced using the Illumina HiSeq2500 platform, which generated 9,210,384 reads (260-bp insert size) totaling 2,219 Mb. The Illumina sequence data were trimmed by Trimmomatic version 0.32 (13) and assembled using SPAdes version 3.5 (14) and ALLPaths-LG version r52488 (15). The final draft assembly for *Rhizobium* sp. strain LCM 4573 consisted of 30 contigs with an *N*<sub>50</sub> contig size of 378.27 kb and 301.9× coverage of the genome. The final assembled genome had a total sequence length of 5,521,535 bp with a G+C content of 61.2%.

The assembled *Rhizobium* sp. strain LCM 4573 genome was annotated via the NCBI Prokaryotic Genome Annotation Pipeline (PGAP), resulting in 5,356 candidate protein-encoding genes, 48 tRNAs, and 2 rRNA regions.

**Accession number(s).** This whole-genome shotgun sequence has been deposited at DDBJ/EMBL/GenBank under the accession number MDDW00000000. The version described in this paper is the first version, MDDW01000000.

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**REFERENCES**


