Title: A unique mitovirus from Glomeromycota, the phylum of arbuscular mycorrhizal fungi

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Abstract

Arbuscular mycorrhizal (AM) fungi that belong to the phylum Glomeromycota associate with most land plants and supply mineral nutrients to the host plants. One of the four viral segments found by deep-sequencing of dsRNA in the AM fungus *Rhizophagus clarus* strain RF1 showed similarity to mitoviruses and is characterized in this report. The genome segment is 2,895 nucleotides in length, and the largest ORF was predicted by applying either the mold mitochondrial or the universal genetic code. The ORF encodes a polypeptide of 820 amino acids with a molecular mass of 91.2 kDa and conserves the domain of the mitovirus RdRp superfamily. Accordingly, the dsRNA was designated as *R. clarus* mitovirus 1 strain RF1 (RcMV1-RF1). Mitoviruses are localized exclusively in mitochondria and thus generally employ the mold mitochondrial genetic code. The distinct codon usage of RcMV1-RF1, however, suggests that the virus is potentially able to replicate not only in mitochondria but also in the cytoplasm. RcMV1-RF1 RdRp showed the highest similarity to the putative RdRp of a mitovirus-like ssRNA found in another AM fungus, followed by RdRp of a mitovirus in an ascomycotan ectomycorrhizal fungus. The three mitoviruses found in the three mycorrhizal fungi formed a deeply branching clade that is distinct from the two major clades in the genus *Mitovirus*. 
Introduction

Arbuscular mycorrhizal (AM) fungi that belong to the phylum Glomeromycota associate with most land plants and supply mineral nutrients, in particular phosphorus, to the host plants through extensive hyphal networks constructed in the soil [15]. The plant-AM fungal symbiosis occurred more than 400 million years ago, and the coincidence of the appearances of early land plants and AM associations suggests that the associations were instrumental in the colonization of land by plants [14]. Although AM fungi have been playing a significant role in terrestrial ecosystems via enhancing P-cycling in the soil, biological characteristics of the fungi have been poorly understood due to their obligate biotrophic nature.

Members of the genus *Mitovirus* in the family Narnaviridae composed of a single genome segment of positive-sense RNA that encodes only RNA-dependent RNA polymerase (RdRp) [3]. Mitoviruses are localized exclusively in mitochondria of the host fungi, except for *Thanatephorus cucumeris* mitovirus that is potentially able to replicate both in the cytosol and mitochondria [6]. The infection of mitoviruses often causes malformation of mitochondria, which leads, in the case of plant pathogenic fungi, to debilitation in virulence [18] due to attenuation of mitochondrial function [12]. Accordingly, their possibility as a biological control agent has been studied extensively [1]. The impact of mitoviruses on AM symbiosis is also of interest, but no mitovirus has been described in the Glomeromycota so far.
One technical limitation for virological study in AM fungi was the difficulty in obtaining a sufficient amount of fungal material for characterization of viral genomes. We have established an open culture system for mass production of AM fungal mycelia and initiated virological studies of the fungi recently, in which four distinct dsRNA viruses, including a new class of virus, were described for the first time in the phylum [5]. In the present study, one dsRNA that was found to be similar to mitoviruses in the previous study is characterized with reference to the members of the genus *Mitovirus*.

**Provenance of the virus material**

*Rhizophagus clarus* (Nicolson & Schenck) Walker & Schüßler strain RF1 (= *Glomus* sp. strain RF1) MAFF520086 was isolated by plant trap culture of *Petasites japonicus* subsp. *giganteus* grown in acidic soil in Hokkaido, Japan in 2005 [5] and has been maintained with sorghum and groundnut grown in a greenhouse. To obtain fungal material, the strain was grown with seedlings of *Lotus japonicus* cv. Miyakojima in the mesh bag-separated open culture system [2], and dsRNA was extracted from extraradical mycelia, purified, and electrophoresed [5]. Four dsRNA segments observed in the gel were excised from the gel, purified, and randomly amplified using the anchored-N6 primer according to Márquez et al [7]. The amplicons were directly sequenced by Roche 454 FLX GS Titanium using a 1/8-scale gasket, and assembled. Among contigs obtained in the sequencing, an ORF of a
2.5-kbp contig showed similarity to RNA-dependent RNA polymerase (RdRp) of mitoviruses.

The nucleotide sequence of the coding region of 2.5-kbp dsRNA was reconfirmed by sequencing three clones for each of two >1-kbp cDNAs obtained by nested RT-PCR, and the extreme ends were determined by sequencing three clones for each of three and two RACE products of the 5' and 3' ends, respectively (Supplementary Table S1 and Fig. S1). The sequences were analyzed and annotated with Artemis (Sangar Institute) and has been deposited in the DDBJ under accession no. AB558120. The amino acid (aa) sequence of predicted ORF was subjected to BLASTp searches and aligned with those of other mitoviruses using MUSCLE implemented in MEGA 5 [17]. Neighbor-joining (NJ) and maximum-likelihood (ML) trees were constructed with MEGA 5 for phylogenetic analysis.

Four well-characterized mitoviruses and an uncharacterized mitovirus-like ssRNA were selected for comparative sequence analysis of the dsRNA of *R. clarus* RF1: TeMV found in the ectomycorrhizal fungus *Tuber excavatum* in Germany [16], CpMV found in a hypovirulent strain of the chestnut blight fungus *Cryphonectria parasitica* in USA [10], TcMV found in a hypovirulent strain of *Th. cucumeris* in USA [6], HmMV1-18 found in the violet root rot fungus *Helicobasidium mompa* in Japan [8], and an uncharacterized mitovirus-like ssRNA found in the AM fungus *Rhizophagus* sp. strain HR1 (= *Glomus* sp. strain HR1 [2]) (RMV-like ssRNA-HR1) in Japan.
Sequence properties

The sequencing of the RACE products revealed that complete genome of the dsRNA was 2,895 nucleotides (nt) in length, which was approx. 400-nt longer than that predicted by the 454 sequencing. Between a 297-nt 5' UTR and a 135-nt 3' UTR, the largest ORF (2,463 nt) was predicted by applying either the mold mitochondrial or the universal genetic code (Supplementary Fig. S2). The predicted ORF encodes a polypeptide of 820 amino acids (aa) with a molecular mass of 91.2 kDa and conserves the domain of mitovirus RdRp superfamily (Pfam PF05919), including the GDD motif (Fig. 1a). Accordingly, the dsRNA was designated as *R. clarus* mitovirus 1 strain RF1 (ReMV1-RF1). Generally functional RdRp in mitoviruses can be translated only if the mold mitochondrial genetic code is invoked [13]. This is because tryptophan residues in mitovirus RdRps are usually encoded either by a UGA or a UGG codon, but the former codon encodes a translation terminator in the universal genetic code (in the cytosol). In fact, 55, 52, and 84% of tryptophan residues are encoded by the UGA codon in the RdRps of TeMV, CpMV, and HmMV1-18, respectively. On the other hand, all tryptophan residues in RcMV1-RF1 RdRp are encoded by the UGG codon (Supplementary Fig. S2, TGG in cDNA) as well as those in TcMV RdRp [6] and putative RdRp of RMV-like ssRNA-HR1 (data not shown), suggesting that functional RdRp could be translated both in the cytosol and in mitochondria. The codons for all tryptophan residues within the conserved domain of the selected mitoviruses are shown in Fig. 1b. The
RdRp aa sequence of RcMV1-RF1 shows high levels of similarity to those found in the two mycorrhizal fungi throughout the ORF: 34% identity to that of RMV-like ssRNA-HR1 at 98% coverage and 28% identity to TeMV RdRp at 96% coverage. Significant similarity to TcMV RdRp in which all tryptophan residues are encoded by the UGG codon, however, was observed only within the conserved domain (43% identity at 23% coverage). The three RdRps of RcMV1-RF1, RMV-like ssRNA-HR1, and TeMV found in the mycorrhizal fungi form a subclade within the *Mitovirus* clade I [3] in the NJ-tree (Supplementary Fig. S3), although the node separating the clades I and II is poorly supported by a low-bootstrap value (28%). Whereas in the ML-tree the three viral sequences form a deeply branching clade with a bootstrap value of 99%, which is distinct from the two major clades (Fig. 2). A similar tree topology was also reported recently [4]. These observations suggest that the mitoviruses from the mycorrhizal fungi is likely to create the third distinct group in the genus.

The first member of *Mitovirus* in the Glomeromycota has been characterized in the present study. It seems likely that the distinct codon usage found in RcMV1-RF1 is a common feature of mitoviruses in AM fungi. The virus is potentially capable of replicating in the cytoplasm as well as in mitochondria. This might be an advantageous trait for horizontal transmission among the fungi, because those that belong to the same anastomosis group can exchange not only nuclei but also cytosol. Given the 400-million-year history of the close association of the fungi with plants, we also consider another possibility that ancestors of RcMV1-RF1 might be able to shuttle between the fungi and the host plant.
during a certain stage of their evolution. This idea is supported by the evidence that RdRps of the members in the genus *Ourmiavirus*, plant ssRNA viruses, are phylogenetically related to those of the members in the Narnaviridae [11], suggesting that mitoviruses and ourmiaviruses diverged from a common ancestor. It is thus expected that more mitoviruses employing the universal genetic code will be found in AM fungi when their sequences become available.

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**Fig. 1 a)** Alignment of the conserved domain of Mitovirus RdRp in RcMV1-RF1. Motifs are labeled according to Poch et al. [2], and the consensus amino acids are written in bold letters. Numbers in parentheses represent the number of amino acid residue in RcMV1-RF1 RdRp. **b)** Codons for the tryptophan residues (W) corresponding to those indicated in the alignment.
Fig. 2 Phyllogenetic position of ReMV1-RF1. Maximum-likelihood tree was constructed based on the amino acid sequences of mitovirus RdRP according to the JTT matrix-based model. Percentage bootstrap values (1000 replications) are indicated at the nodes. Two major clades (I and II) in Mitovirus are labeled according to Hillman and Cai [3]. Accession numbers are given in parentheses.
**Supplementary Fig. S1** Genome structure of *R. claus* mitovirus 1 strain RF1 (RcMV1-RF1). An ORF encoding RNA-dependent RNA polymerase (RdRp) was predicted. Relative positions and size of cDNA clones sequenced for confirmation are drawn below the genome.
Kitahara et al.

Supplementary Fig. S2
cDNA sequence of ReMV1-RF1 and predicted ORF according to either the mitochondrial or the universal genetic code. All tryptophan residues are written in a red letter and their codons are highlighted.
**Supplementary Fig. S3** Phylogenetic position of RcMV1-RF1. The Neighbor-joining tree was constructed based on the JTT matrix-based model. Percentage bootstrap values (1000 replication) are indicated at the nodes. Two major clades (I and II) in *Mitovirus* are labeled according to Hillman and Cai [3]. Accession numbers are indicated in Fig. 2.