Ribosomal DNA sequences of *Glugea anomala*, *G. stephani*, *G. americanus* and *Spraguea lophii* (Microsporidia): phylogenetic reconstruction

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ABSTRACT: The microsporidian species *Glugea anomala*, *G. stephani*, *G. americanus* and *Spraguea lophii* were compared by using sequence data derived from their small subunit rDNA genes which were amplified by polymerase chain reaction and directly sequenced. These sequence data and published data of *G. atherinae* were analyzed and were used to infer a phylogenetic tree. The 5 microsporidian fish parasites appeared to be closely related. The higher sequence similarities demonstrated among *G. anomala*, *G. stephani* and *G. atherinae* suggest that these 3 parasites are in fact only 1 species of *Glugea*. Moreover, the higher sequence similarities between *S. lophii* and *G. americanus* support the transfer of the latter *Glugea* species into the genus *Spraguea*.

KEY WORDS: Sequence - SSUrDNA - Microsporidia - Glugea - Spraguea - Taxonomy

INTRODUCTION

Microsporidia are obligately intracellular parasites and are found in many vertebrates and invertebrates (Sprague & Vavra 1977). However, most Microsporidia are found in insects and fishes. Many species belonging to the genus *Glugea* were described in several species of teleost fishes (Canning & Lom 1986). They have been identified and described mainly on the basis of their tissue preference, morphology and developmental life cycle as demonstrated by electron microscopy (Canning & Lom 1986). Molecular analyses based on rDNA sequence data are increasingly applied to the phylum Microsporidia. The small subunit (SSU) rRNA of several Microsporidia have already been determined for many Microsporidia including *Vairimorpha necatrix* (Vossbrinck et al. 1987), *Ameson michaelis* (Zhu et al. 1993a), *Encephalitozoon cuniculi* (Zhu et al. 1993b), *Septata intestinalis* (Zhu et al. 1993c), *Enterocytozoon bieneusi* (Zhu et al. 1994) and *Glugea atherinae* (Genbank: U15987).

Recently, a phylogenetic analysis based on riboprinting of the SSU and LSU rDNA of 7 microsporidian species has shown that *Glugea anomala*, *G. stephani* and *G. atherinae* are very closely related while *G. americanus* may be more closely related to *Spraguea lophii* than to the other *Glugea* spp. (Pomport-Castillon et al. 1997a). When we consider the species description, their taxonomic status is questionable for several reasons. (1) Experimental infection of stickleback with *G. hertwigi* obtained from the smelt *Osmerus eperlanus*, carried out successfully by Weissenberg (1968), led Canning & Lom (1986) to question the validity of the species *G. anomala* and *G. hertwigi*. (2) Successful experimental transmission of *G. atherinae*, a parasite of the sand smelt *Atherina boyeri*, to the flounder *Platichthys flesus* (Mathieu-Daudé et al. 1992) and to the turbot *Scophthalmus maximus* (Leiro et al. 1993) calls the validity of this species and of *G. stephani* into question. (3) In spite of a very similar parasitism of angler fish (little white cysts in grapes in the nervous system), genus *Lophius*, 2 species have been described: *Spraguea lophii* for the European angler fish (*L. budegassa* and *L. piscatorius*) and *G. americanus*.
for the American angler fish (*L. americanus*) (Loubès et al. 1979, Takvorian & Cali 1986).

In order to improve the taxonomy of microsporidian parasites of fish, in particular of the genera *Glugea* and *Spraguea*, we have sequenced the SSU rDNA of *G. anomala, G. stephani, G. americanus* and *S. lophii*. We report here the SSU rDNA sequences of these 4 Microsporida and a phylogenetic reconstruction based on them.

### MATERIALS AND METHODS

#### Fishes and Microsporidia

The sand smelt *Atherina boyeri* Risso, 1810, host of *Glugea atherinae* (Berrebi, 1979), the stickleback *Gasterosteus aculeatus* L., 1758, host of *Glugea anomala* (Moniez, 1887), the flounder *Platichthys flesus* L., 1758, host of *Glugea stephani* (Hagenmüller, 1889), the American angler fish *Lophius americanus* Valenciennes, 1837, host of *Glugea americanus* (Takvorian and Cali, 1986) and the European angler fish *Lophius piscatorius* L., 1758 and *L. budegassa* Spinosa, 1807, hosts of *Spraguea lophii* (Dollein, 1898) were collected directly from fishermen or from a dealer. Fishes were dissected and xenomas were isolated from parasitized host organs. The spores were recovered from the cysts in a phosphate-buffered saline (PBS) solution (pH 7.2) and purified on a Percoll gradient (25, 50, 75 and 100%). The spor suspensions obtained were stored at −20°C in PBS.

**Microsporidian DNA isolation.** After centrifugation (3000 × g), the spore pellet was resuspended in a solution containing 0.4 ml buffer (0.5 M NaCl, 0.2 M Tris HCl pH 7.5, 0.01 M ethylenediamine tetra-acetic acid and 1% sodium n-lauroyl sarcosine) and 0.4 g glass beads. The DNA was liberated from the spores by shaking in a Mini Bead Beater (Biospec. Product, Bartlesville, OK, USA). The aqueous phase was extracted with Tris-saturated phenol and precipitated with ethanol (Pomport-Castillon et al. 1997a).

**PCR amplification and sequencing.** The SSU rDNA was amplified using the forward primer 5'-GGTGG- ATTCTGCCTGACGT-3' (Baker et al. 1994) and one of the reverse primers 5'-GACGGCCGTGTGTAACAAG-3' or 5'-GGTCCGGTTTCAAGACGG-3' (Vossbrinck et al. 1993). Amplification conditions were 1 min at 94°C, 1.5 min at 55°C and 2 min at 72°C for 30 cycles using Taq DNA polymerase (Boehringer Mannheim, Germany) (Pomport-Castillon et al. 1997a). The initial primer and additional primers were used to sequence the SSU rDNA. Direct sequencing of a purified fragment was accomplished using the 70170 Sequenase™ PCR Product Sequencing kit (Amer sham, United States Biochemicals, Cleveland, Ohio, USA). PCR and sequencing were done 4 times with different isolates to take any variability into account.

GenBank/NCBI accession numbers are: *Spraguea lophii* AF056013, *Glugea americanus* AF056014, *G. stephani* AF056015 and *G. anomala* AF056016. The *G. atherinae* sequence was obtained from GenBank (accession no. U15987).

**Phylogenetic analysis.** Base compositions and GC contents of the SSU rDNA sequences of *Glugea anomala, G. stephani, G. atherinae* and *Spraguea lophii* sequences are reported in Table 1. The base composition appears to be nearly identical for *G. anomala*, *G. stephani* and *G. atherinae* with a small bias to G+C bases, whereas the 2 other species have a bias to A+T bases. The SSU rDNA shows little variation among the taxa analyzed. The *G. anomala* and *G. stephani* sequences are identical and only 1 base was different between these 2 and the *G. atherinae* sequence. By contrast, about 160 bases were different between these 3 sequences and the *G. americanus* and *S. lophii* sequences. The *G. americanus* and *S. lophii* sequences were very similar with only 7 base differences between the 2 sequences (Table 2).

Table 2 shows the percent divergence among the 5 species. The dis-
Table 2. Percent sequence divergence among 5 species of Microsporidia. S. Spraguea, G. Glugea

<table>
<thead>
<tr>
<th></th>
<th>S. lophii</th>
<th>G. americanus</th>
<th>G. anomala</th>
<th>G. stephani</th>
</tr>
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<tbody>
<tr>
<td>S. lophii</td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>G. americanus</td>
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<td>G. anomala</td>
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<td>15.8</td>
<td>0.0</td>
<td></td>
</tr>
<tr>
<td>G. stephani</td>
<td>16.0</td>
<td>15.8</td>
<td>0.1</td>
<td>0.1</td>
</tr>
<tr>
<td>G. atherinae</td>
<td>15.9</td>
<td>15.7</td>
<td>0.1</td>
<td>0.1</td>
</tr>
</tbody>
</table>

Differences between Glugea atherinae and G. anomala/G. stephani and between G. americanus and Spraguea lophii are smaller (0.1 and 0.6%, respectively) than that between G. atherinae and G. americanus/S. lophii (15.7/15.9%).

Fig. 1 illustrates a phylogenetic tree of the 5 species showing 2 clusters, i.e. the first grouping: Glugea anomala, G. stephani and G. atherinae, and the second grouping: G. americanus and Spraguea lophii. Such a tree supports the exclusion of G. americanus from the genus Glugea. Moreover, the analysis of the SSU rDNA sequences using the DOLLOP programs of the PHYLIP package generated a tree showing the phylogenetic relationship between the different fish Microsporidia (Fig. 2).

DISCUSSION

Using conserved primers in the SSU rDNA, we obtained sequences of 4 microsporidian species belonging to the genera Glugea and Spraguea (G. anomala, G. stephani, G. americanus and S. lophii) and have compared these with a sequence of another species of the Glugea genus, G. atherinae.

Our study and some recent riboprinting experiments on several microsporidian species (Pomport-Castillon et al. 1997a) show the following.

The SSU rDNA sequences of the 5 species are closely related since the lowest similarity is about 80.3 % as observed between Glugea anomala, G. stephani and G. americanus. Microsporidian fish parasites constitute a homogeneous group in comparison with other species. Malone & McIvor (1995) have shown a great sequence variability between species of the Nosema genus parasitising different classes of arthropods: 62.4 % similarity was observed between N. bombycis and N. apis.

The analysis of the base composition outlines a linking of some of these species. Pieniazek et al. (1995) have shown a variation of the GC content of 15 microsporidian species with, at the extreme, 34% for Nosema bombycis and 56% for Vavraia oncoperae. However, in the same genus the GC content is very similar, e.g. 34 to 38% for Nosema spp. Malone & McIvor (1995) have demonstrated that, mainly in Protozoa, the ratio G+C/A+T is less than or equal to 1. This is consistent for Glugea americanus and Spraguea lophii, but not for the 3 other species which show a majority of G+C bases.

Glugea anomala, G. stephani and G. atherinae are very closely related while G. americanus appears to be more closely related to Spraguea lophii than to the other 3 Glugea species. Indeed, the G. anomala and G. stephani sequences are identical or almost identical (99.9% similarity) to the G. atherinae sequence. This result is in accordance, for these 3 species, with a well established homogenous life cycle inducing voluminous xenomas encapsulated in connective tissues of their host fishes. In spite of some morphological differ-
ences in the spores, experimental infections have been carried out successfully between some of these species (Mathieu-Daudé et al. 1992, Leiro et al. 1993). These observations and our results suggest that G. anomala, G. stephani and G. atherinae could be only 1 species. However, there is evidence that some Microsporidia are cross-infective between hosts and that the morphology can be altered in different hosts (Canning & Lom 1986). Moreover, Docker et al. (1997) have demonstrated the existence of genetic variability in the rDNA sequences of another microsporidian parasites in the same host species. Indeed, the authors have sequenced 2 samples of Loma salmonae and have shown differences of 2 nucleotides. Such intra-species variability can explain the difference observed between G. anomala, G. stephani and G. atherinae by riboprinting and sequencing, and could be an argument in favour of the existence of only one species.

Glugea americanus and Spraguea lophi, in spite of causing a very similar parasitism of the American and the European angler fishes' nervous systems respectively, produce different spores. G. americanus has oval and uninucleated spores of 2.8 × 1.5 μm while S. lophi has 2 types of spores: oval, uninucleated ones and curved, binucleated ones with a size of 4.2 × 2.5 μm and 3.7 × 1.4 μm, respectively (Loubès et al. 1979, Lom & Dykova 1992). However, it is important to mention that the early proliferative forms and the complete development of G. americanus are unknown and only spores and xenomas have been described (Takvorian & Cali 1986). In our study, these 2 angler fish parasite species have a higher SSU rDNA sequence similarity (98.6%) which is due to only 2 transitions, 2 transversions and 1 deletion. The closely related sequences determined here confirm the result obtained by riboprinting and could support the transfer of G. americanus into the genus Spraguea. Indeed, percent similarities and divergencies have already been used to rename species. Zhu et al. (1993c) and Hartskeel et al. (1995) have justified the transfer of Septata intestinalis to the genus Encephalitozoon by a lower divergence (21.9%) and a high similarity (77%) between S. intestinalis and Encephalitozoon cuniculi. However, it would be of interest to know how many nucleotide substitutions constitute differences between species and genera. Molecular systematics are clearly helpful when there are many nucleotide differences but slight differences create difficulties in interpretation.

The low numbers of differences between gene sequences of these species give little information. The present work is the second step, after riboprinting, in the study of the phylogeny relationships of these 5 microsporidian species and more sequence data have to be obtained. Moreover, further steps should be taken to confirm the hypotheses inferred from the molecular data: (1) an examination of the morphology of Glugea americanus and description of the life cycle to see whether it exhibits sporal dimorphism as is present in Spraguea lophii; (2) the separation of the 2 spore types of S. lophi and analysis of both sequences to confirm that they belong to the same species; (3) cross-transmission experiments if possible among the G. anomala, G. atherinae, G. stephani group; and (4) sequencing of the ITS and the LSU rDNA or another gene more variable among the 5 species under study.

**LITERATURE CITED**


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