ESTIMATES of the number of species of ciliates vary from 3000 to 30,000 (Foissner, 2008; Fokin, 2012). Numerous descriptions of new ciliate taxa from exotic habitats and scantily investigated regions and sometimes places not yet investigated by professional taxonomists (Foissner, 2016; Foissner & Berger, 2021; Krenek et al., 2015; Rossi et al., 2016) underpin the argument that a total of 30,000 species for all Ciliophora is not unrealistic. Ciliates occupy highly diverse ecological niches. Most are free-living, some are epi and endocommensals, parasites of other unicellular organisms and metazoans, and serve as hosts for bacterial epi and endosymbionts, fungi, algae, and other protists (Dziallas et al., 2012; Fokin et al., 2014; Görtz, 2006). Ecological and trophic preferences (e.g. being bacterivores) of ciliates seem to favor endosymbioses with bacteria and a diversity of symbiotic relationships. The biodiversity of prokaryotic symbionts in Ciliophora is fascinatingly rich; and it is not by chance they might harbor potentially pathogenic bacteria.

Most of the studies published over the past decade in the field of symbiosis in ciliated protists address newly discovered or re-investigated systems of ciliate bacteria or have summarized such results (Bright et al., 2014; Fokin, 2012; Görtz, 2014; Potekhin et al., 2021; Schramlhammer & Potekhin, 2020; Serra et al., 2016; Szokoli et al., 2016; Vannini et al., 2014), and some of the investigations were devoted to the molecular aspects of such systems (Boscaro et al., 2017; Dohra et al., 2014; Fujishima, 2009; Garushyants et al., 2018).

**Abstract**

The diversity of prokaryotic symbionts in Ciliophora and other protists is fascinatingly rich; they may even include some potentially pathogenic bacteria. In this review, we summarize currently available data on biodiversity and some morphological and biological peculiarities of prokaryotic symbionts mainly within the genera *Paramecium* and *Euplotes*. Another direction of ciliate symbiology, neglected for a long time and now re-discovered, is the study of epibi- onts of ciliates. This promises a variety of interesting outcomes. Last, but not least, we stress the new technologies, such as next generation sequencing and the use of genomics data, which all can clarify many new aspects of relevance. For this reason, a brief overview of achievements in genomic studies on ciliate's symbionts is provided. Summing up the results of numerous scientific contributions, we systematically update current knowledge and outline the prospects as to how symbiology of Ciliophora may develop in the near future.

**KEYWORDS**

Ciliophora, endosymbiosis, episymbiosis, *Euplotes*, *Holospora*, *Holospora*-like bacteria, morphology, *Paramecium*, phylogeny, symbionts
The ability to gain and retain symbionts varies among the different ciliate groups, as it is in the case of other prokaryotic taxa. Most of the detected prokaryotic symbionts belong to Alphaproteobacteria, but step-by-step symbiotic abilities have been demonstrated in other bacterial groups (Boscaro et al., 2012; Bright et al., 2014; Fokin, 2004, 2012; Rosati et al., 1998; Schrallhammer & Potekhin, 2020).

Holospora and Holospora-like bacteria (HLB) are non-motile Alphaproteobacteria, showing a characteristic life cycle with infectious and reproductive forms, found mainly in the nuclear apparatus of representatives of Oligohymenophorea, Heterotrichea, Armophorea, Phyllopharyngea, and Prostomatea (Fokin, 2012; Fokin & Görtz, 2009). The majority of such cases have not been investigated in depth, and representatives of the Holospora genus were confirmed only in Paramecium. Bacteria associated with bacteriophages were found in species of Heterotrichea and Oligohymenophorea. A rare type of endosymbiotic motile bacteria was found in ciliates of the two different classes: in the cytoplasm of heterotrichs or in the macronucleus and its perinuclear space of Oligohymenophora (Fokin, 2012; Vannini et al., 2014). Some other non-motile endosymbionts were discovered in the perinuclear space of both nuclei of ciliates (Fokin & Karpov, 1995). Episymbionts are more common in Heterotrichea, Armophorea, and Plagiopylea, but recently such bacteria were recorded for Oligohymenophora (Bright et al., 2014; Castelli et al., 2019).

Apparently, Oligohymenophorea and Spirotrichea are the most promising groups of Ciliophora for the investigation of prokaryotic symbioses. A remarkable wealth of symbioses has been observed so far in Paramecium, a member of the first group. They include an infectious type of bacterial symbiosis (parasitism). A similar kind was detected in the related genus Frontonia (Fokin et al., 2019; Schrallhammer & Potekhin, 2020). In this review, particular attention will be given to the HLB group—“Candidatus Gortzia” (from here on, we will use symbiotic bacterial taxa names with abbreviation of the prefix “Candidatus”—a term that indicates that the taxon name does not yet meet all requirements for nomenclature of prokaryotes), “Ca. Hafkinia”, and Preeria as well as to endosymbionts of Euplotes species.

Among spirotrichs, Euplotes is certainly one of the most studied genera for its proneness to establish symbiotic relationships with bacterial organisms of different natures. It often harbors, in its cytoplasm, true microbial consortia (i.e. more than one bacterial species stably present inside the host cell). The best known Euplotes symbiont is Polynucleobacter necessarius, a betaproteobacterium which establishes an obligate mutualistic relationship with the host (Heckmann & Schmidt, 1987; Vannini et al., 2005, 2012). Many other species of bacteria have been found in different species of Euplotes, ranging from Alphaproteobacteria to Gammaproteobacteria and even to verrucomicrobia (Boscaro et al., 2019; Schrallhammer et al., 2013; Serra et al., 2020). It is still unclear which is the role played by the symbiont in these cases.

Another direction of ciliate's symbiology, neglected for long time and now re-discovered, is the study of epibionts of ciliates. The topic promises diverse and interesting insights, as those achieved with epixenosomes in Euplotidium spp. (Rosati et al., 1997, 1998, 1999), and more recently with “Ca. Deinanirea vastatrix” in Paramecium (Castelli et al., 2019), and the episymbionts of Parablepharisma spp. (Campello-Nunes et al., 2020) and Zoothamnium niveum (Bright et al., 2019; Rinke et al., 2006).

Endosymbionts can occupy almost all cellular compartments of ciliated protists, but not much is known about the way in which colonization takes place and how the symbiont communicates with host ciliate cell (Boscaro et al., 2017; Fujishima, 2009; Garushyants et al., 2018; Sabaneyeva et al., 2010; Schrallhammer & Potekhin, 2020). Many groups of ciliates other than Oligohymenophorea and Spirotrichea have been shown to contain symbionts, but they have generally been little investigated. Many of those endosymbionts were found and described in the “pre-molecular era”. It is valuable to find them again, analyze them using molecular approaches, and investigate how they interact with the host cell. In this perspective, new technologies, such as next-generation sequencing and the use of genomics data, can provide essential information to clarify many aspects of the subject. For this reason, also a brief overview of achievements in genomics studies on ciliate's symbionts is provided.

**Holosporaceae Symbionts in Oligohymenophorean Ciliates**

The genus Paramecium (Ciliophora, Oligohymenophorea), which comprises around 20 morphospecies (Kronek et al., 2015; Przybós & Tarcz, 2018), is one of the most studied ciliate genera. Therefore, it is not surprising that the largest number of bacterial symbionts has been reported for these ciliates, some of them having been precisely described from the middle of the 20th century (Fokin & Görtz, 2009; Potekhin et al., 2021).

Most endosymbionts of Paramecium belong to two orders within Alphaproteobacteria, namely, Rickettsiales and Holosporales (Fokin et al., 2019; Schrallhammer & Potekhin, 2020; Serra et al., 2016; Szokoli et al., 2016). We focus on representatives of the Holosporaceae family as they are the most studied in ciliate symbiosis.

Bacteria from the family Holosporaceae are known as obligate endosymbionts of eukaryotes, mostly ciliates. In the last decade, new members of the family were found in various hosts from different environments (Boscaro, Fokin et al., 2013; Fokin, 2012; Fokin et al., 2019; Lanzoni et al., 2015; Serra et al., 2016).
The biology of most Holospora and HLB, as well as their morphological peculiarities, have been described in detail in several reviews (Fokin & Görtz, 2009; Fokin & Sera, 2014; Görtz, 2006, 2014; Schrallhammer & Potekhin, 2020). They have been characterized as immobile, generally host- and compartment-specific infectious bacteria, with reproductive and infectious forms in their life cycles. Members of Holospora have been traditionally defined as bacteria able to invade nuclei of certain Paramecium species only.

Among Paramecia, two species, P. bursaria and P. chlorelligerum, contain autotrophic eukaryotic symbionts, that is, Chlorella or Chlorella-like green algae. We described such a symbiotic complex in P. chlorelligerum, in which the unicellular algae belong to Meyerella genus and inhabit its cytoplasm, while a new endosymbiont, “Ca. Holospora parva” colonizes the host macronucleus (Lanzoni et al., 2015). “Ca. Holospora parva” is a basally branching member of the Holospora but shares almost all characteristics of the genus, except connecting piece formation during the nuclear division. The connecting piece is a median body of the infected macronucleus or of the micronucleus formed during the host cell division process, in which most of the infective forms concentrate (Figure 1a): this peculiar formation is thought to improve the release of the infectious symbionts to the environment (Fokin & Sabaneyeva, 1997). This phenomenon could be considered an adaptation of bacteria that exploit the host cell division machinery to accomplish complex life cycles (Fokin, 2015; Fokin & Sabaneyeva, 1997). This feature has been found only in some HLB species retrieved in the macronuclei of representatives of Paramecium (Figure 1b) and Frontonia species (Beliaevskaya et al., 2020; Boscaro, Fokin, et al., 2013; Fokin et al., 2019; Potekhin et al., 2018; Serra et al., 2016), more in detail, in Holospora species infecting P. caudatum and P. bursaria, “H. bacillata” from P. nephridiautum or P. calkinsi, and “H. curvata” from P. calkinsi (Fokin, 1989a, 2015; Fokin & Sabaneyeva, 1993, 1997). In other words, it was shown that the connecting piece is characteristic for all Holospora species, except “Ca. Holospora parva”. Therefore, the formation of a distinctive connecting piece during host cell division was proposed as a key feature to discriminate between classical Holospora and other HLB (Fokin et al., 1996).

Classical characteristics of Holospora infectious forms are the straight rod shape with differentiated cytoplasmic and periplasmic parts and a recognition tip-like structure. In case of “Ca. Holospora parva” the periplasmic region of the cell always has two different regions, a larger and denser one and another more electronically transparent (Figure 1c). The latter characteristic was never recorded in other Holosporas, but is common for HLB (Figure 1d–f).

Considering both morphological and molecular data obtained for “Ca. H. parva”, we do not claim the presence of the connecting piece as a purely apomorphic feature for all Holospora species. However, apparently, this adaptation originated during co-evolution between Holosporas and Paramecia at a rather early step, being present in five Holospora species and absent only from one, “Ca. H. parva” (Figure 2).

Recently the former Holospora member Holospora caryophila (Preer & Preer, 1982) was reinvestigated (Potekhin et al., 2018). The authors wrote “surprisingly, they are only distantly related to other Holospora species, suggesting that they belong to a new genus within the family Holosporaceae, here described as Preeria caryophila comb. nov.” (Potekhin et al., 2018). Indeed, according to the 16S rRNA gene sequence comparison made, this HLB species is located in a different branch separated from all classical Holosporas and even other investigated HLB (Fokin et al., 2019; Potekhin et al., 2018). Two and a half decades ago, this conclusion had been reached on the basis of the absence of the connecting piece in the life cycle of the endosymbiont (Fokin et al., 1996).

Another feature that could generally separate Holospora from HLB is the relatively low host specificity of the latter. In the case of Preeria caryophila, the endosymbiont could not only infect but also survive in three ciliates of P. aurelia complex: P. biaurelia, P. octaurelia, and P. novaurelia as well in P. caudatum and P. polycaryum (Fokin, 2004; Fokin & Görtz, 2009; Lebedeva, Potekhin and Fokin, unpubl. data; Potekhin et al., 2018). “Ca. Gortzia” has the same tendency: “Ca. G. infectiva” (Boscaro, Fokin et al., 2013) could infect the macronucleus of P. jenningsi (main host) and P. quadecaurelia (another native host); “Ca. G. yakutica” (Beliaevskaya et al., 2020) infected the macronucleus in P. putrinum and P. nephridiatum (Fokin, Lebedeva, Serra, unpubl. data); “Holospora bacillata” (Fokin, 1989a) infected the macronucleus in P. nephridiatum and P. calkinsi. “Ca. Hafkinia simulans” from Frontonia salmastra and “Ca. G. shahrazadis” from P. multimicronucleatum have only been observed once (Fokin et al., 2019; Serra et al., 2016).

Regarding symbionts other than in the genus Holospora, the first complete description of “Ca. Gortzia” (Boscaro, Fokin et al., 2013) was from P. jenningsi (main host) and from P. quadecaurelia from Thailand (Figure 1d). It is a member of the Holospora—“Hafkinia”—“Gortzia” cluster (Figure 2). This bacterium has low host specificity, and it has been identified again in India from P. jenningsi (Serra et al., 2016).

Holospora phylogeny suggests that this genus has co-evolved with Paramecium, occurring in P. caudatum, P. chlorelligerum, and P. bursaria (Fokin et al., 2019; Potekhin et al., 2021). Three other genera of HLB closely related to Holospora include infectious bacteria manifesting similar life cycles and inhabiting the macronuclei of peniculine ciliates. They are: “Ca. Gortzia” (Boscaro, Fokin et al., 2013) and Preeria (Potekhin et al., 2018), which colonize different Paramecium species, and “Ca. Hafkinia” which is so far known only from Frontonia salmastra. Preeria seems to be more divergent from Holospora in molecular analyses (Figure 2).
The number of HLBs might be much higher than the 12 or so members referred to in the literature (see: Fokin et al., 2019). Several species, described in pre-molecular era, such as “H. bacillata”, “H. curvata”, and “Holospora sp.” from the macronucleus of P. putrinum (Fokin, 1989a; Fokin et al., 1999; Fokin & Görtz, 2009; Fokin & Sabaneyeva, 1993), probably belong to the “Ca. Gortzia” genus, considering the type of host and morphological traits (Figure 1b).

The infectious forms of “Ca. Hafkinia simulans”—HLB of the F. salmastra macronucleus have a distinctive appearance. They are present in the nucleus as large (up to 30 × 3.5 µm) spindle-shaped or even skittle-like infectious forms, quite similar in shape with the diatom Phaedactylum tricornutum—a common prey of the ciliate. Some forms
(probably the developing infectious forms) have many periplasmic zones and lack the recognition tip—an otherwise distinctive trait of the infectious form. Conversely, some infectious forms had two recognition tips located at opposite ends of the bacterial cell (Fokin et al., 2019). The last feature possibly improves the probability of infection. Double tips were reported in Preeria caryophila as well (Figure 1e).

In phylogenetic trees, “Ca. Hafkinia simulans” fell inside the Holosporaceae as a sister taxon to the Holospora clade, with strong statistical support (Figure 2). Thus, the biology and morphology of “Ca. Hafkinia simulans” differs more that of from Holospora, than other HLB, but phylogenetically it is more similar to classical holosporas than the “Ca. Gortzia” genus representatives (Fokin et al., 2019). The opposite argument could be made for Preeria caryophila—it is morphologically most similar (among HLB) to classical holosporas, but on phylogenetic reconstruction trees, it is far from Holospora and other HLB (Figure 2; Fokin et al., 2019; Potekhin et al., 2018). Thus, morphological and molecular results in these cases are considerably different.

One representative of the “Ca. Gortzia” genus, “Ca. G. shahrazadis”, detected in P. multimicronucleatum in India (Serra et al., 2016), has an unusual and surprising ability to reproduce in the host cytoplasm as well as in the macronucleus. This pattern has never been reported for any species of Holospora nor for members of “Ca. Gortzia”, Preeria, or “Ca. Hafkinia” (Fokin et al., 2019; Potekhin et al., 2018; Serra et al., 2016). Double infection of the cytoplasm and the macronucleus occurs in “Ca. Paraholospora nuclevisitans” (Eschbach et al., 2009), which is phylogenetically distant from the “Ca. Gortzia” clade (Figure 2; Fokin et al., 2019). As in the case of “Ca. Gortzia shahrazadis”, the endosymbiont is mainly located in the cytoplasm and morphologically does not resemble HLB; we conclude that this life cycle arose independently in these lineages of Holosporaceae.

Currently, the closest relative of “Ca. P. nucleovisitans” is “Ca. Mystax nordicus”, a cytoplasmic symbiont of P. nephridiatum (Korotaev et al., 2020). This endosymbiont has a remarkable appearance, suggesting it is the same as described in the 1980s (Fokin, 1989b). The endosymbiont was never observed inside the macronucleus; instead, it aggregated in clusters close to mitochondria.

New findings of bacteria belonging to major HLB groups and other representatives of Holosporaceae will definitely continue to occur. Some of endosymbions described in the pre-molecular period have been recollected from nature, and the bacteria were identified according to the current rules of bacterial nomenclature. This applies to “Ca. M. nordicus” (Korotaev et al., 2020) and probably to “Ca. G. yakutica”, even though the re-description of the endosymbiont was made without transmission electron microscope investigation (Beliavskiaia et al., 2020). The finding of the endosymbiont in the macronucleus of F. leucas that resemble Preeria (Fokin, unpubl. data) also suggests that more HLB biodiversity will be reported.

As last remarks: first, we recommend that we avoid the practice of naming symbiotic bacteria by the geographic place of their occurrence (Beliavskiaia et al., 2020; Korotaev et al., 2020) because in our opinion, the same endosymbiont may be found in other host populations far from the place of the first description. As an example, “Ca. Gortzia yakutica” (from Yakutia, Sakha Republic, Russia) was first found in Germany (Fokin et al., 1999) and more recently in another host species, P. nephridiatum, on the Baltic Sea coast (Fokin, Lebedeva, Serra, unpubl. data).

The second remark concerns the proposal of H. undulata as the type species of the genus Holospora some time ago (Gromov & Ossipov, 1981). Indeed, it is now clear that this species has considerable morphological plasticity, and its 16S rRNA gene sequence is almost identical to the one of H. elegans and H. recta (Potekhin et al., 2021; Serra et al., 2016; Wackerow-Kouzova & Myagkov, 2021). A recent attempt to solve this issue was made by Wackerow-Kouzova and Myagkov (2021), who proposed to introduce a subspecies level for those three holosporas, namely, Holospora undulata subsp. elegans, H. undulata subsp. recta, and Holospora undulata subsp. undulata. This proposal has, unfortunately, been made based on few molecular data, not sufficient in our opinion to correctly address the problem, and moreover, we would discourage the use of subspecies level for bacterial species.

**ENDOSYMBIONTS IN EUPLOTSES**

*Euplotes* is one of the most studied genera of Spirotrichea (Ciliophora). An increasing number of species is being described (Serra et al., 2020; Syberg-Olsen et al., 2016), with currently more than 40 free-living species. Members of the genus typically feed on bacteria, microalgae, and smaller protists. They are worldwide in distribution and widely present in marine, brackish, freshwater, and terrestrial habitats.

Generally, symbiotic events in *Euplotes* appear to be common and widespread, especially involving bacteria (Boscaro et al., 2019; Görtz, 2006; Heckmann & Schmidt, 1987; Serra et al., 2020; Vannini et al., 2005). Most studies have focused on freshwater or brackish water *Euplotes*, and little is known about endosymbions in marine species (Görtz, 2006; Vannini et al., 2004).

An important event in the evolutionary history of the genus was the origin of the obligate symbiosis among some phylogenetically close species of *Euplotes* and *Polynucleobacter necessarius* or *Polynucleobacter* spp. (Betaproteobacteria,
Burkholderiales), previously known as omicron or omicron-like particles (Heckmann & Schmidt, 1987). This kind of obligate endosymbionts occurs in all *Euplotes* species from brackish or freshwater habitats, belonging to a monophyletic group known as “clade B” (Syberg-Olsen et al., 2016). The obligate relationship with *Polynucleobacter*, essential for *Euplotes*, has been deeply explored, although not all mechanisms or benefits are understood. Thanks to genomic studies, it is clear that *Euplotes* acquired *Polynucleobacter* several times, with a process of free-living bacterial strains taking over existing symbiotic systems and replacing the former symbiotic bacteria (Boscaro et al., 2017).

This kind of plasticity in symbiont replacement was demonstrated also by laboratory experiments, in which endosymbiotic bacteria have been transferred between different *Euplotes* species by microinjection (Fujishima & Heckmann, 1984; Vannini et al., 2017).

Interestingly, if *P. necessarius* is not present in the cytoplasm of the host, another less common *Burkholderiales* bacterium, namely, “*Ca*. Protistobacter hekmanni” can take its place (Boscaro, Petroni, et al., 2013; Vannini et al., 2007, 2012) or, in the case of *E. platysomalharpa*, distantly related endosymbionts, such as “*Ca*. Devosia symbiotica” (*Alphaproteobacteria*, *Hyphomicrobiales*; Boscaro et al., 2019). The presence of this bacterium in a “clade B” *Euplotes* has led to alternative scenarios that explain the initial symbiotic event between this clade of *Euplotes* and their modern symbionts. “*Ca*. Devosia spp.” (*Alphaproteobacteria*, *Hyphomicrobiales*) have been detected in two “clade A”–*Euplotes* (*E. magnicirratus*, *E. enigma*; Boscaro et al., 2019), suggesting that the first symbiosis event involved a common ancestor of clades A and B and an alphaproteobacterium, and the symbiosis with a betaproteobacterium would have arisen later. Further studies and *Euplotes* screening for symbionts are required to solve the issue.

Beside *Betaproteobacteria* endosymbionts, *Euplotes* (from “clades A”, “C”, and “E”) can host a wide range of phylogenetic far-related bacteria, spanning from *Alphaproteobacteria* to *Gamma proteobacteria* and *Verrucomicrobia* (Boscaro et al., 2019; Schrallhammer et al., 2013; Serra et al., 2020).

*Alphaproteobacteria* symbionts are among the most frequently found endosymbionts in the cytoplasm of *Euplotes* spp. Most belong to the *Rickettsiales* (Boscaro, Petroni, et al., 2013; Senra et al., 2016; Vannini et al., 2010) or *Holosporales* (Boscaro et al., 2019) or *Hyphomicrobiales* (Boscaro et al., 2019; Vannini et al., 2004).

Representatives of *Gammaproteobacteria* are found in few species of *Euplotes*, e.g. “*Ca*. Nebulobacter yamunensis” (Boscaro et al., 2012), *Francisella endociliophora* (Schrallhammer et al., 2011), *F. adeliensis* (Vallesi et al., 2019), and “*Ca*. Endonucleario bacter sp.” (Boscaro et al., 2019).

The first verrucomicrobiunm found as an endosymbiont of a ciliate is represented by “*Ca*. Pinguicoccus supinus” (*Verrucomicrobia, Opitutae*), endosymbiont of *E. vanleeuwenhoekii*, a “clade A” *Euplotes* (Serra et al., 2020). This roundish prokaryote is about 1 μm in size and is a very peculiar endosymbiont from morphological and molecular points of view. It has distinctive invaginations of the inner membrane and an ultra-reduced genome, one of the smallest found in symbionts of ciliates, being only 163,218 bp, encoding 205 genes (Serra et al., 2020). The most striking and fascinating feature of this bacterium is that it does not possess any gene for the catalytic subunit of the DNA polymerase. Unfortunately, the role of this symbiont in its host is still unclear.

Another verrucomicrobial endosymbiont was recently found in association with one *Euplotes* sp. from the Organic lake in Antarctica: “*Ca*. Organicella extenuata”, phylogenetically close to “*Ca*. Pinguicoccus supinus” (Williams et al., 2021). The genome of this endosymbiont is even smaller: 158,228 bp (encoding 194 genes). It lacks any capacity for the biosynthesis of amino acids or vitamins. It has a certain capacity for replication, transcription, translation, and protein-folding, but no dedicated DNA polymerase for DNA replication was detected, as in “*Ca*. Pinguicoccus supinus”. Unfortunately, for “*Ca*. Organicella extenuata”, no ultrastructural analyses of the endosymbiont nor fluorescence in situ hybridization (FISH) experiments for localization inside the host have been reported.

Generally speaking, *Euplotes* species have a tendency to harbor bacteria and even true microbial consortia (i.e. two or more species of different prokaryotic species stably associated to the host.). Similar bacterial associations were detected and characterized from different *Euplotes* species. For example, one *E. octocarinatus* strain from Italy has been found infected by six different endosymbionts (including *P. necessarius* and four *Alphaproteobacteria*; Boscaro et al., 2019); one freshwater *E. aediculatus* strain from India harbored *P. necessarius* together with two *Alpha*- and one *Gamma proteobacteria* species in a stable association (Boscaro, Petroni, et al., 2013; Boscaro et al., 2012; Vannini et al., 2012, 2014). One brackish *E. woodruffii* strain from Brazil was reported to host the *Alphaproteobacteria* “*Ca*. Bandiella woodruffii” and *P. necessarius* (Senra et al., 2016). In all these cases, all the endosymbionts occupied the cytoplasm of the host, and *P. necessarius* was present. As they were found in different *Euplotes* species, from different habitats and very distant locations, it suggests that these kinds of associations could be more common than one might expect. The dynamics among these prokaryotic endosymbionts is still far from being understood, although it was hypothesized that *P. necessarius* might play a crucial role in facilitating the establishment of other symbionts in the same ciliate host (Senra et al., 2016).

As general considerations, all the endosymbionts of *Euplotes* were found in the cytoplasm (with or without a symbiosome; i.e. a specialized subcellular compartment of the host that houses an endosymbiont in a symbiotic relationship) and not in organelles, such as the macronucleus, but the reason for this is still unclear (Boscaro et al., 2019). Possibly, it is connected with the particular way that the macronucleus reorganizes before cell division of the ciliate. Moreover,
none seems to be provided of any motile structures (Boscaro et al., 2019). Despite a considerable amount of knowledge on Euplotes’ symbionts having been recently acquired, a lot of work has still to be done to understand how and why these kinds of symbioses take place in Euplotes (so often!) and which cellular metabolic pathways are involved.

EPISYMBIONTS IN CILIATES

The episymbiosis phenomenon in ciliates is a tremendously interesting and diverse topic. Episymbioses have been observed in many ciliate species belonging to different classes of the phylum, involving many kinds of bacteria from Verrucomicrobia to Rickettsiales to Gammaproteobacteria and many others. The nature of the relationship can differ greatly from case to case. The prokaryotic symbiont may be parasitic and highly damaging for the host, as is the case of the episymbiont “Ca. Deianiraea vastatrix” (Alphaproteobacteria, Rickettsiales) in P. primaurelia as it causes loss of cilia and the death of the host (Castelli et al., 2019).

Relationships may be mutualistic, as in the case of the mouthless karyorelictean Kentrophoros and its microbial “kitchen garden” (Fenchel & Finlay, 1989), from which, from time to time the ciliate gains energy as food, phagocytizing a number of bacteria. The symbionts of Kentrophoros are phagocytosed by the ciliates along the whole cell body. The bacteria are sulfur oxidizers (thiotrophs) and assigned to the so called “Ca. Kentron” clade within the Gammaproteobacteria (Seah et al., 2017).

One of the most interesting and peculiar cases of mutualistic symbiosis is the relationship among Euplotidium species with the so-called epixenosomes. These are Verrucomicrobia bacteria distributed along the dorsal surface of the ciliate. They protect the ciliate from predation with an extrusive apparatus that is triggered by external signals which are mediated by membrane receptors (Rosati et al., 1997, 1998, 1999).

Episymbionts can influence the lifestyle of their ciliate hosts, and even their morphology, as demonstrated for Zoothamnium niveum and its thiotrophic ectosymbiont “Ca. Thiobios zoothamnicoli” (Bright et al., 2014; Rinke et al., 2006), another thiotrophic bacterium belonging to the Gammaproteobacteria. Bright et al. (2019) demonstrated that Z. niveum is capable of a striking polymorphism connected to the presence of these ectosymbionts. Symbiotic and aposymbiotic organisms differed significantly in colony growth, form, and fitness (Bright et al., 2019).

Episymbionts of ciliates can also be employed as diagnostic features, such as in case of members of Parablepharisma (Campello-Nunes et al., 2020): P. bacteriophora and P. brasiliensis possess rod-shaped ectosymbiotic bacteria transversally attached to the cortex, while P. granulata shows episymbionts longitudinally attached to the cortex. No molecular studies have been carried out yet on these organisms, and we still do not know if there is a species-specificity among Parablepharisma species and their ectosymbionts.

A fascinating aspect is of episymbionts of ciliates living in anoxic or semi-anoxic conditions, such as members of Plagiopylea and Armophorea, for example, Plagiopyla ramani, P. nasuta (nitla et al., 2019), Metopus contortus, and Caenomorpha levanderi (Fenchel & Ramsing, 1992). Molecular data are not available for these cases, but preliminary FISH experiments suggest the presence of sulfate-reducing microorganisms (Fenchel & Ramsing, 1992).

The examples above reflect a small part of the literature dedicated to episymbionts of ciliates. Despite the remarkable observations, the field is still poorly investigated. Episymbioses is a topic that deserves extensive efforts from the scientific community as they may provide the keys to understanding: (1) how eukaryote cells are colonized by prokaryotes which could later become endosymbionts in a second evolutionary step after the prolonged cell to cell contact of an episymbiotic relationship and/or (2) how prokaryotes evolved a separate adaption path to another different kind of niche (extra vs. intracellular).

GENOMES OF CILIATE SYMBIONTS

Second- and third-generation sequencing techniques allow us to obtain genomes for uncultivable organisms, such as the symbionts in ciliates. To date, around 20 complete genomes of such symbionts have been sequenced and annotated (Table 1). Most belong to the Proteobacteria, in particular the Alphaproteobacteria and Gammaproteobacteria (Table 1), but recently, the first genomes of two methanogenic archaeal endosymbionts have been provided (Lind et al., 2018).

In terms of genome size, symbionts of ciliates range from 158 and 163 kbp of “Ca. Organicella extenuata” and of the E. vanleewihoeki endosymbiont, “Ca. Pinguicoccus supinus” (Serra et al., 2020), to the genomes of 3.31 and 5.02 Mbp long from the symbionts “Ca. Kentron” clade from Kentrophoros (Seah et al., 2019). The latter are unique among thiotrophic symbionts because they do not encode canonical pathways for autotrophic carbon fixation but have a variety of heterotrophic features. They have the potential to oxidize sulfur to provide energy for assimilating organic carbon as the main carbon source for growth (Seah et al., 2019).
| Host                      | Symbiont                           | Symbiont classification                                                                 | Localization          | GC content (%) | Genome size (Mb) | References                  |
|--------------------------|------------------------------------|----------------------------------------------------------------------------------------|-----------------------|----------------|-----------------|-----------------------------|
| Paramecium caudatum      | Holospora obtusa                   | Bacteria; Alphaproteobacteria; Holosporales; Holosporaceae                             | Macronucleus          | 35.2           | 1.33            | Dohra et al. (2014)         |
| Paramecium caudatum      | Holospora elegans                  | Bacteria; Alphaproteobacteria; Holosporales; Holosporaceae                             | Macronucleus          | 36             | 1.27            | Dohra et al. (2014)         |
| Paramecium caudatum      | Holospora undulata                 | Bacteria; Alphaproteobacteria; Holosporales; Holosporaceae                             | Macronucleus          | 36.1           | 1.40            | Dohra et al. (2013, 2014)   |
| Paramecium bursaria      | Holospora curviscula               | Bacteria; Alphaproteobacteria; Holosporales; Holosporaceae                             | Macronucleus          | 37.6           | 1.70            | Garushyants et al. (2018)   |
| Paramecium biaurelia     | "Caelidomonas varicaedens"         | Bacteria; Alphaproteobacteria; Holosporales; Caeidimonadaceae                         | Cytoplasm or macronucleus | 42.1           | 1.68            | Suzuki et al. (2015)        |
| Paramecium sp.           | "Candidatus Fokinia solitaria"     | Bacteria; Alphaproteobacteria; Rickettsiales; Midichloriaceae                         | Cytoplasm             | 35.8           | 0.83            | Floriano et al. (2018)      |
| Paramecium primaurelia   | "Candidatus Deianiraea vastatrix"  | Bacteria; Alphaproteobacteria; Rickettsiales; Deianiraeeae                            | External cell surface | 32.9           | 1.2             | Castelli et al. (2019)      |
| Paramecium tredec Aurelia| "Candidatus Sarmatiella mevalonica"| Bacteria; Alphaproteobacteria; Rickettsiales; Rickettsiaceae                         | Cytoplasm             | 38             | 1.27            | Castelli et al. (2021b)     |
| Paramecium polycaryum    | "Candidatus Gromoviella agglomerans"| Bacteria; Alphaproteobacteria; Holosporales; Holosporaceae                             | Cytoplasm             | 32.2           | 0.589           | Castelli et al. (2021a)     |
| Euplotes petzi           | Francisella adeliensis             | Bacteria; Gamma proteobacteria; Thiotrichales; Francisellaceae                        | Cytoplasm             | 32.6           | 2.05            | Vallesi et al. (2019)       |
| Undescribed Plagiopylea  | "Candidatus Azomaticus ciliaticola"| Bacteria; Alphaproteobacteria                                                          | Cytoplasm             | 24.4           | 0.29            | Graf et al. (2021)          |
| Kentrophoros sp.         | "Candidatus Kentron" clade         | Bacteria; Alphaproteobacteria                                                          | External cell surface | Approximately 50 | 3.3–5.0     | Seah et al. (2019)          |
| Paramecium tretdec Aurelia| Caedibacter taeniosp oralis        | Bacteria; Gamma proteobacteria; Thiotrichales; Fastidiosibacterae                     | Cytoplasm             | 41.3           | 1.32            | Pirritano et al. (2020)     |
| Pseudoblepharisma tenue  | "Candidatus Thiodicyon intracellulare" | Bacteria; Gamma proteobacteria; Chromatiaceae                                        | Cytoplasm             | 64.2           | 2.9             | Muñoz-Gómez et al. (2021)   |
| Trimyema compressum      | Unnamed, strain TC1                | Bacteria; Firmicutes                                                                  | Cytoplasm             | 32.8           | 1.59            | Shinzato et al. (2016)      |
| All Euplotes spp.        | belonging to clade B               | Polynucleobacter spp., including Polynucleobacter necessarius                       | Cytoplasm             | 45.6           | 1.5–1.9         | Boscardo, Felletti, et al. (2013) |
| Euplotes sp. (AntOrgLke) | "Candidatus Organicella extenuata" | Bacteria; Verrucomicrobia                                                            | Unknown               | 32             | 0.158           | Williams et al. (2021)      |
| Euplotes vanleewenhoeki  | "Candidatus Pinguicoccus supinus"  | Bacteria; Verrucomicrobia                                                            | Cytoplasm             | 25.1           | 0.163           | Serra et al. (2020)         |
| Metopos contortus        | Methanocorpusculum sp. MCE         | Archaea; Methanomicrobia; Methanomicrobia                                             | Cytoplasm             | 50.1           | 1.69            | Lind et al. (2018)          |
| Nyctotherus ovalis       | Methanobrevibacter sp. NOE         | Archaea; Methanobacteria; Methanobacteriales                                          | Cytoplasm             | 25.4           | 1.92            | Lind et al. (2018)          |
Complete genome assembly of “Ca. Gromoviella agglomerans” is the smallest reported genome among the order *Holosporales* (589,967 bp) and presents a severely reduced metabolism, both for what concerns biosynthetic pathways and for energy production and conversion (Castelli et al., 2021a).

Some obligate intracellular symbionts are subject to genomic reduction that may be accompanied by extensive gene loss, pseudogenization, a high rate of mutations, and low GC content (Floriano et al., 2018; Garushyants et al., 2018; Lind et al., 2018). The highly reduced genome of “Ca. Azoamicus ciliaticola”, from an undescribed plagiopyloid, has presumably preserved traits that are beneficial to the host and that provide energy from anerobic respiration. Indeed, this symbiont is an obligate endosymbiont that has retained cellular functions that are markedly similar to those of mitochondria, although it did not originate from the mitochondrial line of descent (Graf et al., 2021).

In other cases, genome reduction is much less extreme in *P. necessarius*, the betaproteobacterial endosymbiont of the ciliate *Euplotes* has a genome (1.56 Mbp long) that is approximately as large as its free-living counterparts (Boscaro et al., 2017). Genomic analyses show that intracellular bacteria use strategies to interact with, invade, and exploit their host cell, including secretion systems and effector, such as Type IV and Type VI secretion systems (Castelli et al., 2019), proteins with repeat motifs such as ankyrin repeat motifs (Floriano et al., 2018) and ADP/ATP translocase that directly import ATP from the host (Castelli et al., 2021b; Dohra et al., 2014; Garushyants et al., 2018; Vallesi et al., 2019). This potential genomic adaptation for an intracellular lifestyle is not always so marked. This is shown by the genomes of the two methanogenic endosymbionts of the anerobic ciliates, *Metopus contortus* and *Nyctotherus ovalis*. Their genomes are in an early stage of adaptation toward endosymbiosis, as evidenced by the large number of genes undergoing pseudogenization (Lind et al., 2018).

**PERSPECTIVES**

Genomic studies of ciliate symbionts give excellent opportunities to deepen our knowledge on the processes of symbioses. Currently, this line of research is poorly explored because of the absence of extensive molecular data. Hence, a fundamental challenge will be to increase the number of complete genomes which, in parallel with comparative genomics analyses, will shed light on specific adaptations to symbiosis. A better understanding of host–symbiont interactions will depend on more targeted genomic studies.

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