Colonial choanoflagellate isolated from Mono Lake harbors a microbiome

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ABSTRACT

Choanoflagellates offer key insights into bacterial influences on the origin and early evolution of animals. Here we report the isolation and characterization of a new colonial choanoflagellate species, *Barroeca monosierra*, that, unlike previously characterized species, harbors a stable microbiome. *B. monosierra* was isolated from Mono Lake, California and forms large spherical colonies that are more than an order of magnitude larger than those formed by the closely related *Salpingoeca rosetta*. By designing fluorescence in situ hybridization probes from metagenomic sequences, we found that *B. monosierra* colonies are colonized by members of the halotolerant and closely related *Saccharospirillaceae* and *Oceanospirillaceae*, as well as purple sulfur bacteria (*Ectothiorhodospiraceae*) and non-sulfur *Rhodobacteraceae*. This relatively simple microbiome in a close relative of animals presents a new experimental model for investigating the evolution of stable interactions among eukaryotes and bacteria.

IMPORTANCE

The animals and bacteria of Mono Lake (California) have evolved diverse strategies for surviving the hypersaline, alkaline, arsenic-rich environment. We sought to investigate whether the closest living relatives of animals, the choanoflagellates, exist among the relatively limited diversity of organisms in Mono Lake. We repeatedly isolated members of a single species of choanoflagellate, which we have named *Barroeca monosierra*, suggesting that it is a stable and abundant part of the ecosystem. Characterization of *B. monosierra* revealed that it forms large spherical colonies that each contain a microbiome, providing an opportunity to investigate the evolution of stable physical associations between eukaryotes and bacteria.
A newly identified choanoflagellate species forms large colonies that contain a microbiome

Choanoflagellates are the closest living relatives of animals and, as such, provide insights into the origin of key features of animals, including animal multicellularity and cell biology [1,2]. Over a series of four sampling trips to Mono Lake, California (Fig. 1A; Table S1) we collected single-celled choanoflagellates and large spherical choanoflagellate colonies, many of which were hollow (Fig. 1B) and resembled the blastula stage of animal development. In colonies and single cells, each cell had the typical collar complex observed in other choanoflagellates: an apical flagellum surrounded by a collar of microvilli [1,2]. In these “rosette” colonies, the cells were oriented with the basal pole of each cell pointing inwards and the apical flagellum facing out (Fig. 1). To study the Mono Lake choanoflagellates in greater detail, we established clonal strains from ten independent isolates, two of which were each started from a single-celled choanoflagellate and the remaining eight of which were each started from a single colony (Table S1). The two strains started from single-celled choanoflagellates, isolates ML1.1 and ML1.2, took on the colonial morphology observed in the other isolates after culturing in the laboratory, suggesting that the colonies and single cells isolated from Mono Lake could belong to the same species. We are aware of no prior reports of choanoflagellates having been cultured from any alkaline soda lake, including Mono Lake.

The 18S rDNA genes for six of the Mono Lake isolates were sequenced and found to be >99% identical (Table S1). Phylogenetic analysis confirmed that all of the isolates are members of a single choanoflagellate species (Fig. S1). In further phylogenetic analyses based on 18S rDNA and two protein-coding genes from isolate ML2.1 (Fig. 1C) [3], we found that its closest relatives are the emerging model choanoflagellate S. rosetta [4–8], additional Salpingoeca spp. [9] and Microstomoeca roanoka [3,10]. The phylogenetic distance separating the Mono Lake species from its closest relatives is similar to the distance separating other choanoflagellate genera. Therefore, we propose the name Barroeca monosierra, with the genus name inspired by esteemed choanoflagellate researcher Prof. Barry Leadbeater and the species name inspired by the location of Mono Lake in the Sierra Nevada mountain range. (See Supplemental Methods for further details and a formal species description.)

Although B. monosierra and S. rosetta form rosette-shaped spherical colonies, they differ greatly in size. S. rosetta colonies range from 10-30 µm in diameter while B. monosierra forms among the largest choanoflagellate colonies observed [1,11], with a single culture exhibiting colony sizes spanning from 10-120 µm in diameter (Fig. 1D-F). Unlike the rosettes of S. rosetta, in which the basal poles of cells are closely apposed in the rosette center [5,7,11,12], cells in large B. monosierra rosettes form a shell on the surface of a seemingly hollow sphere. Inside the ostensibly hollow sphere, a branched network of extracellular matrix connects the basal poles of all cells (Fig. S2.)
Upon staining *B. monosierra* with the DNA dye Hoechst 33342, we observed the expected toroidal nuclei in each choanoflagellate cell [12,13], but were surprised to detect Hoechst-positive material in the interior of *B. monosierra* colonies (Fig. 2A, A’). Transmission electron microscopy revealed the presence of 1 µm and smaller cells with diverse morphologies bounded by cell walls in the centers of rosettes (Fig. 2B, B’; Fig. S3). Together, these observations led us to hypothesize that the centers of *B. monosierra* colonies contain bacteria.

By performing hybridization chain reaction fluorescence *in situ* hybridization (HCR-FISH [14–16]) with a broad-spectrum probe of bacterial 16S rRNA, EUB338 [17], we confirmed that the cells in the center of colonies are bacteria (Fig. 2C). A second probe that specifically targeted 16S rRNA sequences from Gammaproteobacteria, GAM42a, revealed that the majority of the bacteria inside the colonies are Gammaproteobacteria (Fig. 2C’)[18]. Finally, by incubating *B. monosierra* cultures with fluorescently labeled D-amino acids, which are specifically incorporated into the cell walls of growing bacteria, we found that the bacteria in *B. monosierra* colonies are alive and growing (Fig. S4)[19]. Therefore, *B. monosierra* contains a microbiome as defined in [20].

To visualize the spatial distribution of choanoflagellate and bacterial cells in a representative colony, we generated a 3D reconstruction from serial sections imaged by TEM. The colony contained 70 choanoflagellate cells that were tightly packed, forming a largely continuous monolayer of cells (Fig. 2D). As observed by immunofluorescence microscopy (Figs. 2A and 2A’), all cells were highly polarized and oriented with their apical flagella and collars extending away from the centroid of the rosette. Many cells were connected by fine intercellular bridges (Fig. S5) that have been previously observed in other colonial choanoflagellates, including *S. rosetta* [5,12].

The 3D reconstruction also revealed at least 200 bacterial cells in the center of the rosette (Fig. 2D’, 2D’’), some of which were physically associated with and wrapped around the choanoflagellate ECM (Fig. S6). A small number of bacterial cells were observed between the lateral surfaces of choanoflagellate cells, although it was not possible to determine whether they were entering or exiting the colony (Fig. 2D’’; Fig. S7). Colonies failed to incorporate bacteria-sized bovine serum albumin (BSA)-coated latex microspheres (0.2 µm and 1 µm) into their centers, suggesting that environmental bacteria may not be capable of passively accessing the centers of *B. monosierra* colonies (Fig. S8).

**Gammaproteobacteria and Alphaproteobacteria in the *B. monosierra* microbiome**

We next sought to identify which bacteria comprise the microbiomes of *B. monosierra*. To identify candidate bacteria for which to design FISH probes, we first sequenced and assembled metagenomes and 16S rDNA sequences from choanoflagellate-enriched samples and from environmental bacteria-enriched samples. These samples were derived from two co-cultures of *B. monosierra* with Mono Lake bacteria, ML2.1E and ML2.1G (Fig. S9), with the enrichment for choanoflagellates or
bacteria performed by centrifugation. A total of 24 different bacterial species were
identified via two complementary bioinformatic approaches (EukRep Metagenomic
Analysis and EMIRGE 16S rRNA Analysis; Table S2), of which 22 species were present
in fractions enriched with B. monosierra colonies (Table S3). The phylogenetic
relationships among these and other bacterial species were determined based on
analysis of highly conserved ribosomal proteins and 16S rDNA sequences (Fig. 2E and
S10).

The 22 bacterial species detected in cultures with B. monosierra may have co-
sedimented with the B. monosierra colonies due their community-structure densities
e.g. biofilms), a transient association with the choanoflagellate colonies (e.g. as prey),
or through a stable association with the choanoflagellate colonies. Upon investigation by
FISH microscopy, we detected ten or eleven of these species in the centers of B.
monosierra colonies (Table S4, Fig. S11). (The uncertainty regarding the precise
number of choanoflagellate-associated bacterial species stems from the inability to
disambiguate 16S rDNA sequences corresponding to one or two of the species.) Of
these microbiome bacteria, nine were Gammaproteobacteria from the families
Oceanospirillaceae (Fig. S11A; OceaML1, OceaML2, OceaML3, OceaML4, OceaML4),
Ectothiorhodospiraceae (Fig. S11B; EctoML1, EctoML2, EctoML3, EctoML4), and
Saccharospirillaceae (Fig. S11C; SaccML), matching our original observation that the
majority of the bacteria were Gammaproteobacteria (Fig. 2C, C'). The remaining
species was a Roseinatronobacter sp. (RoseML; Alphaproteobacteria) (Fig. S11D). The
microbiome bacteria exhibited an array of morphologies, from long and filamentous to
rod shaped (Fig. S3 and Fig. S12). Intriguingly, with the exception of OceaML3, which
was exclusively detected inside B. monosierra colonies of ML2.1E (Fig. S13), all other
microbiome species identified in this study were detected both inside and outside the
colonies.

In animals, the microbiome often contains a core set of host-adapted bacteria
that are present in many or all individuals in a host species, as well as a more flexible
set of bacteria that may be found in only a subset of individuals [21–23]. To identify core
members of the B. monosierra microbiome, we measured the frequency with which
colonies of ML2.1EC contained or lacked a number of representative microbiome
bacteria (Fig. S14A) and estimated the abundance of each species relative to the total
microbiome (Fig. S14B). Only one bacterium tested, OceaML1, was found in the
microbiome of all B. monosierra colonies (Fig. S14A). The other most frequently
observed members of the microbiome were SaccML (93.3% of colonies), EctoML3
(91.8% of colonies) and EctoML1 (82.4% of colonies; Fig. S14A). Two other
Gammaproteobacterial species, OceaML2 and EctoML2, were found in ~50 - 60% of
colonies, while the Alphaproteobacterium RoseML was found in only 13.9% of rosettes.

The most common resident of the B. monosierra microbiome, OceaML1, was also the
most abundant, representing on average 66.4% of the total bacterial load per colony
(Fig. S14B). Other abundant bacteria, some found in >80% of colonies, represented
approximately smaller percentages of the average bacterial biomass in the
microbiomes in which they were found. For example, SaccML was found in 93.3% of
microbiomes but represented only 30.3% of total bacteria in the *B. monosierra* rosettes in which it was found. EctoML1, which was found in 82.4% of *B. monosierra* rosettes represented less than 10% of the bacteria in the bacteria in which was detected. Thus, only OceaML1 appears to be a core member of the *B. monosierra* microbiome. Other symbionts detected in *B. monosierra* (e.g. SaccML and OceaML2-4) are close relatives of OceaML1 and may engage in similar metabolic or developmental functions in their interactions with *B. monosierra* [24].

**Discussion**

Interactions with bacteria are essential to choanoflagellate nutrition and life history. Bacteria are the primary food source for choanoflagellates, and the choanoflagellate *S. rosetta* responds to different secreted bacterial cues to undergo either multicellular developmental or mating [25–28]. We report the isolation and characterization of a new choanoflagellate species, *B. monosierra*, that forms large colonies and contains a microbiome consisting of at least ten different bacterial symbionts. To our knowledge, this is the first example of a stable interaction between choanoflagellates and ectosymbiotic bacteria. Future studies will be important to determine how the *B. monosierra* colonies and their bacterial symbionts interact.

We detected 10 – 11 bacterial species from four different families (*Saccharospirillaceae*, *Oceanospirillaceae*, *Ectothiorhodospiraceae*, and *Rhodobacteraceae*) in the *B. monosierra* microbiome. Comparisons with other symbioses suggest that their interactions with *B. monosierra* may relate to metabolism and detoxification of environmental sulfur. Such functions have been reported for *Oceanospirillaceae* species that are symbionts of diverse marine animals, from corals to snails [29–36] and for *Ectothiorhodospiraceae* that are symbionts of diverse ciliates and animals [37–40].

Although the composition of the animal gut microbiome often varies between individuals in a species, many host species harbor a core set of microbes, with which they participate in stable metabolic interactions and may coevolve [22,23]. Indeed, choanoflagellates express homologs of Toll-like receptors that, in animals, mediate interactions with gut bacteria to maintain homeostasis [10,41]. In *B. monosierra*, OceaML1 was found in 100% of rosettes assessed and was the most abundant bacterium in the microbiome, suggesting that its interactions with *B. monosierra* may be essential for *B. monosierra* biology. Interestingly, the related bacterium *Endozoicomonas* sp. (Order Oceanospirillales), is a core member of the gut microbiome of the ascidian *Ciona intestinalis*, where it represents up to 54% of the bacterial biomass [32].

*B. monosierra* and its associated microbiome provide a unique opportunity to characterize a new symbiotic interaction between a single choanoflagellate species and a microbial community. Due to the phylogenetic relevance of choanoflagellates, this symbiotic relationship has the potential to illuminate the ancestry of and mechanisms
underlying stable associations between animals and bacteria, colonization of organisms by diverse microbial communities, and insights into one of the most complex animal-bacterial interactions, the animal gut microbiome.
**Data Availability:** Genbank accession numbers for bacterial 16S rDNA sequences are listed in Table S6. Sequences for *B. monosierra* 18S rDNA, EFL, and Hsp90 (Fig. 1C) have been assigned GenBank accession numbers MW838180, MW979373 and MW979374, respectively. 18S sequences for different *B. monosierra* strains (Fig. S1, Table S1) have been assigned GenBank accession numbers MZ015010-MZ015015. The assembled *B. monosierra* genome sequence is currently being uploaded to GenBank and will be available under the accession number PRJNA734368. The *B. monosierra* genome, all bacterial genome sequences, and all relevant input and output data from the phylogenetic trees presented in Fig. 1C and Fig. S1 are available via FigShare (https://doi.org/10.6084/m9.figshare.14474214).

**Author contributions:** *P.W. performed metagenomic analysis, K.M. did the transmission electron microscopy, D.L. and P.B. did the TEM-reconstruction, and C.F. helped culture B. monosierra. A.G.D.L.B. imaged the theca. K.H. performed all other experiments and analysis. D.J.R. originally isolated B. monosierra and contributed to manuscript editing and the taxonomic description. J.B. and N.K. contributed to project leadership, experimental design, figure design, writing, and editing.*

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Figure 1
Figure 1. A new colony-forming choanoflagellate isolated from Mono Lake.

(A) Choanoflagellates were collected from two sampling sites (asterisks) near the shore of Mono Lake, California. (Modified from map at monolake.org.) (B) *B. monosierra* forms large colonies (DIC image). Scale bar = 50µm. (C) *B. monosierra* (shown in bold) is a craspedid choanoflagellate closely related to *S. rosetta* and *Microstomoeca roanoka*. Phylogeny based on sequences of 3 genes: 18S rDNA, EFL and HSP90. Metazoa (7 species) were collapsed to save space. Bayesian posterior probabilities are indicated above each internal branch, and maximum likelihood bootstrap values below. (A ‘-’ value indicates a bifurcation lacking support or not present in one of the two reconstructions.) (D-E) Two representative colonies reveal the extremes of the *B. monosierra* colony size range (D, 58 µm diameter; E, 19 µm diameter; scale bar = 20 µm). In *B. monosierra* colonies, each cell is oriented with its apical flagellum (white; labeled with anti-tubulin antibody) and the apical collar of microvilli (red; stained with phalloidin) pointing out. Nuclei (cyan) were visualized with the DNA-stain Hoechst 33342. (F) Colonies of *B. monosierra* span from 10 µm in diameter, a size comparable to that of small *S. rosetta* colonies, to 120 µm, over an order of magnitude larger. Diameters of *B. monosierra* and *S. rosetta* colonies were plotted as a violin plot; median indicated as thick black line. Diameters of representative colonies indicated as colored bars behind violin plot (D, red bar; E, blue bar).
Figure 2
**Figure 2.** *B. monosierra* colonies are filled with bacteria.

(A, A’) The center of a representative *B. monosierra* colony, shown as a maximum intensity projection (A) and optical z-section (A’), contains DNA (revealed by Hoechst 33342 staining; cyan). Apical flagella were labeled with anti-tubulin antibody (white); microvilli were stained with phalloidin (red). Hoechst 33342 staining (cyan) revealed the toroidal choanoflagellate nuclei along the colony perimeter and an amorphous cloud of DNA sitting within the central cavity formed by the monolayer of choanoflagellate cells.

(B-B’) Thin section through a representative *B. monosierra* colony, imaged by transmission electron microscopy (TEM), revealed the presence of small cells in the central cavity. (B’) Inset (box; panel B’) reveals that the interior cells are each surrounded by a cell wall. (C-C’’) The small cells inside *B. monosierra* colonies are bacteria, as revealed by hybridization with a broad spectrum 16S rRNA probe (C, green) and a probe targeting Gammaproteobacteria (C’, red). Choanoflagellate nuclei and bacterial nucleoids were revealed by staining with Hoechst (C”, cyan). (C’’) Merge of panels C – C”. Scale bar for all = 5 μm. (D-D”) 3D reconstruction of a 70-cell *B. monosierra* choanoflagellate colony from transmission electron micrographs of serial ultrathin sections revealed that the bacteria are closely associated with and wrapped around the ECM inside the colony. (D) Whole colony view. (D’) Cut-away view of colony center. Color code: cell bodies (cyan); microvilli (orange); flagella (green); bacteria (red); ECM (white); intercellular bridges (yellow, see also Fig. S5); filopodia (purple). (D’’) Reducing the opacity of the choanoflagellate cell renderings revealed the presence of bacteria positioned between the lateral surfaces of choanoflagellate cells (brackets, see also Fig. S7). (E) Unrooted phylogenetic tree based on 16 concatenated ribosomal protein sequences representing bacterial diversity modified from [42], illustrated to indicate the phylogenetic placement of bacteria co-cultured from Mono Lake with *B. monosierra*. The bacteria belonged to four major classes: Spirochaetia, Alphaproteobacteria, Gammaproteobacteria, and Bacteroidetes, however the bacteria found associated with *B. monosierra* colonies came only from Alphaproteobacteria and Gammaproteobacteria. Circles represent the phylogenetic placement of non-symbionts (white) and symbionts (*Oceanospirillaceae* sp., magenta; *Saccharospirillaceae* sp., green; *Ectothiorhodospiraceae* sp., blue; *Roseinatronobacter* sp., orange). See also Figs. S10 and S11.
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