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ABSTRACT

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2 We investigated how Legionella pneumophila (Lp) JR32 interacts with Anteglaucoma 3 CS11A and Colpoda E6, two ciliates that we isolated from sewage and sink trap sludge, 4 respectively, using a handmade maze device containing a 96-well crafting plate. Our 18S rDNA-based phylogenetic analysis showed that Anteglaucoma CS11A and Colpoda E6 5 6 formed distinct clades. Scanning electron microscopy showed that Anteglaucoma CS11A 7 had a bigger-sized body than Colpoda E6 and, unlike Tetrahymena IB (the reference 8 strain), neither ciliate produced pellets. Fluorescence microscopic observations revealed 9 that although the intake amounts differed, all three ciliates rapidly ingested LpJR32 10 regardless of the presence or absence of the icm/dot virulence genes, indicating that they 11 all interacted with LpJR32. In co-cultures with Anteglaucoma CS11A, the LpJR32 levels 12 were maintained but fell dramatically when the co-culture contained the LpJR32 icm/dot 13 deletion mutant instead. Anteglaucoma CS11A died within 2 days of co-culture with 14 LpJR32, but survived co-culture with the deletion mutant. In co-cultures with Colpoda 15 E6, LpJR32 levels were maintained but temporarily decreased independently of the 16 virulence gene. Concurrently, the Colpoda E6 ciliates survived by forming cysts, which 17 may enable them to resist harsh environments, and by diminishing the sensitivity of 18 trophozoites to Lp. In the *Tetrahymena* IB co-cultures with LpJR32 or Δicm/dot, the Lp 19 levels were maintained, albeit with temporal decreases, and the *Tetrahymena* IB levels 20 were also maintained. We conclude that unlike *Tetrahymena* IB with pellet production, 21 Anteglaucoma CS11A can be killed by LpJR32 infection, and Colpoda E6 can resist 22 LpJR32 infection through cyst formation and the low sensitivity of trophozoites to Lp.

1	Thus, the two ciliates that we isolated had different susceptibilities to LpJR32 infection.
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3	Keywords
4	Legionella pneumophila JR32, Anteglaucoma CS11A, Colpoda E6, infection,
5	susceptibility, co-cultures
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INTRODUCTION

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2 Legionella pneumophila (Lp) is ubiquitous in a wide range of natural environments such 3 as soil or pond water, where it interacts with amoebae [1-3]. Amoebae provide Lp with 4 the intracellular niche required for its replication. In the process of adapting to its cellular 5 environment, Lp acquired a set of virulence genes encoding the Icm/Dot system that 6 deliver effector proteins to support its successful phagocytosis and invasion, thereby 7 favoring its growth inside human cells along with its own growth [4-7]. As a human 8 pathogen, Lp can cause life-threatening atypical pneumonia (legionellosis) in 9 immunocompromised patients when aerosols or mine dust contaminated with it are 10 inhaled [8-10]. Lp prefers to colonize plumbing walls or gravel floors in hot springs, and 11 Legionella outbreaks frequently occur when people take public baths in circulating water 12 systems or in free-flowing hot springs where aerosols containing Lp are frequently 13 formed [11-13]. Therefore, from a public health perspective, the ability to control Lp requires better understanding of its interactions with amoebae. 14

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Various microorganisms such as predatory ciliates with numerous cilia gather in places where amoebae and Lp interactions occur [14-16], and several studies have shown an association of Lp with ciliates [17, 18]. Meanwhile, the role of these ciliates in maintaining and/or killing Lp is not fully understood, although some ciliates (*Paramecium* and *Tetrahymena*) play a role in the symbiotic interaction of Lp [19-23]. With the exception of *Paramecium* and *Tetrahymena*, research on the interactions of other ciliates with Lp has not progressed because the active movements and the global

diversity with low concentration of ciliates make them difficult to isolate [24, 25].

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3 Ciliates are diverse and over 8,000 species have been reported to date [26-28]. As 4 mentioned above, interactions between Lp and Paramecium or Lp and Tetrahymena have 5 been investigated, and both ciliates have been found to support the intracellular growth of 6 Lp, albeit differently [19-23]. Some Paramecium strains can maintain the survival of Lp 7 symbiotically, depending on the lefA gene (Legionella endosymbiosis-modulating factor 8 A), a key factor contributing to the life stage change in Lp from endosymbiosis to host 9 lysis, which enables its escape into the outside environment [19]. It has been observed 10 that Lp inhibits phagosome formation by *Paramecium* through a component of the type I 11 secretion system (a TolC-dependent mechanism) [20]. It is also reported that 12 Tetrahymena can support the survival of Lp [21, 22]. One way in which Tetrahymena 13 does this is by expelling environmental Lp-laden pellets, a finding indicating that it may 14 be a defense mechanism against Lp rather than a symbiotic role [22, 23]. However,

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Here, we assessed the interaction of Lp with two wild ciliates, *Anteglaucoma* CS11A and *Colpoda* E6, which we isolated from sewage and sink trap sludge, respectively, using a handmade maze device fitted with a 96-well crafting plate. We show that unlike *Tetrahymena* IB with pellet production, *Anteglaucoma* CS11A can be effectively killed by Lp infection, whereas *Colpoda* E6 is protected against Lp infection through cyst formation and the low sensitivity of trophozoites to Lp. Thus, these two ciliates are

studies on how Lp interacts with other ciliates are limited in number.

1 differentially susceptible to Lp infection.

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4 **METHODS**

Ciliates

- 6 Anteglaucoma CS11A (isolated from sewage) and Colpoda E6 (isolated from sink strap
- 7 sludge) were used in this study (See below for "Isolation methods for ciliates"). These
- 8 ciliates were maintained in Sonneborn's Paramecium medium (SPM, also known as
- 9 ATCC medium 802) consisting of cerophyl grass powder (Toyotama Healthy Foods Co.,
- 10 Ltd. Japan) in distilled water with live Enterobacter aerogenes (our laboratory stock) at
- 11 22°C [29]. Tetrahymena IB was also maintained in peptone-yeast extract glucose (PYG)
- broth, which contains peptone (BD, Franklin Lakes, NJ), yeast extract (BD), and glucose
- 13 (FUJIFILM Wako Pure Chemical Co., Tokyo, Japan) at 15°C, as described previously
- 14 [30].

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Bacteria

- 17 The Lp originally derived from the Philadelphia-1 progenitor strain [JR32 (wild-type
- strain) and its mutant $(\Delta icm/dot)$] were used in this study [31]. Both strains were
- 19 genetically modified to carry a GFP-expressing plasmid (pAM239GFP) for easy
- 20 visualization. The plasmid, which continuously expressed GFP was introduced to
- 21 LpJR32 and its mutant by electroporation. Also, mCherry-expressing-Escherichia coli
- 22 DH5α (Ec) was constructed by the introduction with a plasmid (pBBR122mCherry)

- 1 (Funakoshi, Tokyo, Japan), which continuously expressed mCherry. Lp and Ec were
- 2 cultured on B-CYE agar (Thermo Fisher Scientific, Waltham, MA) and LB agar plate
- 3 with or without 10µg/mL of chloramphenicol, respectively, at 37 °C for appropriate
- 4 period (2 days for Lp, one day for Ec). The mutant with the plasmid (pAM239GFP) was
- 5 kindly provided by Dr. Nagai (Gifu University, Japan).

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Isolation methods for ciliates

- 8 Ciliates were isolated according to the procedure with the maze device described in Fig.
- 9 S1. To make this device inexpensive and easy to use, we used commercially available
- 10 96-well culture plates in its operation. We simply broke the walls between the plate wells
- with a heated spatula (Fig. S2, a photo of the device that we used). Briefly, sink trap
- sludge samples (n=5) and manhole sewage samples (n=13) were collected from different
- places in our university, because Lp frequently colonizes in sinks or sewages [32-36]. The
- sample solutions, which were diluted in Page's amoeba saline (PAS) [37], were cultured
- with sterilized rice grains at room temperature for 2 weeks. When a large number of
- ciliates had been reared, a concentrated sample containing >1,000 ciliate cells was placed
- into the "sample addition hole" of the maze device, which was filled with PAS except for
- 18 the "sample addition hole", which was partitioned with a cotton plug. After removing the
- 19 cotton plug-containing partition, the ciliates were cultured for several days at room
- 20 temperature. Ciliates captured in the "capture hole" were carefully collected, cultured in
- 21 SPM medium (See above), and isolated by the limiting dilution method. The maze device
- could be reused approximately three times by performing intermittent sterilization (95°C,

1 10 min, twice).

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Direct sequencing and phylogenetic analysis

4 Total DNA was extracted from the isolated ciliates using the High Pure PCR Template Preparation Kit (Roche) according to the manufacturer's instructions. Extracted DNA 5 6 was amplified using Quick Taq HS Dye Mix (TOYOBO, Osaka, Japan) with an universal 7 primer set for each ciliate's 18S rDNA gene (P-SSU-342f; 5'-CTT TCG ATG GTA GTG 8 TAT TGG ACT AC-3', Medlin B; 5'-TGA TCC TTC TGC AGG TTC ACC TAC-3'), as 9 previously reported [38]. The amplified products were electrophoretically separated and 10 then extracted from the agarose gel using the FastGene Gel/PCR Extraction Kit (NIPPON 11 Genetics, Tokyo, Japan) according to the manufacturer's protocol, and then sequenced by 12 Fasmac (Kanagawa, Japan). The 18S rDNA phylogenetic tree containing the isolated 13 ciliates, which also contains representative ciliates, was constructed by the neighbor-joining method (bootstrap replication value, 500) with MEGA X [39]. 14 15 Representative sequences were obtained from the National Center for Biotechnology 16 Information (NCBI: (https://www.ncbi.nlm.nih.gov/). All the nucleotide sequence

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Scanning electron microscopy (SEM)

accession numbers are shown in Fig. 1.

The isolated ciliates were processed according to a previous method [40]. Briefly, they were washed in saline, fixed with 2.5% glutaraldehyde in phosphate-buffered saline for 2 h at room temperature, and then soaked in osmium tetroxide for 1 h at 4°C. The samples

- were then dehydrated in ethanol, freeze-dried, and coated with osmium using a plasma osmium coater. The samples were analyzed by SEM (Hitachi S-4800; Hitachi, Tokyo, Japan).

 Co-culture experiment with imaging

 Our co-culture system was constructed with green fluorescent protein (GFP) or mCherry-expressing Lp (LpJR32 or Δ*icm/dot*) or Ec (10⁷ CFU/mL) with or without each ciliate (*Anteglaucoma* CS11A, *Colpoda* E6, *Tetrahymena* IB) (10³ cell/ml) in 50 mL PAS or CA-1 medium (some experiment), a nutrient-rich medium used for preventing cyst
- or CA-1 medium (some experiment), a nutrient-rich medium used for preventing cyst formation [41]. The cultures were maintained at room temperature. Samples were collected immediately (1, 30, 120 min) or daily over 7 days, and used for extraction of DNA, ciliate's counts or imaging with a conventional fluorescence microscope (BIOREVO BZ-9000, Keyence, Osaka, Japan), an Olympus microscope (OKX41) with a fluorescence unit (U-RFLT50, Tokyo, Japan), or a confocal laser fluorescence

microscope (TCSSP5 TIRF, Wetzlar, Germany). The amount of amplicon targeting Lp

16 *mip* gene and each ciliate was then quantified (see below).

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Pellet production assay

The following experiments were conducted to confirm whether pellet production is a phenomenon peculiar to *Tetrahymena* IB. Briefly, each of the ciliates (*Anteglaucoma* CS11A, *Colpoda* E6, *Tetrahymena* IB) (10³ cell/ml) was mixed with GFP-expressing Lp (JR32 or Δ*icm/dot*) (10⁷CFU/ml) to 10ml of PAS solution in a 15ml-sterilized tube, and

- incubated upright without shaking for one day at 22°C. After incubation, the supernatant containing a 50 µl solution was carefully removed, and the remaining solution was fixed
- 3 with 4% paraformaldehyde. The presence or absence of pellets in the sample was then
- 4 confirmed under a fluorescence microscope (BIOREVO BZ-9000, Keyence).

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Determining Lp amounts using quantitative (q)PCR

7 The samples obtained from co-culture system were treated with freeze-thawing and then

used for DNA extraction [42]. DNA extraction was performed using a Instagene kit

(Bio-Rad, Hercules, CA, USA) according to the manufacture protocol. The Lp and ciliate

amounts in the co-culture system were quantified by CFX Connect (BioRad) with SYBR

Green (KOD SYBR qPCR Mix, TOYOBO) targeting the bacterial mip gene (Lg3: 5'-

12 GCT ACA GAC AAG GAT AAG TTG -3', Lg4: 5'- GTT TTG TAT GAC TTT AAT

13 TCA -3') [43]. The aliquot solution of mixed culture of Lp with ciliates was simply used

as the sample for qPCR, but not for culture. We adopted the qPCR instead of the CFU to

quantify Lp for two main reasons; firstly, it is very difficult to uniformly suspend Lp into

ciliates or expelled pellets; and secondly, bacteria, which is given as food during

subculture, may also grow together on an agar medium.

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Determining ciliate amounts using a cell counting chamber

- 20 Ciliate (trophozoite) numbers and cyst numbers were determined using 10µl of culture in
- 21 a disposable cell counting chamber (Thermo Fisher Scientific, Tokyo, Japan) by the
- 22 naked eye under a microscope.

2	Statistical analysis
3	Data obtained from the co-culture experiments were compared using the Bonferroni
4	correction. P-values of less than p <0.05 were considered significant. The presence of a
5	correlation for each ciliate's (trophozoite) number and cyst number was determined by
6	Pearson's correlation coefficient test. A correlation coefficient value of >0.3 or <-0.3
7	with a <i>P</i> -value of less than 0.05 was considered significant.
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9	Nucleotide sequence accession numbers
10	The 18S rDNA sequences from Anteglaucoma CS11A and Colpoda E6 described in this
11	study have been deposited in the DDBJ GenBank database
12	(https://www.ddbj.nig.ac.jp/index.html) under accession numbers LC573510 and
13	LC573511.
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16	RESULTS
17	Characterization of the ciliates newly isolated from sewage and sink
18	trap sludge by phylogenetic analysis and SEM observations
19	First, we successfully isolated two ciliates (Anteglaucoma CS11A and Colpoda E6).
20	Their taxonomic positions were determined by phylogenetic analysis of their 18S rDNA
21	sequences. As a result, two ciliates, CS11A and E6, were assigned to Anteglaucoma
22	(isolated from sewage) (accession number: LC573510) and Colpoda (isolated from sink

1 trap sludge) (accession number: LC573511), respectively. Both fell into distinct clades 2 and their locations in the phylogenic tree differed from those of Paramecium and 3 Tetrahymena, which are the only ciliates that have been studied for their interactions with 4 Lp [19-23] (Fig. 1). Our SEM observations revealed that unlike *Colpoda* E6 (Fig. 2A and B), Anteglaucoma CS11A possesses a large body with numerous thin cilia projecting 5 6 outwards from it (Fig. 2C and D). Although Tetrahymena IB, which has thicker cilia, has 7 been reported to expel pellets [23] (Fig. 2F, arrows), pellet production was not observed 8 in Colpoda E6 and Anteglaucoma CS11A. To confirm this, we assessed if the pellet 9 production with the packaging of Lp occurred as a phenomenon specific to *Tetrahymena* 10 IB. As a result, in contrast to the other ciliates, the presence of expelled pellets was only 11 observed in the remaining solution of the standing-mixed culture of Tetrahymena IB with 12 Lp at 24 h after co-culture (Fig. S3), indicating the pellet production with the packaging 13 of Lp is unique to Tetrahymena IB among ciliates used for this study. Thus, based on these morphological differences, the two ciliates we isolated may have different 14 15 interactions with Lp.

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Interactions of the newly isolated ciliates with Lp immediately (1 min,

18 30 min, 2 h) or after 24 h in co-culture

- 19 We next determined whether the wild ciliates (Anteglaucoma CS11A and Colpoda E6),
- and the control ciliate (Tetrahymena IB), interact with Lp (the LpJR32 wild-type strain or
- 21 the Δicm/dot mutant) using confocal laser microscopy. All three ciliate genera ingested
- 22 Lp and their features were distinctive at 24 h post-inoculation regardless of whether the

infections involved the wild-type or mutant Lp (Fig. 3). Furthermore, we assessed if these ciliates ingested Lp immediately (1 min, 30 min, 2 h) after co-culture through visualization of Lp in ciliates. As a result, the uptakes of Lp regardless of its mutant in the ciliates (Anteglaucoma CS11A and Colpoda E6) was observed from 1 min after co-culture, although there was a difference in the amount of uptake as the Anteglaucoma CS11A ingested a large amount (Fig. 4). Also, as expected, we confirmed that the uptake of these bacteria into *Tetrahymena* IB normally occurred at 30 min after co-culture (Fig. S4), and as previously reported, the ciliates packaged it in pellets at 24 h after co-culture (Fig. 3C and Fig. S3) [23], thereby confirming that our co-culture conditions were optimal. Notably, Anteglaucoma CS11A ingested a large amount of Lp compared with Colpoda E6. Together, the findings indicate that some interaction between ciliates and Lp occurred in our co-culture system.

Changes in Lp amounts in the co-culture system

We used qPCR to monitor changes in Lp amounts (the LpJR32 wild-type strain or the JR32Δ*icm/dot* mutant) in the ciliates using the co-culture system over 7 days. As a result, the Lp amount in the *Anteglaucoma* CS11A co-culture was constantly maintained over the 7-day period (Fig. 5A), but fell dramatically when the Δ*icm/dot* mutant was substituted for JR32 (Fig. 5B, "+*Anteglaucoma* CS11A"). In contrast with *Anteglaucoma* CS11A, the Lp amount was maintained in the co-culture system with *Colpoda* E6 independently of the virulence genes, but with a temporal decrease occurring one day after starting the culture (Fig. 5, "+*Colpoda* E6"). In addition, the amount of Lp was

- 1 maintained in *Tetrahymena* IB irrespective of the virulence genes (Fig. 5, "+*Tetrahymena*
- 2 IB"). Thus, the findings indicate that the Lp amounts in the co-culture system differed
- 3 between Anteglaucoma CS11A and Colpoda E6, although the change of bacterial
- 4 amount in the culture with the *Colpoda* was similar to those of *Tetrahymena* IB.

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Changes in ciliate amounts during co-culturing

- 7 Using cell counting, we also monitored changes in ciliate amounts in the co-culture
- 8 system with Lp (the LpJR32 wild-type strain or the Δicm/dot mutant) over 7 days.
- 9 Although uptake of LpJR32 and the mutant was separately confirmed, no ciliates were
- observed in the co-culture of *Anteglaucoma* CS11A and LpJR32 (Fig. 6A). While the
- 11 Anteglaucoma CS11A-Δicm/dot co-culture was maintained at levels similar to the
- bacteria alone control during the culture period (Fig. 6B, "+LpJR32 Δicm/dot"), the
- ciliates were completely killed in the presence of LpJR32 by day 2 (Fig. 6B, "+LpJR32").
- 14 Conversely, the morphology of *Colpoda* E6 was maintained even in the presence of
- LpJR32, a finding similar to that of the "ciliates alone" control (Fig. 7A). Although the
- ciliate amounts gradually decreased, there was no significant difference in Colpoda E6
- amounts among the co-cultures, regardless of the presence or absence of the virulence
- genes (Fig. 7B). Interestingly, when the number of ciliates decreased, the number of cysts
- tended to increase, although this finding was not statistically significant (Fig. 8). Also, to
- 20 confirm the sensitivity of Lp to trophozoites, Colpoda E6 was co-cultured with LpJR32
- 21 or Δ*icm/dot* mutant in a nutrient-rich medium, CA-1, which inhibits cyst formation (See
- 22 the Methods), and the number of ciliates was calculated under the condition suppressive

1 of cyst formation. However, the trophozoites of ciliates grew well regardless of the 2 presence or absence of Lp during the culture period (Fig. S5), suggesting that along with 3 resistance to Lp by cyst formation, the low sensitivity of trophozoites to Lp also may 4 contribute to their resistance to Lp. In addition, the Tetrahymena IB amounts were maintained regardless of the presence or absence of Lp during the culture period (Fig. S6). 5 6 Thus, we conclude that unlike *Tetrahymena* IB, which can resist Lp by enwrapping it and 7 expelling it in pellets, Anteglaucoma CS11A can be completely killed by Lp infection, 8 whereas Colpoda E6 resists Lp infection by forming cysts with the low sensitivity of 9 trophozoites to Lp. Therefore, the two ciliates we isolated have distinctive to Lp 10 infection.

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DISCUSSION

As mentioned above, Lp is ubiquitous in a wide range of natural environments such as soil or pond water [1-3]. Lp interacts with various protozoa including amoebae and ciliates, many of which are unknown species [26-28]. More importantly, Lp propagating through such interactions can create the ideal conditions for legionellosis in humans [8-10]. However, few studies have been published on the interactions of Lp with ciliates, and those that have been published are limited to *Paramecium* and *Tetrahymena* [16-23]. Therefore, we investigated the interaction of Lp with two wild ciliates, Anteglaucoma CS11A and Colpoda E6, which we isolated from sewage and sink trap sludge, respectively, using a handmade maze device with a 96-well crafting plate. Our data

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1 clearly shows that unlike Tetrahymena IB, Anteglaucoma CS11A is killed by Lp 2 infection, whereas Colpoda E6 resists Lp infection through cyst formation and the low 3 sensitivity of trophozoites to Lp. 4 Ciliates are not easy to isolate because they move around actively [24, 25]. Therefore, we 5 6 created a simple handmade maze device that can discriminate differences in the moving 7 capacities of other microbes such as motile bacteria. Meanwhile, there are some limitations in using the maze device. First, intermittent sterilization (<95°C, 10 mins, 8 9 twice) of the device deforms it making it unusable after three uses. Second, toxic gas is 10 generated when making the device, so it is necessary to prepare it in a well-ventilated 11 place. Third, because the wall is broken manually, the groove width will subtly differ in 12 each device. Fourth, the separation cannot be done unless the number of ciliates added exceeds $> 10^3$ cells, approximately. 13 14 15 We isolated two morphologically distinct ciliates using this device and assigned them as 16 Anteglaucoma CS11A and Colpoda E6. These ciliates differ from Paramecium and 17 Tetrahymena, which have been extensively studied for their interactions with Lp. 18 Anteglaucoma, which was originally isolated from a farmland pond in China, has been 19 reported as a new ciliate genus in the *Glaucomidae* family, and is recognized by its hectic 20 jerking motion [34]. Anteglaucoma CS11A also has an outstanding athletic ability, 21 possibly affording it a predation advantage. In contrast, a large number of studies have

been published over a long time period on the ecological and biological properties of

1 Colpoda, and these support it as a ubiquitous and prosperous genus among known ciliates

2 [45-47]. With its ubiquitous nature, Colpoda has been used as an indicator of

environmental pollution (e.g., heavy metal pollution) [48, 49]. Because there are

currently no studies on the interactions of these ciliates with Lp, this study is the first to

discover their interactions.

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We found that Anteglaucoma CS11A can be killed by LpJR32 depending on whether or not the icm/dot virulence genes are present. Although it is well known that Tetrahymena ensures bacterial survival by expelling any bacteria (e.g., Salmonella, Mycobacterium, and Lp) packaged in its fecal pellets as a way for ciliates to discard preys [23, 50, 51], no pellet production but the accumulation of Lp at earlier interaction was seen in the Anteglaucoma CS11A-co-cultures, suggesting that the defense mechanism of the ciliates against Lp may be lacking. Also, the exact reason why the Lp infections caused ciliate death in our experiments is not known, but it is possible that some Lp effectors (e.g., those which shut down endoplasmic reticulum-mitochondria interactions), may be involved in this type of cell death [52]. Meanwhile, it was not clear that Lp can replicate inside the ciliates. Thus, Anteglaucoma CS11A could be used as a biological indicator to verify the pollution status of Lp in water supplies or hot springs. We also found that Colpoda E6 can be maintained in the presence of Lp independently of the icm/dot virulence genes through cyst formation. Stimulation of cyst formation is not a sign of Lp infection, because cyst formation is observed even when ciliates are cultured alone. Furthermore, because we observed that despite the presence of Lp, the ciliates mostly consisting of trophozoites

Acknowledgments

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1	grew well in a nutrient-rich condition that inhibits cyst formation (See Fig. S5), it is
2	possible that along with the transitions to cysts in low-nutrient environments, the low
3	sensitivity of trophozoites to Lp may contribute to their resistance to Lp. In addition, it
4	has been reported that starvation can induce production of the cyst coat protein
5	responsible for cyst formation in Colpoda steinii [53].
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7	In conclusion, we successfully isolated two ciliates (Anteglaucoma CS11A and Colpoda
8	E6) from sewage and sink trap sludge, respectively, using a handmade maze device. Each
9	one differs in its susceptibility to Lp infection. Contrasting with Tetrahymena IB with
10	pellet production, Anteglaucoma CS11A can be killed by Lp infection, whereas Colpoda
11	E6 resists Lp infection through cyst formation and the low sensitivity of trophozoites to
12	Lp (Fig. 9). Our new identification of two ciliates with different susceptibilities to Lp
13	infection provides useful public health information towards the control of human
14	pathogenic Lp.
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6 Author contribution

- 7 HY and TO designed the study. AK and TO carried out the experiments. HY, AK, and TO
- 8 analyzed the data. SN carried out imaging analysis. MM provided the Lp strains. JT
- 9 edited the manuscript and made critical suggestions. HY wrote the manuscript.

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11 Conflicts of interest

12 The authors declare that there are no conflicts of interest.

References

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- 2 1. Fields BS, Benson RF, Besser RE. Legionella and Legionnaires' disease: 25 years of
- 3 investigation. Clin Microbiol Rev 2002;15:506-526.
- 4 2. Boamah DK, Zhou G, Ensminger AW, O'Connor TJ. From Many Hosts, One
- 5 Accidental Pathogen: The Diverse Protozoan Hosts of Legionella. Front Cell Infect
- 6 *Microbiol* 2017;7:477.
- 7 3. Caicedo C, Rosenwinkel KH, Exner M, Verstraete W, Suchenwirth R,
- 8 Hartemann P, Nogueira R. Legionella occurrence in municipal and industrial
- 9 wastewater treatment plants and risks of reclaimed wastewater reuse: Review. Water
- 10 Res 2019;149:21-34.
- 4. Swart AL, Harrison CF, Eichinger L, Steinert M, Hilbi H. Acanthamoeba and
- 12 Dictyostelium as Cellular Models for Legionella Infection. Front Cell Infect
- 13 *Microbiol* 2018;8:61.
- 14 5. **Qiu J, Luo ZQ.** *Legionella* and *Coxiella* effectors: strength in diversity and activity.
- 15 Nat Rev Microbiol 2017;15:591-605.
- 16 6. Finsel I, Hilbi H. Formation of a pathogen vacuole according to Legionella
- 17 pneumophila: how to kill one bird with many stones. Cell Microbiol
- 18 2015;17:935-950.
- 19 7. Isaac DT, Isberg R. Master manipulators: an update on Legionella pneumophila
- 20 Icm/Dot translocated substrates and their host targets. Future Microbiol
- 21 2014;9:343-359.
- 22 8. Principe L, Tomao P, Visca P. Legionellosis in the occupational setting. Environ

- 1 Res 2017;152:485-495.
- 2 9. Parr A, Whitney EA, Berkelman RL. Legionellosis on the Rise: A Review of
- 3 Guidelines for Prevention in the United States. J Public Health Manag Pract
- 4 2015;21:E17-26.
- 5 10. Cunha BA, Burillo A, Bouza E. Legionnaires' disease. *Lancet* 2016;387:376-385.
- 6 11. Leoni E, Catalani F, Marini S, Dallolio L. Legionellosis Associated with
- 7 Recreational Waters: A Systematic Review of Cases and Outbreaks in Swimming
- 8 Pools, Spa Pools, and Similar Environments. Int J Environ Res Public Health
- 9 2018;15:1612.
- 10 12. Herwaldt LA, Marra AR. Legionella: a reemerging pathogen. Curr Opin Infect Dis
- 11 2018;31:325-333.
- 12 13. Barker KA, Whitney EA, Blake S, Berkelman RL. A Review of Guidelines for the
- Primary Prevention of Legionellosis in Long-Term Care Facilities. J Am Med Dir
- 14 *Assoc* 2015;16:832-6.
- 15 14. Barbaree JM, Fields BS, Feeley JC, Gorman GW, Martin WT. Isolation of
- protozoa from water associated with a legionellosis outbreak and demonstration of
- intracellular multiplication of Legionella pneumophila. Appl Environ Microbiol
- 18 1986;51:422-424.
- 19 15. King CH, Shotts EB Jr, Wooley RE, Porter KG. Survival of coliforms and
- 20 bacterial pathogens within protozoa during chlorination. Appl Environ Microbiol
- 21 1988;54:3023-3033.
- 22 16. Yli-Pirilä T, Kusnetsov J, Haatainen S, Hänninen M, Jalava P, Reiman M, Seuri

- 1 M, Hirvonen MR, Nevalainen A. Amoebae and other protozoa in material samples
- from moisture-damaged buildings. *Environ Res* 2004;96:250-256.
- 3 17. Tsao HF, Scheikl U, Herbold C, Indra A, Walochnik J, Horn M. The cooling
- 4 tower water microbiota: Seasonal dynamics and co-occurrence of bacterial and protist
- 5 phylotypes. *Water Res* 2019;159:464-479.
- 6 18. Paranjape K, Bédard É, Shetty D, Hu M, Choon FCP, Prévost M, Faucher SP.
- 7 Unravelling the importance of the eukaryotic and bacterial communities and their
- 8 relationship with *Legionella* spp. ecology in cooling towers: a complex network.
- 9 Microbiome 2020;8:157.
- 10 19. Watanabe K, Nakao R, Fujishima M, Tachibana M, Shimizu T, Watarai M.
- 11 Ciliate Paramecium is a natural reservoir of Legionella pneumophila. Sci Rep
- 12 2016;6:24322.
- 13 20. Nishida T, Hara N, Watanabe K, Shimizu T, Fujishima M, Watarai M. Crucial
- Role of Legionella pneumophila TolC in the Inhibition of Cellular Trafficking in the
- 15 Protistan Host *Paramecium tetraurelia*. Front Microbiol 2018;9:800.
- 16 21. Berk SG, Garduño RA. The tetrahymena and acanthamoeba model systems.
- 17 *Methods Mol Biol* 2013;954:393-416.
- 18 22. Fields BS, Shotts EB Jr, Feeley JC, Gorman GW, Martin WT. Proliferation of
- 19 Legionella pneumophila as an intracellular parasite of the ciliated protozoan
- 20 Tetrahymena pyriformis. Appl Environ Microbiol 1984;47:467-471.
- 21 23. Hojo F, Sato D, Matsuo J, Miyake M, Nakamura S, Kunichika M, Hayashi Y,
- 22 Yoshida M, Takahashi K, Takemura H, Kamiya S, Yamaguchi H. Ciliates expel

- 1 environmental Legionella-laden pellets to stockpile food. Appl Environ Microbiol
- 2 2021;78:5247-5257.
- 3 24. Vincensini L, Blisnick T, Bastin P. 1001 model organisms to study cilia and flagella.
- 4 *Biol Cell* 2011;103:109-130.
- 5 25. Finlay BJ. The global diversity of protozoa and other small species. Int J Parasitol
- 6 1998;28:29-48.
- 7 26. Zhao F, Filker S, Xu K, Huang P, Zheng S. Patterns and Drivers of Vertical
- 8 Distribution of the Ciliate Community from the Surface to the Abyssopelagic Zone in
- 9 the Western Pacific Ocean. Front Microbiol 2017;8:2559.
- 10 27. Weisse T. Functional diversity of aquatic ciliates. Eur J Protistol
- 11 2017;61(PtB):331-358.
- 12 28. Vďačný P. Integrative taxonomy of ciliates: Assessment of molecular phylogenetic
- 13 content and morphological homology testing. *Eur J Protistol* 2017;61(Pt B):388-398.
- 14 29. Cassidy-Hanley DM. Tetrahymena in the laboratory: strain resources, culture,
- maintenance, and storage. *Methods Cell Biol* 2012;109:237-276.
- 16 30. Matsuo J, Oguri S, Nakamura S, Hanawa T, Fukumoto T, Hayashi Y,
- 17 Kawaguchi K, Mizutani Y, Yao T, Akizawa K, Suzuki H, Simizu C, Matsuno K,
- 18 Kamiya S, Yamaguchi H. Ciliates rapidly enhance the frequency of conjugation
- between Escherichia coli strains through bacterial accumulation in vesicles. Res
- 20 *Microbiol* 2010;161:711-719.
- 21 31. Rao C, Benhabib H, Ensminger AW. Phylogenetic reconstruction of the Legionella
- 22 pneumophila Philadelphia-1 laboratory strains through comparative genomics. PLoS

- 1 *One* 2013;8(5):e64129.
- 2 32. Vicente D, Marimón JM, Lanzeta I, Martin T, Cilla G. Fatal Case of Nosocomial
- 3 Legionella pneumophila Pneumonia, Spain, 2018. Emerg Infect Dis
- 4 2019;25:2097-2099.
- 5 33. Fleres G, Couto N, Lokate M, van der Sluis LWM, Ginevra C, Jarraud S, Deurenberg
- 6 RH, Rossen JW, García-Cobos S, Friedrich AW. Detection of Legionella Anisa in
- Water from Hospital Dental Chair Units and Molecular Characterization by
- 8 Whole-Genome Sequencing. *Microorganisms* 2018;6:71.
- 9 34. Yamamoto N, Kubota T, Tateyama M, Koide M, Nakasone C, Tohyama M,
- 10 Shinzato T, Higa F, Kusano N, Kawakami K, Saito A. Isolation of Legionella
- anisa from multiple sites of a hospital water system: the eradication of Legionella
- contamination. J Infect Chemother 2003;9:122-125.
- 13 35. Bajrai LH, Azhar EI, Yasir M, Jardot P, Barrassi L, Raoult D, La Scola B,
- Pagnier I. Legionella saoudiensis sp. nov., isolated from a sewage water sample. Int
- 15 *J Syst Evol Microbiol* 2016;66:4367-4371.
- 16 36. Palmer CJ, Tsai YL, Paszko-Kolva C, Mayer C, Sangermano LR. Detection of
- 17 Legionella species in sewage and ocean water by polymerase chain reaction, direct
- fluorescent-antibody, and plate culture methods. Appl Environ Microbiol
- 19 1993;59:3618-3624.
- 20 37. Page FC. A New Key to Freshwater and Soil Gymnamoebae
- 21 (ISBN-10:18711105021) Freshwater Biological Association, Ambleside, UK (1988)
- 22 122 pp.

- 1 38. Karnati SKR, Yu Z, Sylvester JT, Dehority BA, Morrison M, Firkins JL.
- 2 Technical note: Specific PCR amplification of protozoal 18S rDNA sequences from
- 3 DNA extracted from ruminal samples of cows. *J Anim Sci* 2003;81:812-815.
- 4 39. Kumar S, Stecher G, Li M, Knyaz C, Tamura K. MEGA X: Molecular
- 5 Evolutionary Genetics Analysis across Computing. Mol Biol Evol
- 6 2018;35:1547-1549.
- 7 40. Okude M, Matsuo J, Nakamura S, Kawaguchi K, Hayashi Y, Sakai H, Yoshida
- 8 M, Takahashi K, Yamaguchi H. Environmental chlamydiae alter the growth speed
- 9 and motility of host acanthamoebae. *Microbes Environ* 2012;27: 423-429.
- 10 41. Shi T, Shi X, Lu F, Yu F, Fu Y, Hu X. A Colpoda aspera isolate from animal
- faeces: In vitro cultivation and identification. *Eur J Protistol* 2014;50:33-39.
- 12 42. Hayden RT, Uhl JR, Qian X, Hopkins MK, Aubry MC, Limper AH, Lloyd RV,
- 13 Cockerill FR. Direct detection of Legionella species from bronchoalveolar lavage
- and open lung biopsy specimens: comparison of LightCycler PCR, in situ
- 15 hybridization, direct fluorescence antigen detection, and culture. J Clin Microbiol
- 16 2001;39:2618-2626.
- 17 43. Mahbubani MH, Bej AK, Miller R, Haff L, DiCesare J, Atlas RM. Detection of
- Legionella with polymerase chain reaction and gene probe methods. Mol Cell Probes
- 19 1990;4:175-187.
- 44. Pan X, Shi Z, Wang C, Bourland WA, Chen Y, Song W. Molecular Phylogeny and
- Taxonomy of a New Freshwater Hymenostomatid from Northeastern China, with the
- Establishment of a New Genus Anteglaucoma gen. n. (Protista, Ciliophora,

- Oligohymenophorea). *J Eukaryot Microbiol* 2016;64:564-572.
- 2 45. Cutler DW, Crump LM. The Rate of Reproduction in Artificial Culture of
- 3 *Colpidium colpoda. Biochem J* 1923;17:174-86.
- 4 46. Foissner W, Stoeck T, Agatha S, Dunthorn M. Intraclass evolution and
- 5 classification of the Colpodea (Ciliophora). J Eukaryot Microbiol 2011;58:397-415.
- 6 47. Foissner W, Bourland WA, Wolf KW, Stoeck T, Dunthorn M. New SSU-rDNA
- 7 sequences for eleven colpodeans (Ciliophora, Colpodea) and description of
- 8 Apocyrtolophosis nov. gen. *Eur J Protistol* 2014;50:40-46.
- 9 48. Madoni P, Romeo MG. Acute toxicity of heavy metals towards freshwater ciliated
- 10 protists. *Environ Pollut* 2006;141:1-7.
- 49. Janssen MP, Oosterhoff C, Heijmans GJ, Van der Voet H. The toxicity of metal
- salts and the population growth of the ciliate *Colpoda cucculus*. *Bull Environ Contam*
- 13 *Toxicol* 1995;54:597-605.
- 14 50. Rehfuss MY, Parker CT, Brandl MT. Salmonella transcriptional signature in
- 15 Tetrahymena phagosomes and role of acid tolerance in passage through the protist.
- 16 *ISME J* 2011;5:262-273.
- 17 51. Denoncourt AM, Paquet VE, Charette SJ. Packaging of Mycobacterium
- smegmatis bacteria into fecal pellets by the ciliate Tetrahymena pyriformis. FEMS
- 19 *Microbiol Lett* 2017;364(23).
- 20 52. Arasaki K, Mikami Y, Shames SR, Inoue H, Wakana Y, Tagaya M. Legionella
- 21 effector Lpg1137 shuts down ER-mitochondria communication through cleavage of
- 22 syntaxin 17. *Nat Commun* 2017;15:15406.

1	53. Tibbs J. Induction of cyst coat protein synthesis by starvation in the ciliate <i>Colpoda</i>
2	steinii. Biochem J 1971;124:419-426.
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1 Figure legends 2 3 Fig. 1. 18S rDNA-based phylogenetic tree showing the location of Anteglaucoma CS11A 4 and Colpoda E6 among representative ciliates. The red and blue circles show the 5 locations of Anteglaucoma CS11A and Colpoda E6, respectively. Small numbers in the 6 phylogenetic tree indicate local bootstrap probability. Numbers after species names are 7 accession numbers in the NCBI database. 8 9 Fig. 2. Representative SEM images showing the morphological features of three ciliates. 10 A and B, Colpoda E6. C and D, Anteglaucoma CS11A. E and F, Tetrahymena IB. Arrows, 11 expelled fecal pellets. Images in small dashed boxes (A and C) in the left-hand panels are 12 enlarged in the right-hand panels (B and D). White bars show scales for 5µm (B and D), 13 10μm (E and F), and 20μm (A and C). 14

Fig. 3. Representative confocal laser microscopic images showing three the ciliates from
 this study with distinct features at 24 h after co-culture with GFP-expressing Lp. GFP-Lp,
 GFP-expressing L. pneumophila. A, Images show Anteglaucoma CS11A co-cultured
 with JR32 (up) and JR32Δicm/dot (down). B. Images show Colpoda E6 co-cultured with

JR32 (up) and JR32 $\Delta icm/dot$ (down). C. Images show *Tetrahymena* IB co-cultured with JR32 (up) and JR32 $\Delta icm/dot$ (down). Each image shows one of the observed 3-5 fields of

view. White bars show scales for $10\mu m$.

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1 Fig. 4. Representative fluorescence microscopic images showing the uptake of Lp into 2 each of the ciliates [Anteglaucoma CS11A (A) and Colpoda E6 (B)] immediately (1, 30, 3 120 min) after co-culture with mCherry-expressing Lp (the wild-type LpJR32 strain and 4 the $\Delta icm/dot$ mutant) or mCherry-expressing Ec. mCherry-Lp, mCherry-expressing L. pneumophila. mCherry-expressing Ec, mCherry-expressing E. coli. Each image shows 5 6 one of the observed 3-5 fields of view. Black bars show a scale of 10µm. 7 8 Fig. 5. Changes in the amount of amplicon targeting Lp mip gene in the co-culture system 9 of Lp (JR32 strain and Δicm/dot mutant) with Anteglaucoma CS11A and Colpoda E6. 10 The co-culture system was maintained for 7 days. "Day 0" means immediately after 11 co-culture. A. Amount of amplicon targeting Lp *mip* gene in the co-culture with LpJR32. 12 B. Amount of amplicon targeting Lp mip gene in the co-culture system with the 13 LpJR32 $\Delta icm/dot$ mutant. The data (average \pm SD) independently obtained from four experiments were compared using the Bonferroni correction. *, p<0.05 vs. "Day0" of 14 15 each group. 16 17 Fig. 6. Changes in the Anteglaucoma CS11A cell amounts in the co-culture system of Lp 18 (JR32 and Δicm/dot mutant) with Anteglaucoma CS11A. The co-culture system was 19 maintained for 7 days. "Day 0" means 2 h after co-culture. A. Representative images 20 showing the interaction of the ciliate with LpJR32 or $\Delta icm/dot$ during the culture period. 21 Because the ciliates could not be seen at 2 days after co-culture, no images were obtained 22 after the second day. Each image shows one of the observed 3-5 fields of view. Black bars

1 show a scale of 10µm. B. Changes in Anteglaucoma CS11A cell amounts in the 2 co-culture system. The data (average ± SD) independently obtained from three 3 experiments were compared using the Bonferroni correction. *, p<0.05 vs. "Ciliates 4 alone". 5 6 Fig. 7. Changes in *Colpoda* E6 cell amounts in the co-culture system of Lp (JR32 and the 7 Δicm/dot mutant) with Colpoda E6. The co-culture system was maintained for 7 days. "Day 0" means 2 h after co-culture. A. Representative images showing the interaction of 8 9 the ciliate with LpJR32 or mutant LpJR32 during the culture period. Each image shows 10 one of the observed 3-5 fields of view. Black bars show a scale of 10µm. B. Changes in 11 Colpoda E6 cell amounts in the co-culture system. The data (average \pm SD) were 12 independently obtained from three experiments. 13 14 Fig. 8. Changes in the cyst amounts in the co-culture system of Colpoda E6 with Lp 15 (JR32 and $\Delta icm/dot$ mutant). The upper images show the appearances of the cysts from 16 each of the cultures at 7 days post-co-culture. The middle graphs show the changes in cyst 17 numbers over time. "Day 0" means 2 h after co-culture. Black bars show a scale of 10µm. 18 Bottom graphs show the correlations between the ciliate (trophozoites) numbers and cyst 19 numbers in the cultures with or without LpJR32 or the LpJR32\Delta icm/dot mutant. The data 20 for "Ciliate (trophozoites) numbers" used for correlation in this experiment were the same as that for Fig. 7B. "r" shows the correlation values. The data (average \pm SD) were 21 22 independently obtained from three experiments.

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2	Fig. 9. Results summary. Upper illustration shows the fate of Anteglaucoma CS11A after
3	interacting with LpJR32. Whether or not this ciliate dies depends on the presence/absence
4	of the icm/dot virulence genes. Middle illustration shows the fate of Colpoda E6 after
5	interacting with LpJR32. Whether or not this ciliate survives depends on avoiding Lp
6	infection by forming cysts and the low sensitivity of trophozoites to Lp. Bottom
7	illustration shows the fate of Tetrahymena IB after interacting with LpJR32. Whether of
8	not this ciliate survives depends on enwrapping Lp in pellets and excreting it outside of its
9	body, as reported previously [22, 23].
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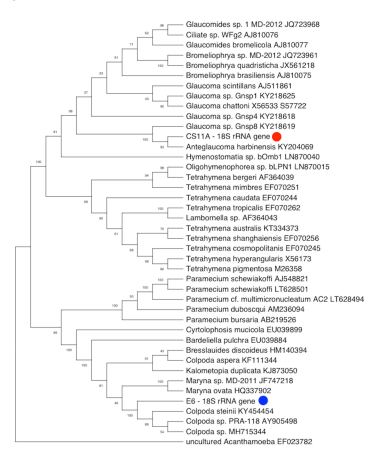
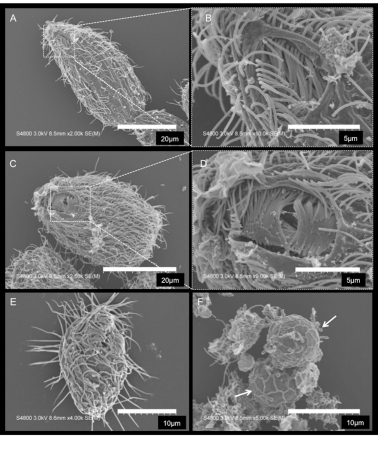
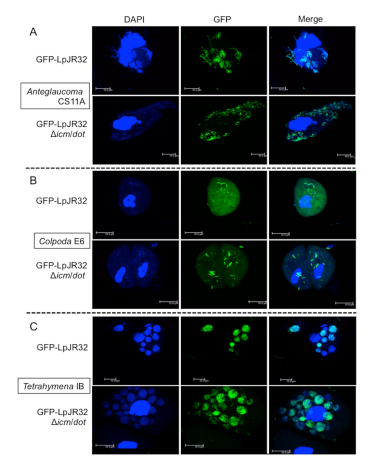
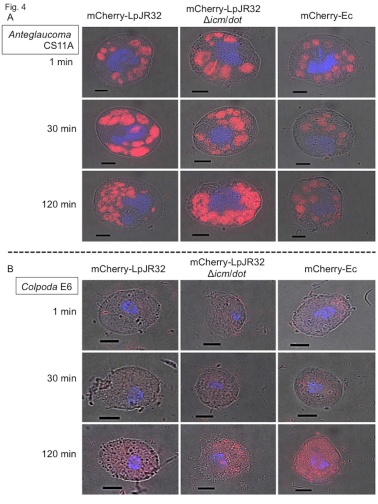
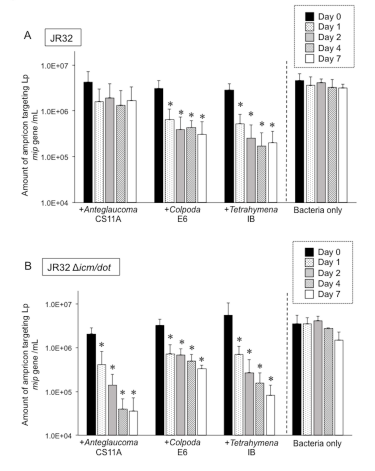


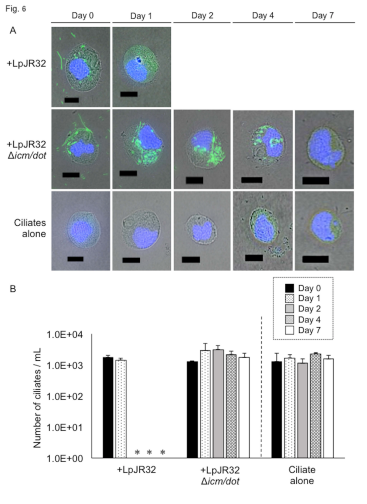
Fig. 2











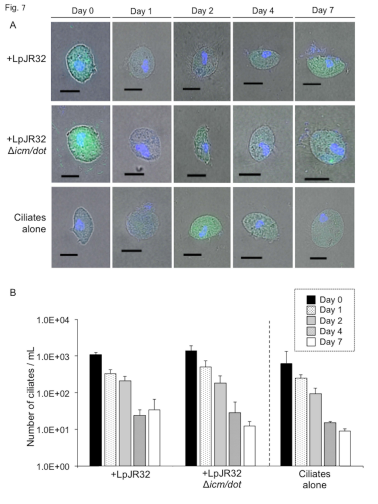


Fig. 8

