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1	Tetrahymena promotes interactive transfer of carbapenemase gene
2	encoded in plasmid between fecal Escherichia coli and environmental
3	Aeromonas caviae
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6	Running title: Ciliates promote bacterial gene transfer
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#### **ABSTRACT**

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Ciliates (*Tetrahymena*) can facilitate plasmid transfer among *Escherichia coli* or from *E*. 34 35 coli to Salmonella Enteritidis via vesicle accumulation. In this study, we assessed 36 whether ciliates promote the interactive transfer of plasmids encoding  $bla_{IMP-1}$  between fecal E. coli and environmental Aeromonas caviae. Both bacteria were mixed with or 37 38 without ciliates and incubated overnight at 30°C. The frequency of plasmid-acquired 39 bacteria was estimated by colony counts using an agar plate containing ceftazidim (CAZ) 40 followed by minimum inhibitory concentration (MIC) assessment. Cultures containing 41 ciliates interactively transferred the plasmid between E. coli and Aeromonas with a frequency of 10<sup>-4</sup> to 10<sup>-5</sup>. All plasmid-acquired bacteria showed a MIC against CAZ of 42 43 >128µg/ml, and the plasmid transfer was confirmed by polymerase chain reaction (PCR) 44 amplification of the bla<sub>IMP-1</sub> gene. Fluorescent observation showed both bacteria accumulated in the same vesicle and that transwell sequestering significantly decreased 45 the transfer frequency. Although ciliates preferentially ingested E. coli rather than A. 46 47 caviae, both bacteria were co-localized into same vesicles of ciliates, indicating that their meeting was associated with the gene transfer. Thus, ciliates interactively promote 48

49	plasmid transfer between E. coli and A. caviae. The results of this study will facilitate
50	control of the spread of multiple-antibiotic resistant bacteria.
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52	Keywords: Aeromonas caviae; ciliates; gene transfer; Escherichia coli; one health
52 53	<b>Keywords:</b> Aeromonas caviae; ciliates; gene transfer; Escherichia coli; one health

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#### **INTRODUCTION**

56 There is currently a pandemic of multidrug-resistant (MDR) Enterobacteriaceae 57 producing extended spectrum β-lactamase (ESBL), such as carbapenemase encoded by 58 plasmids such as  $bla_{NDM}$  or  $bla_{IMP}$ , which resist carbapenem, and their continuing spread 59 is a growing global concern that is becoming an immeasurable threat to hospitals and other healthcare-associated facilities (1-5). Additionally, lack of comprehensive 60 management protocols encompassing livestock, circulating food, public hygiene 62 practices, and monitoring of natural environments has resulted in failure to control 63 dissemination of ESBL-producing bacteria to or from humans through fecal waste, contaminated water, and processed meat products (6-9). Therefore, a "One Health" 64 65 approach requiring the combined efforts of physicians, veterinarians, epidemiologists, 66 public health workers and urban planners worldwide has been proposed to control the emergence of MDR bacteria (10). 67 68 Ciliate protozoa (Tetrahymena) ubiquitously inhabit animal rumens or natural 69 environments such as rivers and ponds in most of the world (11-13). The organisms are at the top of the complex-food chain in the microbial community, where they act as grazers 70

71 that feed on bacteria (14). Moreover, it has been reported that ciliates facilitate cell-to-cell contact among bacteria captured in their vesicles by expelling vesicles with 72 73 undigested-live bacteria to outside environments, thereby contributing to gene transfer 74 (15, 16). Indeed, we previously demonstrated that ciliates strongly promote transfer of the TnphoA, which encodes alkaline phosphatase, in Escherichia coli (SM10  $\lambda pir^+$  pRT733<sup>+</sup> 75 76 with mobRP4) to a clinical isolate of E. coli via vesicle accumulation (17). We also found that ciliates can facilitate the transfer of plasmids encoding bla<sub>NDM-5</sub> between E. coli 77 78 strains or from E. coli to Salmonella Enteritidis (18, 19). These findings suggest that 79 ciliates support effective plasmid transfer among bacteria in natural environments and are 80 involved in their circulation between humans and natural environments. Alternatively, we 81 speculated that ciliates can be a hotspot in natural environments, facilitating bacterial plasmid transfer from human pathogens to environmental bacteria, responsible for 82 83 spreading plasmid into the environments, presumably coming the plasmid back to human pathogenic bacteria. Additionally, Aeromonas caviae could play a role as both the 84 85 recipient and donor of plasmids to human pathogenic bacteria because they are 86 environmental bacteria that inhabit rivers and coastal areas as a fish pathogen, and are

therefore closely connected to human life. (20). However, it is not yet known if ciliates
can promote the transfer of plasmids between fecal *E. coli* and environmental organisms
such as *Aeromonas caviae*.

In this study, we assessed whether ciliates promoted the interactive transfer of plasmids encoding  $bla_{IMP-1}$  between fecal E. coli and environmental A. caviae. We present here the first evidence of the interactive transfer of this plasmid between E. coli and A. caviae in two distinct Tetrahymena.

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#### MATERIALS AND METHODS

# 96 Ciliates and bacteria used for this study

Table 1 shows a list of the traits and sources of ciliates and bacteria used in this study. Two
distinct ciliate protozoa, *Tetrahymena thermophila* (Tth; kindly provided by Dr. Sugai of
lbaraki University, Japan) and *Tetrahymena* sp. (Tsp; gifted from Dr. Tukii of Hosei
University, Japan) were used in this study. Ciliates were maintained in peptone-yeast
extract glucose broth (PYG) containing 0.75% peptone (Difco), 0.75% yeast extract
(Difco), and 1.5% glucose (Wako) at 30°C as previously described (21). *Citrobacter* 

freundii carrying a plasmid encoding  $bla_{IMP-1}$  was originally isolated from Hokkaido University Hospital and Aeromonas caviae no. 86 was kindly gifted from Professor Y. Tamura, Rakuno Gakuen University. Other bacteria (E. coli J53, E. coli DH5α, E. coli ATCC 29213, and Staphylococcus aureus ATCC 25922) were purchased from the American Type Culture Collection (ATCC). E. coli TC170328 strain carrying the plasmid encoding  $bla_{IMP-1}$  was experimentally established by the broth-mating method with E. coli J53 and C. freundii as previously described (22). Moreover, green fluorescent protein (GFP)-expressing E. coli DH5α was constructed according to a previous study (23), then used for imaging analysis (see below).

#### Mixed culture and assessment of gene transfer frequency

The frequency of bacterial gene transfer through protozoa was assessed as previously described (17). Briefly, equal amounts (approximately 10<sup>9</sup> colony-forming units [CFU]) of both donor [ceftazidime (CAZ)-resistant] (0.5 ml) and recipient bacteria (0.5 ml) were mixed with or without 10<sup>6</sup> cells of ciliates (1.0 ml) in Page's amoeba saline (PAS) (24), then incubated for 24 h at 30°C. In addition, an experiment with a transwell (0.4 μm pore

size) was conducted to separate ciliates from donor and recipient bacteria. After incubation, the solution was spread on Trypticase-soy agar (Nissui Pharmaceutical, Japan) containing 50  $\mu$ g/ml of CAZ and 50  $\mu$ g/ml of irgasan (Sigma-Aldrich) for *A. caviae*-derived plasmid-acquired bacteria or Luria-Bertani (LB) agar (Nacalai tesque) containing 50  $\mu$ g/ml of CAZ and 100  $\mu$ g/ml of sodium azide (Wako) for *E. coli* J53-derived plasmid-acquired bacteria. The remaining solution was used for DNA extraction for polymerase chain reaction (PCR). The gene transfer frequency was calculated by the following equation: frequency of transfer events = the number of plasmid-acquired bacterial colonies/total colony numbers (donor plus recipient).

## **DNA** extraction and PCR

DNA was extracted from a single-colony culture of the bacteria (donor, recipient and plasmid-acquired bacteria) by boiling as previously described (17), after which 2  $\mu$ l of boiling template was used for PCR with primers against  $bla_{\rm IMP-1}$  (25). The replicon type of the  $bla_{\rm IMP-1}$ -encoding plasmid was also determined by PCR (26).

135	Imaging
136	To confirm co-localization of <i>E. coli</i> and <i>A. caviae</i> in ciliate vesicles, 24-h-mixed cultures
137	comprising ciliates (T. thermophile or T. pyriformis), GFP-expressing E. coli DH5 $\alpha$
138	(fluorescence color: green,) (22) and vital-stained <i>A. caviae</i> no. 86 (fluorescence color:
139	red) were fixed with 4% formalin and then analyzed by fluorescence microscopy
140	(BioZero, Keyence). Vital staining of bacteria was performed using a PKH-26 labeling
141	kit (PKH-26GL, Sigma) according to the manufacturer's protocols.
142	
143	Plasmid-acquired bacterial determination
144	Bacterial identification was accomplished using a Matrix Assisted Laser
145	Desorption/Ionization (MALDI) Biotyper system (Bruker Daltonics) according to the
146	manufacturer's instructions.
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148	Antimicrobial susceptibility test
149	The minimum inhibitory concentration (MIC) of CAZ against donors, recipients, and
150	plasmid-acquired bacteria in this study was determined by the agar-dilution method (27).

151 E. coli ATCC 25922 and S. aureus ATCC 29213 were used as quality control strains.

## **Grazing assay**

Briefly, equal amounts (approximately 10<sup>9</sup> colony-forming units [CFU]) of *E. coli* DH5α (5 ml PAS) or *A. caviae* no. 86 (5 ml PAS)) were mixed with or without 10<sup>6</sup> cells of ciliates, Tth or Tsp (10 ml PAS), then incubated for 24 h at 30°C. Samples were subsequently collected at 0, 1, 2, 4, 8, and 24 h after incubation, and colony numbers of each bacteria were estimated by distinct colony color (*E. coli*, blue: *A. caviae*, white) on Colimark agar plates (Eiken). Grazing rates were expressed as decreasing bacterial rate (%) at 24 h compared to starting bacterial number (100%) immediately after incubation.

# **Meeting frequency**

Amounts (approximately 10<sup>8</sup> colony-forming units [CFU]) of GFP-expressing *E. coli* DH5α and vital stained *A. caviae* no. 86 (See above) were mixed with 10<sup>5</sup> cells of ciliates, Tth or Tsp in 1 ml of PAS, then incubated for 1 h at 30°C. Following incubation, mixed solutions were fixed with 4% formalin and then analyzed to estimate the meeting

167	frequency by fluorescence microscopy (BioZero, Keyence), which was determined by
168	counting the number of vesicles (green, E. coli alone; red, A. caviae alone; yellow, both)
169	formed into ciliates. Microscopic fields were randomly selected and at least 100 ciliate
170	cells were counted.
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172	Statistical analysis
173	All experiments were repeated at least three times. Statistical analysis was conducted
174	using the Mann-Whitney U-test. A $p$ value <0.05 was considered to be significant.
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177	RESULTS
178	Transfer frequency of the plasmid encoding $bla_{IMP-1}$ between $E.\ coli$ and
179	A. caviae in cultures with ciliates
180	First, we assessed if ciliates were associated with increased transfer of a plasmid
181	encoding <i>bla</i> <sub>IMP-1</sub> of <i>E. coli</i> TC17032828 (Donor) to <i>A. caviae</i> no. 86 (recipient) (Fig. 1A,
182	left). In contrast to the absence of ciliates $(1.0 \times 10^{-7})$ , the gene transfer frequency was

increased in the presence of ciliates to  $1.0 \times 10^{-5}$  (Tth) [p<0.05 vs. ciliate(-)] or  $1.0 \times 10^{-4}$ 183 184 (Tsp) [p < 0.05 vs. ciliate(-)] (Fig. 1B, left). Successful gene transfer was also confirmed 185 by PCR amplification with primers specific to  $bla_{IMP-1}$  (Fig. 1C, left). Next, we assessed whether ciliates facilitated reverse transfer of the A. caviae no. 86<sup>r</sup> plasmid to E. coli J 53 186 (Fig. 1A, right). As expected, the gene transfer frequency was significantly increased in 187 the presence of ciliates with a frequency of  $1.0 \times 10^{-4}$  (Tth) [p<0.05 vs. ciliate(-)] or 188  $1.0 \times 10^{-4}$  (Tsp) [p < 0.05 vs. ciliate(-)] (Fig. 1B, right). Successful plasmid transfer was also 189 190 confirmed by PCR amplification with primers specific to bla<sub>IMP-1</sub> (Fig. 1C, right). Moreover, assessment of MICs revealed that plasmid-acquired bacteria (E. coli 192 TC17032828 and A. caviae no. 86<sup>r</sup>) were resistant to CAZ, similar to those of C. freundii 193 carrying the plasmid encoding  $bla_{IMP-1}$ , an origin of the plasmid (Table 2). These results 194 also confirmed that representative colonies of plasmid-acquired bacteria were correctly 195 identified to the species level using the MALDI Biotyper system (Supporting Information 196 Fig S1). Overall, the results of this study indicate that cultures with ciliates could interactively prompt the transfer of plasmids encoding  $bla_{IMP-1}$  between  $E.\ coli$  and A.197 caviae. In addition, PCR-based replicon typing showed that the bla<sub>IMP-1</sub>-encoding 198

plasmid was incA/C type.

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Co-localization of E. coli and A. caviae in ciliate vesicles, and the influence of separating bacteria from ciliates on gene transfer frequency We evaluated whether the accumulation of two different kinds of bacteria [GFP-expressing E. coli DH5α (green) and vital-stained A. caviae no. 86<sup>r</sup> carrying a plasmid encoding *bla*<sub>IMP-1</sub> with PKH-26 (red)] could occur in the same ciliate vesicle. Co-localization of the two different bacteria in the same ciliate vesicle was observed, regardless of ciliate strains, suggesting that the gene transfer events effectively occurred through bacterial accumulation in the ciliate vesicle (Fig. 2). Interestingly, some vesicles packed with both bacteria were expelled from ciliates, after which these bacteria were released to the outside of disrupted vesicles, implying circulation of bacteria via ciliates in culture (Fig. 2, bottom right panel). To confirm this hypothesis, we estimated the gene transfer frequency of A. caviae no. 86<sup>r</sup> carrying a plasmid encoding bla<sub>IMP-1</sub> to E. coli J53 when ciliates were segregated from these bacteria with a transwell membrane. Segregation significantly diminished the gene transfer frequency when compared to cultures without the transwell, regardless of ciliate strains (Fig. 3).

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Grazing rate and meeting frequency showing both ciliates favor E. coli

rather than A. caviae for digesting their bacteria

To determine why the gene transfer frequency from E. coli (donor) to A. caviae (recipient) was inferior to that from A. caviae (donor) to E. coli (recipient), we estimated the grazing rate (decreasing bacterial number) with meeting frequency of both ciliates on digesting their bacteria. As a result, grazing rates of Tth and Tsp at 24 h after incubation with E. coli were 29.1±48.2% and 1.7±1.5%, respectively (Fig. 4A). Meanwhile, the rates of Tth and Tsp with A. caviae were 171.6±277.6% and 51.5±26.1%, respectively (Fig. 4B). The results showed both ciliates preferred to ingest E. coli rather than A. caviae. To confirm this, we assessed the meeting frequency of the bacteria inside the vesicles of ciliates. As expected, E. coli alone was more frequently seen in the vesicles than A. caviae, regardless of ciliate strains used for this study [Fig. 5A (green, E. coli; red, A. caviae; yellow, both together) and B (white bar)]. Moreover, both bacteria were found to co-localize into the same vesicles of ciliates (Fig. 5B, see black bars), indicating that their

231 meeting prompted gene transfer.

#### **DISCUSSION**

Multidrug-resistant Enterobacteriaceae producing ESBL such as carbapenemase encoded by  $bla_{\rm IMP}$  or  $bla_{\rm NDM}$ , which are resistant to carbapenem, pose a severe threat to hospitals and other healthcare facilities (1-5). Therefore, it is necessary to develop comprehensive management measures encompassing livestock, circulating food, public hygiene practices, and monitoring of natural environments to control the emergence of these bacteria (6-10). In this study, we demonstrated interactive transfer of the plasmid encoding  $bla_{\rm IMP-1}$  between  $E.\ coli$  and  $A.\ caviae$  in ciliates, indicating a pathway responsible for plasmid transfer among bacteria underlying the circulation of MDR bacteria.

Enterobacteriaceae producing carbapenemase encoded by  $bla_{IMP-1}$  have been widely detected in patients and hospital facilities, as well as livestock and aquatic environments, including sewers and rivers, which has led to circulation of these bacteria between

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humans and the environment (28-31). E. coli is a representative fecal bacterium belonging to *Enterobacteriaceae* that can widely adapt to a variety of mammalian bodies, including animal or human intestines. These bacteria contain plasmids that encode drug resistant genes, resulting in the bacterial constant release to natural environments such as rivers or soil. In fact, many studies have revealed the actual release of these bacteria such as Salmonella or E. coli into natural environments (32-35). Moreover, Aeromonas spp. including A. caviae comprise a representative natural environmental bacterium widely distributed in coastal areas and rivers as a fish pathogen, as well as in many areas in which people live (36, 37), that serve as a potential reservoir of the plasmid. Therefore, these two bacteria and  $bla_{IMP-1}$  as a factor gene were selected as the donor and recipient to assess gene transfer via Tetrahymena cells. The gene transfer frequency between E. coli and A. caviae was significantly higher in the presence of ciliates. However, when compared with the transfer from E. coli TC170328 to A. caviae no. 86, ciliates weakly promoted the transfer of the plasmid from A. caviae no. 86<sup>r</sup> to E. coli JR53. Because both ciliates favored E. coli rather than A. caviae, the preferential ingestion of E. coli rather than A. caviae in both ciliates is rigidly

263 involved in the difference in gene transfer frequency depending on the direction of plasmid transfer between E. coli and A. caviae. Moreover, sequence analysis revealed that 264 the plasmid was assigned to a type with incA/C, which can effectively mobilize into the 265 266 Enterobacteriaceae family (37). Imaging analysis revealed co-localization of *E. coli* and *A. caviae* in ciliate vesicles. 267 268 Because of unsuccessful establishment of fluorescence-expressing bacteria, live labeling 269 of A. caviae was conducted by staining with PKH-26, a hydrophobic fluorescence dye. 270 After staining, the bacteria were still viable, indicating that the staining protocol had minimal effects on bacterial survival. After mixed-culture, ciliates that ingested the 272 live-stained A. caviae became red. Spread of the dye to the ciliate's cytoplasm revealed 273 obvious ingestion of A. caviae. Interestingly, ciliates expelled pellets containing bacteria, 274 which subsequently released the bacteria to the culture, further increasing the frequency 275 of encounter with other environmental bacteria. These findings are concordant with the 276 results of previous studies (15, 16). Meanwhile, because of an increase in the pellet 277 amounts expelled by ciliate (Tsp), it is likely to be changed depending on the ciliate's strain, presumably indicating the presence of optimal sets with bacteria and ciliate strains 278

279 on accelerating gene transfer frequency. Because several studies have shown that living 280 bacteria such as Legionella and non-pathogenic E. coli packaged into multi-membrane vesicles can be released outside of protozoa such as *Tetrahymena* or *Acanthamoeba*, such 282 bacterial expulsion may involve the inability to digest bacteria responsible for host 283 cellular protection (38-40). Therefore, in this study, we defined the pellets as being 284 actively expelled, rather than excreted, from *Tetrahymena*. 285 Segregation using a transwell membrane significantly diminished gene transfer 286 frequency, regardless of ciliate strains. Overall, these findings and those presented above 287 indicated that transfer occurred via a series of processes comprising attachment, digestion 288 and accumulation of both bacteria into ciliates. It should be noted that an increasing trend 289 of the baseline of gene transfer frequency was observed the cultures with transwell, 290 although this was not significant. It is well known that filter-associated mating between bacteria is very effective (41, 42); therefore, it is possible that bacterial accumulation on 291 292 the pore membrane slightly promoted gene transfer frequency. 293 In conclusion, we demonstrated that ciliates promote transfer of plasmids between E. 294 coli and A. caviae. While further study is needed to clarify these results using other

295	bacteria, the information presented herein will be useful to control of the spread of
296	multiple-antibiotic resistant bacteria.
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310	DISCLOSURE
311	The authors declare that they have no conflicts of interest associated with this work.
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313	submission of this study.
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# Figure legends

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Figure 1 Gene transfer frequency of E. coli and A. caviae containing the plasmid encoding  $bla_{IMP-1}$  in cultures with ciliates. A. Direction of plasmid transfer between E. coli and A. caviae. A1 and A2 show the direction of plasmid transfer form E. coli to A. caviae and from A. caviae to E. coli, respectively. **B.** Plots show frequency of transfer of a plasmid encoding *bla*<sub>IMP-1</sub> from *E. coli* TC170328 (Donor) to *A. caviae* no. 86 (Recipient) (B1) and from A. caviae no. 86<sup>r</sup> back to E. coli J 53 (B2). Gene transfer frequency was estimated as the number of plasmid-acquired bacteria for each recipient (see text). Data shown are the averages of gene transfer frequency  $\pm$  standard deviations. Black and white boxes show the range from the mean to 75% and 25% values, respectively. \*, p < 0.05versus values for control without ciliates. C. Representative images showing the results of agarose gel electrophoresis of the  $bla_{IMP-1}$  PCR products amplified from the recipient (R) and plasmid-acquired bacteria (Pa). C1 and C2 show the direction of plasmid transfer form E. coli to A. caviae and from A. caviae to E. coli, respectively. Pa1–6 show the PCR results of representative plasmid-acquired bacterial colonies. N, negative control

473 [DNAase-free water (sigma)]. M, DNA ladder marker.

**Figure 2** Co-localization of *E. coli* DH5 $\alpha$  and *A. caviae* no. 86<sup>r</sup> in ciliate vesicles. Upper and lower sections of each of the four panels show mixed cultures of both bacteria with the ciliates, *T. thermophile* (Tth) and *T. pyriformis* (Tsp), respectively. Green indicates GFP-expressing *E. coli* DH5 $\alpha$ , while red shows live stained *A. caviae* no. 86<sup>r</sup>. Squares surrounded by dashed lines are enlargements of the same image (lower right panel). Scale, 50 μm.

**Figure 3** Influence of separating bacteria ( $A.\ caviae$  no.  $86^{\rm r}$  and  $E.\ coli$  J53) from ciliates with a transwell on gene transfer frequency. Upper images show the direction of plasmid transfer and images of the experiment. Plots show the gene transfer frequency of  $A.\ caviae$  no.  $86^{\rm r}$  (donor) containing a plasmid encoding  $bla_{\rm IMP-1}$  to  $E.\ coli$  J53 (recipient). Gene transfer frequency was estimated as the number of plasmid-acquired bacteria for each recipient (see the text). Data represent the average gene transfer frequency  $\pm$  standard deviation. Black and white boxes show the range from the mean to 75% and

489	25% values, respectively. *, $p$ <0.05 versus values for control without ciliates of each
490	experiment.
491	
492	Figure 4 Grazing rate of both ciliates when incubated with E. coli and A. caviae. The
493	rates were monitored for 24 h after incubation as described in the methods. A. Grazing
194	rate of Tth with E. coli and A. caviae. <b>B.</b> Grazing rate of Tsp with E. coli and A. caviae.
495	Tth, <i>T. thermophile</i> . Tsp, <i>Tetrahymena</i> . Ec DH5 $\alpha$ , <i>E. coli</i> DH5 $\alpha$ . Ae 86, A. caviae no. 86.
496	*, $p$ < 0.05 versus values for control without ciliates at each of the time points.
497	
498	<b>Figure 5</b> Meeting frequency of <i>E. coli</i> and <i>A. caviae</i> inside vesicles of both ciliates. The
499	frequency was estimated at 1 h after incubation as described in the methods. A.
500	Co-localization of these bacteria inside both ciliates. Green, GFP-expressing <i>E. coli</i> DH5
501	$\alpha$ . Red, vital stained A. caviae no. 86 with PKH-26 dye. Yellow, both bacteria
502	co-localized inside the vesicles of ciliates. Magnification, ×400. B. Assessing the
503	frequency of <i>E. coli</i> and <i>A. caviae</i> inside vesicles of both ciliates.
504	

# **Supporting Information**

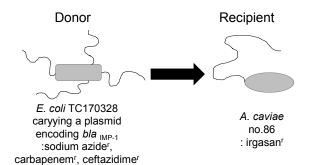
- Figure S1. Representative plasmid-acquired bacteria (E. coli TC170328 and A. caviae no.
- 507 86) of colonies identified to the species level using the MALDI Biotyper system.

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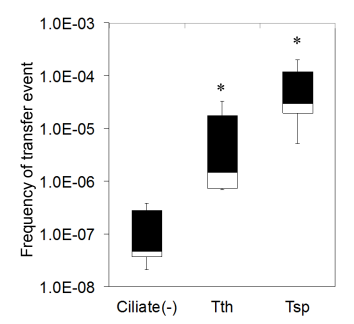
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509	List of Abbreviations:
510	ATCC, American Type Culture Collection; CAZ, ceftazidime; CFU, colony-forming
511	units; DNA, deoxyribonucleic acid; ESBL, extended spectrum β-lactamase; GFP, green
512	fluorescent protein; LB, Luria-Bertani; MALDI, matrix assisted laser
513	desorption/ionization; MDR, multidrug-resistant; MIC, minimum inhibitory
514	concentration; PAS, Page's amoeba saline; PCR, polymerase chain reaction; PYG
515	peptone-yeast extract glucose
516	
517	

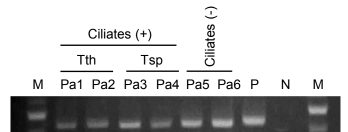
**A1** 



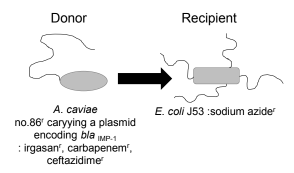
B1



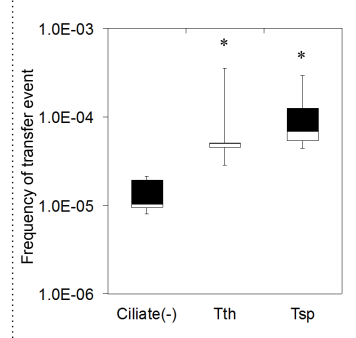
C1



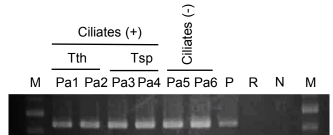
A2



B2



C2



GFP PKH26 Mixed culture of ciliates (Tth) with GFP-expressing E. coli DH5a and PKH26-stained A. caviae 50 µm Merge 50 µm 50 µm GFP PKH26 Mixed culture of ciliates (Tsp) with GFP-expressing E. coli DH5α and PKH26-stained A. caviae  $50\,\mu$  m Merge

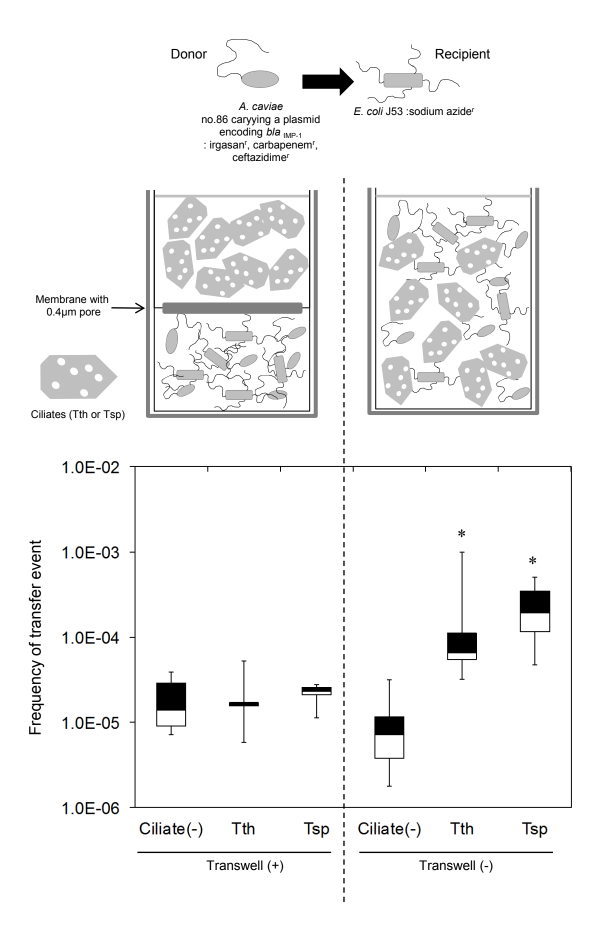
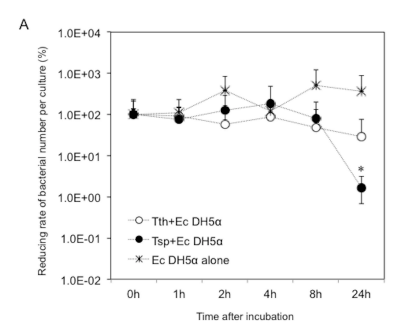


Fig. 4



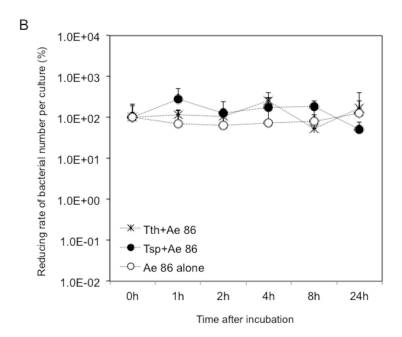
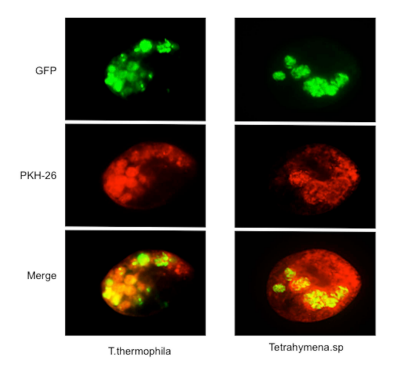


Fig. 5 A



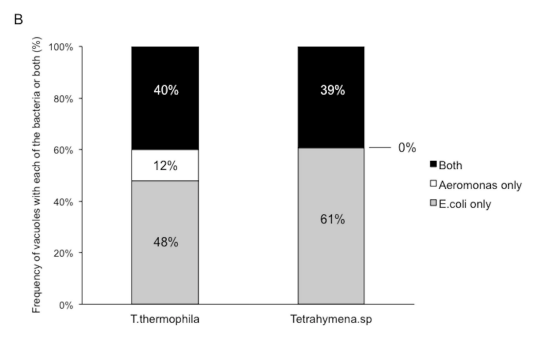


Table 1 Bacteria and protozoa used for this study

Bacteria and protozo	Purpose	Characteristics or sources
<b>Protozoa</b> Ciliates		
T. thermophila (Tth)	Host cells / Imaging	Gifted from Dr. Sugai of Ibaragi University, Japan
Tetrahymena sp. (Tsp)	Host cells / Imaging	Gifted from Dr. Tukii of Hosei University, Japan
Bacteria		
C. freundii	Donor (original)	Clinical isolate, carrying a plasmid encoding <i>bla</i> IMP-1 :carbapenem <sup>r</sup> , ceftazidime <sup>r</sup>
E. coli J53	Recipient	Sodium azide <sup>r</sup> , purchased from ATCC
E. coli TC170328	Donor	Carrying a plasmid encoding <i>bla</i> <sub>IMP-1</sub> :sodium azide <sup>r</sup> , carbapenem <sup>r</sup> , ceftazidime <sup>r</sup>
		*Established by mating with <i>C. freundii</i> caryying a plasmid encoding <i>bla</i> <sub>IMP-1</sub>
A. caviae no.86	Recipient /Imaging	Environmental isolate (river), irgasan <sup>r,</sup> gifted from Dr. Tamura of Rakno University, Japan
E. coli DH5α	Imaging	For expressing GFP
E. coli ATCC 29213	Control for MIC	purchased from ATCC
S. aureus ATCC 25922	Control for MIC	purchased from ATCC

Table 2 MICs of bacteria against CAZ

Bacteria	Classification	MIC against CAZ (μg/ml)
C. freundii	Donor (original)	>128
E. coli TC170328	Transconjugant/Donor	>128
A. caviae no.86	Recipient	1
<i>A. caviae</i> no.86 <sup>r</sup>	Transconjugant/Donor	>128
E. coli J53	Recipient	0.5
E. coli ATCC 29213	Control for MIC	0.5
S. aureus ATCC 25922	Control for MIC	16