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## 1 Permissivity of insect cells to Waddlia chondrophila, Estrella lausannensis and

## 2 Parachlamydia acanthamoebae

- 3 Carole Kebbi-Beghdadi, Morgane Fatton, Gilbert Greub\*
- 4 Center for Research on Intracellular Bacteria (CRIB), Institute of Microbiology, University
- 5 Hospital Center and University of Lausanne, Lausanne, Switzerland
- 6
- 7 \*Corresponding author:
- 8 Prof. Gilbert Greub
- 9 Institute of Microbiology
- 10 Rue du Bugnon 48
- 11 1011 Lausanne
- 12 Switzerland
- 13 Tel: 0041 21 314 4979
- 14 Fax 0041 21 314 4060
- 15 e-mail: <u>gilbert.greub@chuv.ch</u>

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#### 19 Abstract

20 Recent large scale studies questioning the presence of intracellular bacteria of the

- 21 Chlamydiales order in ticks and fleas revealed that arthropods, similarly to mammals,
- reptiles, birds or fishes, can be colonized by *Chlamydia*-related bacteria with a
- 23 predominant representation of the *Rhabdochlamydiaceae* and *Parachlamydiaceae*
- families. We thus investigated the permissivity of two insect cell lines towards *Waddlia*
- 25 chondrophila, Estrella lausannensis and Parachlamydia acanthamoebae, three bacteria
- representative of three distinct families within the *Chlamydiales* order, all documented in
- ticks and/or in other arthropods. We demonstrated that *W. chondrophila* and *E.*
- 28 lausannensis are able to very efficiently multiply in these insect cell lines. E.

29 *lausannensis* however induced a rapid cytopathic effect, which somehow restricted its

- replication. *P. acanthamoebae* was not able to grow in these cell lines even if inclusions
- 31 containing a few replicating bacteria could occasionally be observed.
- 32
- *Keywords:* Intracellular bacteria; *Chlamydia*-related bacteria; Cell permissivity; Host
   range; Reservoir; Vector
- 35

## **1. Introduction**

37 All bacteria belonging to the *Chlamydiales* order are strict intracellular organisms sharing a biphasic life cycle that involves two distinct bacterial forms, an infectious Elementary 38 Body (EB) capable of attaching to and entering into its host cell, but that cannot replicate 39 40 before it differentiates into a metabolically active form, called Reticulate Body (RB). At the end of the multiplication phase, RBs differentiate back into infectious particles and 41 42 lyse their host cells to start a new cycle [32]. Chlamydiales bacteria have been isolated from samples of multiple origins such as mammals, birds, fishes, reptiles or protozoa 43 revealing the large and probably still largely underestimated diversity of their ecological 44 niches (reviewed in [22, 36]). Besides the well-studied human and animal pathogens 45 belonging to the Chlamydiaceae family (such as Chlamydia pneumoniae, C. trachomatis 46 or *C. abortus*), this clade also comprises 8 other family-level lineages whose members 47 48 are very diverse being either emerging pathogens able to grow in cells of various origins (Waddlia chondrophila, Simkania negevensis) or harmless environmental species only 49 replicating in amoebae (Protochlamydia amoebophila, Neochlamydia hartmanellae). 50 51 Recent large scale studies questioning the presence of *Chlamydiales* bacteria in ticks and fleas [13], [Pilloux et al. submitted 2015] revealed that arthropods may be colonized 52 by these bacteria and thus could serve as reservoir and vectors for potential novel 53 54 pathogens. In this context, we investigated the permissivity of two insect cell lines, Aedes albopictus larva cells and Spodoptera frugiperda ovary cells (Sf9), towards 55 Waddlia chondrophila, Estrella lausannensis and Parachlamydia acanthamoebae, three 56 Chlamydia-related bacteria representative of three different families, all belonging to the 57

- *Chlamydiales* order, documented in fleas and/or ticks and whose pathogenic potential for humans and cattle is confirmed or highly suspected [2-6, 9, 15, 16, 23, 34, 37, 14].
- 61 2. Materials and methods
- 62 2.1. Cell culture and bacterial strains
- 63 Aedes albopictus clone C6/36 larva cells (ATCC<sup>®</sup> CRL-1660<sup>™</sup>) and Spodoptera
- 64 *frugiperda* ovarian epithelial cells (Sf9) (ATCC<sup>®</sup> CRL-1711<sup>™</sup>) were routinely maintained
- respectively at 28°C and 5%CO<sub>2</sub> in Dulbecco's modified essential medium (DMEM;
- 66 Gibco Invitrogen, Basel, Switzerland) supplemented with 10% foetal calf serum
- 67 (Biochrom, Berlin, Germany) or at 27°C in Grace insect medium (GIM; Gibco Invitrogen,
- Basel, Switzerland) supplemented with 10% foetal calf serum.
- 69 W. chondrophila strain WSU 86-1044 (ATCC VR-1470), E. lausannensis strain CRIB 30
- and *P. acanthamoebae* strain Hall's coccus were grown at 32°C within *Acanthamoeba*
- *castellanii* strain ATCC 30010, as described in [19].

72

#### 73 2.2. Infection procedure

Insect cells were seeded at  $1 \times 10^5$  cells per well in 24-wells microplates (Corning) the day before infection. Infection was performed at 28°C or 27°C, as described in [24], with a 5 days-old culture of bacteria in A *castellanii* diluted, if not otherwise described, 1:1000 for *W. chondrophila* (MOI: 1-20) and *P. acanthamoebae* (MOI: 0.1-1) and 1:2000 for *E. lausannensis* (MOI: 1-10)

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80 2.3. Immunofluorescence and confocal microscopy

At different time points after infection, immunofluorescence was performed on cells 81 82 cultivated on glass coverslips following precisely the protocol described in [27] except that rabbit anti-P. acanthamoebae and mouse anti-E. lausannensis were used 83 respectively at a 1:1000 and 1:500 dilution. Secondary antibody was diluted 1:500 and 84 mixed with a 1/50 dilution of Concanavalin A (Molecular Probe) and 150 ng/ml DAPI 85 (Molecular Probe, Eugene, Oregon, USA). Cells were observed under an epifluorescent 86 microscope (Axioplan 2, Zeiss, Feldbach, Switzerland) or a confocal microscope 87 (AxioPlan 2 LSM 710, Zeiss). 88

89

#### 90 2.4. Electron microscopy

One T25 flask of *Aedes albopictus* or Sf9 cells was infected as described above with *W. chondrophila* and *E. lausannensis* diluted 1/200. 24 hours post-infection, cells were
washed once with PBS and harvested. After one centrifugation step of 10 min at 1'000
rpm, cells were resuspended in phosphate buffer (19ml of 0.2 M NaH<sub>2</sub>PO<sub>4</sub> + 81ml of
0.2M Na<sub>2</sub>HPO<sub>4</sub>, pH 7.4) containing 0.2% glutaraldehyde (Fluka Biochemika, Buchs,
Switzerland) and 4% paraformaldehyde (Electron Microscopy Sciences, Hatfield, USA)

and incubated overnight at 4°C. After two more washing steps with phosphate buffer,
cells were prepared as described previously [10]. Thin sections on grids were examined
with a transmission electron microscope Philips CM 100 (Philips, Eindhoven, The
Netherlands).

101

102	2.5.	Quantitative	PCR
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103 Cells were harvested at different time points after infection, genomic DNA was extracted

104 following manufacturer's instructions (Wizard SV Genomic DNA purification kit,

105 Promega, Madison, WI) and qPCR was performed as described in [24]. To account for

variation in the inocula used for infection, bacterial growth is expressed as a fold

increase in the number of bacteria after normalization using time point "0h" as reference.

108

109 2.6. Cell viability

110 Cell viability was determined with a propidium iodide assay as described in [14] except

that propidium iodide was added immediately after infection. Positive control was

obtained by incubating non-infected cells 5 minutes with MetOH at

-20°C. Results were normalized considering the positive control as 100% mortality.

114

#### 115 **3. Results**

116 3.1. Growth kinetic in insect cells

117 Two insect cell lines, Aedes albopictus larva cells and Sf9 (S. frugiperda ovarian

epithelial cells) were infected with either *W. chondrophila, E. lausannensis* or *P.* 

acanthamoebae and bacterial growth was monitored using specific quantitative PCRs

120 (qPCRs) developed in our laboratory [11, 16, 30]. In parallel, bacterial multiplication was

also assessed by immunofluorescence and confocal microscopy using in-housepolyclonal antibodies and by electron microscopy.

123

#### 124 3.1.1. W. chondrophila and E. lausannensis

125 Results shown in Fig. 1 indicated that W. chondrophila and E. lausannensis are both able to efficiently replicate in insect cells, leading to an increase of bacterial genomic 126 DNA copies of about 3 logs in 48 hours for W. chondrophila (Fig. 1 panels a and b) and 127 of about 2 logs in 48 hours for *E. lausannensis* (Fig. 1 panels c and d). During the first 8 128 hours following infection, EBs enter their insect cell host, differentiate into RBs and 129 130 create a replicative niche able to support their exponential multiplication. Inclusions containing dividing bacteria can already be observed by immunofluorescence and 131 confocal microscopy 24 hours post infection (Fig. 2A, panels a,b,c,and d). Similarly to 132 what has been described in other cell lines [24, 25], W. chondrophila then exponentially 133 replicate until 48 hours p.i. and ultimately lyse their host to release infectious particles 134 ready to infect new insect cells (data not shown). Replication of W. chondrophila and E. 135 lausannensis was also documented in Sf9 cells by electron microscopy 24 hours post 136 infection (Fig. 2B). The *W. chondrophila*-containing vacuoles were surrounded by tightly 137 138 associated mitochondria, a feature that was also described in other cell lines [12, 25].

139

140 3.1.2. P. acanthamoebae

No growth of *P. acanthamoebae* could be detected by qPCR or immunofluorescence in
the two insect cell lines tested (Fig.1 and 2A, panels e and f) even after 7 days.
However, when cells were infected with a 10 fold higher bacterial load (MOI of 1-10),
rare inclusions containing replicating bacteria could be observed by confocal microscopy

in *Aedes albopictus* cells (Fig. 2A, panel g). This bacterial replication is very limited and
 restricted to about 3% of all infected cells.

147

148 3.2. Cytophatic effect

149 Direct examination by confocal microscopy of *E. lausannensis* infection revealed that these bacteria, like *W. chondophila*, efficiently multiplied in insect cells but these cells 150 were more rapidly lysed than those infected with W. chondrophila. This observed 151 cytopathic effect was confirmed in a host cell viability assay based on propidium iodide 152 incorporation that showed, 48 hours post infection, 100% mortality of Aedes albopictus 153 cells infected with E. lausannensis versus 50% in cells infected with W. chondrophila 154 and 23% in non-infected cells (Fig. 3). Results were normalized using methanol-treated 155 cells at time of infection as a positive control of 100% mortality. Mortality rate over 100% 156 are thus explained by an increase, at later time points, of the total number of cells that 157 ultimately died. We could observe a weak cytopathic effect of P. acanthamoebae 158 infection that probably resulted more from the lower fitness of infected cells than from 159 the few replicating bacteria. 160

In Sf9 cells, a similar cytopathic effect of *E. lausannensis* was observed by confocal
 microscopy (data not shown), however, due to interferences of the culture medium with
 the propidium iodide assay, this increased cell mortality could not be precisely
 quantified.

165

166 **4. Discussion** 

In the present study, we demonstrated that *W. chondrophila* and *E. lausannensis* are able to enter and efficiently multiply in two different insect cell lines. *E. lausannensis* growth is however slightly less efficient than *W. chondrophila* growth, a difference that can be explained by the rapid cytopathic effect of *E. lausannensis*, which restricts the number of replication rounds possible.

W. chondrophila is an emerging pathogen for humans and animals and it is able to 172 readily grow at temperatures ranging from 25°C to 37°C in mammalian cell lines of 173 diverse origins [12, 21, 25, 28], in fish cells [24] as well as in protozoa [17, 31]. E. 174 lausannensis, whose pathogenic potential is currently under investigation, was first 175 176 isolated from an environmental water sample and is also able to grow at different temperatures in protozoa, fishes and mammalian cells [14, 24, 30]. Strikingly, these two 177 organisms that display such a broad host range, both encode an extended family of 178 OmpA proteins with beta-barrel structure that were recently shown to be dominant in the 179 outer membrane of *W. chondrophila* and to play a role in adhesion of this bacteria to its 180 host [1, 7, 8, 26, 29]. In addition, Simkania negevensis, another Chlamydia-related 181 bacteria that also possess a large family of 37 MOMP-like proteins in its outer 182 membrane similarly displays a wide host range [1, 27]. Based on these observations, we 183 184 recently hypothesized that the large diversity of the OmpA protein family is linked to the wide host range of these bacteria [26]. In addition, ability to grow in multiple and diverse 185 eukaryotic cells correlates in most cases with virulence towards mammals [27]. 186 187 *P. acanthamoebae* only encodes one homolog of MOMP and its efficient growth seems to be restricted to amoebal hosts [18, 20]. Several studies have reported a limited 188 replication of this bacteria in host cells such as bone-marrow derived macrophages, fish 189 or insect cell lines [24, 33, 35]. This present work also demonstrated that growth of P. 190

acanthamoebae in insect cells is limited and restricted to a few cells. Sixt et al. also 191 192 reported a limited P. acanthamoebae replication in Aedes albopictus, Sf9 and Drosophila S2 cells and they observed a bacteria-induced programmed cell death in the 193 last two cell lines [35]. We did not observe nuclear fragmentation in P. acanthamoebae-194 infected cells and cell viability was stable during the first 72 hours post infection, which is 195 not in agreement with apoptosis. Furthermore, preliminary experiments performed in 196 Aedes albopictus cells in presence of a pan-caspase inhibitor did not demonstrate an 197 enhanced multiplication of *P. acanthamoebae* (A. Croxatto, unpublished). 198 In conclusion, we demonstrated the permissivity of insect cells to Waddla chondrophila 199 and Estrella lausannensis and further highlighted the very broad host range of these 200 possible pathogens and the potential role of insects as reservoir or vectors for these 201 strict intracellular bacteria. 202

203

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## 212 6. Conflict of interest

The authors have no conflict of interest.

214

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## 318 **7. Figure legends**

#### 319 Fig. 1 Bacterial growth within insect cells

- 320 W. chondrophila (panels a and b), E. lausannensis (panels c and d) and P.
- 321 *acanthamoebae* (panels e and f) replication measured by qPCR in *Aedes albopictus*
- cells (panels a, c and e) and in Sf9 cells (panels b, d and f). Values are normalized to
- 323 the number of bacteria at 0h post infection (p.i). Results are the mean +/-SD of at least
- 324 four independent experiments performed in duplicates.
- 325

## **Fig.2 Bacterial replication assessed by confocal and electron microscopy**

- A. Immunofluorescence and confocal microscopy of *W. chondrophila* (panels a and
- b), *E. lausannensis* (panels c and d) and *P. acanthamoebae* (panels e and f) in
- Aedes albopictus cells (panels a, c and e) and in Sf9 cells (panels b, d and f) 24
- hours post-infection. Panel g displays one rare inclusion of *P. acanthamoebae* in
- Aedes albopictus cells 4 days post infection. Bacteria (green) are stained with
- 332 species-specific polyclonal antibodies and insect cells (red) are stained with Texas
- Red-conjugated Concanavalin A. Scale bar 10  $\mu$ m.
- B. Electron micrographs of *W. chondrophila* (panel a) and *E. lausannensis* (panel b)
  in Sf9 cells 24 hours post infection. Scale bar: 1 μm.
- 336

#### 337 Fig. 3 Cell viability

Aedes albopictus cells viability was determined with a propidium iodide assay at different
 time points after infection. Results are the mean +/- SD of 2 independent experiments
 performed in triplicates.



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В





## Figure 3

