The obligate intracellular bacterium *Orientia tsutsugamushi* differentiates into a developmentally distinct extracellular state

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*Orientia tsutsugamushi* (Ot) is an obligate intracellular bacterium in the family Rickettsiaceae that causes scrub typhus, a severe mite-borne human disease. Its mechanism of cell exit is unusual amongst Rickettsiaceae, as Ot buds off the surface of infected cells enveloped in plasma membrane. Here, we show that Ot bacteria that have budded out of host cells are in a distinct developmental stage compared with intracellular bacteria. We refer to these two stages as intracellular and extracellular bacteria (IB and EB, respectively). These two forms differ in physical properties: IB is both round and elongated, and EB is round. Additionally, IB has higher levels of peptidoglycan and is physically robust compared with EB. The two bacterial forms differentially express proteins involved in bacterial physiology and host-pathogen interactions, specifically those involved in bacterial dormancy and stress response, and outer membrane autotransporter proteins ScaA and ScaC. Whilst both populations are infectious, entry of IB Ot is sensitive to inhibitors of both clathrin-mediated endocytosis and macropinocytosis, whereas entry of EB Ot is only sensitive to a macropinocytosis inhibitor. Our identification and detailed characterization of two developmental forms of Ot significantly advances our understanding of the intracellular lifecycle of an important human pathogen.
**Orientia tsutsugamushi** (Ot) is an obligate intracellular Gram-negative bacterium that causes the mite-transmitted human disease scrub typhus. This condition is one of the most severe rickettsial diseases; it is associated with a median 6% mortality in untreated patients. Scrub typhus is a leading cause of severe febrile illness in the Asia-Pacific, a region containing two-thirds of the world’s population. Recent reports from Latin America, the Middle East, and Africa suggest that the disease is more globally distributed. Knowledge of the fundamental biology of this organism lags behind that of pathogens of equivalent prevalence and severity. Historical underestimations of Ot incidence, its classification as a biosafety level 3 pathogen, its fastidious growth in cultured cells, and the fact that it remains genetically intractable likely account for the paucity of studies focused on this pathogen.

The order Rickettsiales contains two major families: Anaplasmataceae and Rickettsiaceae. Members of the Anaplasmataceae, which include *Anaplasma* spp. and *Ehrlichia* spp., reside within membrane-bound vacuoles inside host cells. The developmental cycle of certain Anaplasmataceae family members is biphasic and involves switching between a noninfectious intracellular replicative form (reticulate body) and an infectious extracellular form (elementary body). While members of the Rickettsiaceae, namely *Rickettsia* spp. and Ot, do not appear to differentiate into distinct forms during their intracellular lifecycle, Ot is unique amongst the characterized Rickettsiaceae in budding off the surface of infected host cells in a manner reminiscent of enveloped viruses. This observation led us to investigate whether budded Ot is in a distinct developmental stage compared with intracellular counterparts.

Ot has been shown to enter host cells using the zipper-like invasion mechanism, which is common to many Rickettsiales members. This process is initiated by binding of bacterial surface adhesins to receptors expressed on mammalian cells. Adhesins such as TSA56, ScaA, and ScaC6–8 bind to host cell fibronectin and syndecan-4; these proteins then interact with integrin α5β1, leading to activation of intracellular signaling that culminates in actin remodeling and clathrin-mediated endocytosis. It is important to emphasize that these studies were entirely focused on intracellular Ot, as they were performed on bacteria isolated from host cells by mechanical lysis of infected cells. However, the reentry mechanism utilized by bacteria that have budded out of cells may be influenced by the presence of an enveloping host cell membrane and/or its differentiation into a distinct state.

Here, we show that Ot exists in distinct intracellular (IB) and extracellular (EB) bacterial populations. We report the characterization of these two states in terms of their physical properties, metabolic activity, and gene expression profiles and demonstrate that they enter host cells using different mechanisms.

**Results**

**IB and EB forms of Ot differ in physical properties and metabolic activity.** Ot is typically described as a coccobacillary bacterium. Its small size and propensity to aggregate make it difficult to obtain clear images of this bacterium using conventional light microscopy. To circumvent this challenge, we labeled Ot with an antibody against the abundant surface protein TSA56 and employed stochastic optical reconstruction microscopy (STORM) for high-resolution imaging. We first investigated the shape of bacteria located inside and outside host cells (Fig. 1A, B and Supplementary Fig. 1). Since Ot begins to appear in the supernatant of infected L929 cells at high levels 5 days post-infection (Supplementary Fig. 2) we focused on the shape of EB Ot beginning 5 days post-infection, whilst we measured the shape of IB Ot throughout the infection cycle. We found that EBs were approximately round in shape with a diameter of about 1.2 µm, and they were often aggregated into clusters. In contrast, IBs adopted diverse round and elongated shapes and varied in dimensions; some bacteria were 3–4 µm in length. Additionally, IB size varied during the course of infection. Initially, and up to 1-day post-infection, IBs were similar in length to EBs. At later stages, i.e., 2–7 days following infection, IBs were characterized by longer lengths. These kinetic changes in shape correlated with Ot growth dynamics whereby replication starts 24–48 h post-infection. These differences in shape provide evidence for the existence of two separate bacterial forms.

We next investigated potential differences in bacterial cell wall composition. We previously showed that Ot builds a low-abundance peptidoglycan-like structure. Thus, we assessed relative peptidoglycan levels in IB and EB Ot forms using a NOD1 reporter assay (Fig. 1C). We detected significantly more peptidoglycan in IB cells compared with EB cells. This finding may account for the observed differences in shape (Fig. 1A) as peptidoglycan is required to build elongated cells.

In related structural studies, we capitalized on our previous finding that the abundant surface protein TSA56 is involved in forming disulfide bonds with itself or other proteins, leading to the formation of aggregates. These complexes can be identified as smears following sample preparation in the absence of reducing agents and evaluation by immunoblot analyses. In this study, we found evidence for substantially higher disulfide bond formation in EB Ot than IB Ot, as determined by the presence of a high molecular weight smear (Fig. 1D). We hypothesize that this crosslinked structure confers EB Ot with additional structural rigidity in the absence of peptidoglycan. These experiments were performed in the presence of iodoacetamide, an agent that covalently modifies cysteine residues and prevents them from forming new bonds, thus ensuring that the cross-links were not formed during sample preparation and processing.

Our next goal was to assess potential changes in genome copy number as, during rapid growth, bacteria such as *E. coli* carry multiple genome copies within a single cell. To address this issue, we determined bacterial cell numbers using fluorescence microscopy and bacterial genome copies using qPCR. We found no differences: Ot IB and EB harbored just over one estimated genome copy per cell (Fig. 1E). This measurement is an approximation that suffers limitations associated with absolute quantification using qPCR and microscopy, but any such errors are expected to be constant between the IB and EB measurements.

Finally, we asked whether IB and EB Ot forms differed in their metabolic activity. Ot is an obligate intracellular bacterium with limited central metabolic and biosynthetic capability and a reliance on host cells for carbon and nitrogen sources and metabolic intermediates. These observations led us to hypothesize that EB bacteria are metabolically inactive owing to their extracellular location. To test this, we used a microscopy-based assay whereby growing bacteria are exposed to a clickable methionine analog L-Homopropargylglycine (HPG) that is subsequently labeled with a fluorophore. Using this methodology, we observed that 95% of IB were translationally active 5 days post-infection, compared with 0% of EB (Fig. 2B). It is possible that EB Ot undergo some residual metabolic activity that is not detected by our assay of protein translation. The fact that not all IB were labeled with HPG indicated that this result did not simply reflect a difference in bacterial permeability between the IB and EB forms. These results provide an additional line of evidence to support the existence of two fundamentally different Ot states.

**IB and EB forms of Ot exhibit different gene and protein expression profiles.** We next investigated whether the differences
in physical properties and metabolic activity observed in IB and EB forms of Ot were accounted exclusively by physical location relative to host cells. Exposure of EB forms to an osmotically different extracellular environment combined with the lack of nutrient availability might account for our observations. However, a second possibility is that EB forms arise by differentiation of IB into a biochemically distinct state. We reasoned that bacteria in the same developmental stage but differing in position...
relative to host cells would exhibit almost identical protein profiles. To test this, we performed shotgun mass spectrometric analyses on isolated IB and EB Ot forms. We found that whilst the correlation coefficient of proteins detected in IB and EB isolated four days post-infection was strong (correlation coefficient = 0.843) (Supplementary Fig. 3) there were several proteins that accumulated to different levels in the two populations. A complete list of the relative levels of proteins detected in IB and EB is provided in Supplementary Data 1. Pathway analyses revealed that IB Ot was enriched in proteins involved in protein synthesis (Fig. 3A). In contrast, EB Ot displayed high expression of SpoT and RpoH, two proteins involved in the stringent response (Fig. 3B). In order to validate these results, we generated antibodies against four proteins that were present at different levels in IB and EB Ot: SpoT, RpoH, RpoD and CtrA. We carried out immunoblot analysis of IB and EB Ot and found that both SpoT and RpoH were present at higher levels in EB than IB, whilst RpoD and CtrA were present at higher levels in IB Ot (Fig. 3C), consistent with the trends observed in our proteomics analysis.

We next investigated whether EB and IB Ot exhibited differences in transcript expression levels. These experiments required the propagation of large amounts of EB because this form of Ot is metabolically inactive and thus harbors very low mRNA levels. We selected 11 genes, three of which (curA, ttxB, sodB) were expressed at similar levels in our proteomic studies comparing EB and IB. Additionally, we quantified mRNA expression of four genes encoding proteins enriched in IB Ot (yccA, yuaB, dapA, scaA) and four genes enriched in EB Ot (obgE, spoT, tsA56, rpoH). We isolated RNA from IB and EB bacteria 4 days post-infection and assessed mRNA levels by qRT-PCR (Fig. 3D). Whilst we observed differences in relative mRNA levels between IB and EB for most genes, we frequently found no correlation between mRNA and protein levels for a particular gene. For example, rpoH transcript levels were similar in IB and EB Ot, but protein levels were higher in EB. These observations provide additional evidence supporting the existence of molecularly distinct bacterial forms and suggest that transcriptional and post-transcriptional regulatory mechanisms participate in the differentiation process.

Surface proteins are present at different relative levels on IB and EB forms of Ot. Our differential proteomics analyses revealed that IB and EB have different levels of specific surface proteins (Fig. 4A). Surface proteins play a central role in Ot function for two reasons. First, unlike most Gram-negative bacteria, Ot lacks lipopolysaccharide, which generally serves as a membrane shield. Second, Ot is a cytosolic bacterium, and its surface is in direct contact with cytoplasmic host components. Ot harbors genes encoding four widely conserved autotransporter proteins: ScaA, ScaC, ScaD, and ScaE. Using our proteomics data, we detected ScaA and ScaC, but not ScaD or ScaE. ScaA was slightly enriched in IB relative to EB Ot (1.5X), while ScaC was enriched in EB (1.87X). Ot also harbors genes encoding three immunogenic surface proteins that dominate the humoral response in human patients, TSA22, TSA47, and TSA56 [16]. We found that TSA22 was enriched in IB relative to EB Ot (3.2X) while TSA47 and TSA56 were not present at significantly different levels (1.14X and 1.32X increase in EB compared with IB, respectively). To verify our observations, we generated polyclonal antibodies against ScaA and ScaC and determined their relative levels in IB and EB bacteria using immunoblot analyses (Fig. 4B and Supplementary Fig. 4). Our experiments up to this point demonstrate that EB Ot exhibits a broader repertoire of surface proteins, which could explain the differences in virulence and host response observed between IB and EB forms of Ot.
point were performed in mouse fibroblasts (L929) because Ot grows readily in these cells. Moreover, L929 fibroblasts are suitable for characterizing bacterial properties that are not affected by the cellular environment. However, for our validation studies, we also investigated bacteria grown in primary human umbilical vein endothelial cells (HUVEC) as a more physiologically relevant model of Ot infection in humans. Our immunoblot studies in both L929 and HUVECs confirmed enrichment of ScaA in IB Ot and higher levels of ScaC in EB compared to IB Ot, in agreement with our mass spectrometric analyses.

Fig. 4 Autotransporter proteins ScaA and ScaC are differentially expressed on IB and EB forms of Ot. A Volcano plot showing differential expression of surface proteins in proteomics analysis of IB and EB forms of Ot. We observe higher expression of ScaC, TSA56, and TSA47 in EB whilst IB Ot expresses higher ScaA and TSA22 levels. Statistical significance was determined using a t test. B Immunoblot analysis of IB and EB Ot from infected L929 and HUVEC cells confirming differential expression of ScaA, and ScaC between IB and EB forms. Loading was normalized by measuring bacterial genome copy number using qPCR. Uncropped immunoblot images with molecular weight markers are given in Supplementary Fig. 4. C Quantification of translational activity, as determined by a positive confocal microscopy signal observed with the clickable methionine probe HPG. HUVEC cells were infected with Ot for 5 days, then fixed and labeled with HPG and anti-ScaA/anti-ScaC antibodies. ScaA/ScaC-positive bacteria were scored as being positive or negative for HPG labeling. D Confocal micrographs of HUVEC cells infected with Ot for 5 days and labeled for DNA (Hoechst, blue), translational activity (HPG, green), ScaA (anti-ScaA antibody, magenta), and ScaC (anti-ScaC antibody, red). Scale bar = 10 μm. Examples of ScaC-positive extracellular bacteria are shown with a white arrow. More images and equivalent experiments in L929 cells are shown in Supplementary Figs. 5–7. E Quantification of experiments described in D. The percentage of IB and EB Ot at 5 days post-infection that were positively labeled with anti-ScaA- and -ScaC antibodies was counted. F Quantification of ScaA- and ScaC-positive IB bacteria inside HUVEC cells one- and three hours post-infection, showing an increase in ScaA labeling and a decrease in ScaC labeling. In (B, E, F), n = 30 randomly imaged fields across three independent experiments. Graphs show individual values and means. Statistical significance was calculated using a two-sided unpaired t test with Graphpad Prism software. *p value ≤ 0.05; **p value ≤ 0.01; ***p value ≤ 0.001; ****p value ≤ 0.0001.
ScaA and ScaC as preliminary markers of EB and IB forms of Ot. Our observation that 95% of IB is translationally active 5 days post-infection, compared with 0% of EB, combined with the fact that these Ot forms display opposite relative levels of ScaA and ScaC led us to investigate whether these surface proteins can be used as bona fide markers of metabolically active IB and metabolically quiescent EB Ot forms, respectively. We used immunofluorescence microscopy to identify bacteria expressing ScaA and ScaC and then quantified the number of translationally active ScaA- and ScaC-positive bacteria by measuring incorporation of HPG. We found (Fig. 4C and Supplementary Fig. 7A) that ScaA-positive bacteria, in contrast to ScaC-positive Ot, were mostly translationally active. We then quantified the number of ScaA/ScaC-positive bacteria inside and outside cells and confirmed that ScaC and ScaA levels were higher in EB and IB Ot, respectively (Fig. 4D, E and Supplementary Figs. 5, 6, 7B). Together, these results show that ScaA and ScaC are present at different relative levels depending on bacterial location relative to host cells: metabolically active IB exhibit higher levels of ScaA relative to EB Ot. Conversely, levels of ScaC in metabolically inactive EB is high relative to that of IB forms of Ot. We therefore propose that ScaA and ScaC can be used as markers to distinguish between IB and EB forms of Ot. However, it is worth noting that the correlation between expression of these proteins and bacterial position relative to host cells is not absolute and it is likely that more robust markers of the different forms of Ot may be uncovered in future studies.

The observation that EB exhibits high scaA transcript levels (Fig. 3D) but low levels of ScaA protein led us to hypothesize that ScaA is translated into protein soon after entry into host cells, using pre-formed mRNA. We quantified the number of IB Ot expressing ScaA and ScaC at 1- and 3 h post-infection and found a decrease in ScaC levels and an increase in ScaA levels as the infection proceeded (Fig. 4F and Supplementary Fig. 7C). These results show that ScaA accumulates during the first 3 h after infection, suggesting that scaA transcripts are made in EBs in anticipation of new infections. These results point to the existence of regulatory mechanisms that control levels of ScaA and ScaC on the surface of Ot cells, depending on bacterial location and stage of host infection.

IB and EB Ot populations are infectious but differ in physical stability and mechanisms of entry into host cells. Previous studies have shown that certain Anaplasmataceae species differentiate into distinct intracellular and extracellular forms and that only the extracellular form is effective at infecting host cells17. This observation led us to investigate whether the ability to infect host cells differs in IB compared with EB Ot. Growth curve measurements revealed that both bacterial forms could infect L929 and HUVEC cells and replicate with similar dynamics (Fig. 5A and Supplementary Fig. 8). However, IB appears to retain infectivity more efficiently than EB following physical and osmotic challenges. Specifically, when we subjected IB and EB Ot to physical stress using a bead mill for 1–2 min, we found that EB displayed a reduction in growth that was dependent on the time of exposure to stress; in contrast, IB was unaffected by this treatment (Fig. 5B). Furthermore, osmotic shock resulted in a significantly greater reduction in viability in EB compared with IB (Fig. 5C). Together, these results demonstrate that IB is physically more robust than EB; this property is likely due to differences in peptidoglycan content, as described above.

To investigate whether EB and IB forms of Ot enter host cells using the same invasion mechanism, we pre-treated HUVEC cells with the clathrin-mediated endocytosis inhibitor dynasore or the macropinocytosis inhibitors amiloride and zoniporide. We then infected the cells with IB or EB Ot and quantified the number of intracellular bacteria 3 h post-infection using confocal microscopy (Fig. 6A, B). The entry of IB Ot into the cells was blocked by dynasore, thus confirming the participation of clathrin-mediated endocytosis in this process. IB entry was also inhibited by amiloride and zoniporide, suggesting additional involvement of macropinocytosis in bacterial uptake. In contrast, EB Ot entry into cells was not affected by dynasore treatment but was completely inhibited by pre-incubation with amiloride or zoniporide. To confirm that EB Ot did not use clathrin-mediated endocytosis to enter cells, we treated HUVEC and L929 cells with siRNA against clathrin. Control microscopy experiments demonstrated that clathrin-targeting siRNA reduced the levels of clathrin foci in uninfected cells (Supplementary Fig. 9). We found that entry of IB form Ot required clathrin, whilst entry of EB form Ot did not (Fig. 6C, D and Supplementary Fig. 10). Together these results demonstrate that IB form Ot uses clathrin-mediated endocytosis to enter cells whilst EB form Ot does not.

To further demonstrate that IB and EB Ot were entering host cells using distinct mechanisms, we examined the role of the surface adhesins TSA56 and ScaC on bacterial entry. These are
outer membrane proteins that have been shown to be involved in entry of Ot into host cells, in experiments carried out using what we now describe as IB Ot stage bacteria \(^6,7\). We used antibodies targeting ScaC and TSA56 and determined whether these antibodies would block the entry of Ot into host cells. Our results with IB Ot replicated published studies and showed that blocking ScaC and TSA56 using antibodies led to a strong reduction in bacterial entry (Fig. 6E). With EB Ot, however, only a minimal effect was observed (Fig. 6E). This result is consistent with a model in which IB Ot enters host cells using adhesin-induced clathrin-mediated endocytosis, whilst in EB Ot the surface proteins are shielded by host derived membrane and the bacterium enters cells through macropinocytosis.

**Discussion**

Our studies have unveiled a new feature of the obligate intracellular bacterium Ot. This organism’s unique exit strategy, which involves budding off the surface of infected cells, is associated with differentiation into a molecularly and functionally distinct developmental state (Fig. 7). While unusual in the family Rickettsiaceae, this property is reminiscent of the biphasic lifecycle of multiple species in the sister family Anaplasmataceae \(^4\) and the unrelated obligate intracellular bacteria *Chlamydia* spp \(^18\). However, in contrast to these organisms, Ot resides in the host cytosol in naked form rather than in a membrane-bound vacuole. To our knowledge, this is the first description of an intracellular bacterium living freely in the host’s cytosol and having a biphasic lifecycle. Our findings have important implications for our understanding of Ot biology and unveil opportunities to develop better therapeutic strategies to treat scrub typhus.

The physical properties of prokaryotic cell walls are crucial for survival. These stiff structures maintain cellular shape, protect the interior, counteract osmotic pressure, and provide structural strength through peptidoglycan layers. Our studies revealed that EB and IB forms of Ot differ in peptidoglycan content. Lower levels in EB compared with IB make this form more susceptible to physical and osmotic stress. There is a long-standing question regarding the role of peptidoglycan in obligate intracellular bacteria. These organisms reside in an osmotically protective niche and are under selective pressure to reduce the abundance of immunogenic pathogen-associated molecular patterns (PAMP)s \(^19\). In the case of pathogenic *Chlamydia* species, it has been shown that peptidoglycan is retained only at the septum and is required for cell division \(^20\). Higher peptidoglycan levels in IB Ot vs. EB Ot point to the retention of a peptidoglycan-like structure for growth and division rather than structural integrity maintenance.

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**Fig. 6 IB and EB Ot enter host cells using distinct mechanisms.** Images (A) and quantification (B) of the effect of endocytosis inhibitors on entry of IB and EB Ot into HUVEC cells. IB and EB Ot were seeded onto fresh HUVEC cells that had been treated with inhibitors of clathrin-mediated endocytosis (100 \(\mu\)M dynasore) or macropinocytosis (50 \(\mu\)M amiloride or 100 \(\mu\)M zoniporide). Cells were fixed 3 h post-infection. Dynasore inhibits entry of IB but not EB, whilst amiloride and zoniporide inhibited entry of both IB and EB. Red = actin (phalloidin), blue = Hoechst (DNA), green = Ot (anti-TSA56 antibody). Scale bar = 10 \(\mu\)m. Similar results were found in L929 cells (Supplementary Fig. 10). Images (C) and quantification (D) of the effect of 0.2 \(\mu\)M siRNA against clathrin on entry of EB and IB Ot into HUVEC cells. Experiments were carried out as in A, B. In B, D, \(n = 30\) fields randomly imaged over 3 independent experiments. In each field the percentage of infected host cells was counted. Individual values and means are shown here. Statistical significance in B was determined by comparing all groups using one-way ANOVA, followed by a Dunnett’s multiple comparison tests with the control within the IB or EB group, and in D by performing a Mann–Whitney test comparing the scramble control and the clathrin siRNA treated condition for IB and EB separately. E Blocking the bacterial surface adhesins TSA56 and ScaC with anti-TSA56 and anti-ScaC antibodies strongly inhibits entry of IB but not EB Ot in HUVEC cells. Bacterial entry in the presence of vehicle control or antibodies was determined by quantifying the number of intracellular bacteria per host cell 30 min post-infection using immunofluorescence microscopy of bacteria labeled with HPG. Statistical significance was determined by comparing all groups using one-way ANOVA, followed by a Dunnett’s multiple comparison tests with the control within the IB or EB group. ***\(p \leq 0.0001\); ns not significant. *\(p \leq 0.05\); **\(p \leq 0.01\); ***\(p \leq 0.001\); ****\(p \leq 0.0001\).
The observed differences in physical properties between EB and IB forms are complemented by measurable differences in the accumulation of certain proteins, suggesting that each form performs unique functions or responds to specific challenges. Proteomics analyses in EB Ot revealed increased levels of heat shock sigma factor RpoH and the alarmone (p)ppGpp synthetase SpoT associated with the stringent response. These results suggest that the EB form of Ot shares features with the dormant state of other bacteria. Notably, Ot is the only reported member of the Rickettsiaceae to encode a full-length spoT gene. This exclusive feature is consistent with the fact that developmental differentiation of Ot is highly unusual in members of this family. The changes associated with differentiation into the EB state likely serve one or both of the following purposes: ensuring survival in the extracellular state and priming the bacteria for reinfection of new cells. Indeed, the observation that ScaA protein is synthesized early upon entry into host cells supports the idea that EB Ot is primed for infection.

Comparing the protein and RNA transcript levels of selected genes revealed a poor correlation for many genes. A potential explanation is that RNA levels are low in the metabolically inert EB form, which does not reflect newly translated proteins. Ot strain UT76 encodes only 8 predicted transcription factors and two sigma factors, RpoD and RpoH. Thus, it is possible that gene expression is not strictly regulated at the transcriptional level but that other forms of regulation are prevalent in Ot. We recently showed high antisense transcription levels in Ot, especially in genes associated with the rampant integrative conjugative element RAGE, which dominates the Ot genome. Notably, Ot is the only reported member of the Rickettsiaceae to encode a full-length spoT gene. This exclusive feature is consistent with the fact that developmental differentiation of Ot is highly unusual in members of this family. The changes associated with differentiation into the EB state likely serve one or both of the following purposes: ensuring survival in the extracellular state and priming the bacteria for reinfection of new cells. Indeed, the observation that ScaA protein is synthesized early upon entry into host cells supports the idea that EB Ot is primed for infection.

In summary, our finding that Ot differentiates into molecularly and functionally distinct forms provides a novel framework to study the infection cycle of this important obligate intracellular pathogen and opens avenues to investigate the physiological significance of the distinct populations characterized.

Methods

Cell culture, bacterial propagation, and reagents. Proteomics and qRT-PCR experiments were performed using the mouse fibroblast cell line L929 kindly provided by Dr. Blacksell at the Mahidol Oxford Tropical Medicine Research Unit, Bangkok, Thailand. For all other experiments, we used L929 cells (ATCC CCL-1), Primary Human Umbilical Vein Endothelial Cells (HUV-EC) (ATCC PCS-100-010) or monkey kidney epithelial Vero cells (ATCC CRL-1586) all purchased from ATCC. We cultured cells in either DMEM (L929) or Large Vessel Endothelial Supplement (LVES, HUV-EC) supplemented with 10% FBS (Gibco 16140071) at 37 °C and a 5% CO2 atmosphere. The Karp-like clinical isolate strain of Ot UT76 was used throughout the study. For routine propagation, bacteria were grown in 25- or 75-cm2 culture flasks, as described previously. We extracted DNA using alkaline lysis buffer, and then assessed bacterial quantities by determining genome copy numbers relative to a standard curve, using Biorad CFX Maestro manager software v3.1. We used a primer/probe set to amplify the single copy gene tsa47 (tsa47F 5′-TCCAGATTTAATGAGAATTAGGAC-3′, tsa47R 5′-TTAGTAATTACATCTCCAGGAGCAA-3′), the 16S gene (Rick16SF 5′-TTCGACATTGTGCTGCAGATCCTTC[TAM]-3′ or the 16S gene [Rick16SF 5′-GCTACACGGCTGCTACAATGG-3′, Rick16SR 5′-TGGTGAACGGCCGAGAACG-3′, Rick16SF 5′-6FAM] ATCGCTAGTAATCGCGGATCAGCA TGCC [TAM]-3′). For drug inhibition assays, dynasore (Sigma D7693) and amiloride (Sigma A3085) were added to L929 or HUVEC cells 1 h before infection with Ot.

Fluorescence microscopy

Immunofluorescence labeling and confocal microscopy. Following fixation, cells were permeabilized by incubation in 0.5% Triton X-100 for 30 min on ice followed by blocking in 1XPBS (pH7.4) containing 1% mg/ml BSA or sera (diluted 1:20) matching the labeled antibody’s host, for 30 min at room temperature. Incubations with primary antibodies took place for 1 h at 37 °C. These studies were conducted using in-house-generated antibodies (i.e., rabbit monoclonal antibody against TSA56 diluted 1:20; rabbit polyclonal antibody against Ot ScaC diluted 1:200) or commercial products (i.e., MA1-065, a ThermoScientific monoclonal mouse antibody against clathrin heavy chain diluted 1:200). The fixed cells were washed twice with PBS-BSA and were then incubated with appropriate secondary antibodies (1:1000 dilutions) for 30 min at 37 °C in the dark. We used the following secondary antibodies: goat anti-mouse IgG, SuperSonic Alexa Fluor 647 conjugate (ThermoFisher A28180), goat anti-rat Alexa Fluor 488 (ThermoFisher A11006), and...
goat anti-rabbit Alexa Fluor 594 (Thermo Fisher 11012). The nuclear stain Hoechst (1:1000 dilution) was supplemented to incubations with secondary antibodies. All antibodies used included Alexa Fluor 594 (1:200, Thermo Fisher Scientific A12381) was used to detect host cell. The cells were washed with 3X PBS before supple-
mentation of mounting medium [20 mM Tris (pH 8.0); 0.5% N-propyl-gallate; 90% glycerol]. Imaging was performed with a Zeiss Observer Z1 LSM700 confocal microscope equipped with an HBO 100 illuminating system equipped with a 63×/1.4 Plan-APOCHROMAT objective lens (Carl Zeiss, Germany) and 405 nm and 555 nm laser lines or a Leica TCS SP8 confocal microscope (Leica Microsystems, Germany) equipped with a 63×/1.4 Plan-APOCHROMAT objective lens with a 1.4-mm working distance and 405 nm, 488, 552, and 638 nm laser lines acquired using Leica TCS SP8 software.

Fluorescence intensity measurements were performed manually using Fiji ImageJ software. 30 (Fig.4C) or 100 (Fig.2B); individual bacteria were quanti-
ed by measuring the brightest pixel intensity within the bacterial area. Images from individual experiments were acquired under identical laser intensity and acquisition conditions. Length and width measurements were performed manually using Fiji ImageJ software.

STORM imaging and analysis. We acquired 3D-STORM data using a homebuilt multi-color Total Internal Reflection Fluorescence (TIRF) microscope that is based on an inverted microscope (Nikon, Ti-E), a high NA objective (Nikon, CFi-Apo 100X, NA 1.49), a 405 nm photoactivation laser (Coherent, OBI 405 100x), a 488 nm excitation laser (MPR, 2R-VPFL-P-20000642-BIR), and an EMCCD camera (Andor, iXon Ultra-888). Samples immunlabelled with Alexa 488 were mounted on the microscope, transferred to STORM buffer [0.05M Phosphate buffer (Sigma), 5.25 mM cysteine-
tetraetrae (COT)], and illuminated with a strong (408 nm) laser to record single-molecule fluorescence blinking events at a frame rate of 25 Hz. Illumina-
tion at 405 nm was intercalated with pulsed 488 nm excitation to enhance photoactivation of Alexa 488 without exacerbating background signal. A cylindrical lens (Thorlabs, L1516×-Å) was placed in front of the camera to determine the z-coordinates of fluorescent molecules through the elongated point spread function (PSF) methods. Each STORM image was reconstructed from single-molecule localization data obtained by analysing 30,000 frames with custom Matlab codes. Briefly, bright single fluorescence spots were identified by thresholding methods, and their 2-D coordinates were determined by 2-D elliptical Gaussian fitting along with a pre-calibrated conversion table between eccentricity and z-coordinate. Focus drift was actively stabilized with Perfect Focus System (Nikon) while acquiring the data, whereas xy-drift was post-
corrected using a redundant cross-correlation algorithm. Each localization event was represented as a 2-D Gaussian of 20 nm standard deviation in STORM images. For the analysis of bacterial morphology, we selected cells that showed clear cell boundary, manually adjusted the z-coordinate to locate the cell mid-
plane and measured long- and short-axes length.

Click labeling. Our metabolic click labeling was based on the Click-IT HPG Alexa Fluor protein synthesis Assay Kit (Click-IT HPG Alexa Fluor Protein Synthesis Assay Kit, ThermoFisher C10428) strategy. To incorporate L-homopropargylglycine (HPG), we incubated infected cells with minimal medium (Dubelco’s Modified Eagle Medium, DMEM, ThermoFisher 21031) lacking L-methionine and containing 25 µM HPG, for 30 min at 37 °C. Labeled bacteria were precipitated with PBS containing formaldehyde, and subsequently permeabilized with 0.5% Triton X-100 for 20 min on ice. After further washing with PBS containing 1 mg/ml BSA, we supplemented the Click-IT reaction cocktail (Click-IT HPG Alexa Fluor Protein Synthesis Assay Kit, ThermoFisher C10428) and incubated the cells for 30 min at room temperature in a light protected environment. The Azide dye (Alexa Fluor 488) was used at a final concentration of 5 µM. Following completion of the click reaction, the cells were subjected to immunofluorescence labeling and imaging as above.

Gene expression analysis by RT-qPCR. L929 cells were infected with Ot using multiple MOIs to a target MOI of 400:1 (quantitative PCR on freshly isolated Ot showed variation of actual MOIs ranging from 200:1 to 672:1 across three inde-
pendent infections). Very high MOIs were used due to the low levels of EB bacteria combined with the fact that RNA levels in EB Ot were very low. Infected cells were harvested 4 days post-infection and were stored in RNAProtect Bacteria Reagent (Qiagen 76506) at −80 °C. Total RNA was extracted using the QIAGEN RNeasy Plus Mini Kit (Qiagen 74136) according to the manufacturer’s instructions. One µg of total RNA was incubated with DNaseI (Thermo Fisher Scientific, AM2238) for 16 h at 37 °C or 37 °C with 1 U of RNase-free DNA-free RNase-free DNaseI (90 min) at room temperature in the dark, and digested with endonucleotides (Lysozyme; LysC: protein ratio = 1:50) overnight at 37 °C and, subsequently, with trypsin (trypsin: protein ratio = 1:1000, overnight with trypsin). The whole protein was used in each replicate without being normalized. For each sample, prior to the fact that the RNAs and EBs were isolated from different environment and therefore have diverse amounts of co-puri-
ed host protein. Samples were acidified with Trifluoroacetic acid (TFA) to 1% final concentration and centrifuged to remove insoluble materials. Trypsinized samples were desalted using an Oasis™ HLB 1cc Silica Gel cartridge (Waters) and then reverse-transcribed using the Advantage cDNA synthesis kit (clontech) and equilibrated with 500 µl of 0.5% acetic acid, we loaded samples and then washed the cartridge with 1 ml MS-grade water. Peptides were eluted with 500 µl of 80% acetonitrile in 0.5% acetic acid, then transferred to low-binding microtubes, and dried using a SpeedVac. Each peptide was resuspended with 20 µl of MS-grade water. We estimated peptide content using 5% of each sample, so-called Mini-runs. This was to check whether we need to adjust or normalize sample loadings in the actual runs. Briefly, individual aliquots in 1 µl of peptides

**Primer name**

| Primer name | Primer sequences (5’ to 3’) |
|-------------|----------------------------|
| spoT forward | GCCGACATATGTAAGCCATTAC |
| cyclC reverse | GAAATCTTGAAGCCCTGGCAG |
| rhoP forward | CTCGCGAGGAAGGATAATAGG |
| oglA reverse | CCCCTCGGAAATCTAAAACAA |
| ssdB forward | GCAAAATGTTAGAGCTGGT |
| dolA reverse | GAATCAGCAGCAATGTTG |
| miP2 forward | GTTGCAGCTTTGACCTACCC |
| miP2 reverse | TGATAGCTGACGATCCAG |

**Proteomics sample preparation.** We used Ot to infect L929 mouse fibroblasts at a 4:101 MOI in six independent culture flasks. High MOIs were used to ensure sufficient material for proteomics analysis. Despite the high MOI the L929 cells remained intact and attached to the surface of the flask at the point of harvesting the samples. We replaced the culture medium at 1-day post-infection (dpi) to remove bacteria that failed to enter the host. Intracellular bacteria were isolated at four dpi; two separate flasks were harvested as one replicate, yielding a total of three replicates. L929 cells were grown in 75-cm² plastic flasks at 37 °C in a 5% CO₂ atmosphere, using DMEM medium (Gibco 1196092) supplemented with 10% FBS (Sigm-Aldrich F7524). At four dpi, the media were replaced with 50 ml conical tubes for extracellular bacteria isolation. Culture flasks were rinsed with 15 ml of PBS, and cells were scraped up 2 ml of fresh media per flask. The detached cells were transferred to ml Safe-lock tubes (Eppendorf 0030120094) and homogenized for 3 min using a Bullet blender® homogenizer (Next Advance BBB24M) with speed control setting at 8. Host cell debris was removed by centrifugation at 300 × g for 3 min; we then incubated extracellular bacteria (EB) by an additional 3 min centrifugation at 20,000 × g. Before isolation of extracellular bacteria (EB), we removed possible detached host cells containing IB through a pre-spin step (1000 × g, 5 min, 4 °C). No visible pellet was observed indicating that there were negligible amounts of detached host cells potentially contaminating the EB prep with IB. Following transfer of supernatants to new conical tubes, we harvested extracellular bacteria by centrifugation at 3000 × g for 20 min at 4 °C. IBs and EBs were washed with 0.3 M sucrose three times, resuspended in PBS, and lysed with 1% Triton X-100 for 30 min at 4 °C. Bacterial proteins were precipitated by addition of 80% ice-cold acetone and overnight incubation at −20 °C. Protein pellets were washed in acetonitrile and resuspended in 10 µl Tris buffer (pH 8.0) to a final concentration and centrifuged to remove unsoluble materials. Trypsinized samples were desalted using an Oasis™ HLB 1cc Silica Gel cartridge (Waters) and then reverse-transcribed using the Advantage cDNA synthesis kit (clontech) and equilibrated with 500 µl of 0.5% acetic acid, we loaded samples and then washed the cartridge with 1 ml MS-grade water. Peptides were eluted with 500 µl of 80% acetonitrile in 0.5% acetic acid, then transferred to low-binding microtubes, and dried using a SpeedVac. Each peptide was resuspended with 20 µl of MS-grade water. We estimated peptide content using 5% of each sample, so-called “Mini-runs”. This was to check whether we need to adjust or normalize sample loadings in the actual runs. Briefly, individual aliquots in 1 µl of peptides.
were supplemented with 4 µl of 1% (v/v) acetonitrile containing 0.06% (v/v) trifluoroacetic acid and 0.5% (v/v) acetic acid, transferred to a 96-well autosampler plate, and then injected (3 µl) into an Orbitrap Fusion™ Mass Spectrometer. After we screened out host proteins from the Mini-runs result, the frequency distribution of Ot protein abundances were plotted. Medians of IB and EB replicates showed abundance values with maximum difference of 1 log scale, suggesting a single normalization process may be performed during data analysis. We decided to fractionate the rest of the samples without the need of normalizing sample loadings. The remaining 95% of each sample (19 µl in MS-grade water) was mixed with 20 µl of 10 mM ammonium formate (pH 10) and centrifuged at 20,000 x g for 3 min to eliminate insoluble components. Each peptide sample was then fractionated through a step-wise desalting process on C18 stage tips using 10, 15, 25, and 50% acetonitrile in 10 mM ammonium formate (pH 10). Fractionated samples were washed with 60% acetonitrile containing 0.1% formic acid twice to eliminate residual ammonium formate by evaporation before tandem MS analysis.

**LC-MS/MS.** Dried samples were resuspended in 10 µl of 2% (v/v) acetonitrile containing 0.06% (v/v) trifluoroacetic acid and 0.5% (v/v) acetic acid, and 1 µl was loaded onto an autosampler plate. Online chromatography was performed in an EASY-nLC 1000 (ThermoScientific) instrument using a one-column setup and 0.1% formic acid in water/0.1% formic acid in acetonitrile as mobile phases. The samples were injected and separated on a reversed-phase C18 analytical column (EASY-Spray LC Column, 75 µm inner diameter x 50 cm, 2 µm particle size; ThermoScientific) maintained at 50 °C and using a 2–23% (v/v) acetonitrile gradient over 60 min, followed by an increase by 50% of 60% acetonitrile for 60 min, and then to 90% over 5 min. The final mixture was maintained on the column for 5 min to elute all remaining peptides. Total run duration for each sample was 90 min at a constant flow rate of 300 nl/min.

Data were acquired using an Orbitrap Fusion Lumos™ mass spectrometer (ThermoScientific) using the instrument’s data-dependent mode and using Orbitrap Fusion Lumos Tune Application (v3.3.2782.23). Samples were ionized using 2.5 K and 300 °C at the nanospray source; positively-charged precursor MS1 signals were detected using an Orbitrap analyser set to 60,000 resolution, automatic gain control (AGC) target of 400,000 ions, and maximum injection time (IT) of 50 ms. Precursors with charges 2–7 and having the highest ion counts in each MS1 scan were further fragmented using collision-induced dissociation (CID) at 35% normalized collision energy. MS2 signals were analysed by ion trap at an AGC of 15,000 and maximum IT of 50 ms. Precursors used for MS2 scans were excluded for 90 s to avoid re-sampling of high abundance peptides. The MS1–MS2 cycles were repeated every 3 s until completion of the run.

**Proteomic data analysis.** Proteins were identified using Proteome Discoverer™ (v2.3, ThermoScientific) instrument using the one-column setup and 0.1% formic acid in water/0.1% formic acid in acetonitrile as mobile phases. The samples were injected and separated on a reversed-phase C18 analytical column (EASY-Spray LC Column, 75 µm inner diameter x 50 cm, 2 µm particle size; ThermoScientific) maintained at 50 °C and using a 2–23% (v/v) acetonitrile gradient over 60 min, followed by an increase by 50% of 60% acetonitrile for 60 min, and then to 90% over 5 min. The final mixture was maintained on the column for 5 min to elute all remaining peptides. Total run duration for each sample was 90 min at a constant flow rate of 300 nl/min.

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For label-free quantitation (LFQ), features were extracted from the raw data files using the Minora Feature Detector processing workflow node on Proteome Discoverer. Chromatographic matched ion feature picking was performed with a tolerance of 10 min to align the retention times and enable quantitation of the precursors. No pre-normalization was performed using the Proteome Discoverer workflow, and raw results were manually normalized using the Median normalization method. A protein was classified as detected if at least two peptides were detected in a minimum of two biological replicates; the mean LFQ across the three replicates was used for further analyses. There was no normalization during protein treatments and peptide injection as described earlier. Normalization between samples was performed only in the data analysis process. Briefly, frequency distributions of Ot protein abundances from six separate runs were plotted (excluded co-purified proteins). And the median from each plot was used to normalize the whole set, prior to data trimming, making all dataset scaled to the same median reference value of 1. Normalized abundance values were then averaged from independent biological replicates (Supplementary Data 1). Statistical significance (t test) was determined using GraphPad Prism software 8.4.3. Subsequently, proteins detected in both IBs and EBs for at least twice were selected for protein abundance comparison, described as Ext/Int ratio. Proteomic data is available from the jPOST repository with the identifier PXD028218.

**Immunoblot analyses.** L929 and HUVEC cells were grown in 24-well plates and infected with Ot at an MOI of 12:1. Noninfectious bacteria were removed by replacing the media 24 h post-infection. EB and IB populations were prepared 4 days post-infection. EB was put to an Eppendorf tube, pelleting at 16,873 x g, then resuspending in 100 µl PBS containing 1x protease inhibitor cocktail (ThermoFisher 78425). For TSAs, immunoblots, samples were prepared in the presence of 1 mM iodoacetamide (Sigma-Aldrich catalog number h6125). IB was prepared by removing adherent cells using trypsinization, spinning the cell pellet, and resuspending it in PBS supplemented with protease inhibitors; host cells were lysed using a bead mill for 1 min. Host cell debris was removed by centrifugation at 5000 x g for 5 min; the supernatant was transferred to a clean tube, spun at 16,873 x g for 5 min, and resuspended in 100 µl PBS supplemented with protease inhibitors. Bacterial protein copy number was quantified by qPCR using tsa47 primers, as described above. EB and IB samples were diluted to ensure equal loading of bacterial cells on gels. Samples were diluted in 3X SDS loading buffer with and without β-mercaptoethanol and heated at 95 °C for 5 min. Samples were subjected to electrophoresis on a 4–20% gradient gel (BioRad 4561094) and then transferred to a nitrocellulose membrane. Following blocking in PBST containing 5% BSA and 5% skim milk for 30 min, we incubated the blots with primary antibody (1:1000) in PBST-BSA for 1 h at room temperature. Membranes were washed, and immunoreactive proteins were visualized using a LiCor Odyssey CLX imager and analyzed using Image Studio V3.1.

**Quantification of genome copy number per cell.** The genome copy number per cell was calculated by determining the number of bacterial genome copies in a sample of purified EB and IB Ot, using tsa47 qPCR, and dividing by the number of bacterial cells in the same sample, as measured by confocal microscopy. Three µl of purified bacteria were spread evenly across a well of a multi-well slide, dried, and

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| Antibody name | Epitope used for immunization | Description of epitope | Species and type |
|---------------|-----------------------------|-----------------------|-----------------|
| TSA56         | KLTPPOQTIMPS1ADRDF—KLH     | Synthetic peptide from strain Karp conjugated to KLH carrier protein | Rat monoclonal  |
| RpoD          | MASLRKNNKNCNKKTCNNTVNEALSNVFDESE5KLSS5DSDIN | Recombinant protein M1-N103 from strain Karp | Rabbit polyclonal |
| Ctrl          | KGHEAVIRGVHGLSINLDRS5VSMYNCDQKNPIHNLTKYKILE | Recombinant protein K120-E145 from strain Karp | Rabbit polyclonal |
| SpoT          | LAAKKGTVLTKEMLSHSGLYDEPIEKIDFVCKRLKEKASNGINF | from strain Karp | Rabbit polyclonal |
| RpoH          | ETPWVGYRIGAEFSKNDNNSVNNVVEE | Recombinant protein E556-V661 from strain Karp | Rabbit polyclonal |
| ScaA          | NNTTESSDNQGDTGSGATSNPSNPSNPSEIGA5TPATGASDTGD | Recombinant protein N50-GT7 from strain Gillam | Rabbit polyclonal |
labeled for immunofluorescence microscopy using TSA56, as described above. The bacteria in a selected field were manually counted to determine the number of bacterial cells in the sample.

Quantification of relative peptidoglycan levels using NOD1 reporter assay
HEK-Blue hNOD1 cells (Invitrogen, hkb-hnodi, USA) were grown in DMEM containing 4.5 g/l glucose, 10% heat-inactivated FBS, 100 U/ml penicillin, 100 mg/ml streptomycin, 100 mg/ml Normocin, 2 mM L-glutamine, and the selective antibiotics blasticidin (30 µg/ml) and Zeocin (100 µg/ml). The cells were seeded on clear-bottom black 96-well plates (Corning 29444-008) 2 days before treatment. Bacterial aliquots from stocks kept in sacrose phosphate glutamate (SPG) at −80 °C were heat-inactivated at 90 °C for 30 min before supplementation to host cells. These studies were carried out in triplicate.

− were seeded on clear-bottom black 96-well plates (Corning 29444-008) 2 days 1 day prior to infection. To measure the effect of perturbing clathrin-mediated growth rate of IB and EB Ot from frozen stock were

Growth curve measurements and bead mill treatment
To determine the growth rate of IB and EB Ot from frozen stock were first grown in two T25 flasks containing a monolayer of L929 cells. At 6 days post-infection, IB and EB were prepared and used to inoculate fresh L929 cells and the subsequent bacterial growth was measured. 


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Author contributions

S.A., J.W., and S.G. contributed equally to this work and carried out the majority of experiments in this work. S.A. and S.G. carried out multiple experiments throughout the work whilst J.W. performed all proteomics experiments and analysis. K.S. carried out qRT-PCR experiments and Y.J. and S.C. performed Ot growth curve experiments. T.C. and H.H. carried out STORM microscopy experiments, whilst S.H.I. supervised that work and analyzed that data. L.C.W. worked with J.W. on proteomics sample preparation and analysis, whilst RMS supervised the proteomics work and assisted with the analysis. J.S. designed experiments, analyzed data, carried out experiments, and wrote the paper.

Competing interests

The authors declare no competing interests.

Additional information

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