



Development of a Plasmid Shuttle Vector System for Genetic Manipulation of Chlamydia psittaci

Kensuke Shima,^a Mary M. Weber,^b Christiane Schnee,^c Konrad Sachse,^d Nadja Käding,^a Matthias Klinger,^e Jan Rupp^{a,f}

^aDepartment of Infectious Diseases and Microbiology, University of Lübeck, Lübeck, Germany ^bDepartment of Microbiology and Immunology, University of Iowa Carver College of Medicine, Iowa City, Iowa, USA Institute of Molecular Pathogenesis, Friedrich-Loeffler-Institut (Federal Research Institute for Animal Health), Jena, Germany ^aRNA Bioinformatics and High-Throughput Analysis, Faculty of Mathematics and Computer Science, Friedrich-Schiller-Universität Jena, Jena, Germany elnstitute of Anatomy, University of Lübeck, Lübeck, Germany fGerman Center for Infection Research (DZIF), partner site Hamburg-Lübeck-Borstel, Germany

ABSTRACT The obligate intracellular bacterium Chlamydia psittaci is a known avian pathogen causing psittacosis in birds and is capable of zoonotic transmission. In human pulmonary infections, C. psittaci can cause pneumonia associated with significant mortality if inadequately diagnosed and treated. Although intracellular C. psittaci manipulates host cell organelles for its replication and survival, it has been difficult to demonstrate host-pathogen interactions in C. psittaci infection due to the lack of easy-to-handle genetic manipulation tools. Here, we show the genetic transformation of C. psittaci using a plasmid shuttle vector that contains a controllable gene induction system. The 7,553-bp plasmid p01DC12 was prepared from the nonavian C. psittaci strain 01DC12. We constructed the shuttle vector pCps-TetmCherry using the full sequence of p01DC12 and the 4,449-bp fragment of Chlamydia trachomatis shuttle vector pBOMB4-Tet-mCherry. pCps-Tet-mCherry includes genes encoding the green fluorescent protein (GFP), mCherry, and ampicillin resistance (AmpR). Target genes can be inserted at a multiple cloning site (MCS). Importantly, these genes can be regulated by a tetracycline-inducible (tet) promoter. Using the pCps-Tet-mCherry plasmid shuttle vector, we show the expression of GFP, as well as the induction of mCherry expression, in C. psittaci strain 02DC15, which belongs to the avian C. psittaci 6BC clade. Furthermore, we demonstrated that pCps-Tet-mCherry was stably retained in C. psittaci transformants. Thus, our C. psittaci plasmid shuttle vector system represents a novel targeted approach that enables the elucidation of host-pathogen interactions.

IMPORTANCE Psittacosis, caused by avian C. psittaci, has a major economic impact in the poultry industry worldwide and represents a significant risk for zoonotic transmission to humans. In the past decade, the tools of genetic manipulation have been improved for chlamydial molecular studies. While several genetic tools have been mainly developed in Chlamydia trachomatis, a stable gene-inducible shuttle vector system has not to date been available for C. psittaci. In this study, we adapted a C. trachomatis plasmid shuttle vector system to C. psittaci. We constructed a C. psittaci plasmid backbone shuttle vector called pCps-Tet-mCherry. The construct expresses GFP in C. psittaci. Importantly, exogeneous genes can be inserted at an MCS and are regulated by a tet promoter. The application of the pCps-Tet-mCherry shuttle vector system enables a promising new approach to investigate unknown gene functions of this pathogen.

KEYWORDS Chlamydia psittaci, Gram-negative bacteria, intracellular bacteria, plasmid shuttle vector, transformation

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Address correspondence to Kensuke Shima. kensuke.shima@uksh.de.

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The obligate intracellular Gram-negative bacterium *Chlamydia psittaci* (*C. psittaci*) is an important zoonotic pathogen that can be encountered in more than 400 bird species, as well as in sheep, cattle, swine, horses, goats, and cats (1–3). When avian *C. psittaci* strains infect humans, they can cause severe atypical pneumonia, with even a fatal outcome in some cases. In contrast, nonavian *C. psittaci* strains might be excluded as potential zoonotic risk factors since the case reports of human psittacosis usually include an avian source (1–3).

In host cells, chlamydiae undergo a unique developmental cycle that alternates between two distinct bacterial forms, infectious elementary bodies (EBs) and the replicating reticulate bodies (RBs). The developmental cycle is completed within the confines of a membrane-bound vacuole termed the inclusion (4). During infection, chlamydiae interact with various host cell organelles to acquire host-derived nutrients for their survival (5–7).

Genome sequence analysis of *C. psittaci* revealed a circular 1.2-Mb chromosome and an approximately 7.5-kb plasmid harboring seven to eight putative coding DNA sequences (CDS) (1, 8–10).

The lack of tools for genetic manipulation of chlamydiae hampered molecular research progress for many years. However, Binet and Maurelli first demonstrated transformation of *C. psittaci*, showing successful allelic exchange on the bacterial chromosome (11). Afterward, Wang et al. demonstrated a stable targeted genetic modification method of *C. trachomatis* using a plasmid shuttle vector (12). Their system has been modified for other *Chlamydia* species (spp.) including *C. pneumoniae* and *C. muridarum*, but not *C. psittaci* (12–14). To replicate efficiently, the plasmid shuttle vector usually has to match the chlamydial plasmid with the target host species due to the presence of replications barriers (13–15). In this study, we established a stable targeted genetic modification system for *C. psittaci* infections.

RESULTS

Construction of a plasmid shuttle vector for transformation of *C. psittaci.* Plasmid DNA sequence comparison of a nonavian isolate *C. psittaci* 01DC12 (GenBank: HF545615.1) shares 99.97% identity to the avian isolate *C. psittaci* 6BC (GenBank: CP002587.1) (Fig. S1 in the supplemental material). In the amino acid sequence of protein coding regions, only one amino acid is different between pCps6BC and p01DC12 (Fig. S1). Due to the sequence similarity and biosafety considerations, the p01DC12 plasmid derived from *C. psittaci* 01DC12 was selected for construction of the shuttle vector.

We constructed a 12,008-bp pCps-Tet-mCherry shuttle vector using a fragment of pBOMB4-Tet-mCherry, including genes for the green fluorescent protein (GFP), mCherry, and ampicillin resistance (AmpR) (16), along with the full sequence of the p01DC12 plasmid derived from *C. psittaci* 01DC12 (Fig. 1A). mCherry can be replaced with various target genes at a multiple cloning site (MCS) using restriction digestion. Target genes can be regulated by a tetracycline-inducible (tet) promoter (Fig. 1B).

The expected plasmid size of 12,008 bp was confirmed by digestion of pCps-TetmCherry with Notl (Fig. S2). Sequencing of the pCps-Tet-mCherry plasmid shuttle vector revealed that the full sequence of p01DC12 was 100% identical to the original sequence (GenBank: HF545615.1). The origin of replication in pCps-Tet-mCherry was identical to pGFP::SW2 (12). The amino acid sequence of mCherry was identical to that of the pMCherry-C1 vector (TaKaRa Bio, Saint-Germain-en-Laye, France). Another region of the pBOMB4-Tet-mCherry fragment was 100% identical to the literature data (Gen-Bank: KF790910.1) (16).

Since our plasmid shuttle vector was constructed from a plasmid of *C. psittaci* 01DC12, we selected the same strain as a control to perform our initial transformation. After transformation of *C. psittaci* 01DC12 with pCps-Tet-mCherry and infection in epithelial cells, a strong GFP signal was detected in *C. psittaci* 01DC12 (*C. psittaci* 01DC12-pCps-Tet-mCherry) inclusions at 24 and 48 h postinfection (hpi) (Fig. S3A). Furthermore, mCherry was successfully induced by both 10 and 100 ng/ml anhydro-





FIG 1 Map of the *C. psittaci*-derived shuttle vector pCps-Tet-mCherry. (A) The CDSs of p01DC12 are shown in orange. GFP, mCherry, and the Tet repressor are shown in green, red, and light blue, respectively. AmpR, pUC ori, and MCIP are shown in light gray. The full sequence of p01DC12 was amplified from *C. psittaci* 01DC12. Another fragment was amplified from pBOMB4-tet-mCherry. mCherry can be induced by a tetracycline-inducible promoter. (B) An MCS containing Notl, Pstl, Kpnl, and Sall restriction sites in pCps-Tet-mCherry.

tetracycline hydrochloride (aTC) treatment, and we did not detect mCherry expression in the absence of aTC (Fig. S3A and S3B).

The bovine strain *C. psittaci* 02DC15 and the psittacine isolate *C. psittaci* 6BC are categorized in the same *ompA* genotype A. Genetically, *C. psittaci* 02DC15 belongs to the psittacine *C. psittaci* 6BC clade (1, 17).

Due to their genome sequence similarity and biosafety considerations, we next investigated whether *C. psittaci* strain 02DC15 could be transformed with pCps-Tet-mCherry. Following transformation, we observed not only expression of GFP, but also

induction of mCherry in *C. psittaci* 02DC15 (*C. psittaci* 02DC15-pCps-Tet-mCherry) inclusions using 10 and 100 ng/ml of aTC (Fig. 2A and B).

Impact of pCps-Tet-mCherry on the growth of *C. psittaci.* We investigated whether the transformation of *C. psittaci* using pCps-Tet-mCherry affected chlamydial growth and morphological characteristics (Fig. 3 and Fig. S4). While penicillin (PEN) treatment slightly reduced the growth of *C. psittaci* 02DC15-pCps-Tet-mCherry compared to wild-type *C. psittaci* 02DC15, *C. psittaci* 02DC15-pCps-Tet-mCherry showed similar growth characteristics compared to wild-type *C. psittaci* 02DC15 and 100 ng/ml of aTC did not inhibit the growth of *C. psittaci* 02DC15-pCps-Tet-mCherry (Fig. 3A). Moreover, we confirmed both 10 and 100 ng/ml of aTC did not inhibit the growth of *C. psittaci* 02DC15-pCps-Tet-mCherry (Fig. 3A). The same trend was observed using the control strain *C. psittaci* 01DC12 (Fig. S4A). Immunofluorescence analysis revealed similar inclusion morphology between wild-type and transformed *C. psittaci* regardless of the presence or absence of aTC (Fig. 3B and Fig. S4B). These data indicate that pCps-Tet-mCherry itself and aTC treatment at concentrations used in this study do not interfere with chlamydial growth characteristics.

Plasmid retention in transformed *C. psittaci.* The average of the endogenous plasmid copy number is 2.0 \pm 0.2 per chromosome in the wild-type *C. psittaci* 02DC15 at 48 hpi (Fig. S5A). In contrast, *C. psittaci* 02DC15-pCps-Tet-mCherry harbors 3.6 \pm 0.1 of the pCps-Tet-mCherry plasmid shuttle vector per chromosome (Fig. S5B). Transformation resulted in 98 \pm 0.1% (n = 3) reduction of endogenous wild-type plasmid in *C. psittaci* 02DC15-pCps-Tet-mCherry. A similar trend was observed using the control strain *C. psittaci* 01DC12 (Fig. S5A and B). While the endogenous plasmid copy number is 1.1 \pm 0.2 per chromosome in the strain *C. psittaci* 01DC12 at 48 hpi (Fig. S5A), *C. psittaci* 01DC12-pCps-Tet-mCherry harbors 3.0 \pm 0.3 pCps-Tet-mCherry plasmid shuttle vector per chromosome (Fig. S5B) and shows a decrease of 96 \pm 0.8% of the endogenous plasmid compared to the wild-type strain.

Plasmids are generally prone to being lost without selection pressure. We next investigated the shuttle vector/plasmid stability in transformed *C. psittaci*. Plasmid shuttle vector pCps-Tet-mCherry was stably maintained in both *C. psittaci* 02DC15-pCps-Tet-mCherry and the control strain *C. psittaci* 01DC12-pCps-Tet-mCherry in the presence of PEN (Fig. 4A and Fig. S6A). Although copy numbers were similar in the presence or absence of PEN at initial culture, pCps-Tet-mCherry was significantly reduced in the absence of PEN from passage 1 (Fig. 4A and B, Fig. S6A and B). This indicates that the pCps-Tet-mCherry plasmid could be stably retained with selection pressure for long-term maintenance.

Impact of aTC on host cell metabolism. It is known that a long-term treatment of tetracycline inhibits host cell metabolism, especially mitochondrial function (18). Therefore, we investigated the impact of aTC on mitochondrial activity.

aTC (10 ng/ml) at the concentration used in this study attenuated host cell mitochondrial activity, as indicated by basal respiration, ATP production, and maximal respiration (Fig. 5A). This indicates that we have to take into account the effect of aTC on host cell functions. However, when we compared mitochondrial activity in aTCtreated uninfected control cells to aTC-treated *C. psittaci* 02DC15-pCps-Tet-mCherry infected cells, significantly upregulated mitochondrial activity was observed in aTCtreated *C. psittaci* 02DC15-pCps-Tet-mCherry infected cells (Fig. 5B and C). This upregulation was similar to that observed in wild-type *C. psittaci* 02DC15 inclusion-bound mitochondria (Fig. S7A and B).

DISCUSSION

C. psittaci has the capability of causing severe disease in animals and humans (1–3). However, due to the lack of genetic manipulation tools for this important pathogen, molecular studies of *C. psittaci* are lagging far behind that of other human chlamydial pathogens, such as *C. trachomatis*.

Development of chlamydial shuttle vectors and stable transformation of *C. trachomatis*, *C. muridarum*, and *C. pneumoniae* have been established only in the last decade (12–14, 19). Bauler and Hackstadt further developed a novel plasmid shuttle vector





FIG 2 GFP expression and mCherry induction in pCps-Tet-mCherry-transformed *C. psittaci* strain 02DC15. Transformed *C. psittaci* 02DC15 bacteria were grown in epithelial cells with 1U/ml PEN for 24 and 48 h. (A) GFP fluorescence of chlamydial inclusions was visualized in living cells without fixing and staining. aTC (10 and 100 ng/ml) was added at 1 hpi to induce mCherry expression. Images are representative of three independent experiments. White arrows show chlamydial inclusions. White scale bars represent 10 μ m. (B) mCherry was analyzed by Western blotting and densitometric analyses at 24 and 48 hpi. mCherry protein amounts were normalized to chlamydial HSP60. GAPDH was used as a loading control. (*n* = 4; meen ± SEM; Sidak's multiple comparison: ***, *P* ≤ 0.001).





FIG 3 One-step growth curve and inclusion morphology of pCps-Tet-mCherry-transformed and untransformed *C. psittaci* strain 02DC15. (A) *C. psittaci* strain 02DC15 transformed or untransformed with pCps-Tet-mCherry was grown in epithelial cells with or without 10 U/ml PEN and 0, 10, or 100 ng/ml of aTC. Recoverable *C. psittaci* 02DC15 IFUs were determined at 5, 24, 48, and 72 hpi. The numbers of recoverable *C. psittaci* 02DC15 under each condition (II to VIII) at the indicated time were compared to those of untransformed *C. psittaci* 02DC15 without PEN (I). (n = 3 to 6; mean \pm SEM; Sidak's multiple comparison: **, $P \le 0.01$; ***, $P \le 0.001$). (B) Representative immunofluorescence images of pCps-Tet-mCherry-transformed *C. psittaci* 02DC15 at 5, 24, 48, and 72 hpi. Chlamydial inclusions were stained by FITC-labeled monoclonal chlamydial-LPS antibodies. Evans blue counterstaining of host cells was used for better characterization of intracellular inclusions. Images are representative of 3 to 6 independent experiments. White scale bars represent 10 μ m.

pBOMB4-Tet-mCherry, which encodes the Tet-inducible promoter system for the expression of recombinant target proteins in *C. trachomatis* (16). Moreover, Weber et al. were able to identify 10 novel inclusion membrane proteins in *C. trachomatis* infection using this shuttle vector system (20).

Considering there are some barriers to plasmid replication (13–15) and the advantages of pBOMB4-Tet-mCherry reporter system, the *C. psittaci* backbone plasmid shuttle vector pCps-Tet-mCherry was constructed from the full sequence of p01DC12 and the





FIG 4 The pCps-Tet-mCherry plasmid can be stably maintained by *C. psittaci* strain 02DC15 and expresses GFP. (A) pCps-Tet-mCherry-transformed *C. psittaci* 02DC15 bacteria were subcultured in epithelial cells in the presence or absence of 10 U/ml PEN every 2 days over 5 passages. Quantitative PCR was performed using primers specific to genomic DNA or to the pCps-Tet-mCherry plasmid. Copy numbers of pCps-Tet-mCherry were normalized to genomic DNA at each passage in the presence or absence of PEN, and were compared to passage 0 of *C. psittaci* 02DC15 in the presence of PEN. (*n* = 3; mean \pm SEM, Sidak's multiple comparison: *, *P* \leq 0.05; **, *P* \leq 0.01; ***, *P* \leq 0.001). (B) Representative GFP and immunofluorescence images of pCps-Tet-mCherry-transformed *C. psittaci* 02DC15 cells 48 hpi at passage 5. After a GFP signal was detected by fluorescence microscopy, the cells were fixed by methanol. Then, chlamydial inclusions were stained by FITC-labeled monoclonal chlamydial-LPS antibodies. Evans blue counterstaining of host cells was used for better characterization 02DC15 that lost pCps-Tet-mCherry during passages. White scale bars represent 20 μ m.

fragment of *C. trachomatis* shuttle vector pBOMB4-Tet-mCherry. We showed that GFP was expressed after transformation of the control strain *C. psittaci* 01DC12 using pCps-Tet-mCherry. Since avian *C. psittaci* strains cause severe lung diseases in humans, it is important to assess whether pCps-Tet-mCherry system works in avian *C. psittaci* strains, including *C. psittaci* 6BC. However, conducting the experiment using *C. psittaci* 6BC is generally troublesome due to stricter regulations for the biosafety level. Therefore, we used *C. psittaci* 02DC15 as a substitute for avian *C. psittaci* 6BC. While whole-genome analysis using UpSetR (21) revealed a total of 984 CDS in *C. psittaci* 6BC





FIG 5 Analysis of mitochondrial activity in cells treated with aTC and infected with *C. psittaci* 02DC15pCps-Tet-mCherry. (A) The effect of aTC (10 ng/ml) on mitochondrial activity in noninfected epithelial cells. Mitochondrial activity shown by oxygen consumption rate (OCR) was measured by a Mito Stress test kit at 24 hpi. I, II, and III indicate basal respiration, ATP production, and maximal respiration, respectively. (B) Phase contrast image of control cells and image overlay of phase contrast and GFP or mCherry in pCps-Tet-mCherry-transformed *C. psittaci* 02DC15-infected cells with 10 ng/ml of aTC. White scale bars represent 20 μ m. (C) Mitochondrial activity in control cells and pCps-Tet-mCherry-transformed *C. psittaci* 02DC15-infected cells with 10 ng/ml aTC at 24 hpi. Arrows show sequential injection of different chemical compounds: O, oligomycin; F, FCCP; A & R, antimycin A plus rotenone. I, II, and III indicate basal respiration, ATP production, and maximal respiration, respectively. Images are representative of three independent experiments. (A: n = 13 from five independent experiments, C: n = 13 from five independent experiments; mean \pm SEM, Student's *t* test: ***, $P \le 0.001$).

and 991 CDS in *C. psittaci* 02DC15 genomes (Fig. S8), we were able to show that all of the genes present in *C. psittaci* 6BC are encountered in *C. psittaci* 02DC15 as well, thus indicating their genetic equivalence. Due to the genome similarity between *C. psittaci* 6BC and *C. psittaci* 02DC15, *C. psittaci* 02DC15 is widely used for a number of immunological and cell biological studies, as well as animal infection trials (22–28). Reinhold et al. established a bovine respiratory model of acute *C. psittaci* infection using strain 02DC15 (27). Koch-Edelmann et al. demonstrated the importance of lipid metabolism in *C. psittaci* 02DC15 infection (25). Dutow et al. elucidated the role of effector of the complement (C3a) in *C. psittaci* 02DC15 infection (22). Most recently, Radomski et al. reported that *C. psittaci* 02DC15 infection of dendritic cells enhances exosome release, resulting in strong induction of IFN- γ production by natural killer cells and enhancement of apoptosis (24). Therefore, successful transformation using *C. psittaci* 02DC15 is particularly meaningful with respect to the biology of *C. psittaci* infection.

In addition to transformation of *C. psittaci* 02DC15, the induction of mCherry expression was controlled by aTC treatment. In this study, mCherry was used as an exogenous gene, but it is possible to insert other target genes into the MCS. Therefore, the pCps-Tet-mCherry is a useful plasmid shuttle vector to investigate the function of specific proteins during *C. psittaci* infection.

A tetracycline-inducible system is beneficial for biological research. However, it has been demonstrated that >100 ng/ml of tetracycline or its derivatives attenuates mitochondrial activity (18, 29, 30). Since mitochondrial function is essential for chlamydiae to sustain their life (8, 31, 32), we investigated the impact of aTC on mitochondria in *C. psittaci* infection.

Oxygen consumption of *C. psittaci* 02DC15-pCps-Tet-mCherry-infected cells was measured by a Seahorse XF analyzer with solid state sensor probes containing polymer embedded fluorophores (33). The peak absorption and emission of the oxygen sensor are 530 and 650 nm, respectively. Although *C. psittaci* 02DC15-pCps-Tet-mCherry expresses GFP and mCherry, intracellular dye fluorescence does not interfere with optical sensors given enough distance to the object, as various groups have demonstrated (34, 35).

Similar to previous studies (18, 29, 30), 10 ng/ml of aTC attenuated the mitochondrial oxygen consumption rate in noninfected epithelial cells. However, we could show that mitochondrial activity was significantly upregulated in *C. psittaci* 02DC15-pCps-Tet-mCherry-infected cells under aTC treatment, similar to that observed in wild-type *C. psittaci* 02DC15 infection in the absence of aTC. These data indicate that pCps-TetmCherry can be a beneficial tool for further biological studies in *C. psittaci* infection, though we need to take into account the impact of aTC on host cell functions.

Recently, gene knockout techniques such as the TargeTron system have also been established in *C. trachomatis* (36, 37). Our plasmid shuttle vector pCps-Tet-mCherry and the future development of a knockout system are promising new approaches to elucidate chlamydial protein functions during *C. psittaci* infection.

Taken together, our findings highlight that pCps-Tet-mCherry can be used for *C. psittaci* transformation and functional studies. This system will enable the identification of novel virulence factors and improvement of treatment strategies for *C. psittaci* infection.

MATERIALS AND METHODS

Bacterial strains, epithelial cells, chemicals, and antibodies. Animal-derived isolates *C. psittaci* 01DC12 (*ompA* genotype E, pigeon clade) and *C. psittaci* 02DC15 (*ompA* genotype A, psittacine clade) were provided by the National and OIE Reference Laboratory for Chlamydioses at FLI Jena (1). HeLa cells (ATCC CCL-2)/HEp-2 cells (ATCC CCL-23) were used for the growth of *C. psittaci*. All chemicals were purchased from Sigma-Aldrich (Deisenhofen, Germany). Rabbit anti-mCherry and mouse anti-heat shock protein 60 (HSP60) were purchased from Thermo Fisher Scientific (Waltham, MA). Rabbit anti-glyceraldehyde-3-phosphate dehydrogenase (GAPDH), horseradish peroxidase (HRP)-horse anti-mouse IgG and HRP-goat anti-rabbit IgG were purchased from Cell Signaling Technology (Frankfurt am Main, Germany).

Cell culture medium. Dulbecco modified Eagle medium (DMEM) was supplemented with 10% fetal bovine serum (FBS) (Invitrogen), 1 mM sodium pyruvate (Pan-Biotech GmbH, Aidenbach, Germany) and 30 mM HEPES. RPMI 1640 medium was supplemented with 5% FBS (Invitrogen), nonessential amino acids (Gibco/Thermo Fisher Scientific), and 2 mM L-glutamine (Lonza, Walkersville, MD).

Construction of *C. psittaci***-derived plasmid shuttle vector.** The plasmid vector pCps-Tet-mCherry was constructed using NEBuilder HiFi DNA assembly cloning kit (New England BioLabs [NEB], Ipswich, MA). Using Phusion Hot Start II High-Fidelity DNA polymerase (Thermo Fisher Scientific), the full sequence of p01DC12 was amplified from DNA of *C. psittaci* 01DC12 (7,553 bp, GenBank: HF545615.1). The fragment, including the tet promoter, mCherry, MCS (NotI, PstI, KpnI and Sall), AmpR, pUC origin of replication (ori), the meningococcal class I protein promoter (MCIP) derived from *Neisseria meningitidis* MC50, and GFP, was amplified from pBOMB4-Tet-mCherry (16). The plasmid vector pBOMB4-Tet-mCherry was kindly provided by Ted Hackstadt (National Institutes of Allergy and Infectious Disease, NIH, Hamilton, MT, USA). These two segments were assembled following the NEB manufacturer's instructions, resulting in pCps-Tet-mCherry. Primer sets are listed in Table S1.

Genetic transformation. Transformation was performed as described in a previous study (13). pCps-Tet-mCherry was extracted from a Dam- and Dcm-methylase-deficient strain, *E. coli* GM2163, using a Qiagen plasmid mega kit (Hilden, Germany) (12). PEN (1U/ml) was used for the selection of transformed *C. psittaci*. A mixture of *C. psittaci* 01DC12 or 02DC15 1 × 10⁸ inclusion forming units (IFU) and 15 μ g of pCps-Tet-mCherry were incubated in 200 μ l calcium chloride buffer (10 mM Tris, 50 mM calcium chloride, pH 7.4) for 30 min at room temperature. Then, 200 μ l of *C. psittaci* mixed with pCps-Tet-mCherry were incubated 8 × 10⁶ epithelial cells in calcium chloride buffer for 20 min at room temperature with mild agitation.

The total volume (400 μ l) of the mixture of epithelial cells, pCps-Tet-mCherry, and *C. psittaci* 01DC12 or 02DC15 was distributed over 4 wells (each 100 μ l) containing 2 ml DMEM medium and 1 μ g/ml cycloheximide in a 6-well plate and incubated at 37°C under 5% CO₂. At 24 hpi, 1U/ml PEN was added

to each well and further incubated for 48 h at 37° C under 5% CO₂. Afterward, infected epithelial cells were scraped and lysed with glass beads.

Freshly prepared epithelial cells were infected with transformed *C. psittaci* in 6-well plates containing 5 ml DMEM medium with 1 μ g/ml cycloheximide and 1U/ml PEN. Passages were performed every 2 to 3 days. GFP expression was observed by passage 2.

Induction of mCherry. A total of 2×10^5 epithelial cells in 24-well plates were infected with 2×10^5 IFU of transformed *C. psittaci* in DMEM medium. At 1 hpi, 10 or 100 ng/ml of aTC was added to induce mCherry.

Western blot analysis. To determine the amount of mCherry protein, 1×10^6 cells were seeded in 6-well plates (Greiner bio-one, Frickenhausen, Germany) and infected with *C. psittaci* 02DC15-pCps-Tet-mCherry or *C. psittaci* 01DC12-pCps-Tet-mCherry (1 IFUs/cell). At 1 hpi, 10 or 100 ng/ml of aTC was added to induce mCherry. *C. psittaci*-infected cells were lysed by 8 M urea supplemented with 325 U/ml Benzonase nuclease (Sigma-Aldrich) at the indicated times. Cell lysates were diluted into Laemmlie buffer (50 mM Tris-HCl pH 6.8, 2% SDS, 1% 2-mercaptoethanol, 10% glycerol, 0.1% bromophenol blue). Samples were analyzed by Western blot analysis (Bio-Rad, Hercules, CA) and visualized using enhanced chemiluminescence (ECL) reagent (Millipore and Thermo Fisher Scientific). Images were acquired by Fusion FX7 (Vilber Lourmat, Eberhardzell, Germany) and the density of each band was measured by Bio-1D software (Vilber Lourmat).

Recovery assay. A total of 5×10^4 epithelial cells in 1 ml DMEM medium were seeded into 24-well plates (Greiner bio-one) and cultured overnight at 37°C under 5% CO₂. For the infection, 2×10^5 IFUs of either untransformed or pCps-Tet-mCherry-transformed *C. psittaci* strains as well as cycloheximide (1 µg/ml) were added to each well. When appropriate, PEN (10 U/ml) and 0, 10, or 100 ng/ml of aTC were added at 1 hpi. The plate was centrifuged at 700 × g for 1 h at 35°C and incubated for 5, 24, 48, and 72 h. After the indicated time, the cells were subsequently cultured for 48 h for determination of the recoverable *C. psittaci*.

Real-time quantitative PCR. Determination of plasmid copy number was performed as demonstrated previously (16). Wild type or pCps-Tet-mCherry-transformed *C. psittaci* were cultured in epithelial cells in the presence or absence of 10 U/ml PEN. Afterward, isolated *C. psittaci* were boiled in 20 mM dithiothreitol (DTT) for 15 min and the supernatant was collected to obtain the genomic and plasmid DNA mixture. One primer set was designed to detect the *hctA* gene (GenBank, AEG87858.1; CPS0B_0937) in the genome. Another primer set was designed to detect the *GFP* gene for quantitation of the pCps-Tet-mCherry plasmid shuttle vector. For the endogenous plasmid, primers were designed to detect CDS5 or the CDS1-CDS2 junction, which is separated by the fragment of pBOMB4-Tet-mCherry. Real-time quantitative PCR was performed using LightCycler 480 SYBR green I Master on LightCycler 480 II (Roche Molecular Biochemicals, Mannheim, Germany). Plasmid copy number was calculated using the threshold cycle ($2^{\Delta \Delta C7}$) method (38) relative to genomic DNA.

Plasmid stability. A total of 3×10^5 epithelial cells in 1 ml DMEM medium were initially infected with pCps-Tet-mCherry-transformed *C. psittaci* at 0.5 IFUs/cell in the presence or absence of 10 U/ml PEN. The infected cells were subcultured every 2 days for 5 times. From the second passage, epithelial cells were infected with serial dilutions of *C. psittaci* at 0.5 to 0.8 IFUs/cell. To determine plasmid copy number, extracted genomic and plasmid DNA mixture was analyzed as described above. At passage 5, the GFP signal was detected by fluorescence microscopy and cells were fixed by methanol. Afterward, chlamydial inclusions were stained by fluorescein isothiocyanate (FITC)-labeled monoclonal chlamydial lipopolysac-charide (LPS) antibodies.

Fluorescence microscopy. A fluorescence microscope Keyence BZ-9000 (Keyence, Osaka, Japan) was used to detect the fluorescence signal of GFP or mCherry expressed in *C. psittaci*. In addition, untransformed or pCps-Tet-mCherry-transformed *C. psittaci*-infected cells were analyzed by immunofluorescence staining with mouse anti-chlamydial LPS antibody (green), which stained chlamydial inclusions. Evans blue was used for counterstaining of host cells (red).

Transmission electron microscopy. *C. psittaci* 02DC15-infected cells were fixed with 2% paraformaldehyde and 2.5% glutaraldehyde in 0.1 M cacodylate buffer for 1 h. Postfixation was performed with 1% OsO_4 in 0.1 M cacodylate buffer for 2 h. Samples were dehydrated with a graded ethanol series and embedded in araldite (Fluka, Buchs, Switzerland). Ultrathin sections were stained with uranyl acetate and lead citrate and were examined with a JEOL 1011 transmission electron microscope (TEM) (JEOL, Tokyo, Japan).

Plasmid sequencing. Extracted pCps-Tet-mCherry plasmid was sequenced at Eurofins Genomics (Ebersberg, Germany). Primer sets are listed in Table S1.

Plasmid maps. Plasmid maps were made by SnapGene. Using CLUSTALW in GenomeNet (http:// www.genome.jp/), p01DC12 plasmid sequences were compared to pCps6BC.

Genome comparion of *C. psittaci* **02DC15 and** *C. psittaci* **6BC.** UpSetR package was used in the genome comparison (21). A total of 13 genomes (12 species of *Chlamydia*) and their homologous gene groups identified with genome analysis pipeline RIBAP were selected as input for UpSetR.

Metabolic analysis. A total of 2 × 10⁴ HeLa cells in RPMI 1640 were treated with 10 ng/ml aTC. In addition, cells were infected with untransformed or pCps-Tet-mCherry-transformed *C. psittaci* without cycloheximide in 24-well XF plates (Agilent, Santa Clara, CA). Plates were centrifuged at 700 × *g* for 1 h at 35°C and incubated for 24 h. Afterward, Mito Stress test kits were used following Seahorse Bioscience manufacturer's instructions with chemical concentrations as follows: oligomycin (0.5 μ M), FCCP (0.2 μ M), and antimycin A (1 μ M) plus rotenone (1 μ M). Before the assay of pCps-Tet-mCherry transformed-*C. psittaci* infected cells, expression of GFP and mCherry was detected by fluorescence microscopy BZ-9000 (Keyence).



Statistics. Data are indicated as means \pm standard error of the mean (SEM). Statistical analyses were performed by GraphPad Prism 7 statistical software. When three or more groups were compared in the experiment, Sidak's multiple comparison was used in cases where one-way analysis of variance showed statistical significance (*P* values \leq 0.05). Data between two groups were evaluated using Student's *t* test. In Sidak's multiple comparison and Student's *t* test, *P* values of \leq 0.05 were considered statistically significant.

Data availability. The sequence of the pCps-Tet-mCherry plasmid shuttle vector reported is available in the DDBJ/EMBL/GenBank databases under accession number LC548057.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, TIF file, 0.3 MB. FIG S2, TIF file, 0.1 MB. FIG S3, TIF file, 1.5 MB. FIG S4, TIF file, 2.4 MB. FIG S5, TIF file, 0.2 MB. FIG S6, TIF file, 1.6 MB. FIG S7, TIF file, 1.1 MB. FIG S8, PDF file, 0.1 MB. TABLE S1, XLSX file, 0.01 MB.

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