Molecular characterization of the Dot/Icm-translocated AnkH and AnkJ eukaryotic-like effectors of Legionella pneumophila

Fabien Habyarimana, Chris T Price, Marina Santic, Souhaila Al-Khodor and Yousef Abu Kwaik

1Department of Microbiology and Immunology, Room MS-410, College of Medicine,
University of Louisville, KY 40292.

Running title: L. pneumophila AnkH and AnkJ effectors

Key Words: Dot/Icm, Legionnaires’, Anaplasma, Coxiella

* For correspondence: Tel (502) 852-4117, Fax (502) 852-7531
e-mail: abukwaik@louisville.edu
ABSTRACT

While most Dot/Icm-translocated effector of Legionella pneumophila are not required for intracellular proliferation, the eukaryotic-like Ankyrin effectors, AnkH and AnkJ are required for intracellular proliferation. In this report, we show that the IcmSW chaperones are essential for translocation of AnkJ but not AnkH. The 10 C-terminal residues and the ANK domains of AnkH and AnkJ are required for translocation. Our data indicate that the two ANK domains of AnkH are critical domains that required for the function of the effector in intracellular replication of L. pneumophila. The ankH and ankJ mutants are severely defective in intrapulmonary proliferation in mice. Expression of AnkH and AnkJ fusions within HEK293 cells show a punctuate distribution in the cytosol but no association with endocytic vesicles, the Golgi apparatus or the ER. Interestingly, the defect in intracellular proliferation of the ankH or ankJ mutants is rescued in HEK293 cells expressing the respective protein. We conclude that AnkH and AnkJ are effectors translocated by the Dot/Icm system by distinct mechanisms and modulate distinct cytosolic processes in the host cell. This is the first demonstration of a trans-complementation of an effector mutant of L. pneumophila through expression of the respective effector in the host cell.
**Introduction**

The gram-negative intracellular bacterial pathogen *L. pneumophila* is found ubiquitously in aquatic environments where it replicates within a wide range of protozoan hosts (2, 18). Once inhaled by humans in aerosolized contaminated water, *L. pneumophila* replicates in human alveolar macrophages and causes Legionnaires’ disease or a less severe flu-like symptoms designated Pontiac fever (19, 29).

*L. pneumophila* is equipped with many sophisticated mechanisms that allow it to survive and replicate within the host cell by creating a specialized ER-like compartment known as the *Legionella*-containing vacuole (LCV) (16, 23, 26, 42, 43). Within the LCV, *L. pneumophila* utilizes its specialized Dot/Icm type IVB secretion machinery to export a cohort of >200 effectors into the host cell cytosol that are essential to modulate various cellular processes such as interception of ER-to-Golgi vesicle traffic, evasion of endocytic traffic, and triggering pro- and anti-apoptotic processes (1, 13, 15, 16, 27, 28, 32, 40, 45). Likewise, *L. pneumophila* is able to translocate into the host cell cytosol specific Dot/Icm substrates before its internalization (12, 35). The IcmS and IcmW chaperones facilitate translocation of few Dot/Icm effectors (7, 10, 12, 36).

*In silico* analyses of 4 *L. pneumophila* genomes (Corby, Paris, Lens and Philadelphia-1) have revealed the presence of many eukaryotic-like genes, which have been suggested to be acquired through horizontal gene transfer (11, 14). Among the genes encoding eukaryotic-like proteins in *L. pneumophila* is a family of at least 11 proteins containing ankyrin eukaryotic-like domains (Ank) (4, 11, 14, 22). The ankyrin domain (ANK) is a 33-amino acid structural motif, and is the most common protein domain in the eukaryotic kingdom, where it functions as a scaffold to mediate protein-protein interactions that play essential roles in various eukaryotic cellular processes, ranging from regulation of transcription, signaling, cytoskeleton, and cell cycle regulation (3, in press, 5, 8, 34). Therefore, it is predicted that the
*L. pneumophila* ankyrin proteins may mimic or interfere with various cellular processes to remodel the host cell into a proliferative niche.

So far, most of Dot/Icm-exported substrates reported have little or no detectable role in intracellular proliferation, suggesting a possible functional redundancy among them (16). Strikingly, of the known Dot/Icm effectors, only loss of SdhA, SidJ or AnkB effectors results in a severe intracellular growth defect. The *sdhA* mutant is defective only in macrophages (31) but the *sidJ* mutant (32) and *ankB* mutant (4) are defective in human macrophages and protozoa. Furthermore, two *L. pneumophila* ankyrin proteins (AnkH and AnkJ) play a significant role in intracellular replication of *L. pneumophila* in human macrophages and in protozoa (22), indicating that they modulate cellular processes that are highly conserved through evolution from protozoa to mammals.

The AnkH and AnkJ proteins that possess 2 and 3 ANK domains, respectively, have been reported to be delivered into host cytosol (13, 37). However, their mechanism of translocation and more importantly the role of the eukaryotic-like ANK domains of AnkH and AnkJ in the intracellular proliferation of *L. pneumophila* and in translocation of AnkH and AnkJ proteins remains unknown.

In this study, we show that three of the *L. pneumophila* ankyrins are delivered into infected cells by a IcmSW-dependent mechanism. The AnkH and AnkJ are essential *in vivo* for in intrapulmonary proliferation in the mouse model. The ANK domains and the last 10 C-terminal residues of AnkH and AnkJ are required for translocation and for intracellular replication. Importantly, expression of AnkH and AnkJ in HEK293 cells rescues the intracellular growth defect of the respective effector mutant, indicating modulation of distinct cytosolic processes by the two effectors to enable intracellular proliferation.
Materials and Methods

Bacterial strains, plasmids, primers and media

The parental *L. pneumophila* serogroup 1 strain AA100/130b (ATCC BAA-74) and its isogenic *dotA, icmS, and icmW* mutant strains have been described previously (46). *Escherichia coli* strain DH5α was used as surrogate to clone the Cya fusion constructs. *L. pneumophila* were grown from frozen stocks on buffered charcoal-yeast extract (BCYE) agar at 37°C or in buffered yeast extract (BYE) broth at 37°C with shaking (17) for 3 days. The plates and broth used for the cultivation of the *L. pneumophila* WT expressing Cya fusion proteins were supplemented with 5 µg/ml of chloramphenicol, whereas the mutants expressing Cya fusion proteins were supplemented with 5 µg/ml of chloramphenicol and 50 µg/ml of kanamycin. The plates used for the cultivation of *E. coli* strains were supplemented with 50 µg/ml of chloramphenicol on Luria-Bertani (LB) agar plates or broth at 37°C with 5% of CO₂ or in LB broth at 37°C with shaking.

DNA manipulations and Cya reporter constructs

Transfections, restriction enzyme digestions, and DNA manipulation were performed as previously described (38). Restriction enzymes and T4 DNA ligase were purchased from Promega (Madison, WI). *L. pneumophila* chromosomal DNA was prepared by using the Puregene DNA isolation kit (Gentra Systems, Minneapolis, MN). Plasmid preparations were performed with the Bio-Rad Quantum miniprep kit. Electroporations were performed with a Bio-Rad Gene Pulser, as recommended by the manufacturer’s specifications. Purification of DNA fragments from agarose gels for subcloning was carried out with a QIAquick gel purification kit (Qiagen, Valencia, CA). Primers (Table 2) used to amplify the coding sequences of *L. pneumophila ank* genes utilized to engineer Cya-Ank or GFP-Ank
constructs by PCR were from Integrated DNA Technologies, Inc. (Coralville, IA). The coding sequences of *L. pneumophila* ank genes were fused to the C-terminal of adenylate cyclase (Cya) of *B. pertussis* using plasmid *pcya-ralF*, which is a derivative of pMMB207M45NT (35). All of the *L. pneumophila* ank gene PCR products were digested with BamHI and PstI. The pCya-RalF plasmid was digested with BamHI and PstI to release the ralF gene. Digested products were ligated using T4 DNA ligase. Resulting Cya-Ank reporter constructs are displayed in Table 2. For in-frame deletion of the ANK domains and C-terminal of AnkH or AnkJ we used *L. pneumophila* ank genes cloned into the plasmid vector pBC-SK+ as previously described (22). To generate domain mutant alleles of *ankH* and *ankJ*, an inverse PCR strategy was employed using pBCSK+ harboring the *ankH* and *ankJ* genes as a template. Briefly, phosphorylated primers (Table 1) were designed to hybridize adjacent to DNA encoding the ANK domains or C-termini and then the entire plasmid lacking the domain of interest was PCR-amplified using Phusion DNA polymerase (Finnzymes). The resulting PCR product was then treated with DpnI restriction to remove residual template DNA from the reaction and then allowed to religate using T4 DNA ligase. The ligation products were transformed into *E. coli* DH5α. Domain deletions of *ankH* and *ankJ* were verified by DNA sequencing to ensure the integrity of the *ankH* and *ankJ* reading frames. Recombinant plasmids were then electroporated into the *L. pneumophila*. The various pBCSK+ vectors harboring the mutant *ankH* and *ankJ* alleles were used as templates to generate Cya fusions. Primers listed in Table 2 were used in PCR to amplify the mutant *ankH* and *ankJ* alleles and the resulting PCR products were cloned into pCR2.1 via topoisomerization as described by the manufacturer (Invitrogen Corp, Carlsbad, CA). The mutant *ankH* and *ankJ* alleles were then subcloned into the BamHI-PstI sites of *pcya-ralF* (Nagai et al., 2005), resulting in replacement of the ralF gene with *ankH* and *ankJ* alleles in frame with cya. Recombinant plasmids were electroporated into the *L. pneumophila*. 
To generate the mammalian fusion constructs, ORF of *L. pneumophila* ankH and ankJ were cloned into the mammalian expression vectors pAcGFP (Sigma) or p3XFlag-CMV (Sigma) using primers displayed in table 3 to generate GFP or 3XFlag-AnkH or AnkJ fusion constructs.

**Translocation assay**

For adenylate cyclase (Cya) activity assays, differentiated U937 cells monolayers grown in 24-well plates were infected with various strains of *L. pneumophila* at MOI of 50 for 1 h at 37°C. For cytochalasin D treatment assay, U937 cells were treated with 1 µg/ml of cytochalasin D for 30 minutes. Cells were then infected for 30 minutes, washed 3 times extensively with 1X PBS to remove extracellular bacteria and subsequently lysed in 200 µl of 0.25% dodecyltrimethylammonium bromide in assay buffer. Cell lysates were processed for the detection of intracellular cAMP using Amersham cAMP Enzyme Immunoassay kit (GE Healthcare, Piscataway, NJ), as recommended by the manufacturer.

**Macrophage culture and *L. pneumophila* intracellular replication analysis**

Isolation and preparation of the human monocyte-derived macrophages (hMDMs) was carried out as we previously described (41). The hMDMs and U937 cells were maintained in RPMI-1640 tissue culture medium (Gibco BRL) supplemented with 10% heat-inactivated fetal bovine serum (FBS) (Gibco BRL). The cells were cultured under a humidified atmosphere containing 5% CO₂ and 95% air at 37°C. For intracellular proliferation studies, infections of macrophages were performed as we described previously (22). Briefly, cells were infected with various strains of at a multiplicity of infection (MOI) of 10 for 1 h followed by treatment with 50µg/ml gentamicin for 1 h to kill extracellular bacteria, and this was considered the zero time point. The intracellular proliferation was assessed by plating and
enumerating the colony-forming units (CFU) at 24h and 48h post-infection. Alternatively, the intracellular replication was examined after 10h post-infection by single cell analysis using laser scanning microscopy as previously described (22).

**Infection of A/J mice with *L. pneumophila***

Female pathogen-free, 6- to 8-week-old A/J mice were used for infection by intratracheal inoculation as described previously (6, 20). The *L. pneumophila* strain AA100, ankH and ankJ mutants were grown on BCYE agar plates for 72 h. Three mice per strain were used for intrapulmonary proliferation assay and 3 mice per dose were used for survival assay. The mice were inoculated intratracheally with 50 µl containing the bacterial dose of $10^6$ for intrapulmonary proliferation assay and $10^7$, $10^8$, $8 \times 10^8$ or $10^9$ for survival assay as described previously (6, 20). At 2 h, 24 h, and 48 h, 72h and 7 days post-inoculation, mice were humanely euthanized, the lungs were removed, and the bacteria were cultured on BCYE agar for 72 h as described previously (6, 20). For the survival assay, the mice were observed for 3 days post-inoculation.

**Expression of AnkH and AnkJ in mammalian cells**

The human renal epithelial cell line HEK293 cells or HEK293T cells were grown on circular glass cover slips (Fisher) pretreated with 0.1mg/ml of poly-D-lysine in 24-well culture plates at concentration of $5 \times 10^4$ cells/ml in DMEM containing 10% FBS overnight. The sub-confluent culture of HEK293T and HEK293 cells were transiently transfected using calcium phosphate method with the mammalian expression vectors pAcGFP (Sigma) or BAP3XFlag (Sigma) and constructs of AnkH or AnkJ fusion GFP or 3XFlag for 18 h. In this study, HEK293T cells were transiently transfected with GFP or 3XFlag constructs to examine their sub-cellular distribution. Since HEK293T are not suitable for
generating stable transfected cells, therefore we used HEK293 cells. Transiently transfected HEK293 cells were used to generate stably transfected HEK293 cells. Briefly, 1/10 diluted transiently transfected HEK293 cells were grown in presence of 1.4mg/ml of geneticin (G418) (Sigma) for 15 days and every other 3 days the media was replaced by new one containing 1.4mg/ml of geneticin. After 15 days, clones were picked and screened by confocal laser scanning microscopy and western blot analysis for expression of the AnkH or AnkJ fusion protein. To study colocalization of AnkH and AnkJ with late endosomal, nuclear, lysosomal, cis and trans Golgi apparatus and ER markers, we labeled transfected cells with the primary antibodies anti-Lamp2, anti-Cathepsin D, anti-GM130 (cis), anti-Golgi 58k (trans) or anti-KDEL. The anti-LAMP-2 (H4B4) monoclonal antibody (developed by J. T.August and J. E. K.Hildreth) was obtained from the Developmental Studies Hybridoma Bank (University of Iowa, IA). To label the lysosomes, transfected cells were incubated with mouse monoclonal anti-cathepsin D antibody (BD Transduction, Franklin Lakes, NJ). Mouse anti-KDEL monoclonal antibody was purchased from StressGen Biotechnologies (Ann Arbor, Michigan). Rabbit anti-GM130 was obtained from Santa Cruz Biotechnology, Inc. and monoclonal anti-Golgi 58k was purchased from Sigma. The polyclonal anti-COX (mitochondria marker) was obtained from Cell Signaling, monoclonal antibodies anti-actin and anti-tubulin were obtained from Sigma. Primary antibodies were detected by Alexa Fluor 555-conjugated donkey anti-mouse or anti-rabbit IgG (Molecular Probes). The cells were examined using an Olympus Fluoview1000 laser scanning confocal microscopy as described previously (41). On average, 8–15 0.2 mm serial Z sections of each image were captured and stored for further analyses, using Adobe Design Premium CS3.

Western blot assay
To detect the Cya-Ank hybrid proteins in *L. pneumophila* WT, *dotA*, *icmS* or *icmW* strains harboring empty vector or Cya-Ank constructs, bacteria were grown for 3 days on BCYE in presence of appropriate antibiotics prior to infection of U937 cells. Total of $5 \times 10^8$ bacteria were isolated by centrifugation and lysed with B-per bacterial protein extraction reagent (Thermo Scientific, Waltham, MA) followed by boiling in SDS-PAGE sample buffer. Proteins were transferred onto nitrocellulose membrane and the Cya-Ank hybrid proteins were detected by using monoclonal anti-M45 antibody at 1:50 dilution (4). After incubation with a secondary antibody conjugated to horseradish peroxidase, signals were detected by the supersignal west femto maximum sensitivity substrate (Thermo Scientific, Waltham, MA).

**Statistical analysis.**

All experiments were performed in triplicate at least three times, and the data shown are representative of one experiment. To analyze for statistical significant differences between different sets of data, a two-tailed Student $t$ test was used, and the $P$ value was obtained.
Results

Potential translocation of *L. pneumophila* ankyrins upon bacterial attachment to the macrophage

Potential role of the IcmSW chaperon in translocation of the Ank proteins was evaluated using adenylate cyclase assays. These data showed that the *L. pneumophila* AnkD, AnkG and AnkJ are translocated by the Dot/Icm machinery in IcmSW-dependent manner while the translocation of AnkH, AnkI, AnkK and AnkN is independent of the IcmSW complex (data not shown). Western blots confirmed equivalent expression of all the proteins in the *icmS* and *IcmW* mutants (data not shown).

Some Dot/Icm effectors, such as LepA and LepB are translocated into the host cell upon contact of *L. pneumophila* to the host cell prior to bacterial internalization (12, 35). We have recently shown that AnkB is exported into host cell by extracellular bacteria (39, in presse) Therefore, we examined whether the other Ank proteins are translocated into macrophages prior to bacterial internalization. Prior to infection, U937 cells were treated for 30 minutes with 1µg/ml of Cytochalasin D to prevent phagocytosis. The cells were infected for 30 minutes with the WT strain or the dotA mutant harboring Cya-Ank fusion constructs or the empty vector. In addition, we infected cells with the WT strain or the dotA mutant expressing Cya-RalF or Cya-AnkB fusion protein as positive controls. As expected, the WT strain translocated significantly the RalF into untreated cells compared to the empty vector (student t-test, p <0.0001) (Fig. 1). The results showed that the AnkB controls was efficiently translocated by attached extracellular bacteria as indicated by comparable level of cAMP in Cytochalasin D-treated and untreated cell (student t-test, p >0.5) (Fig. 1). In contrast, the levels of cAMP were significantly different between Cytochalasin D-treated and untreated cells infected with WT strain harboring any of the tested Cya-Ank reporters (student t-test, p <0.0001) (Fig. 1).

The C-terminus of *L. pneumophila* AnkH and AnkJ are essential for translocation
In addition to the existence of a C-terminus secretion motifs, Nagai et al. have shown that some but not all Dot/Icm substrates harbor conserved hydrophobic residues in the C-terminus (30, 33, 35). Since AnkH and AnkJ bear hydrophobic residues at the positions -4 or -5 in the C-terminus (Fig. S1), we tested whether the C-terminal region of AnkH and AnkJ proteins were required for their translocation. In-frame deletion of the last 10 residues of AnkH and AnkJ were constructed and fused to Cya (Fig. 2A and Fig. 3A). U937 cells were infected with the WT strain or the dotA mutant expressing the truncated AnkH and AnkJ reporter fusions. Our results showed a significant low levels of cAMP in cells infected with the WT strain expressing C-terminal deletion of AnkH and AnkJ with > 10-fold less compared to cells infected with the WT strain expressing full length AnkH and AnkJ (student t-test, \( p < 0.0001 \)) (Fig. 2C and Fig. 3C). Thus, the C-terminus of AnkH and AnkJ is required for translocation into the host cell.

The ANK domains of \textit{L. pneumophila} are essential for translocation of AnkH and AnkJ

To test whether the ANK domains of AnkH and AnkJ proteins could be required for translocation of the two proteins into macrophages, single, double or triple in-frame deletions of the ANK domains of AnkH and AnkJ were fused to Cya (Fig. 2A and Fig. 3A). Immunoblot analysis revealed equivalent expression of the truncated proteins (Fig. 2B and Fig. 3B). The U937 cells were infected for 1h with the WT strain or the dotA isogenic mutant expressing each of the truncated AnkH or AnkJ fusion proteins and intracellular cAMP level was determined in cell lysates. The level of cAMP in cells infected with \textit{L. pneumophila} harboring the ANK domains deletion was significantly lower with >10-fold for truncated AnkH and >7-fold less for truncated AnkJ compared to the full length fusion or the positive control (student t-test, \( p < 0.0001 \)) (Fig. 2C and 3C). Taken together, these data indicate that the ANK domains of AnkH and AnkJ are essential for their translocation into the host cell. This is the
first example of the role of the eukaryotic-like domains in translocation of Dot/Icm effectors into the host cell. It is also possible that the reduced translocation may be due to a mild reduction in the protein level of the variant proteins.

Role of the ANK domains of AnkH and AnkJ in intracellular growth of *L. pneumophila*

We have previously shown that AnkH and AnkJ are required for intracellular replication of *L. pneumophila* (22). The aforementioned role of ANK domains of AnkH and AnkJ in translocation of AnkH and AnkJ proteins into host cytosol prompted us to examine their role in intracellular replication. Therefore, we performed single, double or triple in-frame deletion of the ANK domains of AnkH and AnkJ. The *ankH* and *ankJ* mutants were trans-complemented with the full length genes or the corresponding engineered ANK domain in-frame deletion constructs. The hMDMs were infected with *L. pneumophila* WT strain, *dotA, ankH* or *ankJ* mutant and *ankH* and *ankJ* mutants trans-complemented with the various constructs. At 24 h post-infection, there was an increase in the number of bacteria in all strains ranging from 10-fold to 65-fold with an increase in number of bacteria for the WT strain and the *ankH* mutant harboring all different constructs compared to *dotA* and the *ankH* mutant (student t-test, \( p < 0.001 \)) (Fig. 4). However, at 48 h post-infection there was a significant reduction in the cfus of the *ankH* mutant trans-complemented with AnkH\( \Delta A1 \), AnkH\( \Delta A2 \) or AnkH\( \Delta A1 \Delta A2 \) compared to mutant complemented with the full length gene with > 100-fold increase (student t-test, \( p < 0.0001 \)). As expected, the *dotA* and the *ankH* mutants did not grow in hMDMs (Fig. 4). Because of inconsistent results in multiple experiments, data from the *ankJ* mutant trans-complemented with engineered ANK deletion constructs are not shown. Our results demonstrate that the two ANK domains of AnkH play a vital role in intracellular proliferation of *L. pneumophila*. It is also possible that the reduced intracellular growth may be due to a reduced translocation of the truncated proteins.
The AnkH and AnkJ effectors are essential for intracellular proliferation of *L. pneumophila* within human monocyte-derived macrophages (hMDMs) and protozoa (22), but whether the two effectors are required for the infection *in vivo* in animal models is not known. Moreover, the role of Dot/Icm-translocated effectors in the pulmonary infection in animal models has never been examined for any Dot/Icm effector. To determine whether the mutation in *ankH* or *ankJ* caused a decrease in mortality in the A/J mouse model, we infected mice intratracheally (6, 9, 20) with doses of $10^7$-$10^9$ cfus. By the first day post-infection with a high dose of $10^9$ cfus there was a mortality rate of 80% in mice infected by the WT strain compared to the mortality rate of 20% in mice infected by *ankB*, *ankH* or *ankJ* mutant with the similar dose (Fig. 5A) (student *t*-test, *p* <0.001). The AnkB attenuated mutant was used as a negative control (39, in press). These data show that the two Ank effectors contribute the lethality of Legionnaires’ disease in the mice model of the disease, consistent with their role in intracellular proliferation within cultured macrophages.

To investigate whether AnkH and AnkJ are required for intrapulmonary proliferation of *L. pneumophila*, we infected A/J mice with $10^6$ of the *L. pneumophila* WT strain, the *ankH*, or the *ankJ* mutant. Multiplication of *L. pneumophila* in lungs of infected mice was assessed by CFU enumeration after 24h, 48h, 72h and 7 days post-infection. At 48h and 72h post-infection, the CFUs of *L. pneumophila* recovered from mice infected with the *ankH* or the *ankJ* mutant were significantly lower than the WT strain (student *t*-test, *p* <0.001) with at least a 1000-fold and 100-fold less bacteria, respectively, were recovered from the lungs for both mutants (Fig. 5B). The *ankB* attenuated mutant (39, in press), which was used as a control, was severely defective in intra-pulmonary replication. These data
show that the \( \text{ankH} \) and \( \text{ankJ} \) mutants are defective in intrapulmonary proliferation and mice mortality, consistent with their \textit{in vitro} intracellular growth defect in cultured macrophages (22). Taken together, our data show that the two Ank effectors are the first effectors of \textit{L. pneumophila} shown to be required for intrapulmonary proliferation and lethality of \textit{L. pneumophila} in the mice model of Legionnaires’ disease.

**Expression and trafficking of AnkH and AnkJ in mammalian cells**

Ectopic expression of bacterial proteins in eukaryotic cells has been an important strategy to study localization of bacterial effector proteins translocated into host cell and may provide key insights into the function of the effectors (13, 30, 37). Therefore, GFP-tagged or 3XFlag-tagged AnkH and AnkJ were constructed and transiently or stably expressed in HEK293T or HEK293 cells to study their subcellular distribution and trafficking in mammalian cells. Constructs of GFP or bacterial alkaline phosphatase (BAP) fusion was used as a negative control. Transient transfection of HEK293T cells with plasmids harboring GFP-tagged or 3xFlag-tagged \( \text{ankH} \) and \( \text{ankJ} \) using calcium phosphate yielded 70%-75% transfection efficiency. Transient or stable expression of AnkH and AnkJ were not toxic to the HEK293 cells (data not shown). Transient or stable expression of GFP-tagged or 3XFlag-tagged AnkH and AnkJ were distributed in the cytoplasm with punctuate-like distribution but neither of the two effectors were detected in the nucleus (Fig. 6), suggesting that AnkH and AnkJ were associated with host cell vesicles. Nevertheless, these punctuate-like structures did not colocalize with \textit{L. pneumophila}.

We utilized laser scanning confocal microscopy to determine trafficking and potential co-localization of the two effectors with endosomal, lysosomal, Golgi, ER, microfilament, microtubules, mitochondria, and nuclear compartments using specific markers Lamp2, cathepsin D, GM130 or P58-k, KDEL, actin, tubulin, mitochondrial protein, and nuclear dye (DAPI), respectively. The data showed
that there were no significant differences in association of the above markers with GFP-AnkH or GFP-AnkJ fusion proteins compared to GFP negative control (13-20%) (student t-test, \( p > 0.5 \)), indicating that these proteins are not associated with endosomal, lysosomal, ER, or Golgi vesicles (Fig. 7A and B). We conclude that despite the punctuate distribution of the two Ank effectors in mammalian cells, they do not co-localize with any sub-cellular compartment.

*L. pneumophila* ankH and ankJ mutants are rescued in HEK 293 cells expressing AnkH and AnkJ-GFP fusion proteins

Since *L. pneumophila* ankH and ankJ mutants exhibited intracellular growth defect and the AnkH and AnkJ are translocated into host cells by the Dot/Icm system, we examined whether stable HEK293 cells expressing *L. pneumophila* AnkH or AnkJ fusion proteins could rescue the respective mutant for the defect in intracellular proliferation. Therefore, HEK293 cells with stable expression of GFP, GFP-AnkH or GFP-AnkJ were infected with *L. pneumophila* WT, dotA, ankH or ankJ mutant. After 10h post-infection, infected cells were labeled with anti-*L. pneumophila* antibody to evaluate the intracellular replication by single cell analysis using confocal microscopy. The data showed that by 10h post-infection, ~70% of the WT strain-infected cells harbored more than 7 bacteria/cell in HEK293 cells expressing GFP, GFP-AnkH or GFP-AnkJ. In contrast, the dotA mutant did not replicate with 1-3 bacteria/infected cell. Similar to the WT-infected cells expressing GFP-AnkH or GFP-AnkJ, ~60% of the ankH mutant-infected cells and ankJ mutant-infected cells harbored more than 7 bacteria (Fig. 8A and B). In contrast, the ankH and ankJ mutants did not replicate in HEK293 cells expressing GFP alone with >60% of the ankH and ankJ mutant-infected cells harbored ≤3 bacteria/cell (Fig. 8A and B). The ankH mutant was not rescued in cells expressing ankJ, or vice versa (data not shown). These data show that the ankH and ankJ mutants are rescued in mammalian cells expressing *L. pneumophila* AnkH and
AnkJ, respectively. Collectively, our data indicate that AnkH and AnkJ modulate distinct processes in the host cell cytosol, consistent with their distinct structure and mode of export. This is the first example of a trans-complementation of a Dot/Icm effector mutant of \textit{L. pneumophila} through expression of the effector in the host cell.
Discussion

The ANK domains are the most abundant domain in the eukaryotic kingdom, where they function as scaffold to mediate protein-protein interactions required for various eukaryotic cellular processes, ranging from regulation of transcription, signaling, cytoskeleton, and cell cycle regulation (5, 8, 34). Recently, genomic analyses have shown that *L. pneumophila* genome encodes a large family of eukaryotic-like ankyrin proteins (4, 11, 14, 22). It is thought that these proteins have been acquired by *L. pneumophila* through horizontal gene transfer through co-evolution with its natural protozoan hosts to perhaps mimic or interfere with host cell processes to establish a replicative niche within host cells. In addition to AnkB, the AnkH and AnkJ proteins play a significant role in intracellular replication of *L. pneumophila* in human macrophages and protozoa (22), indicating that these Dot/Icm-translocated effectors modulate cellular processes that are highly conserved through evolution from protozoa to mammals. Importantly, our data show that the AnkH and AnkJ Dot/Icm-translocated effectors are essential for intrapulmonary proliferation *in vivo* in A/J mice. These data are consistent with the role of the two effectors in intracellular growth of *L. pneumophila* in macrophages (22). To our knowledge, this is the first demonstration for an essential role of Dot/Icm-translocated effectors in intrapulmonary proliferation in animal models.

In several other pathogens, such as *Agrobacterium*, *Bordetella*, *Helicobacter*, *Anaplasma*, *Coxiella* and *Brucella*, the TFSS is essential for delivery of host cell-modulating effectors. In agreement with two different studies (13, 37), our data show that 7 *L. pneumophila* Ank proteins are delivered into the host cytosol. In contrast to de Filipe et al. study (13), using a different strategy our data show that AnkG/LegA7 is part of the cohort of Dot/Icm-translocated effectors. Consistent with the *L. pneumophila* Ank proteins being translocated by the Dot/Icm secretion system, there is a recent report of 13 Dot/Icm-translocated ankyrin proteins of *Coxiella burnetii* when expressed in *L. pneumophila* as a surrogate host.
Moreover, *Anaplasma phagocytophilum* encodes an ankyrin protein (AnkA), which is translocated by the TFSS into the host cell cytosol and nucleus (21, 24, 25). Given the rising number of translocated effector proteins by *L. pneumophila* into the host cell and its large spectrum of environmental hosts, it is possible that *L. pneumophila* selectively deploy a specific set of effectors that best promote its survival and proliferation within a specific host cell in the environment or in humans during infection.

The components of the Dot/Icm TFSS engage some of its effector proteins through a recognition of a translocation signal predicted to be at the C-terminus of Dot/Icm substrates (10, 35). Our data show that deletion of the last 10 residues of AnkH and AnkJ abrogates their translocation, indicating that their translocation signal is located at C-terminus (4, 12, 35).

The deletion of the ANK domains in AnkH and AnkJ reduced substantially their translocation, suggesting that the ANK domains actively participate in their translocation. Two explanations appear to validate the involvement of the ANK domains in AnkH and AnkJ proteins delivery into host cytosol. First, the ANK domains may participate in folding or unfolding of the AnkH or AnkJ proteins for suitable presentation of their C-terminal translocation signal to the Dot/Icm components. Second, the ANK domains may be involved in interaction between AnkH or AnkJ proteins and the Dot/Icm components in order to be properly delivered into host cells, where they interfere with host cell processes to facilitate bacterial proliferation. However, it is possible that the ANK deletion proteins are translocated into host cell and the enzymatic activity of adenylate cyaclase (Cya) is inhibited due to misfolding of the truncated protein, thereby Cya become inaccessible to the activating host calmodulin.

In addition, possible misfolding of the in-frame deletion of the Ank protein may have rendered it unrecognizable to be delivered by the Dot/Icm system.
Interestingly, expression of AnkH and AnkJ in mammalian cells show a punctate distribution throughout the cytosol, but our data indicate no association of AnkH or AnkJ with endosomal, lysosomal, ER, mitochondria and Golgi vesicles, or actin and tubulin. These data are consistent with our previous findings that the phagosomes harboring the \textit{L. pneumophila ankH} and \textit{ankJ} mutants are trafficked in similar manner as the ones harboring the WT strain (22), suggesting that these proteins are not involved in trafficking of LCV or recruitment of the ER to LCV. It is unlikely that the punctuate distribution of proteins in mammalian cells is due to protein aggregation, since the ectopically expressed protein is functional in trans-rescue of the mutants for the defect in intracellular proliferation. Sub-cellular localization as well as the host cell targets of the Ank effectors still to be identified.

The complementation of the \textit{ankH} mutant by in frame-deletions of the ANK domains of AnkH does not restore its intracellular growth defect (22). However, deletion of the ANK domains of AnkH abrogates its translocation, indicating that the role of AnkH in intracellular replication requires its ANK domains that are also indispensable for its translocation. Whether translocation of the in-frame deletion of the Ank protein would render it functional in the host cell cytosol is not known.

Remarkably, our data indicate that \textit{ankH} and \textit{ankJ} mutants exhibit a severe intrapulmonary replication defect resulting in less mortality compared to the WT strain. This is consistent with our previous \textit{ex vivo} results in human macrophages and alveolar epithelial cells (22) and corroborates with high rate of survival of animal infected by the \textit{ankH} or \textit{ankJ} mutants. This is the first demonstration for the role of Dot/Icm effectors in the development of Legionnaires’ disease in animal models.

Interestingly, when HEK293 cells expressing AnkH and AnkJ are infected with the \textit{ankH} and \textit{ankJ} mutants, the intracellular growth defect of the respective mutant is rescued. These data completely support the findings that the AnkH and AnkJ are translocated into the host cell cytosol to modulate
distinct cytosolic processes needed to sustain the intracellular proliferation of the ankH and ankJ mutants. This is the first finding that ectopic expression of a Dot/Icm effector in mammalian cells can rescue the growth of an effector mutant.

In summary, our data show that 3 L. pneumophila Ank proteins are delivered into the host cells in an IcmSW complex-dependent manner, and none of the 7 translocated Ank proteins tested are delivered into the host cell by attached extracellular bacteria. Furthermore, our data indicate that the ANK domains and the C-terminus of the AnkH and AnkJ are indispensable for translocation into the host cell, which is essential for intracellular proliferation of L. pneumophila. Ectopic expression in mammalian cells and co-localization studies show that AnkH and AnkJ proteins are distributed in punctuate-like structures throughout the cytosol and are not associated with nuclear, endosomal, lysosomal, Golgi or ER compartments. Our data show that the L. pneumophila ankH and ankJ mutants are rescued for their intracellular growth defect in HEK293 cells expressing the respective effector.
Table 1. List of ANK deletion constructs and primers used to generate ANK deletions in *ankH* and *ankJ*.

<table>
<thead>
<tr>
<th>Constructs</th>
<th>Primer Sequences (5'-3')</th>
</tr>
</thead>
</table>
| ankH∆A1 | F: CCAGACGTACAGGACGC  
R: TTCATCGATATCATCCAAG |
| ankH∆A2 | F: TACACTCGTAATGGGTTTG  
R: GACGTCTGGGTTGTATAT |
| ankH∆1391-1401 | F: TTAATTAGGATTAATCCCAACTCATCCAGAATT  
R: TAAAGGTAAGAAATAATT |
| ankJ∆A1 | F: TGATGCTATTTTCATTTC  
R: TACGCCCCCCCCACTTATG |
| ankJ∆A2 | F: GGGGGGGGCGTACTGATT  
R: CCTATCAACTCATCAAAAG |
| ankJ∆A3 | F: ATAGTTGAGAGAGATAC  
R: CTGTTGATAGACCAATAAT |
| ankJ∆797-807 | F: TTATCTTCTCAAAACGACTCTCTGGAAC  
R: GGAGAAATACCTCCTTCAAGAA |

All primers are 5’-phosphorylated
Table 2. List of Cya-Ank reporter constructs and primers used to generate ank fusions.

<table>
<thead>
<tr>
<th>Constructs</th>
<th>Primer Sequences (5’-3’)</th>
<th>Restriction sites</th>
</tr>
</thead>
<tbody>
<tr>
<td>pCya-ankB</td>
<td>F: GGATCCTTATGAAAAAGAATTTTTTTCTG</td>
<td>BamHI PstI</td>
</tr>
<tr>
<td></td>
<td>R: CTGCAGTTAACAACACGACCTTGT</td>
<td></td>
</tr>
<tr>
<td>pCya-ankC</td>
<td>F: CCCGGATCTTATGGATTGTTAGTGAATGGAATG</td>
<td>BamHI PstI</td>
</tr>
<tr>
<td></td>
<td>R: CCCCTGCAGTTATTTTTAGGCAAACCTCTG</td>
<td></td>
</tr>
<tr>
<td>pCya-ankD</td>
<td>F: CCCGGATCTTTATGGACTCTCCGCTGCC</td>
<td>BamHI PstI</td>
</tr>
<tr>
<td></td>
<td>R: CCCCTGCAGTTAGCCTGAGGATTTCTTTTA</td>
<td></td>
</tr>
<tr>
<td>pCya-ankG</td>
<td>F: CCCGGATCTTTCTGAAATTTATAAGGATGAC</td>
<td>BamHI PstI</td>
</tr>
<tr>
<td></td>
<td>R: CCCCTGCAGTTATTTCAAACCAAAACGAG</td>
<td></td>
</tr>
<tr>
<td>pCya-ankH</td>
<td>F: CCCGGATCTTTATGAGTATTGCAAAC</td>
<td>BamHI PstI</td>
</tr>
<tr>
<td></td>
<td>R: CCCCTGCAGTTATAGGCTGCAAACAGGA</td>
<td></td>
</tr>
<tr>
<td>pCya-ankI</td>
<td>F: CCCGGATCTTTATGATTATTTTATATGATT</td>
<td>BamHI PstI</td>
</tr>
<tr>
<td></td>
<td>R: CCCCTGCAGTTAAATAGGCTGCAAACAGGA</td>
<td></td>
</tr>
<tr>
<td>pCya-ankJ</td>
<td>F: CCCGGATCTTTATGATTATTTTATATGATT</td>
<td>BamHI PstI</td>
</tr>
<tr>
<td></td>
<td>R: CCCCTGCAGTTAAATAGGCTGCAAACAGGA</td>
<td></td>
</tr>
<tr>
<td>pCya-ankK</td>
<td>F: CCCGGATCTTTATGATTATTTTATATGATT</td>
<td>BamHI PstI</td>
</tr>
<tr>
<td></td>
<td>R: CCCCTGCAGTTAAATAGGCTGCAAACAGGA</td>
<td></td>
</tr>
<tr>
<td>pCya-ankN</td>
<td>F: CCCGGATCCTTATGCTTATGCG</td>
<td>BamHI PstI</td>
</tr>
<tr>
<td></td>
<td>R: CCCCTGCAGTTATTTTTTTATAGGCTGCAAACAGGA</td>
<td></td>
</tr>
<tr>
<td>pCya-ankQ</td>
<td>F: CCCGGATCCTTATGCTTATGCG</td>
<td>BamHI PstI</td>
</tr>
<tr>
<td></td>
<td>R: CCCCTGCAGTTATTTTTTTATAGGCTGCAAACAGGA</td>
<td></td>
</tr>
</tbody>
</table>

...
Table 3. List of GFP/3xFlag-Ank fusion constructs and primers used to generate *ank* fusions.

<table>
<thead>
<tr>
<th>Constructs</th>
<th>Primer Sequences (5’-3’)</th>
<th>Restriction sites</th>
</tr>
</thead>
</table>
| p3xflag-ankH    | F: AGATCTGATGATTTGCAAACGATA  
R: GGAATCTTTATAGGCCTGCGCAACAGGATT                                                   | BglII BamH       |
| p3xflag-ankJ    | F: AAGCTTGTGATTAAAAATGGGTAGA  
R: AGATCTTTAAGTGGCGTTTTTAGGGGTATC                                                    | HindIII BglII    |
| pAcGFP-ankH     | F: AGATCTGATGATTTGCAAACGATA  
R: TCTAGATTATAGGCCTGCGCAACAGGATT                                                     | BglII XbaI       |
| pAcGFP-ankJ     | F: AGATCTGATGATTTGCAAACGATA  
R: TCTAGATTAAAGTGGCGTTTTTAGGGGTATC                                                   | BglII XbaI       |
References


Figure Legends

Fig. 1 Potential translocation of *L. pneumophila* Ankyrin proteins by attached extracellular bacteria. Untreated or cytochalasin D-treated U937 cells were infected with the wild-type strain of *L. pneumophila* expressing the indicated Cya hybrid proteins. After 30 minutes of infection, cultured cells were lysed and cAMP was quantified by ELISA and the amount of cAMP is indicated as fmol/well. The experiment was performed three times. The data points are the average of cAMP concentration for one representative experiment performed in triplicate. Error bars represent standard deviations of triplicate samples.

Fig. 2. The ANK repeats and the C-terminus of *L. pneumophila* AnkH are required for delivery into host cell

(A) Organization of the AnkH and different ANK domain or C-terminus deletions. Each protein size and truncation are displayed.

(B) Immunoblots of whole-cell bacterial extracts expressing indicated Cya hybrid proteins from wild-type (WT), probed with monoclonal antibody specific to the M45 epitope and reprobed with anti-CAT as loading control. The numbers represent the different truncated Cya-AnkH hybrid proteins: Cya-AnkH (1), Cya-AnkHΔA1 (2), Cya-AnkHΔA2 (3), Cya-AnkHΔA1ΔA2 (4), Cya-AnkHΔ457-467 (5).

(C) U937 cells were infected with the wild-type strain or the *dotA* mutant of *L. pneumophila* expressing the indicated Cya hybrid proteins. After 1 h of infection, cultured cells were lysed and cAMP was quantified by ELISA and the amount of cAMP is indicated as fmol/well. The experiment was performed three times and the data are the average of cAMP concentration for one representative experiment performed in triplicate. Error bars represent standard deviations of triplicate samples.
Fig. 3. The ANK repeats and the C-terminus of *L. pneumophila* AnkJ are required for delivery into host cell

(A) Organization of the AnkJ and different ANK repeats or C-terminus in-frame deletions. Each protein size and different truncation are displayed.

(B) Immunoblots of whole-cell bacterial extracts expressing the indicated Cya hybrid were probed with monoclonal antibody specific to the M45 epitope and reprobed with anti-CAT as a loading control. The numbers represent the different Cya-AnkJ truncated hybrid proteins: Cya-AnkJ (1), Cya-AnkJΔA1 (2), Cya-AnkJΔA2 (3), Cya-AnkJΔA3 (4), Cya-AnkJΔA1ΔA2 (5), Cya-AnkJΔA1ΔA3 (6), Cya-AnkJΔA2 ΔA3 (7), Cya-AnkJΔA1ΔA2 ΔA3 (8), Cya-AnkJΔ259-269 (9).

(C) U937 cells were infected with the wild-type strain or the dotA mutant of *L. pneumophila* expressing the indicated Cya hybrid proteins. After 1 h of infection, cultured cells were lysed and cAMP was quantified by ELISA and the amount of cAMP is indicated as fmol/well. The experiment was performed three times and the data are the average of cAMP concentration for one representative experiment performed in triplicate. Error bars represent standard deviations of triplicate samples.

Fig. 4. The ANK repeats of AnkH are indispensable for intracellular growth of *L. pneumophila*

Monolayers of hMDMs were infected with *L. pneumophila* WT strain, dotA or ankH mutants complemented with full length AnkH or constructs with in-frame deletion of the ANK domains. The infection was carried out in triplicate with an MOI of 10 for 1h followed by 1h gentamicin treatment to kill extracellular bacteria. At 24h and 48h, *L. pneumophila*-infected cells were lysed and plated onto BCYE plates for CFU enumeration. The WT strain was used as a positive control and dotA mutant strain as a negative control. The results are represented as fold increase (T/T0). The experiment was performed...
three times. The data points are the average of one representative experiment performed in triplicate. Error bars represent standard deviations of triplicate samples.

Fig. 5. The *L. pneumophila* ankH and ankJ mutants are defective in the A/J mice model

(A.) Groups of 30 mice were infected with $10^7$, $10^8$, $8 \times 10^8$ or $10^9$ cfus of *L. pneumophila* WT strain, ankH or ankJ mutant compared to the ankB mutant. After 1, 2, 3, 4 and 5 days post-infection mortality was determined. After 3 days there was no mortality.

(B.) A/J mice were infected with $10^6$ of *L. pneumophila* WT strain, ankH or ankJ mutants compared to the ankB mutant. After 1, 2, 3 and 7 days post-infection, 3 mice were sacrificed and lungs were collected for CFU enumeration.

Fig. 6. Subcellular localization of AnkH and AnkJ

HEK293T cells were transiently transfected with the empty vector pAcGFP, the pAcGFP-AnkH or pAcGFP-AnkJ fusion constructs or with pBAP-3XFlag, and pAnkH-3XFlag or pAnkJ-3XFlag fusion constructs for 18 h. After transfection, localization of AnkH and AnkJ fusion proteins was examined by confocal laser scanning microscopy. Similar results were obtained in stable transfections (data not shown).

Fig. 7. Colocalization of AnkH and AnkJ with endosomal, lysosomal, Golgi and ER compartments

Stable transfected HEK293 cells expressing GFP, the GFP-AnkH or GFP-AnkJ fusion proteins were fixed and stained with anti-Lamp2, anti-cathepsinD, anti-KDEL, anti-GM130 and anti-p58k antibodies. Their association with endosomal, lysosomal and ER compartments was assessed by confocal laser scanning microscopy (A) and quantitation of colocalization is shown in (B). The results
shown are representative of three independent experiments performed in triplicate. The data represent means ± standard deviation.

Fig. 8. *The* *L. pneumophila* *ankH* or *ankJ* mutants are rescued within HEK293 cells expressing AnkH or AnkJ-GFP fusion proteins, respectively. HEK293 cells with stable expression of GFP, AnkH and AnkJ-GFP fusion proteins were infected with the wild-type strain, the *dotA, ankH* or *ankJ* mutant at MOI of 5. After 10h post-infection, cells were stained for laser scanning confocal microscopy analysis (A). Percentage of infected cells harboring ≤3, 4-7 and >7 bacteria per cell were determined based on analyses of 100 infected cells shown in (B). The results shown are representative of three independent experiments performed in triplicate. The data represent means ± standard deviation.
Fig. 5

A.

% mortality

Days

Strain

1
1
1
2
2
2
3
3
3
WT
ankB
ankH
ankJ
AA100
ankB
ankH
ankJ
AA100
ankB
ankH
ankJ

1.00E+06
1.00E+07
1.00E+08
8.00E+08
1.00E+09

B.

Log_{10} CFU/ml

2h
24h
48h
72h
7days

WT
AnkB
AnkH
AnkJ
Fig. 7
A. 

<table>
<thead>
<tr>
<th></th>
<th>GFP</th>
<th>GFP-AnkH</th>
<th>GFP-AnkJ</th>
<th>untrasfected cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td><img src="WT-GFP.png" alt="Image" /></td>
<td><img src="WT-GFP-AnkH.png" alt="Image" /></td>
<td><img src="WT-GFP-AnkJ.png" alt="Image" /></td>
<td><img src="WT-untrasfected.png" alt="Image" /></td>
</tr>
<tr>
<td>dotA</td>
<td><img src="dotA-GFP.png" alt="Image" /></td>
<td><img src="dotA-GFP-AnkH.png" alt="Image" /></td>
<td><img src="dotA-GFP-AnkJ.png" alt="Image" /></td>
<td><img src="dotA-untrasfected.png" alt="Image" /></td>
</tr>
<tr>
<td>ankJ</td>
<td><img src="ankJ-GFP.png" alt="Image" /></td>
<td><img src="ankJ-GFP-AnkH.png" alt="Image" /></td>
<td><img src="ankJ-GFP-AnkJ.png" alt="Image" /></td>
<td><img src="ankJ-untrasfected.png" alt="Image" /></td>
</tr>
</tbody>
</table>

B. 

![Bar chart](Bar_chart.png)

Fig. 8