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The secreted protein kinase CstK from *Coxiella burnetii* influences vacuole development and interacts with the GTPase-activating host protein TBC1D5

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Running title: CstK affects C. burnetii vacuole biogenesis

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The authors declare that they have no conflicts of interest with the contents of this article.

**ABSTRACT**

The intracellular bacterial pathogen *Coxiella burnetii* is the etiological agent of the emerging zoonosis Q fever. Crucial to its pathogenesis is type 4b secretion system (T4SS)-mediated secretion of bacterial effectors into host cells that subvert host cell membrane trafficking, leading to the biogenesis of a parasitophorous vacuole for intracellular replication. The characterization of prokaryotic Serine/Threonine Protein Kinases (STPKs) in bacterial pathogens is emerging as an important strategy to better understand host-pathogen interactions. In this study, we investigated CstK (for *Coxiella* ser/thr Kinase), a protein kinase identified in *C. burnetii* by in silico analysis. We demonstrate that this putative protein kinase undergoes autophosphorylation on Thr and Tyr residues, and phosphorylates a classical eukaryotic protein kinase substrate *in vitro*. This dual Thr-Tyr kinase activity is also observed for a eukaryotic dual-specificity Tyr phosphorylation-regulated kinase class. We found that CstK is translocated during infections and localizes to *Coxiella*-containing vacuoles (CCVs). Moreover, a CstK- overexpressing *C. burnetii* strain displayed a severe CCV development phenotype, suggesting that CstK fine-tunes CCV biogenesis during the infection. Protein-protein interaction experiments identified the Rab7 GTPase-activating protein (GAP) TBC1D5 as a candidate CstK-specific target, suggesting a role for this host GAP in *Coxiella* infections. Indeed, CstK colocalized with TBC1D5 in non-infected cells, and TBC1D5 was recruited to CCVs in infected cells. Accordingly, TBC1D5 depletion from infected cells significantly affected CCV development. Our results indicate that CstK functions as a bacterial effector protein that interacts with the host protein TBC1D5 during vacuole biogenesis and intracellular replication.

Signal transduction is an essential and universal function that allows all cells, from prokaryotes to eukaryotes, to translate environmental signals to adaptive changes. By this mechanism, extracellular inputs propagate through complex signaling networks whose activity is often regulated by reversible protein phosphorylation. Signaling mediated by Serine/Threonine/Tyrosine protein phosphorylation has been extensively studied in eukaryotes, however, its relevance in prokaryotes has only begun to be appreciated. The recent discovery that bacteria also use Ser/Thr/Tyr kinase-based signaling pathways has opened new perspectives to study environmental adaptation, especially in the case of bacterial pathogens, with respect to host infection (1). Thus, advances in genetic strategies and genome sequencing have revealed the existence of "eukaryotic-like" serine/threonine protein kinases (STPKs) and phosphatases in a number of prokaryotic organisms...
(2), including pathogens such as, *Streptococcus* spp. (3-6), *Mycobacteria* (7-12), *Yersinia* spp. (13,14), *Listeria monocytogenes* (15,16), *Pseudomonas aeruginosa* (17), *Enterococcus faecalis* (18) or *Staphylococcus aureus* (19,20). Consequently, the study of STPKs in human bacterial pathogens is emerging as an important strategy to better understand host-pathogen interactions and develop new, targeted antimicrobial therapies. However, if on one hand it is clear that STPKs and phosphatases regulate important functions in bacterial pathogens, their signal transduction mechanism remains ill-defined and restricted to a limited number of microbes.

Importantly, STPKs expressed by pathogenic bacteria can either act as key regulators of important microbial processes, or be translocated by secretion systems to interact with host substrates, thereby subverting essential host functions including the immune response, cell shape and integrity (21). Phosphorylation of host substrates has been demonstrated for some bacterial STPKs, whereas others seem to require their kinase activity but their phosphorylated substrates remain to be identified (21). Therefore, biochemical mechanisms of these pathogen-directed targeted perturbations in the host cell-signaling network are being actively investigated and STPKs are proving to be molecular switches that play key roles in host-pathogen interactions (21).

Among emerging human pathogens, *Coxiella burnetii* is a highly infectious bacterium, responsible for the zoonosis Q fever, a debilitating flu-like disease leading to large outbreaks with a severe health and economic burden (22-24). The efficiency of infections by *C. burnetii* is likely associated with the remarkable capacity of this bacterium to adapt to environmental as well as intracellular stress. Indeed, outside the host, *C. burnetii* generates pseudo-spores that facilitate its airborne dissemination. *C. burnetii* has developed a unique adaptation to the host, being the only bacterium that thrives in an acidic compartment containing active lysosomal enzymes. Upon host cell invasion, bacteria reside within membrane-bound compartments that passively traffic through the endocytic maturation pathway, progressively acquiring early and late endocytic markers such as Rab5 and Rab7, respectively (25). Fusion of *Coxiella*-containing vacuoles (CCVs) with late endosomes and lysosomes is accompanied by the acidification of the endosomal environment, which is required to activate the translocation of bacterial effector proteins by a Dot/Icm Type 4b Secretion System (T4SS) (26). Some of these effectors modulate important signaling pathways of infected cells, including apoptosis and inflammasome activation (27-29), whereas others are essential for the development of the intracellular replicative niche. Among these CvpB and CvpF have been recently implicated in the manipulation of autophagy for optimal vacuole development (30-33). *C. burnetii* genome analysis revealed a close homology to the facultative intracellular pathogen *Legionella pneumophila*, in particular at the level of Dot/Icm core genes (34). *In silico* analysis identified over 100 candidate effector proteins encoded in the *C. burnetii* genome, some of which have been validated for secretion using either *C. burnetii* or *L. pneumophila* as a surrogate model (26,35,36).

In this study, we investigated the candidate effector CBU_0175, which encodes a unique putative *Coxiella* Ser/Thr kinase (CstK). We demonstrated CstK translocation by *C. burnetii* during infection and we reported its localization at CCVs. *In vitro* kinase assays revealed that CstK undergoes autophosphorylation on Thr and Tyr residues, and displays a ‘bona fide’ kinase activity towards a test substrate of eukaryotic protein kinases. Furthermore, the identification of the Rab7 GTP-activating protein (GAP) TBC1D5 as a CstK interactor suggests that this protein might be involved during infection to facilitate CCVs biogenesis. Indeed, TBC1D5 is actively recruited at CCVs during *Coxiella* infections and TBC1D5-targeting siRNAs significantly affect CCVs development. Our data provide the first evidence that a *C. burnetii* secreted kinase might control host cell infection.

**RESULTS**

*Coxiella burnetii* genome encodes a single putative protein kinase. *In silico* analysis of the virulent *C. burnetii* strain RSA493 NMI genome revealed only one gene encoding a putative Serine/Threonine Protein Kinase (STPK). To date, no STPKs have been characterized in this organism. This gene was named *cstK* for *C. burnetii* serine threonine Kinase and encodes a 246 amino acids protein with an estimated molecular mass of 31 kDa. The gene coding for *cstK* is flanked by genes CBU_0174 (which encodes an hypothetical protein) and CBU_0176, a gene coding for the serine protease domain-containing protein degP1. Of note, these genes are not part of an operon (Fig. 1A). InterProScan analysis of CstK revealed the presence of most of the essential amino acids and sequence subdomains characterizing the Hanks family of eukaryotic-like protein kinases (37). CstK shares a common eukaryotic protein kinases
superfamily fold with two lobes and a Gly-rich loop. These include the central core of the catalytic domain, and the invariant lysine residue in the consensus motif within subdomain II, which is usually involved in the phosphate transfer reaction and required for the autophosphorylating activity of eukaryotic STPKs (Fig. 1A) (37-39). The activation loop in the catalytic domain is particularly short in CstK, and the DFG motif is substituted by a GLG motif. Interestingly, the transmembrane domain usually present in classical prokaryotic STPKs is lacking in CstK, thus it is a so-called cytoplasmic STPK.

**CstK is a Dot/Icm effector protein.** Bioinformatics analysis using the prediction software S4TE 2.0 (40) indicated that CstK harbors features corresponding to secreted effector proteins, including a promoter motif typically found in effector proteins from intra-vacuolar bacterial pathogens, suggesting that CstK is indeed a *Coxiella* effector protein (Fig. 1A). Consistently, previous studies by Chen and colleagues have shown that CstK is secreted in a T4SS-dependent manner by the surrogate host *L. pneumophila*, albeit with low efficiency (36). In order to validate CstK secretion in *C. burnetii*, we engineered plasmids encoding, either CstK or CvpB (a known *C. burnetii* effector protein) (30), fused to Beta-Lactamase (BLAM) and expressed in wt *Coxiella Tn1832* (a *C. burnetii* transposon mutant expressing GFP and that phenocopies wild-type *C. burnetii*) or the Dot/Icm-defective *dotA::Tn* mutant, also expressing GFP. By means of a BLAM secretion assay, we could observe that BLAM-CstK was secreted by wt *Coxiella* at 48 and 72 hours, but not at 24 hours post-infection (Fig. 1B). Secretion of BLAM-CvpB or BLAM-CstK was not detectable in cells infected with the *dotA::Tn* strain, indicating that both CvpB and CstK are *C. burnetii* Dot/Icm substrates (Fig. 1B). Next, the intracellular localization of CstK was investigated by ectopically expressing HA-tagged CstK either in non-infected or wt *C. burnetii*-infected U2OS cells. In non-infected cells, CstK localized at intracellular compartments that were negative for the lysosomal marker LAMP1, whereas it was recruited at CCVs (as revealed by the co-localization with LAMP1) in infected cells (Fig. 1C).

**CstK displays autokinase and protein kinase activities.** To determine whether CstK is a functional protein kinase, this protein was overproduced in *E. coli* and purified as a recombinant protein fused to glutathione S-transferase (GST) tag. The purified tagged CstK protein (Fig. 2A, upper panel) was then assayed for autokinase activity in presence of the phosphate donor [\(\gamma^{33}\text{P}\)]ATP. As shown in Fig. 2A (lower panel), CstK incorporated radioactive phosphate from [\(\gamma^{33}\text{P}\)]ATP, generating a radioactive signal corresponding to the expected size of the protein isoform, strongly suggesting that this kinase undergoes autophosphorylation. To confirm CstK autophosphorylation and exclude that contaminant kinase activities from *E. coli* extracts might phosphorylate CstK, we mutated the conserved Lys55 residue present in subdomain II into CstK by site-directed mutagenesis. Indeed protein sequence analysis revealed that Lys55 in CstK is similarly positioned as a conserved Lys residue usually involved in the phosphotransfer reaction and also required for the autophosphorylating activity of eukaryotic-like STPKs (37,38). Thus, Lys55 was substituted by a Met residue, the mutated form of CstK, CstK_K55M, was purified as described above (Fig. 2A, upper panel) and it was then tested for autophosphorylation in presence of [\(\gamma^{33}\text{P}\)]ATP. As expected, no radioactive signal could be detected (Fig. 2A, lower panel), thus establishing that CstK displayed autophosphorylation activity. A kinetic analysis of CstK phosphorylation was next performed to determine the initial CstK phosphorylation rate (Fig. 2B). Incorporation of \(\gamma\)-phosphate occurred rapidly, reaching about 50% of its maximum rate within 5 min of reaction. This autokinase activity was dependent on bivalent cations such as Mg\(^{2+}\) and Mn\(^{2+}\) in the range of 5mM, thus in correlation with concentrations required for canonical STPK activity, as shown in Fig. 2C, and abolished by addition of 20 mM EDTA chelating all the divalent cations available (data not shown).

The recombinant CstK protein was further characterized by studying its ability to phosphorylate exogenous proteins and was thus assayed for *in vitro* phosphorylation of the general eukaryotic protein kinase substrate, myelin basic protein (MBP), in the presence of [\(\gamma^{33}\text{P}\)]ATP. MBP is a commonly used substrate for both Ser/Thr and Tyr kinases. A radiolabeled signal at the expected 18kDa molecular mass of MBP was detected, thus demonstrating that CstK phosphorylates protein substrates such as MBP (Fig. 2A). As expected, the CstK_K55M mutant did not phosphorylated MBP. Altogether, these data indicate that *in vitro*, CstK possesses intrinsic autophosphorylation activity and displays kinase functions for exogenous substrates.

**Identification of CstK autophosphorylation sites.** To determine the specificity of this kinase, we next identified its autophosphorylation sites. A mass
spectrometry approach was used since this technique allows precise characterization of post-translational modifications including phosphorylation (41,42). NanoLC/nanospray/tandem mass spectrometry (LC-ESI/MS/MS) was applied for the identification of phosphorylated peptides and for localization of phosphorylation sites in CstK. This approach led to 97% of sequence coverage, while the remaining residues uncovered that did not include Ser, Thr or Tyr residues.

As detailed in Table 3, analysis of tryptic digests allowed the characterization of three phosphorylation sites in CstK. Surprisingly, unlike classical Ser/Thr or Tyr kinases, CstK was phosphorylated on two Tyr residues (Y14 and Y209), in addition to one Thr site (T232). Since protein sequence analysis did not reveal a classical activation loop in this kinase, the contribution of T232, Y14 and Y209 to CstK kinase activity was individually assessed. Hence, these residues were mutated either to phenylalanine to replace tyrosine residues or alanine to replace threonine residue, generating the single mutants CstK_Y14F, CstK_Y209F, and CstK_T232A as well as the CstK_Y14F/Y209F/T232A triple mutant (CstK_FFA). Next, in vitro kinase assays with [γ-32P]ATP were carried out and revealed that maximum loss in CstK autophosphorylation activity was observed in the CstK_Y14F mutant (Fig. 2D), suggesting that this site is central for CstK activation. In contrast, the CstK_Y209F mutant exhibited a slight hyperphosphorylation, which might indicate that Y209 only plays an accessory role in controlling CstK autophosphorylation (Fig. 2D). Finally, the CstK_T232A mutant showed a reduced CstK phosphorylation and displayed diminished kinase activity towards the exogenous substrate MBP (Fig. 2D). Note that mutating all three autophosphorylation sites fully abrogated CstK kinase activity (Fig. 2D). These results indicate that Y14 and T232 are the major phosphorylation sites in CstK and strongly suggest that CstK might be a dual specificity (Thr/Tyr) kinase.

CstK activity and phosphorylated state affect its intracellular localization.

Next, we ectopically expressed HA-tagged CstK, CstK_K55M and CstK_FFA derivatives in non-infected and C. burnetii infected U2OS cells, to investigate its intracellular localization. CstK mainly localized at vesicular compartments in non-infected cells, and accumulated at CCVs upon C. burnetii infection, suggesting an active role in the biogenesis of this compartment (Fig. 1C). Interestingly, the inactive CstK_K55M mutant localized at vesicular structures positive for the lysosomal marker LAMP1 but not at CCVs while the non-phosphorylated CstK_FFA displayed a diffuse localization in the cytosol of transfected cells (Fig. 2E). Overall, these data suggest that the kinase activity and phosphorylated state of CstK play an important role in its localization in cells.

CstK regulates vacuole development and C. burnetii replication within infected cells. As a first step towards the understanding of CstK functions in the course of infection, and to appreciate the extent to which this kinase is required for growth and viability of C. burnetii, we attempted to inactivate the corresponding chromosomal gene. Unfortunately, after several attempts we were unable to generate a null mutant suggesting that cstK might be essential. However, we had previously isolated a C. burnetii mutant (Tn2496) carrying a transposon insertion allowing GFP expression at position 156783, 32 bp upstream of the starting codon of cstK (43) (Fig. 1A). To determine the effect of this transposon insertion on cstK gene expression, we assessed the expression level of cstK mRNA from wt C. burnetii and Tn2496 strains. Surprisingly, cstK expression was significantly upregulated in the mutant strain, suggesting that the transposon insertion may have released a transcriptional negative regulation (Fig. 3A). This suggested that a putative transcriptional regulator might bind the cstK promoter and control its activity during host invasion.

We next examined the effects of CstK overexpression on C. burnetii infections by challenging Vero cells either with wt C. burnetii, the Dot/Icm-defective dotA::Tn mutant or the Tn2496 mutant. Intracellular growth of the CstK overexpressing strain was significantly reduced over 7 days of infection with an intermediate phenotype between wt and the dotA::Tn mutant (Fig. 3B). Accordingly, multiparametric phenotypic profile analysis of the Tn2496 mutant indicated that this strain exhibited a major defect in CCV development as compared to wt C. burnetii (Fig. 3C & D). To further investigate the effects of CstK overexpression on C. burnetii infections, GFP-expressing C. burnetii were transformed with plasmids expressing HA-tagged wt CstK or its corresponding mutants (CstK_K55M and CstK_FFA) under the control of an IPTG promoter. U2OS cells expressing cytoplasmic mCherry were challenged with the three C. burnetii strains in the presence or absence of IPTG. After 6 days of infection, cells were fixed, labelled with Hoechst and anti-LAMP1 antibody to visualize host cells nuclei and CCVs respectively, and
processed for automated image analysis. In all cases, the overexpression of CstK was detrimental for CCVs biogenesis and bacterial replication (Fig. 3E). Next, U2OS cells were challenged either with wt C. burnetii, the GFP-expressing Tn2496 mutant strain or a combination of the two for 6 days (Fig. 3F). Cells were then fixed and labelled with an anti-C. burnetii antibody to label both bacteria strains and incubated with Hoechst to visualize host cells nuclei (Fig. 3F). Automated image analysis was then used to determine the effects of CstK overexpression on the replication of wt bacteria, in trans. Co-infections resulted in a significant increase in the size of Tn2496 colonies, indicating that wt C. burnetii can partially restore the growth of the CstK-overexpressing strain (Fig. 3G). However, a significant decrease in the size of bacterial colonies labelled by the anti-C. burnetii antibodies indicated that CstK-overexpression has a detrimental effect in trans, on the development of wt bacteria (Fig. 3G). Of note, vacuoles harboring wt or mutant colonies alone were never observed in co-infection experiments. Therefore, we concluded that CstK participates in the formation of the C. burnetii replicative vacuole, and that its expression must be finely tuned for optimal intracellular replication.

**CstK specifically interacts with host cell proteins.** Since CstK is a secreted protein, we assume that this kinase would interfere with host cell signal transduction pathways to subvert host cell defenses to the benefit of the bacteria. To identify host cell proteins that could interact with CstK, we made use of the model amoeba Dictyostelium discoideum. *D. discoideum* is a eukaryotic professional phagocyte amenable to genetic and biochemical studies. Lysate from cells overexpressing CstK tagged with a C-terminal FLAG epitope (CstK-FLAG) was incubated with beads coupled to an anti-FLAG antibody. Beads were extensively washed, and bound proteins were separated by SDS-PAGE before mass-spectrometry analysis. Among the putative interactants of CstK identified by this approach, some were discarded on the basis of their intracellular localization while other retained candidates were mostly involved in the endocytic pathway (Table S1). Among these, the Rab GAP/TBC domain-containing protein, DDB_G0280253 (UniProtKB - Q54VM3), is a 136.4 kDa protein homologous to mammalian TBC1D5 (http://dictybase.org), a GTPase-activating protein (GAP) for Rab7a and Rab7b (44-46) that acts as a molecular switch between the endosomal and the autophagy pathway (47). Given the recently reported implication of TBC1D5 in the biogenesis of *L. pneumophila*-containing vacuoles (48) and the role of autophagy in the biogenesis of CCVs (25,31,49), we aimed at validating the interaction between human TBC1D5 (Hs-TBC1D5) and CstK in HEK-293T cells. Cells co-expressing Hs-TBC1D5-GFP and HA-CstK or CstK mutants were used for immunoprecipitation using anti-HA beads. Wild-type and CstK derivatives were detected as co-immunoprecipitated in presence of Hs-TBC1D5-GFP, thus confirming that Hs-TBC1D5 is a bona fide CstK interactant (Fig. 4A). Significantly higher levels of TBC1D5 were co-immunoprecipitated by the CstK mutants suggesting that the interaction might be increased in the absence of phosphorylation turnover of the kinase (Fig 4A, lower panel). Interestingly, the interaction is not dependent on the phosphorylation status of CstK as the K55M mutant or the triple FFA mutant are still able to interact. Other candidates identified by mass spectrometry are currently being investigated.

**TBC1D5 is recruited at CCVs and regulates their biogenesis.** Given its recently reported role in the development of *L. pneumophila*-containing vacuoles, we investigated the localization of TBC1D5 in U2OS cells infected either with wt *C. burnetii*, the CstK-overexpressing strain Tn2496 or the Dot/Icm-defective mutant dotA::Tn. Consistently with previous studies demonstrating a role in the activation of Rab7a and b, TBC1D5 seems to accumulate specifically at CCVs in a Dot/Icm-independent manner as the eukaryotic protein was found at CCVs generated by all *C. burnetii* (wt, the Tn2496 and the dotA::Tn Fig. 4B and Supplementary Fig.1). Next, we used siRNA to deplete cells of TBC1D5 prior to *C. burnetii* infection, to investigate a possible role in CCVs development and intracellular bacterial replication. Indeed, vacuole development was significantly reduced in cells exposed to Hs-TBC1D5-targeted siRNAs as opposed to cells treated with non-targeting siRNA oligonucleotides (Fig. 4C).

**TBC1D5 is not phosphorylated in vitro by recombinant CstK.** We assessed whether CstK might phosphorylate the recombinant Hs-TBC1D5. Despite the *in silico* prediction of several Ser/Thr and Tyr phosphorylatable residues in Hs-TBC1D5, we failed to detect Hs-TBC1D5 phosphorylation using several *in vitro* kinases assays (Supplementary Fig.2). In addition, Hs-TBC1D5 phosphorylation status was also investigated upon transfection with CstK or its inactive derivative (K55M) followed by Hs-TBC1D5 immunoprecipitation. No
phosphorylation could be detected in our experimental conditions.

**DISCUSSION**

Bacterial Ser/Thr/Tyr kinases expressed by pathogenic bacteria can either act as key regulators of important microbial processes, or be translocated by secretion systems to interact with host substrates, thereby our results provide the first biochemical analysis of the secreted *C. burnetii* kinase CstK and its involvement in the process of infection and CCVs development. Importantly, CstK presents important differences as compared to classical Ser/Thr kinases. In particular, we provided evidence that CstK is a dual kinase able to autophosphorylate on Thr and Tyr residues. Moreover, the observation that a transposon insertion 32 bp upstream of the *cstK* starting codon leads to an increase in the levels of *cstK* mRNAs was indicative of the presence of a negative transcriptional regulation of gene expression, suggesting a fine tuning of the levels of CstK. Indeed, the *Tn*2496 mutant displays a severe CCV biogenesis defect when used to challenge U2OS cells, highlighting the importance of regulating *cstK* expression during *C. burnetii* infections. Accordingly, inducing the expression of wt CstK in wt *C. burnetii* severely impairs CCVs development and bacterial replication. Co-infection experiments demonstrated that CstK overexpression can also act in trans, by perturbing the intracellular replication of wt *C. burnetii*. The identification of candidate eukaryotic interactors of CstK further corroborated a role of the bacterial kinase in subverting host functions during infection. Here we confirmed that CstK interacts with TBC1D5, but we failed to detect phosphorylation of the eukaryotic target by CstK.

Indeed TBC1D5 associates the retromer VPS29 subunit involved in endosomal trafficking, and upon autophagy induction, the autophagy ubiquitin-like protein LC3 can displace VPS29, thus orienting TBC1D5 functions towards autophagy instead of endosomal functions (47). It is thus tempting to propose that CstK might interfere with this tight regulation between TBC1D5, LC3 and VPS29, and redirect TBC1D5 functions to support efficient *C. burnetii* intracellular replication. Further work will need to be carried out to decipher how CstK recognizes these host substrates and how they participate in the establishment of *C. burnetii* parasitophorous vacuoles. Another perspective of this work is the opening of a new field of investigation for future drug development to fight this pathogen. Because CstK seems to be essential, specific inhibitors of this kinase preventing *C. burnetii* growth would be extremely useful for the development of new therapies.

**EXPERIMENTAL PROCEDURES**

**Bacterial strains and growth conditions.** Bacterial strains and plasmids are described in Table 1. Strains used for cloning and expression of recombinant proteins were *Escherichia coli* TOP10 (Invitrogen) and *E. coli* BL21 (DE3)Star (Stratagene), respectively. *E. coli* cells were grown and maintained at 25 °C in LB medium supplemented with 100 µg/ml ampicillin when required. *Coxiella burnetii* RSA439 NMII and transposon mutants *Tn*1832, *Tn*2496, and *dot*A::*Tn* were grown in ACCM-2 (45) supplemented with chloramphenicol (3 mg/ml) in a humidified atmosphere of 5% CO2 and 2.5% O2 at 37 °C.

**Cloning, expression and purification of CstK derivatives.** The *cstK* (*CU*0175) gene was amplified by PCR using *Coxiella burnetii* RSA439 NMII chromosomal DNA as a template with the primers listed in Table 2 containing a *Bam*HI and *Hind*III restriction site, respectively. The corresponding amplified product was digested with *Bam*HI and *Hind*III, and ligated into the bacterial pGEX(M) plasmid, that includes a N-terminus GST-tag, thus generating pGEX(M)·cstK. pGEX(M)·cstK derivatives harboring different mutations were generated by using the QuikChange Site-Directed Mutagenesis Kit (Stratagene, La Jolla, CA), and resulted in the construction of plasmids detailed in Table 2. For overexpression assays, *cstK* and its derivatives were cloned in the pJA-LACO-4xHA vector (30) using *Kpn*I and *Bam*HI restriction sites. All constructs were verified by DNA sequencing.
Transfected E. coli BL21 Star cells with pGEX(M)\textsuperscript{cstK} derivatives were grown at 16 °C in LB medium containing 1 g/liter of glucose and 100 µg/ml of ampicillin and protein synthesis induced with 0.5 mM IPTG overnight. Bacteria were disrupted by sonication (Branson, digital sonifier) and centrifuged at 14,000 rpm for 25 min. Purifications of the GST-tagged recombinants were performed as described by the manufacturer (GE Healthcare). \textit{cstK} coding sequence was also optimized for mammalian cell expression (GenScript), amplified by PCR and cloned into pDXA-3C (50) containing a FLAG-tag for C-terminal fusion. After sequencing, the plasmid was linearized by the restriction enzyme Scal and transfected in \textit{D. discoideum} as described (51). Clone selection was made with 10 mg/ml of G418, and protein expression was assayed by Western blot analysis of \textit{D. discoideum} crude extract with an anti-FLAG rabbit polyclonal antibody (GenScript, USA). For ectopic expression assays, \textit{cstK, cstK.Marshal} and \textit{cstK.FLAG} with optimized codons (IDT) were cloned in pRK5-HA using the primer pair CstKopt primers. H\textit{sTBC1D5} coding sequence has been recombined into pEGFP-N1 RfC Destination vector by the same method (MGC Platform Montpellier). Thus generating pEGFP-\textit{HsTBC1D5} fusion. Human \textit{TBC1D5} coding sequence was obtained from \textit{hORFeome v8.1} and recombined with \textit{HsTBC1D5} ORFs in Gateway Entry clones ready for transfer to Gateway-compatible expression vectors). "HsTBC1D5" coding sequence has been recombined into pEGFP-N1 RfC Destination vector by Gateway reaction (MGC Platform Montpellier) thus generating pEGFP-N1_HsTBC1D5 coding for H\textit{sTBC1D5} with a C-terminal GFP Tag. pmCH_Hs-TBC1D5-mCherry has been generated by the same method (MGC Platform Montpellier).

\textbf{Cloning, expression and purification of TBC1D5 derivatives.} The \textit{D. discoideum} GFP-tagged TBC1D5 was previously generated (48). Cells were grown at 22 °C in HL5 medium as previously described (51). Human TBC1D5 coding sequence was obtained from the hORFeome v8.1 (ORF 2659, Q92609, fully-sequenced cloned human ORFs in Gateway Entry clones ready for transfer to Gateway-compatible expression vectors). H\textit{sTBC1D5} coding sequence has been recombined into pEGFP-N1 RfC Destination vector by Gateway reaction (MGC Platform Montpellier) thus generating pEGFP-N1_HsTBC1D5 coding for H\textit{sTBC1D5} with a C-terminal GFP Tag. pmCH_Hs-TBC1D5-mCherry has been generated by the same method (MGC Platform Montpellier).

\textbf{RNA extraction and quantitative RT-PCR (qRT-PCR).} Fifty ml of \textit{C. burnetii} culture was harvested, resuspended in 600 µL of RNA protect reagent (Qiagen) and incubated 5 min at room temperature. Bacteria were centrifuged and resuspended in 200 µL TE (10 mM Tris-HCl, 1 mM EDTA, pH 8) containing 1 mg/ml lysozyme. Bacterial suspension was incubated at room temperature for 5 min and bacteria were disrupted by vigorous vortexing for 10 sec every 2 min. 700 µL of RLT buffer from RNA easy kit (Qiagen), were added to the bacterial suspension and disruption was completed by vortexing vigorously. RNA was purified with the RNA easy kit according to manufacturer’s instructions. DNA was further removed using DNase I (Invitrogen). cDNA was produced using Superscript III reverse transcriptase (Invitrogen). Controls without reverse transcriptase were done on each RNA sample to rule out possible DNA contamination. Quantitative real-time PCR was performed using LightCycler 480 SYBR Green I Master (Roche) and a 480 light cycler instrument (Roche). PCR conditions were as follows: 3 min denaturation at 98 °C, 45 cycles of 98 °C for 5 sec, 60 °C for 10 sec and 72 °C for 10 sec. The \textit{dotA} gene was used as internal control. The sequences of primers used for qRT-PCR are listed in Table 2. The expression level of \textit{cstK} in the wild type strain was set at 100 and the expression levels of \textit{cstK} in the \textit{Tn2496} mutant were normalized to the wild type levels.

\textbf{In vitro kinase assays.} \textit{In vitro} phosphorylation was performed with 4 µg of wild-type CstK or CstK derivatives in 20 µl of buffer P (25 mM Tris-HCl, pH 7.0; 1 mM DTT; 5 mM MnCl\textsubscript{2}; 1 mM EDTA; 50 µM ATP) with 200 µCi ml\textsuperscript{-1} (65 mM) \(\gamma\textsuperscript{32}P\text{ATP}\) (PerkinElmer, ref: NEG 602H250UC, 3000 Ci mmol\textsuperscript{-1}) for 30 min at 37 °C. For substrate phosphorylation, 4 µg of MBP (Myelin Basic Protein) (Sigma) and 4 µg of CstK were used. Each reaction mixture was stopped by addition of an equal volume of Laemmli buffer and the mixture was heated at 100 °C for 5 min. After electrophoresis, gels were soaked in 16% TCA for 10 min at 90 °C, and dried. Radioactive proteins were visualized by autoradiography using direct exposure to films.

\textbf{Mass spectrometry analysis.} For mass spectrometry analysis, CstK was phosphorylated as described above, except that \(\gamma\textsuperscript{32}P\text{ATP}\) replaced with 5 mM cold ATP. Subsequent mass spectrometry analyses were performed as previously reported (52,53). Briefly, samples were submitted to trypsin digestion and analyzed using an Ultimate 3000 nano-RSLC (Thermo Scientific, San Jose California) coupled on line with a quadrupole-orbitrap Q Exactive HF mass spectrometer via a nano-electrospray ionization source (Thermo Scientific, San Jose California). Samples were injected and loaded on a C18 Acclaim PepMap100 trap-column (Thermo Scientific) and separated on a C18 Acclaim Pepmap100 nano-column (Thermo Scientific). MS data were acquired in a data dependent strategy selecting the fragmentation events based on the 20 most abundant precursor ions in the survey scan (350-1600 Th). The resolution of the survey scan was 60,000 at m/z 200 Th and for MS/MS scan the resolution was set to 15,000 at m/z 200 Th.
Peptides selected for MS/MS acquisition were then placed on an exclusion list for 30s using the dynamic exclusion mode to limit duplicate spectra. Data files were then analyzed with Proteome Discover 2.2 using the SEQUEST HT algorithm against the Uniprot Dictyostelium discoideum to which was included the sequence of CstK.

C. burnetii infections. U2OS epithelial cells were challenged either with Coxiella burnetii RSA439 NMII, the transposon mutants Tn1832, dotA::Tn or Tn2496 as previously described (38,49). For co-infection experiments, cells were challenged with a 1:1 ratio of Coxiella burnetii RSA439 NMII and Tn2496 transposon mutant. For gene silencing, U2OS cells were seeded at 2,000 cells per well in black, clear-bottomed, 96-well plates in triplicate and transfected with siRNA oligonucleotides 24 h later by using the RNAiMAX transfection reagent (Thermo Fisher Scientific) according to the manufacturer’s recommendations. At 24 h post transfection, cells were challenged with C. burnetii (MOI of 100) and further incubated for 5 d. Cells were then fixed and processed for immunofluorescence. Where appropriate, anti-LAMP1 antibodies were used to label lysosomes and CCVs as previously described (54). Samples were imaged with a Zeiss Axio Imager Z1 epifluorescence microscope (Carl Zeiss, Germany) connected to a CoolSNAP HQ2 CCD camera (Teledyne Photometrics, Tucson, AZ). Images were acquired with 40x oil immersion objectives (Teledyne Photometrics, Tucson, AZ). For phenotypic screening, samples were imaged with an ArrayScan VTI Live fluorescence automated microscope (Cellomics) equipped with an ORCA-ER CCD camera (Hamamatsu). 25 fields per well were acquired for image analysis. ImageJ and ICY software were used for image analysis and quantifications. Phenotypic profiles (expressed as z-scores) were calculated using CellProfiler, from triplicate experiments as previously described (54) following median based normalization of 96-well plates. Plates effects were corrected by the median value across wells that are annotated as control.

Beta-lactamase translocation assay. Effector proteins translocation assays were performed as previously described (30). Brieﬂy, Coxiella burnetii Tn1832 (wt) and dotA::Tn were transformed either with pXDC-Blam (negative control), pXDC-Blam-CvpB (positive control) or pXDC-Blam-CstK. Each strain was used to infect U2OS epithelial cells. After 24, 48 or 72 hours of infection, cells were loaded with the fluorescent substrate CCF4/AM (LiveBLAzer-FRET B/G loading kit; Invitrogen) in HBSS (20 mM HEPES pH 7.3) containing 15 mM probenecid (Sigma). Cells were incubated in the dark for 1 h at room temperature and imaged using an EVOS inverted fluorescence microscope. Images were acquired using DAPI and GFP filter cubes. The image analysis software CellProfiler was used to segment and identify all cells in the sample (GFP), positive cells (DAPI) and to calculate the intensity of fluorescence in each channel. The percentage of positive cells versus the total number of infected cells was then calculated and used to evaluate effector translocation.

Immunoprecipitation from D. discoideum lysates. For immunoprecipitation assays, 2x10^5 cells were lysed in lysis buffer (50 mM Tris-HCl pH 7.5, 300 mM NaCl, 0.5% NP40, protease inhibitors (Roche)) and cleared by centrifugation for 15 min at 14 000 rpm in a microfuge. Lysate supernatants were incubated overnight at 4 °C with anti-flag monoclonal antibody coated on agarose beads (GenScript). The beads were then washed five times in wash buffer (50 mM Tris-HCl pH 7.5, 300 mM NaCl, 0.1% NP40) and once in PBS. Bound proteins were migrated on SDS-PAGE and analyzed by LC-MS/MS.

Cell culture, heterologous expression and anti-HA immunoprecipitation. HEK-293T cells were grown in DMEM (Gibco) containing 10% (vol/vol) FBS, 1% glutamax (Gibco, 200 mM stock), 0.5% Penicillin/Streptomycin (Gibco, 10000 U/mL stock) and maintained under standard conditions at 37 °C in a humidified atmosphere containing 5% CO2. Cells were transiently transfected using the jetPEI Transfection Reagent (Polyplus-Transfection Inc.) to express either Hs-TBC1D5-GFP, CstK_HA derivatives, or each CstK derivatives with TBC1D5_GFP protein. Cells were used 24h after transfection for immunoprecipitation assay. Transfected cells were washed two times in cold PBS and lysed in lysis buffer (50 mM Tris, 150 mM NaCl, 0.5 mM EDTA, 0.5 % NP40, protease and phosphatases inhibitors (Roche)). Cleared lysate (950 µL, approximately 1 mg total proteins) were incubated with anti HA magnetic Beads (Pierce) for 30 min at room temperature under gentle rotation. The beads were washed three times in lysis buffer, boiled in 2X Laemmli sample buffer and loaded on ExpressPlus™ PAGE Gels (GenScript). The eluted proteins were visualized by Western Blotting with the following antibodies: anti-HA from Chromotek, anti-GFP from Torrey Pines, donkey anti-rat or anti-rabbit from Jackson ImmunoResearch.
Densitometry.
Regions of Interest (ROIs) were obtained from each band of interest and the intensity was measured using ImageLab (From Biorad). For each band, the same ROI was used for background calculation and removal from areas adjacent to each band.

DATA AVAILABILITY STATEMENT
All the data are contained in this manuscript.

ACKNOWLEDGMENTS
We wish to thank the Montpellier RIO imaging facility at the University of Montpellier, member of the national infrastructure France-BioImaging, that is supported by the French National Research Agency (ANR-10-INBS-04, «Investments for the future»). We acknowledge the contribution of the Protein Science Facility of the SFR Biosciences (UMS3444/CNRS, USB/Inserm, ENS de Lyon, UCBL), especially Frédéric DELOLME and Adeline PAGE who performed the mass spectrometry analysis. This work was supported by grants from the ATIP/AVENIR Program for V.M. and M.B., the Region Occitanie for S. B., a Marie Curie CIG to E.M. (grant n. 293731), the Agence Nationale de la Recherche (ANR) Grant ANR-14-CE14-0012-01 (project AttaQ) to M.B.

CONFLICT OF INTEREST
The authors declare that they have no conflicts of interest with the contents of this article.

REFERENCES


FIGURE LEGENDS

Figure 1. CstK is a Dot/Icm effector protein. (A) Schematic representation of CstK. A conserved promoter-binding domain is predicted between aa -227 and -213 (Apm) and the transposon insertion site for Tn2496 is indicated. The kinase domain is predicted between aa 26 and 174 and is shown in red. The conserved lysine residue is indicated, and the phosphorylated sites are indicated by an asterisk. (B) BLAM secretion assay. U2OS cells were infected for 24, 48, or 72h (black, grey and white bars, respectively) with wt C. burnetii or the dotA::Tn mutant transformed with vectors expressing Beta-Lactamase alone (BLAM, negative control), BLAM-CvpB (positive control) or BLAM-CstK. Protein translocation was probed using the CCF-4 substrate. The average percentage of cells positive for cleaved CCF-4 as compared to the total number of cells was assessed. Values are mean ± SD from 3 independent experiments. ns = non-significant; *** = P<0.0001, one-way ANOVA, Dunnett’s multiple comparison test. CstK localization. (C) U2OS cells ectopically expressing HA-tagged CstK (red) were either left untreated (top panels) or challenged with wt C. burnetii expressing GFP (green) for 3 days. Lysosomal compartments were labelled using anti-LAMP1 antibodies (blue). Scale bars 10 μm.

Figure 2. (A) Biochemical characterization of CstK. Recombinant CstK derivatives were overproduced, purified on glutathione-Sepharose 4B matrix and submitted to gel electrophoresis and stained with Coomassie Blue (upper panel). In vitro phosphorylation assays were performed with [γ-33P]ATP for 30 min and the eukaryotic substrate myelin basic protein (MBP) when required. Proteins were analyzed by SDS-PAGE, and radioactive bands were revealed by autoradiography (lower panel). The upper bands illustrate the autokinase activity of CstK, and the lower bands represent phosphorylated MBP. Standard proteins of known molecular masses were run in parallel. (B) A kinetic analysis was performed by incubation of CstK with [γ-33P]ATP over different times. Proteins were analyzed by SDS-PAGE, and radioactive bands were revealed by autoradiography. (C) Effects of cations on autokinase activity of CstK. Purified CstK protein was subjected to in vitro autophosphorylation assays in the presence of 50 μM ATP [γ-33P]ATP and 5 mM of Mg2+ or Mn2+. Phosphoproteins were separated by SDS-PAGE and then revealed by autoradiography. (D) In vitro phosphorylation of CstK mutant derivatives. The mutated variants CstK_Y14F (harboring a Tyr to Phe substitution at Y14), CstK_Y209F (harboring a Tyr to Phe substitution at Y209), CstK_T232A (harboring a Thr to Ala substitution at T232), or CstK_Y14F/Y209F/T232A (harboring all three substitutions) were incubated in presence of [γ-33P]ATP with or without the eukaryotic substrate myelin basic protein (MBP). Samples were separated by SDS-PAGE, stained with Coomassie Blue (upper panel), and visualized by autoradiography (lower panel). The upper bands illustrate the autokinase activity of CstK, and the lower bands represent phosphorylated MBP. Standard proteins of known molecular masses were run in parallel (kDa lane). (E) U2OS cells ectopically expressing either HA-tagged CstK_K55M or CstK_FFA (red) were challenged with wt C. burnetii expressing GFP (green) for 3 days. Lysosomal compartments were labelled using anti-LAMP1 antibodies (blue). Scale bars 10 μm. Inset magnification scale bars 4 μm.
Figure 3. CstK regulates vacuole development and *C. burnetii* replication within infected cells. (A) Expression of cstk was measured by qRT-PCR from cultures of either wt *C. burnetii* or the Tn2496 mutant strain. Values are means ± SEM of 4 independent experiments. ***, *p < 0.001*, unpaired t-test. (B) Vero cells were challenged either with wt *C. burnetii*, the dotA::Tn mutant or the Tn2496 mutant strains, all expressing GFP. Intracellular replication was monitored over 7 days of infection by monitoring the GFP fluorescence using a microplate reader. Values are mean ± SD from 3 independent experiments. (C) U2OS cells were challenged either with wt *C. burnetii* or the Tn2496 mutant strain, both expressing GFP (green). 7 days post-infection, cells were fixed and labelled with an anti-LAMP1 antibody (red) and Hoechst (blue) to visualize CCVs and host cell nuclei, respectively. Scale bars 10 μm. (D) An average of 50000 cells were automatically imaged and analyzed from triplicate experiments for each condition illustrated in C, and the phenotypic profile of the Tn2496 mutant was compared to that of wt *C. burnetii* and expressed as z-scores over 15 independent features. (E) GFP-expressing *C. burnetii* transformed with plasmids encoding either wt CstK, the K55M or the FFA mutants, all under the regulation of an IPTG-inducible promoters were used to infect U2OS cells for 6 days, in the presence or absence of IPTG. Cells were then fixed and labelled as in C, and an average of 50000 cells were automatically imaged and analyzed for each condition. The mean size of *C. burnetii* colonies and CCVs was calculated (red bars). (F) U2OS cells were challenged either with wt *C. burnetii*, the GFP-expressing mutant Tn2496 or a combination of the two strains for 6 days. Cells were then fixed and labelled with anti-C. burnetii antibodies (anti-NMII) to label both strains. (G) An average of 50000 cells were automatically imaged and analyzed for each condition illustrated in F and the mean size of *C. burnetii* colonies was calculated (red bars). n.s., non-significant; ****, *p < 0.0001*, one-way ANOVA, Dunnett’s multiple comparison test.

Figure 4. CstK interacts with TBC1D5. (A) The interaction between CstK and the human TBC1D5 has been confirmed after cotransfection of HEK-293T cells to express each HA-tagged CstK derivatives with GFP-tagged Hs-TBC1D5. Cells have been lysed and HA-tagged CstK derivatives have been trapped on anti-HA magnetic beads (Pierce). Beads were washed, eluted by boiling, and bound proteins were revealed by Western blot analysis. Anti-HA antibody confirms the immunoprecipitation of CstK derivatives. For the densitometry graph, Regions of Interest (ROIs) were obtained from each band of interest and the intensity was measured (B) Hs_TBC1D5 localization during infection was monitored in U2OS cells expressing mCherry-tagged Hs-TBC1D5 (red) challenged for 4 days either with wt *C. burnetii*, the Dot/Icm-defective dotA::Tn mutant or the CstK-overexpressing mutant Tn2496, all expressing GFP (green). Lysosomal compartments were labelled with anti-LAMP1 antibodies (blue). Scale bars 10 μm. Inset magnification scale bars 4 μm (C) The role of TBC1D5 in *C. burnetii* infections was investigated using siRNA to deplete Hs-TBC1D5 in U2OS prior to challenge with GFP-expressing wt *C. burnetii*. The size of CCVs was automatically calculated over an average of 150000 cells per condition. Red bars indicate medians. ***, *p < 0.0001; **, *p < 0.001; *, *p < 0.01; one-way ANOVA, Dunnett’s multiple comparison test. Scale bars 10 μm.
### Tables

#### Table 1. Strains and plasmids used in this study

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<td><strong>Coxiella burnetii strains</strong></td>
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**Dictyostelium discoideum plasmids**

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Table 2. Primers used in this study

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<tr>
<td>CvpB-KpnI-Fw</td>
<td>GATGTTACCAGCACAGGCGATCCATTG</td>
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</table>

RT-PCR PRIMERS

<table>
<thead>
<tr>
<th>Primers</th>
<th>Sequence</th>
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</thead>
<tbody>
<tr>
<td>DotA-F</td>
<td>GCTCCAGCATCATCATCGT</td>
</tr>
<tr>
<td>DotA-R</td>
<td>GGCACTTACAGCCCTCAT</td>
</tr>
<tr>
<td>CstK-F</td>
<td>GGCAAGGTATAGGCGGAA</td>
</tr>
<tr>
<td>CstK-R</td>
<td>GGGATTCACCATTGGCCT</td>
</tr>
</tbody>
</table>

<sup>a</sup> Restriction sites are underlined and specified into brackets.

<sup>b</sup> Mutagenized codons are shown in bold.
Table 3. Phosphoacceptors identified after phosphorylation of *C. burnetii* CstK. Sequences of the phosphorylated peptides identified in CstK as determined by mass spectrometry following tryptic digestion are indicated, and phosphorylated residues (pT or pY) are shown in bold.

<table>
<thead>
<tr>
<th>Phosphorylated tryptic peptide sequence of CstK</th>
<th>Number of detected phosphate groups LC/MS/MS</th>
<th>Phosphorylated residue(s)</th>
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<tbody>
<tr>
<td>[8-18] LSVADFpYDLKK</td>
<td>1</td>
<td>Y14</td>
</tr>
<tr>
<td>[228-237] EQFNpTAGHLR</td>
<td>1</td>
<td>T232</td>
</tr>
<tr>
<td>[209-215] pYNPHIK</td>
<td>1</td>
<td>Y209</td>
</tr>
</tbody>
</table>
Figure 1

A

---

B

---

C

---
Figure 2

A. Coomassie and autoradiogram of CstK and MBP.

B. Time course of CstK degradation.

C. Effect of metal ions on CstK and MBP.

D. Merged images of wt C. burnetii and HA-CstK_K55M.

E. GFP and LAMP1 localization in wt C. burnetii and HA-CstK_FFA.
**Figure 3**

A. Colony area distribution for wild-type (wt) and Tn2496 strains.

B. Relative fluorescence intensity over days post-infection for wild-type, dotA::Tn, and Tn2496 strains.

C. Immunofluorescence images showing Hoechst staining, C. burnetii, LAMP1, and merged channels.

D. Z-score distribution for various parameters including colony area, CCV area, CCV perimeter, CCV circularity, CCV density, multivacuolar total cell counts, infection foci area, infected cells lys count, non-infected cells lys count, and non-infected cells area.

E. Scatter plots showing colony size (pixel) and normalized CCV area for different strains and conditions.

F. Diagram illustrating the infection process with wild-type (wt), Tn2496, and co-infection conditions.

G. Graph showing colony size distribution with levels of significance indicated.

Legend:
- wt: wild-type
- Tn2496: Tn2496 strain
- co-infection: co-infection of wt and Tn2496
**Figure 4**

A. Transient expression of HA- and GFP-tagged constructs shows their distribution and interactions with C. burnetii. Western blotting with anti-GFP and anti-HA antibodies confirms the expression of the constructs. Immunoprecipitation (IP) with anti-HA antibody and western blotting with anti-GFP antibody reveals the association of CstK with the TBC1D5-GFP complex.

B. Confocal images of C. burnetii with mCherry-TBC1D5 (mCH-TBC1D5), LAMP1, and merged views. The images show the localization of C. burnetii within the host cell and the interaction with LAMP1. The scale bars represent 25 µm.

C. Graphical representation of the log10 of the area (pixels²) of C. burnetii CCVs in cells treated with Ctrl siRNA or TBC1D5 siRNA. The data is presented as a scatter plot with lines indicating the mean. The asterisks denote statistical significance:** P < 0.001, **P < 0.01, *P < 0.05.