Structural analyses of *Legionella* LepB reveal a new GAP fold that catalytically mimics eukaryotic RasGAP

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Rab GTPases are emerging targets of diverse bacterial pathogens. Here, we perform biochemical and structural analyses of LepB, a Rab GTPase-activating protein (GAP) effector from *Legionella pneumophila*. We map LepB GAP domain to residues 313-618 and show that the GAP domain is Rab1 specific with a catalytic activity higher than the canonical eukaryotic TBC GAP and the newly identified VirA/EspG family of bacterial RabGAP effectors. Exhaustive mutation analyses identify Arg444 as the arginine finger, but no catalytically essential glutamine residues. Crystal structures of LepB\(^{313-618}\) alone and the GAP domain of *Legionella drancourtii* LepB in complex with Rab1-GDP-AlF\(_3\) support the catalytic role of Arg444, and also further reveal a 3D architecture and a GTPase-binding mode distinct from all known GAPs. Glu449, structurally equivalent to TBC RabGAP glutamine finger in apo-LepB, undergoes a drastic movement upon Rab1 binding, which induces Rab1 Gln70 side-chain flipping towards GDP-AlF\(_3\) through a strong ionic interaction. This conformationally rearranged Gln70 acts as the catalytic cis-glutamine, therefore uncovering an unexpected RasGAP-like catalytic mechanism for LepB. Our studies highlight an extraordinary structural and catalytic diversity of RabGAPs, particularly those from bacterial pathogens.

**Keywords**: *Legionella*; Rab GTPases; GTPase-activating protein (GAP); type IV secretion system; TBC GAP; membrane trafficking

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**Introduction**

Rab GTPases constitute the largest family of small GTP-binding proteins that act as molecular switches in regulating vesicular transport in eukaryotic cells [1, 2]. Rab GTPases are lipid-modified at their C termini and localized to distinct membrane-bound compartments to ensure corrected delivery of different cargos [3]. Like other small GTPases, Rabs cycle between the GDP-bound “off” and GTP-bound “on” states, the latter of which can interact with downstream effector proteins for appropriate functioning. The conformational states and activities of Rab GTPases are subjected to precise regulation by several classes of Rab modulators [4, 5]. GTPase-activating proteins (GAPs) catalyze GTP hydrolysis and thereby inactivate Rabs on the membrane. Guanine nucleotide dissociation inhibitors (GDIs) extract GDP-bound Rab from the membrane and complex with them in the cytosol. For activation, Rabs are released from RabGDI by the GDI-displacement factor to allow for membrane re-association and subsequent exchange of GDP for GTP, catalyzed by guanine nucleotide exchange factors (GEFs).

The importance of modulating Rab function has also been appreciated by bacterial pathogens. Recent intensive studies on *Legionella pneumophila*, a successful intracellular bacterial pathogen responsible for Legionnaires’ disease, have uncovered a series of stepwise manipulations of Rab1 function by *Legionella* Dot/Icm type
IV secretion system effectors. *L. pneumophila* establishes a replication niche called *Legionella*-containing vacuole (LCV) in host macrophages, which requires recruitment of endoplasmic reticulum (ER)-derived vesicles [6-8]. In line with the fact that the Dot/Icm type IV secretion system is required for maturation and intracellular trafficking of the LCV, several *Legionella* type IV effectors are endowed with activities of manipulating Rab1 cycle and function. SidM/DrrA, one of the Dot/Icm-secreted effectors, recruits Rab1 from the cytosol to the LCV and uses its unique GEF activity to activate Rab1 from the GDI complex [9-14]. The N-terminal domain of SidM bears an additional adenyllyl transferase domain that adenyllylates Rab1-GTP on a switch II tyrosine residue, which prevents GAP-catalyzed GTP hydrolysis and prolongs the duration of Rab1-GTP for activating LidA, a *Legionella*-supplied Rab effector. SidM-induced adenyllylation is enzymatically reversed by another Dot/Icm effector called SidD [16, 17]. This de-adenyllylation modification allows for Rab1 GTP hydrolysis and thereby recycling back to the cytosol, the former of which is catalyzed by LepB, a *Legionella*-encoded RabGAP effector whose mechanism remains unknown [18].

Very recently we discovered a Rab1-specific GAP activity of the VirA and EspG family of type III secretion system effectors from non-vacuole pathogens *Shigella flexneri* and enteropathogenic *E. coli* (EPEC) [19]. These effectors function to counteract autophagy and cytokine secretion-mediated host defenses during *S. flexneri* and EPEC infection, respectively. Notably, the VirA/EspG family of bacterial GAPs adopts a completely different three-dimensional architecture from the host TBC GAPs for Rab GTPases [20], but employs a TBC-like arginine and glutamine dual-finger catalytic mechanism [19]. Inspired by this study, here we perform biochemical and structural analyses of LepB RabGAP activity. We map LepB GAP domain to residues 313-618 and identify Arg444 as the candidate arginine finger [19].

In vitro GAP assay showed that bacterially purified LepB (904-1 294) and LepB (811-1 294) fragments harbored no activities of catalyzing GTP hydrolysis in Rab1 (Figure 1A). In contrast, LepB (1-903) displayed a robust GAP activity comparable to that of full-length LepB. Further systematic truncation analyses identified residues 313-618 of LepB (LepB313-618) as the minimal domain capable of catalyzing GTP hydrolysis in Rab1 (Figure 1A). Notably, the activity of LepB313-618 was even higher than that of full-length LepB or LepB (1-903), suggesting a possible autoinhibitory mechanism.

Both eukaryotic TBC-domain RabGAPs [20] and the VirA/EspG family of bacterial RabGAPs [19] feature an arginine and glutamine dual catalytic finger. To identify the catalytic finger residues, nearly all the arginine (Arg395, 430, 431, 444, 573, 583, 596, and 608) and glutamine residues (Glu373, 453, 468, 486, 488, 506, 511, and 533) in LepB313-618 were individually mutated into lysine and alanine, respectively. Among the 16 mutant proteins, only LepB313-618 R444K was found to be deficient in promoting GTP hydrolysis (Figure 1B). The GAP protein can form a stable complex with the GTPase substrate in the presence of aluminium fluoride (AlF₃) and GDP, which approximates the GTP hydrolysis transition state intermediate [19-22]. As expected, purified LepB313-618 co-eluted with Rab1-GDP from a gel filtration column in the presence of AlF₃ and omission of AlF₃ prevented the LepB313-618-Rab1-GDP complex formation (Figure 1C). In contrast, LepB313-618 R444K and Rab1-GDP eluted separately from the gel filtration column even in the presence of AlF₃ and their elution profile corresponded to that of the two proteins loaded alone. These data suggest that Arg444 is the catalytic arginine finger for LepB GAP activity.

The GAP activity of LepB is highly robust and specific to Rab1

We then profiled the activity of LepB313-618 towards a panel of 30 mammalian Rabs and determined the corresponding kinetic parameters, as we and others have previously performed with Gyp1p and the VirA/EspG fami-
Compared with Gyp1p and the VirA/EspG family, LepB showed a much narrower substrate spectrum and acceleration of GTP hydrolysis was only observed with Rab1, Rab9, Rab13 and Rab35 (Figure 2).
The activity of LepB313-618 on Rab1 (k_{cat}/K_{m}, 7.24 \times 10^5 M^{-1}S^{-1}) was higher than those on other three Rabs (k_{cat}/K_{m}, 5.40 ~ 7.60 \times 10^3 M^{-1}S^{-1}) by more than two orders of magnitude (Figure 2). The catalytic efficiency of LepB313-618 on Rab1 was also higher than those of Gyp1p on Rab1 and Rab33 (≤1 \times 10^5 M^{-1}S^{-1}) [20] as well as that of VirA/EspG on Rab1 (2 ~ 3 \times 10^5 M^{-1}S^{-1}) [19]. These enzymatic analyses establish LepB as a highly selective and most potent GAP protein for Rab1, a feature consistent with an essential role of LepB in mediating rapid inactivation and removal of Rab1 from the LCV during infection [18].

Crystal structure of LepB RabGAP domain

LepB313-618 shows no sequence homology to all known GAP proteins including TBC-domain RabGAPs. To reveal the structural mechanism of LepB RabGAP activity, we crystallized LepB (residues 1-618) that covers the GAP domain (LepB313-618) and solved the 3.2-Å structure by single wavelength anomalous diffraction (SAD) method (Table 1). The structure contains two LepB molecules packing against each other in an asymmetric unit. The interaction between the two molecules likely resulted from crystal packing as purified LepB was exclusively a monomer in solution according to its gel filtration chromatography behavior. No meaningful structural difference was observed between the two molecules in the same asymmetric unit, and chain A was subjected to further analysis. Compared with the N-terminal region, the GAP domain has a higher-quality electron density map and contains residues 326-391, 400-519 and 530-617.

Table 1 Data collection and refinement statistics

| Crystal statistics | LepB313-618 SeMet | LepBdrancourti-Rab1 |
|--------------------|-------------------|-------------------|
| **Data collection statistics** | | |
| Space group | C222, | P6122 |
| Wavelength (Å) | 0.9792 | 0.9792 |
| a, b, c, (Å) α, β, γ, (°) | 101.53, 159.42, 181.15, 90, 90, 90 | 95.56, 95.56, 197.72, 90, 90, 120 |
| Resolution range (Å)* | 20-3.15 (3.20-3.15) | 20-2.80 (2.85-2.80) |
| No. of unique reflections | 25525 (1163) | 14058 (681) |
| Completeness (%) | 99.5 (94.7) | 100 (99) |
| Redundancy | 13.5 (8.7) | 10.5 (10.7) |
| I/σI | 28.87 (2.00) | 17.56 (3.45) |
| Rmerge (%) | 12.8 (92.6) | 15.8 (77.2) |
| **Refinement statistics** | | |
| Rwork/Rfree (%)† | 26.40/28.30 | 22.91/26.91 |
| No. of protein atoms | 6894 | 3625 |
| No. of waters | 5 | 4 |
| RMSD bond lengths (Å) | 0.010 | 0.016 |
| RMSD bond angles (°) | 1.367 | 1.419 |
| **Ramachandran plot statistics** | | |
| Most favored regions (%) | 92.60 | 95.94 |
| Additional allowed regions (%) | 6.70 | 4.06 |
| Outlier region (%) | 0.7 | |

*The data for the highest resolution shell are shown in parentheses.
†R_{free} is calculated by omitting 5% of the total number of reflections in model refinement.
Other residues including 317-325, 392-399 and 520-529 are missing and likely in flexible loop regions.

LepB GAP domain adopts a wide “V”-shape architecture, consisting of 10 α-helices and no β-strands (Figure 3A). The more elongated left arm of the “V” shape is mainly formed by a four-helix bundle (α2, α3, α5 and α6) packed by α1 and α4 on opposite sides. α1 at the top end of the left arm connects the GAP domain to the N-terminal domain (not shown in the picture) through a linking loop; α4 is situated in the valley bottom. The right arm is made up of four helices (α7-α10) organized into a more globular shape. α7 and α8, projecting perpendicular to the plane of the V shape, is located closer to the valley bottom; α9 and α10, parallel to each other, cross α7 and pack against α7/α8 from outside. Notably, the overall shape and structural architecture of LepB GAP domain shows no similarity to that of eukaryotic TBC domain (illustrated using Gyp1p TBC domain) [20] (Figure 3A and 3B). LepB GAP domain is also structurally distinct from the VirA/EspG family of bacterial RabGAPs that adopt an α/β fold despite that they are also of prokaryotic origin and highly specific for Rab1. The structural divergence of LepB and the VirA/EspG family from eukaryotic TBC GAPs shall bring insights into the evolution of bacterial virulence activity.

The candidate arginine finger residue Arg444 is located in the loop between α3 and α4 on the concave surface of the V-shaped valley; its side chain is solvent exposed and projects from the valley bottom upwards to the open space (Figure 3A). Structural superimposition of LepB GAP domain with VirA-Rab1 or Gyp1p-Rab33 complex (see below), which places Arg444 in a position equivalent to that of the catalytic arginine in VirA/Gyp1p and amenable to interacting with GDP and AlF₃, revealed no structural clashes but instead a well accommodation of Rab1/33 in the open space of the V-shaped valley. These structural analyses lend further support to Arg444 as the canonical arginine finger for LepB GAP activity. It is also worth noting that LepB Arg444 is situated in a loop region rather than supplied by a helix as observed with those in Gyp1p and VirA/EspG (Figure 3A and 3B), highlighting an additional structural difference between

Figure 3 Crystal structure of LepB313-618 and comparison with the VirA/EspG-family bacterial RabGAP and the TBC domain of Gyp1p. (A) Overall structure of LepB313-618 in comparison with those of VirA (PDB ID: 4FMB) and Gyp1p (PDB ID: 2G77). Secondary structure elements are drawn with α-helices as coils, β-strands as flat arrows and loops as lines. LepB313-618, VirA and Gyp1p are colored blue, orange and green, respectively. (B) Superimposition of LepB313-618 structure onto those of VirA and Gyp1p using Arg444/Glu449 in LepB313-618, Arg188/Gln280 in VirA and Arg343/Gln378 in Gyp1p as the references. Shown on the right is a close-up view with marked reference residues in stick models. α-helices are shown as cylinders colored as that in A. (C) Structural comparison of Arg444/Glu449 in LepB313-618 with Arg343/Gln378 in Gyp1p. GDP and AlF₃ in the Gyp1p-Rab1 complex structure are shown as yellow sticks. Mg²⁺ and the nucleophilic water are depicted as magenta and red sphere, respectively. Polar interactions between GDP-AlF₃ and Arg343/Gln378 in Gyp1p are denoted by black dashed lines.
LepB and known RabGAPs.

Glu449, occupying the structural position of TBC GAP glutamine finger in apo-LepB, is critical for LepB GAP activity

Different from GAPs for Rho and Ras families, both eukaryotic TBC GAPs and the VirA/EspG family of bacterial GAPs feature a catalytic glutamine finger in addition to the canonical arginine finger [19, 20]. The glutamine side chain is at a distance of ~6 Å from the guanidine group of the arginine finger; its carbonyl amide mediates bipartite hydrogen-bonding interactions with an equatorial fluoride ion and the axial water in the AlF₆⁻ complex. In catalysis, the glutamine carbonyl acts as a general base to deprotonate and orientate the water molecule for nucleophilically attacking on the γ-phosphate in GTP. Consistent with the fact that none of the glutamines in LepB³₁₃₋₆₁₈ are essential for its GAP activity (Figure 1B), no glutamine residues could be found within the 8-Å distance from Arg444, confirming that LepB does not employ a TBC-like catalytic glutamine finger. Notably, we found a glutamate residue (Glu449) whose side-chain carboxyl carbon is ~4 Å away from the carbon atom of the guanidine group of Arg444 (Figure 3B). When LepB GAP domain was structurally superimposed onto Gyp1p TBC domain and VirA using Arg444/Glu449 in LepB and the catalytic arginine/glutamine fingers in Gyp1p/ VirA as the reference, presentation and stereochemistry of Glu449 side chain appeared to be analogous to those of the catalytic glutamine in TBC and TBC-like GAPs despite that the carboxyl group of Glu449 is slightly closer to AlF₆⁻ (Figure 3C). In fact, mutation of Glu449 to alanine, glutamine or aspartate either completely abolished or largely attenuated the activity of LepB³₁₃₋₆₁₈ in

![Figure 4](https://example.com/figure4.png)

**Figure 4** Glu449 is required for the RabGAP activity of LepB as well as AlF₆⁻-mediated LepB-Rab1-GDP complex formation. (A) Effects of Glu449 mutations on the RabGAP activity of LepB. Time courses of GTP hydrolysis for Rab1 (40 µM) catalyzed by LepB³₁₃₋₆₁₈ or its E449A, E449D and E449Q mutants are shown as color indicates. (B) Summary of the kinetic parameters of LepB Glu449 and VirA Gln280 mutants. The Michaelis-Menten kinetic parameters (kcat/Km) determined by using the Lineweaver-Burk plot for LepB³₁₃₋₆₁₈ and VirA or their indicated mutants are listed. Data shown are mean values ± standard deviation (error bar) from two independent experiments. (C) Effects of Glu449 mutations on AlF₆⁻-mediated LepB-Rab1-GDP complex formation. The assay was performed similarly as that in Figure 1C. Shown on the left are the gel filtration chromatograms colored accordingly. Peak 1, 2, and 3 correspond to LepB³₁₃₋₆₁₈-Rab1-GDP-AlF₆⁻ complex, free LepB³₁₃₋₆₁₈ and free Rab1, respectively. Elution fractions in the indicated range were analyzed by SDS-PAGE and Coomassie blue-stained gels are shown on the right.

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promoting GTP hydrolysis in Rab1 (Figure 4A and 4B). Consistently, both LepB313-618-E449Q and E449D mutants were deficient in forming an AlF₃-dependent complex with Rab1-GDP (Figure 4C). Prompted by this observation, we also examined the TBC-like glutamine finger in VirA and found that substitution of Q280 by either a glutamate or an asparagine could also largely abolish VirA GAP activity by more than 90% (Figure 4B). These analyses establish a critical role of Glu449 for the GAP activity of LepB effector.

Crystal structure of LepB<sub>drancourtii</sub> GAP domain-Rab1-GDP-AlF₃ complex

To understand the exact mechanism of LepB recognizing Rab1 and catalyzing GTP hydrolysis, we efforted to determine the structure of LepB GAP domain in complex with Rab1, but unfortunately we were unable to obtain diffractable crystals. We then turned to LepB homologues from other Legionella species. LDG_7216 from <i>L. drancourtii</i> strain LLAP12, a strictly intracellular amoebal pathogen [23], contains 1,189 amino acids and shares about 55% overall sequence similarity to LepB from <i>L. pneumophila</i>. Residues 316-620 at the N-terminal region of LDG_7216 harbored 54% sequence homology to the GAP domain of <i>L. pneumophila</i> LepB. For convenience, a recombinant fragment containing residues 316-620 of LDG_7216 was referred as LepB<sub>drancourtii</sub> GAP domain hereafter. Biochemical assays confirmed that LepB<sub>drancourtii</sub> GAP domain (residues 316-620) harbored a robust GAP activity using Rab1 as the substrate, comparable to that of <i>L. pneumophila</i> LepB (Figure 5A). LepB<sub>drancourtii</sub> GAP domain could also form an AlF₃-dependent complex with Rab1-GDP (Figure 5B). Also similarly to that observed with <i>L. pneumophila</i> LepB, the RabGAP activity and formation of the transition state-approximating ternary complex required Arg447 and Glu452 in LepB<sub>drancourtii</sub> GAP domain (equivalent to Arg444 and Glu449 in <i>L. pneumophila</i> LepB) (Figure 5A and 5B).

We succeeded in obtaining diffraction-quality crystals of LepB<sub>drancourtii</sub> GAP domain in complex with Rab1-GDP and AlF₃ and determined 2.8-Å structure of the ternary complex (Figure 6A and Table 1). The structure contains one LepB<sub>drancourtii</sub> and one Rab1 in an asymmetric unit. Rab1 is cradled by the concave surface of LepB<sub>drancourtii</sub> GAP domain along the V-shaped valley. Consistent with the fact that LepB GAP domain has a completely different fold from both the eukaryotic TBC domain and the VirA/EspG family of bacterial RabGAPs, the Rab1-bind-
ing mode of LepB\textsuperscript{drancourtii} is unique and shows no similarity to those of the two known classes of RabGAPs (Figure 6A). Rab1-bound LepB\textsuperscript{drancourtii} GAP domain is structurally similar to apo-LepB GAP domain with RMSD value of 2.642 for 233 selected Cα atoms (Figure 6B). The overall structure of Rab1 is also nearly identical to those of VirA-bound Rab1 and Gyp1p-bound Rab33 (Figure 6C, right). The interaction of LepB\textsuperscript{drancourtii} with Rab1 engages residues from helices α3, α4 and more importantly the connecting loop in LepB\textsuperscript{drancourtii} GAP domain. The interaction involves a network of extensive hydrogen bonds and hydrophobic interactions (Figure 6C). Specifically, the backbone carbonyl oxygen of Met445 in LepB\textsuperscript{drancourtii} forms a hydrogen bond with the backbone amide of Rab1 Ser42. Rab1 Ser39 has two hydrogen bonds with LepB\textsuperscript{drancourtii} GAP domain; its hydroxyl group donates one hydrogen to the side-chain carbonyl of LepB\textsuperscript{drancourtii} Gln603 and its backbone carbonyl oxygen accepts one hydrogen from the backbone amide of LepB\textsuperscript{drancourtii} Gly601. Rab1 Ile41 contributes to hydrophobic
Figure 7 Structural basis for a RasGAP-like catalytic mechanism of LepB. (A) Sigma-A weighted 2mFo-DFc map contoured at 1.5σ for the modeled GDP, AlF$_3$, Mg$^{2+}$ and putative nucleophilic water in the LepB$^{drancourt}$ GAP domain-Rab1 complex. (B) Structural comparison of GDP, AlF$_3$, and the catalytic arginine and glutamine fingers among LepB$^{drancourt}$ GAP domain-Rab1 complex (left), RasGAP-Ras complex (middle, PDB ID: 1WQ1) and Gyp1p-Rab1 complex (right, PDB ID: 2G77). The catalytic arginine and glutamine and other key residues are shown in sticks and colored the same as their residing chains. Mg$^{2+}$ and the nucleophilic water are shown as green and red spheres, respectively. Polar interactions are indicated by black dashed lines. (C) Conformational changes of Gln70 in Rab1 and Glu449/452 in LepB and comparison with related structural elements in the RasGAP-Ras complex. Shown on the left is structural alignment of key residues in the catalytic sites of VirA-bound Rab1 and Gyp1p-bound Rab33 with those in the LepB$^{drancourt}$-Rab1 complex. The right panel shows the alignment of free Ras with RasGAP-Ras complex. Alignments were performed using Cα chains of Rabs or Ras as the reference. In the left panel, apo-LepB GAP domain was superimposed onto the LepB$^{drancourt}$-Rab1 complex using Cα atoms of LepB Arg444 (Arg447 in LepB$^{drancourt}$) and E449 (Glu452 in LepB$^{drancourt}$) as the reference. Free Ras, RasGAP-bound Ras and RasGAP are colored in magenta, orange and teal blue, respectively. Coloring of LepB$^{drancourt}$-Rab1 complex and LepB GAP domain alone follows that in Figure 6A and Figure 3A, respectively. Rotational movement of the cis-glutamine in Rab1 and Ras and the critical Glu452 in LepB$^{drancourt}$ are denoted by black curved arrows. (D) Effects of mutation of Gln70 in Rab1 on LepB- and VirA-catalyzed GTP hydrolysis. The indicated concentrations of purified VirA and LepB$_{313-618}$ were subjected to in vitro GAP assays using Rab1 Q70E (Left) and Q70N (Right) as the substrate. (E) In vitro GAP assay of LepB$_{313-618}$ using H-Ras as the substrate.
the distinct mode of Rab recognition by LepB\textsuperscript{drancourtii} GAP domain.

\textit{LepB employs a RasGAP-like catalytic mechanism}

In the refined map of the LepB\textsuperscript{drancourtii} GAP domain-Rab1 complex, clear electron density is observed for GDP, Mg\textsuperscript{2+}, a putative nucleophilic water and AlF\textsubscript{3} (Figure 7A). AlF\textsubscript{3} is coordinated by an axial oxygen from GDP and occupies the site of γ-phosphate in the GTP-bound state [24]. The conformational arrangement and chemical geometry of GDP, Mg\textsuperscript{2+} and AlF\textsubscript{3} are similar to those observed in other GAP-GTPase complex including the Gyp1p-Rab33 and VirA-Rab1 complexes (Figure 7B), confirming a GTP-hydrolysis transition state-approximating intermediate. As expected, Arg447 in LepB\textsuperscript{drancourtii} mediates bidentate polar interactions with an oxygen from β-phosphate of GDP and an equatorial fluoride ion. This structural feature also resembles those described for other GAP-GTPase complexes [19, 20, 25], reassuring Arg447 being the arginine finger in catalyzing GTP hydrolysis.

To our surprise, the side chain of Glu452 in LepB\textsuperscript{drancourtii} rotates away from its original position in the apo-LepB structure by ~120 °C (Figure 7C), which no longer occupies the structural position equivalent to the glutamine finger in TBC and TBC-like RabGAPs. Most notably, Glu452 in the Rab1-GDP-AlF\textsubscript{3} complex now bears a strong hydrogen-bond or ionic interaction with Gln70 in Rab1, and this interaction renders a drastic conformational change of the side chain of Gln70 (Figure 7C). Such structural rearrangement of the glutamine residue, though more dramatic, echoes what has been observed in the RasGAP-Ras complex, which is induced by interaction of Gln61 in Ras with the backbone carbonyl oxygen of RasGAP arginine finger (Arg789) [26] (Figure 7C). In fact, Gln61 in Ras serves as the catalytic cis-glutamine to facilitate GTP hydrolysis. Similar to Gln61 in the RasGAP-Ras complex, Gln70 in the LepB\textsuperscript{drancourtii} GAP domain-Rab1 complex also mediates bipartite polar interactions with equatorial fluoride ion and the putative nucleophilic water (Figure 7B and 7C). The importance of Gln70 was supported by results of the in vitro GAP assay, in which Q70E or Q70N mutants of Rab1 did not significantly affect the activity of VirA but were completely resistant to LepB-catalyzed GTP hydrolysis (Figure 7D). These observations clearly distinguish LepB RabGAP from eukaryotic TBC GAPs and the VirA/ EspG family of TBC-like RabGAPs that both supply the catalytic glutamine finger in trans (Figure 7B). Thus, the structural and chemical similarities between LepB\textsuperscript{drancourtii} Arg447/Rab1 Gln70 and RasGAP Arg789/Ras Gln61 establish that \textit{Legionella} LepB effector does not harbor the TBC RabGAP-like dual catalytic fingers but instead employs a RasGAP-like catalytic mechanism in promoting GTP hydrolysis in Rab1. RasGAP has extensive specific interactions with Ras for productive substrate binding [26]. Consistently, LepB did not catalyze GTP hydrolysis when Ras was used a substrate (Figure 7E).

The drastic side-chain movement of Rab1 Gln70 towards GDP-AlF\textsubscript{3} and therefore serving as the catalytic cis-glutamine is unexpected. Given that Glu449/Glu452 in LepB plays an absolutely essential role for LepB to form an AlF\textsubscript{3}-dependent ternary complex with Rab1 and catalyze GTP hydrolysis (Figures 4 and 5), the strong interaction observed between Gln70 in Rab1 and Glu452 in LepB\textsuperscript{drancourtii} is likely a determining structural factor for Gln70 being able to play the catalytic role. Gln70 in Rab1, compared with Gln61 in Ras, is more towards the GTPase inside and relatively difficult to move out. These analyses together help to explain why TBC or TBC-like RabGAPs have to supply the glutamine finger as they do not harbor a glutamate equivalent to Glu449 in LepB.

\textbf{Discussion}

In this study, we map the RabGAP domain of the \textit{Legionella} Dot/Icm type IV effector LepB and determine crystal structures of the GAP domain alone and in complex with Rab1-GDP-AlF\textsubscript{3}. LepB represents a novel RabGAP fold and employs a distinct GTPase-recognition mode. Combining extensive biochemical analyses together with the structural characterization, we identify the canonical arginine finger residue responsible for LepB to catalyze GTP hydrolysis. As a RabGAP, the VirA/EspG family of bacterial effectors shares the arginine and glutamine dual catalytic fingers, but no primary sequence and three-dimensional structural similarities with eukaryotic TBC-domain RabGAPs [19]. Notably, the overall structure of LepB GAP domain is also clearly distinct from those of TBC domain and the VirA/EspG family. The three different classes of RabGAPs all employ a common catalytic arginine finger, representing a striking example of convergent evolution of bacterial virulence activity. The sequence and structural divergence of the three classes of RabGAPs also suggests a possible existence of other structural classes of RabGAPs in eukaryotes that have escaped from identification by conventional sequence and domain analyses. \textit{S. flexneri} is free living after gaining access to host cytosol and VirA inactivation of Rab1 contributes to suppression of autophagy-mediated host defense; GTP hydrolysis and inactivation of Rab1 by EspG and the consequent disruption of the general secretory pathway is important for inhibition of cytokine secretion by EPEC that resides...
extracellularly; intracellular *L. pneumophila* relies on LepB-mediated inactivation of Rab1 to achieve sophisticated manipulation of Rab1 cycle and decorate LCV into an ER-like structure. The three pathogens have completely different pathogenesis and life cycle, but all have evolved a virulence activity of catalyzing GTP hydrolysis in Rab1 for their own purposes. These observations are insightful in deepening and broadening our understanding of hijacking Rab function by bacterial pathogens as well as the associated diverse bacterial virulence mechanism. Therefore, future studies will likely identify more bacterial virulence factors that are RabGAP proteins either similar to or distinct from the above three classes.

Our analyses also identify a glutamate in the catalytic site of LepB that is indispensable for its RabGAP activity and formation of the AlF₃-dependent complex with Rab1-GDP. In apo-LepB, this glutamate occupies the structural position of the catalytic glutamine finger in eukaryotic TBC GAPs and the VirA/EspG family of TBC-like bacterial RabGAPs. Interestingly, this residue, upon LepB binding to GTPase substrate, moves away from contacting AlF₃ and bears a strong ionic interaction with Gln70 in Rab1. The interaction is accompanied by a drastic side-chain flipping of Rab1 Gln70 towards GDP-AlF₃. Such conformational changes provide the structural basis for Gln70 acting as the catalytic *cis*-glutamine in promoting GTP hydrolysis. The catalytic mechanism of LepB indeed mirrors that of RasGAP [26] and also represents the first example that Rabs can supply a catalytic glutamine finger in GAP-mediated GTP hydrolysis [20]. This unexpected finding breaks the boundary of GAP classification and highlights an extraordinary catalytic diversity of GAP proteins for small GTPases.

While this manuscript was being prepared for publication, Goody and their colleagues also reported the structure of *L. pneumophila* LepB in complex with Rab1 [27]. The Goody structure is overall highly similar to our structure of LepB<sup>drancourtii</sup> -Rab1 complex. However, beryllium fluoride (rather than aluminum fluoride) is used as the γ-phosphate surrogate in their study, and therefore the critical nucleophilic water is missing in the catalytic center and direct evidences for Gln70 acting as the *cis*-glutamine are not provided. In fact, the beryllium fluoride complex structure more mimics the ground state of GAP-catalyzed GTP hydrolysis while our structure, obtained by using aluminum fluoride, approximates the subsequent transition-state intermediate.

**Materials and Methods**

*Plasmids and reagents*

DNA for LepB was amplified from *L. pneumophila* genomic DNA and cloned into the pQE-80L vector (QIAGEN) for recombinant expression in *E. coli*. DNA encoding the GAP domain of Lep<sup>drancourtii</sup> (residues 316-620 of ZP_09620807 from *Legionella dran-courtii* LLAP12 strain) was artificially synthesized. Human Rab1a (residues 1-177) were PCR cloned into pET28a vector with an N-terminal SUMO fusion. Truncation and deletion mutants were constructed by standard PCR cloning strategy. All point mutations were generated by QuickChange™ Site-Directed Mutagenesis Kit (Stratagene) and the plasmids were verified by DNA sequencing. Bacterial expression constructs for 30 Rabs were kindly provided by Dr David G Lambright (University of Massachusetts Medical School, USA). All chemicals were Sigma-Aldrich products.

**Recombinant protein expression and purification**

*E. coli* BL21 (DE3) strain (Novagen) was used as the host for expression of recombinant proteins. Protein expression was induced for 16 h at 16 °C or 22 °C with 1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) after OD₆₀₀ reached 0.6–0.8. N-terminal 6× His-tagged LepB and its mutants in the pQE-80L vector were purified by using Ni-NTA agarose beads (QIAGEN) followed by Hitrap Q HP ion exchange chromatography (GE Healthcare). Recombinant proteins were kept in a buffer containing 25 mM HEPES (pH 7.4) and 150 mM NaCl. LepB<sup>drancourtii</sup> GAP domain and Rab1 proteins were also overexpressed in BL21 (DE3) followed by Ni-affinity purification. The SUMO-fusion parts were removed by Ulp1 protease digestion and the proteins were further purified by Hitrap Q ion exchange column. Protein concentrations were determined spectrophotometrically using the theoretical molar extinction coefficients at 280 nm and protein purity was examined by Coomassie blue staining of SDS-PAGE gels. For crystallization, LepB (residues 1-618) were also cloned into the pQE-80L vector for bacterial expression, and proteins were further purified by Mono Q ion exchange and Sephadex 75 gel filtration chromatography.

**GAP assay**

The GAP assay using the EnzChek Phosphate Assay Kit (Invitrogen) and determination of kinetics were carried out by strictly following the previously described procedure [19]. The absorbance at 360 nm was monitored on a UV-2450 spectrometer (Shimadzu).

**Crystallization and data collection**

The LepB<sub>R. leg</sub> crystals were grown by the hanging-drop vapor diffusion method at 22 °C by mixing equal volumes (1 µl each) of the protein and the reservoir solution. Diffraction-quality crystals appeared at the condition of 15 mg/ml of LepB (residues 1-618) with a reservoir solution containing 0.1 M citrate (pH 6.5) and 20% MPEG 2000. For cryo-protection, crystals were transferred into the cryoprotectant solution containing 0.1 M citrate (pH 6.5), 25% MPEG 2000 and 5% glycerol, soaked for several seconds, and then flash-cooled in liquid nitrogen. X-ray diffraction data were collected at Spring-8 beamline BL41XU at the wavelength of 0.9792 Å. To crystallize the LepB<sup>drancourtii</sup> GAP domain-Rab1 complex, LepB<sup>drancourtii</sup> GAP domain was mixed with GDP-loaded Rab1 at the molar ratio of 1:1:1 in a buffer containing 20 mM Tris-HCl (pH 8.0), 150 mM NaCl, 20 mM NaF, 2 mM AlCl₃, 5 mM MgCl₂ and 2 mM DTT, and the mixture was incubated on ice overnight to obtain the LepB<sup>drancourtii</sup> GAP domain-Rab1-GDP-AlF₃ complex. Crystallization of the complex was performed by hanging-drop va-
por diffusion method using the concentrated mixture at 24 mg/ml. The crystals were grown at 20 °C for one week in 0.2 M ammonium acetate, 0.1 M Tris-HCl pH 8.5, and 25% v/v polyethylene glycol 3350. Crystals were coated by cryoprotectant oil containing 50% v/v Parabar 10312 (Hampton Research) and 50% v/v paraffin oil, and then subjected to flash freezing in liquid nitrogen. Data were collected at Shanghai Synchrotron Radiation Facility Beamline 17U.

**Structural determination and refinement**

Raw diffraction data of LepB_{trans} were processed with the HKL package [28]. The phase was solved using selenium single wavelength anomalous dispersion method [29] in the program SHARP (http://www.globalphasing.com/sharp/). Initial experimental map was density-modified in DM [30]. The model was automatically built in Phenix.autobuild [31] and Buccaneer [32], and manually adjusted in COOT [33]. The structure was refined by using the program phenix.refine in PHENIX. Diffraction data of the LepB_{trans} GAP domain-Rab1 complex were processed with HKL2000 and scaled with Scalepack [28]. The complex structure was solved by molecular placement using VirA-bound Rab1 and sculptured LepB-GAP domain [34] as searching models in the program Phaser [35]. The complete model was manually built in COOT [33] and refined in reflmac [36]. All the structural figures were generated using PyMol (www.pymol.org). The atomic coordinates of L. pneumophila LepB (residues 1-618) and LepB_{trans} GAP domain-Rab1 complex have been deposited in the Protein Data Bank with the accession codes 4JVS and 4JW1, respectively.

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