

RESEARCH ARTICLE



transport

Periplasmic binding proteins Bug69 and Bug27 from Bordetella pertussis are in vitro high-affinity quinolinate binders with a potential role in NAD biosynthesis

Leonardo Sorci¹ (b), Gabriele Minazzato², Adolfo Amici³, Francesca Mazzola³ (b) and Nadia Raffaelli²

1 Division of Bioinformatics and Biochemistry, Department of Science and Engineering of Matter, Environment and Urban Planning, Polytechnic University of Marche, Ancona, Italy

2 Department of Agricultural, Food and Environmental Sciences, Polytechnic University of Marche, Ancona, Italy

3 Department of Clinical Sciences, Polytechnic University of Marche, Ancona, Italy

Keywords

Bordetella Uptake Gene; molecular docking; NAD biosynthesis; periplasmic binding protein; guinolinate phosphoribosyltransferase; sequence similarity network

Correspondence

L. Sorci, Division of Bioinformatics and Biochemistry, Department of Science and Engineering of Matter, Environment and Urban Planning, Polytechnic University of Marche, Via Ranieri 67 (Polo Montedago), 60131 Ancona Italy Tel: +39 071 2204678 E-mail: I.sorci@staff.univpm.it

Bordetella's genome contains a large family of periplasmic binding proteins (PBPs) known as Bugs, whose functions are mainly unassigned. Two members, Bug27 and Bug69, have previously been considered potential candidates for the uptake of small pyridine precursors, possibly linked to NAD biosynthesis. Here, we show an in vitro affinity of Bug27 and Bug69 for quinolinate in the submicromolar range, with a marked preference over other NAD precursors. A combined sequence similarity network and genome context analysis identifies a cluster of Bug69/27 homologs that are genomically associated with the NAD transcriptional regulator NadQ and the enzyme quinolinate phosphoribosyltransferase (QaPRT, gene *nadC*), suggesting a functional linkage to NAD metabolism. Integrating molecular docking and structure-based multiple alignments confirms that quinolinate is the preferred ligand for Bug27 and Bug69.

Leonardo Sorci and Gabriele Minazzato shared first authorship equally.

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Periplasmic binding proteins (PBPs), also known as Beyond their canonical role in nutrient uptake, PBPs substrate binding proteins or extracytoplasmatic solute exhibit versatility as initial receptors, participating in receptors, bind and deliver specific target molecules, signal transduction and channel gating [1,2]. Currently, encompassing ions, vitamins, amino acids, carbohy-PBP-dependent multicomponent three drates, and peptides to a membrane transporter, systems have been documented: (a) the ATP-binding which, in turn, imports the solute into the cytoplasm. cassette transporters (ABCs), (b) the tripartite

Abbreviations

Bug, Bordetella Uptake Gene; Ca, citric acid; DHAP, dihydroxyacetone phosphate; Na, nicotinic acid; NAD, nicotinamide adenine dinucleotide; Nam, nicotinamide; NaMN, nicotinic acid mononucleotide; NaMNAT, nicotinic acid mononucleotide adenylyltransferase; NMN, nicotinamide mononucleotide; Pa, picolinic acid; PBP, periplasmic binding protein; PRPP, 5-phosphoribosyl-1-pyrophosphate; Qa, quinolinic acid; QaPRT, quinolinic acid phosphoribosyltransferase; R5P, ribose-5-phosphate; TTT, tripartite tricarboxylate transporter.

ATP-independent periplasmic transporters (TRAPs), and (c) the tripartite tricarboxylate transporters (TTTs). ABC systems, categorized as primary transporters, utilize ATP hydrolysis to energize the transport against concentration gradients [3]. In contrast, TRAP and TTT systems facilitate transport by coupling it to an electrochemical gradient, thus functioning as conventional secondary transport systems [4]. An ABC transport system typically comprises a substrate binding protein, a transmembrane (TM) protein, and a nucleotide-binding protein. Both the membrane and nucleotide-binding components are usually organized as homo- or heterodimers. Such a dimeric structure is crucial for the proper functioning of ABC transporters, enabling coordinated ATP hydrolysis and substrate translocation [5]. Members of the TTT and TRAP families comprise a conserved 12-TM protein (prototyped by TctA in TTT systems) acting as a symport protein, a 4-TM protein (TctB) with unknown function, and a PBP protein (TctC) playing the highaffinity substrate binding role. Recently, the cryo-EM structure of the sialic acid TRAP transporter SiaQM and the soluble substrate binding "P-subunit" (SiaP) from Photobacterium profundum have been reported, suggesting an 'elevator-with-an-operator' transport mechanism where SiaP plays the role of the 'operator' of the elevator [6]. TTT transport system is the most recent and least characterized family of transporters using a PBP. Intriguingly, certain bacteria exhibit a redundancy of TctC homologs without apparent membrane counterparts. With 434 tctT genes, the genome of a-proteobacterium Rhodoplanes sp. Z2-YC6860 represents the largest over-representation of the TTT protein family reported to date, a gene expansion likely spurred by multiple environmental adaptations [7]. In the genome of pathogenic Bordetella pertussis, an over-representation of TctC 'orphan' paralogsreferred to as Bordetella Uptake Genes (Bug)-has also been observed [8].

Here, we elucidate the role of two orphan TctCs from *Bordetella pertussis*, namely Bug27 and Bug69, as *in vitro* quinolinic acid-binding proteins. In most bacteria, quinolinate (Qa) is an intermediate of the *de novo* NAD biosynthesis pathway implemented by genes *nadB*, *nadA*, and *nadC* [9,10]. This pathway leads to the formation of nicotinate mononucleotide (NaMN) intermediate from aspartate and dihydroxyacetone phosphate (DHAP). Once formed, NaMN is converted to NAD by two consecutive reactions [11]. In contrast, most bordetellae lack a fully functional *de novo* NAD biosynthesis pathway, rendering them auxotrophic for the NAD salvage pathway precursors nicotinic acid (Na) and nicotinamide (Nm) (see Ref. [12] and Fig. 1). To meet this nutritional in vitro growth requirement, Bordetella spp. are conventionally cultured in a standard Stainer Scholte (SS) defined medium supplemented with 30 µM Na (or Nm) [13]. However, Bordetellae, akin to Streptococcus pyogenes and other pathogenic bacteria [14,15], retain a functional terminal component of the de novo NAD pathway, the Qa phosphoribosyltransferase (QaPRT) enzyme encoded by gene nadC. This enzyme converts quinolinic acid and phosphoribosyl pyrophosphate (PRPP) to NaMN, thus allowing bacterial growth on Qa as the sole NAD precursor [16]. Of significance is the observation of Brickman et al. regarding the presence of the gene bug69 downstream of nadC in Bordetella species. They hypothesized a direct role of this periplasmic binding protein in the transport of Qa. However, the genetic inactivation of bug69 in Bordetella bronchiseptica did not compromise the bacterium's ability to grow on Qa. This finding led the authors to infer that Bug69 is either not implicated in Qa uptake or that other proteins may be capable of assuming such a function [16]. In B. pertussis, a previous study identified Bug27, a paralog of Bug69 sharing 55% sequence identity, as capable of in vitro binding to Na and Nam [17].

The present study demonstrates for the first time that *Bordetella pertussis* Bug69 and Bug27 are *in vitro* high-affinity Qa binders. This experimental result is supported by a comprehensive evolutionary and *in silico* structural analysis. Finally, comparative genomics revealed a potential functional association of these orphan Bugs with a quinolinate salvage pathway for NAD biosynthesis in Bordetellae and other bacterial species.

Experimental procedures

Cloning, expression, and purification of Bug69 and Bug27

Genes were amplified from the genomic DNA of *Bordetella pertussis* Tohama I for cloning into pET28c vector. For *bug27*, the forward and reverse primers were designed as 5'-TAT<u>CCATGG</u>TTGCCACCGGCGACTTCCCG-3' and 5'-GCC<u>AAGCTTGTCCACTTTCAGGCCGATCTGCC3'</u> to introduce *Nco*I and *Hind*III restriction sites, respectively. For *bug69*, primers were 5'-TAT<u>CCATGG</u>TTGCGCCGG CCACCTATCCC-3' and 5'-AAACTCGAGGTCGGCG CAGCCCA-3' carrying *Nco*I and *Xho*I sites, respectively. Constructs were used to transform BL21(DE3) competent *Escherichia coli* cells for protein expression. Cells were grown at 37 °C in Luria Bertani medium supplemented with kanamycin (50 µg·mL⁻¹). After reaching an OD₆₀₀ of 0.7, the cultures were shifted at room temperature for 20 min, and protein expression was induced with 1 mM

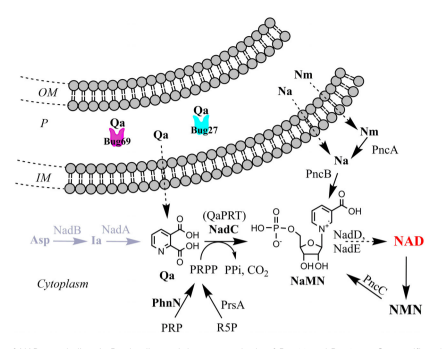


Fig. 1. Reconstruction of NAD metabolism in Bordetellae and the proposed role of Bug69 and Bug27 as Qa-specific solute binding proteins. Pathway reactions are denoted by black arrows. The initial two steps of *de novo* NAD biosynthesis, missing in most Bordetellae, including *Bordetella pertussis*, are depicted in gray. The proposed role of Bug69 and Bug27 as Qa binders is shown. Uptake of pyridines (dotted black arrows) is by unknown mechanisms. Metabolites and enzyme abbreviations (not introduced in the main text). NadD (NaMNAT), nicotinate adenylyltransferase [43]; NadE, NAD synthetase [44]; NMN, nicotinamide mononucleotide; PncC, nicotinamide mononucleotide deamidase [45]; R5P, ribose-5-phosphate; PrsA, PRPP synthetase; Other abbreviations: IM, inner membrane; OM, outer membrane; P, periplasm.

isopropyl b-D-1-thiogalactopyranoside. After 15h of induction, cells were harvested by centrifugation at 5000 g for 8 min at 4 °C. All the following steps were performed at 4 °C. The pellets were resuspended in one-twentieth of the original culture volume of lysis buffer (50 mM sodium phosphate, pH 8.0, 0.3 M NaCl, 5 mM imidazole) freshly supplemented with 1 mM phenylmethylsulfonyl fluoride and protease inhibitor cocktail. Cells were disrupted by passage through a French pressure cell (18 000 psi), and crude extracts were clarified by centrifugation at $20\,000\,g$ for 25 min at 4 °C. Both His-tagged Bug proteins were purified by NiNTA affinity chromatography. The clarified supernatants were loaded onto 1 mL NiNTA columns (Qiagen) equilibrated with the lysis buffer. After washing with 20 mm imidazole in the same buffer, recombinant proteins were eluted with an imidazole concentration of 250 mm. Purification was monitored by SDS/PAGE, according to Laemmli [18]. Protein concentration was determined by Bradford's method [19], and about 9 mg of pure Bug27 and 5 mg of pure Bug69 were obtained starting from 250 mL of cultures.

Fluorescence-based thermal shift binding assay

The assay was adapted from a previously published method [20], and analyses were performed after proteins' denaturation and renaturation steps to remove endogenous ligands, as

described for Bug27 [17]. Briefly, the assay was carried out in a RotorGene 3000 real-time PCR in a final assay volume of 25 μ L. The assay mixture contained 1 μ M of proteins, different concentrations of ligands (dissolved in water), and 2.5× Sypro orange (Invitrogen, Waltham, MA, USA) in 40 mM Tris/HCl, pH 8.0, 250 mM NaCl. The mixtures were pre-heated at 35 °C for 5 min, then heated to 40 °C for 15 s, and finally heated from 40 to 99 °C with a heating rate of 0.1 °C·s⁻¹. The fluorescence intensity was measured with Ex/Em: 470/585 nm. The melting temperature (T_m) was determined by taking the maximum point of the slope-derivative peak.

Intrinsic tryptophan fluorescence quenching assay

The dissociation constants (K_D) of Qa for Bug27 and Bug69 were determined using tryptophan fluorescence titration. The fluorescence quenching was carried out in a cuvette with a path length of 1 cm. The Bug proteins were prepared following the same procedure as for the thermal shift assay, with a final concentration of 0.1 µM. The intrinsic fluorescence spectra of Bug27 and Bug69 were recorded by a fluorescence spectrophotometer (Perkin Elmer LS-45, Waltham, MA, USA) at 20 °C using excitation at 295 nm and emission ranging from 290 to 450 nm. Notably, Qa itself induced negligible changes in fluorescence under the same experimental conditions. Tryptophan fluorescence spectra were collected before and after titration with varying concentrations of Qa. The dissociation constant of Qa to Bug27 and Bug69 was determined by fitting the normalized fluorescence intensity at 328 nm using the following equation [21]:

$$f = \left[(Pt + Lt + K_{\rm D}) - \sqrt{(Pt + Lt + K_{\rm D})^2 - 4PtLt} \right] / 2Pt$$

where f is the fractional change, K_D is the dissociation constant, and Pt and Lt are the total concentration of protein and Qa, respectively.

Statistical analysis

Statistical analyses were performed using PRISM 7 (GraphPad software, San Diego, CA, USA). The thermal shift binding assays and the tryptophan quenching assays were performed in triplicate. The $K_{\rm D}$ values \pm Std. Error are determined using nonlinear regression analysis as reported above.

Bug69-Bug27 sequence similarity network

We used the "Sequence BLAST or option A" within the EFI-Enzyme Similarity Tool (available at https://efi.igb. illinois.edu/efi-est/index.php) to perform all-by-all blast analysis of protein homologs of query Bug69 from Bordetella pertussis Tohama I (Uniprot ID: Q7W019). Applying the default cutoff e-value of 10⁻⁵ for Uniprot BLAST retrieval and SSN edge calculation, we limited the retrieval of homologs to a maximum of one thousand (default option). This dataset contained 928 unique sequences. The full network 10⁻¹²⁰ SSN was generated by applying an *e*-value cutoff of 10^{-120} to the full network. Ninety-four paralogs of the TctC family are present in Bordetella pertussis. With these stringent search parameters, we collected a set of Bug69 homologs, including Bug27, the closest paralog in B. pertussis with a sequence identity of 59%, but excluded all other more distant paralogs. An 85% identity Representative node network was visualized on the Cytoscape platform [22], i.e., all connected sequences that share 85% or more identity are grouped into a single meta node.

In silico binding analysis and other bioinformatics tools

Visualization, molecular docking, 2D and 3D ligand-pocket interaction analysis were performed with tools integrated in Molsoft ICM-PRO 3.9-2c [23]. The X-ray structure of Bug27 (PDB code: 2QPQ) served as the template for docking pyridine derivatives. Prior to docking, the protein was prepared by removing waters, optimizing the hydrogen bonding network, and the orientation of His, Pro, Asn, Gln, and Cys residues. We used the ICM Pocket Finder algorithm [24] to identify the ligand binding pocket of Bug27 (PDB: 2qpq). One of the identified pockets, which partially overlapped with the co-crystallized citrate ligand, was used as the template for docking simulations. The co-crystallized citrate ligand was removed before docking. In independent docking tests, confidence in the docking pose was achieved by progressively increasing the simulation length (using the 'Effort' parameter, ranging from 1 to 10 in ICM-PRO) until convergent results were obtained. Structure-based multiple sequence alignments were generated with PROMALS3D [25] and rendered with Espript 3.0 [26].

Results

Sequence similarity networks of *B. pertussis* Bug69 and Bug27

To investigate the sequence-function relationship of the two protein families typified by *B. pertussis* Bug69 and Bug27, we adopted an integrated strategy leveraging sequence similarity networks (SSN) and genome context analysis tools available at https://efi.igb. illinois.edu/ [27-29]. Sequence similarity networks visually represent pairwise sequence relationships within clusters of homologous proteins [30]. In these networks, each node represents a protein, and edges connect pairs of nodes with pairwise relationships better than a user-defined threshold. These pairwise relationships correspond to BLAST alignment associated with an E-value. When properly selected cutoff values are applied, proteins with identical functions tend to segregate into distinct clusters. Using Cytoscape, we examined the SSN for 928 top homologs of B. pertussis Bug69 (Option A in the EFI-EST tool) at an e-value of 10^{-120} , which corresponds to a median sequence identity of approximately 67%, yielding a total of 156 discrete clusters (Fig. 2A). Bug27 is a closely related paralog of Bug69, with a 59% identity. Hence, this approach also allowed for a comprehensive retrieval of Bug27 homologs. Bug69 and Bug27 were located in separate clusters, while the remaining 92 "Bugs" from Bordetella pertussis were excluded from the initial dataset because they were more divergent (refer to the Methods sections for parameters used for generating the initial dataset of sequences).

Genome context analysis and phylogenetic distribution of Bug69 and Bug27

To aid functional inference, we integrated the SSN with genome neighborhood information using the EFI-GNT tool (Fig. 2A) and assessed the overall phylogenetic distribution of Bug69 and Bug27 homologs.

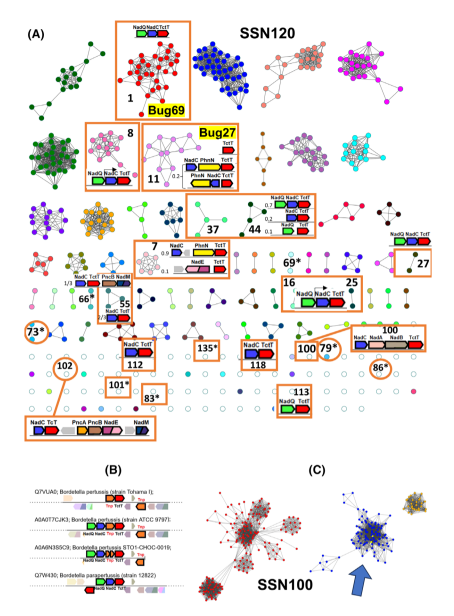


Fig. 2. Sequence similarity networks and genome context analysis of *Bordetella pertussis* Bug69 and Bug27 homologs. (A) SSN displayed at an alignment score of 120 (~67% sequence identity) generated with *B. pertussis* Bug69 (Uniprot: Q7VUA0) as the input sequence. Clusters with the TctC homolog in a NAD cofactor-related genomic context are boxed, and the gene cluster is shown inside. The relative frequency is indicated on the left when different gene contexts are present within the same SSN cluster. Clusters with an asterisk possess the same gene context as in 1 (not repeated for clarity). Taxonomy of sequences in NAD-related clusters: *Bordetella* spp., *Achromobacter* spp., and *Alcaligenes xylosoxydans* (1); *Cupriavidus* spp. (7); *Comamonas* spp. (8); *Bordetella* spp., *Variovorax* spp., *Jezberella montanilacus*, *Alcaligenes faecalis*, *Achromobacter pestifer* (11); *Delftia* spp. (16); *Algalinigenes* spp. (25); *Orrella dioscoreae*, *plant metagenome* (27); *Pollutimonas* spp., *Cupriavidus* sp. *USMAA2-4*, *Alcaligenaceae bacterium* (37); *Pseudorhodoferax* spp., *Lampropedia cohaerens* (44); *Vandammella animalimorsus*, *Franklinella schreckenbergeri* (55); *Alcaligenaceae bacterium* (66, 69, 73, 79); *Pusilimonas* sp. *ANT_WB101* (79); *Pseudorhodoferax* sp. *Leaf267* (83); *Comamonas kerstersii* (86); *Comamonas* sp. *BIGb0124* (113); *Lampropedia hyalina DSM 16112* (100); *Paenalcaligenes hominis* (101); *Verticiella sediminum* (118); *Pusilimonas* sp. *TS35* (135). (B) Genome neighborhood diagrams of *B. pertussis* strains show transposable elements in the context of TctC. (C) A partial view of the SSN is displayed at an alignment score of 100. The blue cluster contains all NAD-related nodes (boxed) of SSN120.

Our analysis revealed that in all 80 members of the Bug69-containing cluster, almost equally represented by *Bordetella* and *Achromobacter* spp. of the

Alcaligenaceae family, *bug69* is in an operon downstream of the NAD transcriptional regulator *nadQ* [31] and *nadC* (Cluster 1 in Fig. 2A). The only exception is B. pertussis Tohama I, where chromosomal rearrangement has translocated the gene elsewhere (Fig. 2B). A close inspection of the gene neighborhood in all sequences showed a prevalent association of TctC with *nadC* and *nadO* in many other clusters populated by genera of Alcaligenaceae, Comamonadaceae, and Burkholderiaceae (Fig. 2A). Occasionally, tctC is adjacent to only *nadC* or *nadO*, with the third gene located elsewhere. In addition to their physical proximity, genes *nadC* and *tctC* are also predicted to be co-regulated by NadQ transcriptional regulator as shown for the examples of Delftia acidovorans SPH-1 (Cluster 16 in Fig. 2A) and Comamonas testosteroni KF-1 (Cluster 8) (https://regprecise.lbl.gov/index.jsp). Overall, our comprehensive analysis corroborates and extends the functional association of QaPRT with Bug69, suggesting that Bug69 might have evolved to recruit the QaPRT substrate quinolinic acid.

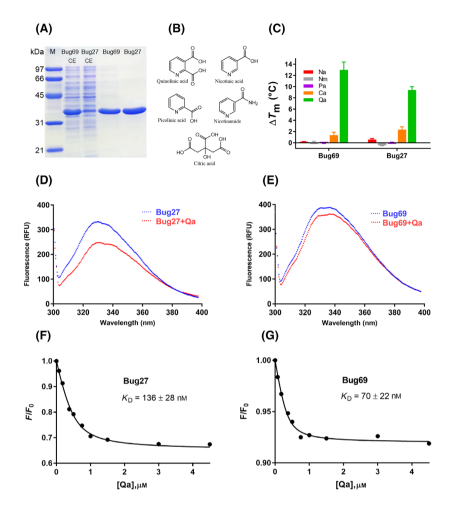
Bordetella pertussis Bug27 is found in cluster 11 of the SSN (Fig. 2A). The structural and functional characterization of *B. pertussis* Bug27 showed its capacity to bind Na and Nam, two precursors of NAD, despite not being in a genome context with NAD metabolic genes [17]. However, an inspection of the gene neighborhood of orthologs from Variovorax spp. (Alcaligeneaceae) and the more distant Jezberella montanilacus (Comamonadaceae) shows that bug27 is adjacent to nadC and phnN. The latter gene encodes the enzyme ribose 1,5-bisphosphate phosphokinase, which converts ribose 1,5-biphosphate (PRP), a crucial component in the metabolism of ribonucleotides and phosphonates. into phosphoribosyl pyrophosphate (PRPP) [32,33]. PRPP, in turn, serves as the phosphoribosyl donor in the QaPRT-catalyzed reaction (see Fig. 1). This finding suggests a role for Bug27 in NAD metabolism as an alternative binder of quinolinic acid, distinct from Bug69.

Furthermore, our gene neighborhood analysis has revealed, in more distant species, alternative genetic arrangements of TctC homologs with the *de novo nad-BAC* genes, salvage *pncAB* genes, and downstream NAD biosynthesis genes *nadE* or *nadM* (see clusters 7100, 102). This provides additional evidence of the involvement of Bug69 and Bug27 in NAD metabolism and allows us to formulate a hypothesis on the evolutionary scenario of NAD-related Bug proteins. These proteins could have initially emerged as promiscuous binders of NAD precursors as Qa, Na, Nm, and Asp and subsequently specialized into more specific Qa scavengers under selective pressure.

Lastly, applying a more relaxed alignment score threshold of 100, we generated a sequence similarity network in which all TcTs found in NAD metabolic contexts in SSN120 segregate in a single, allegedly iso-functional cluster (Fig. 2C).

In vitro characterization of *B. pertussis* Bug69 and Bug27

As inferred from our bioinformatics analysis, the most plausible hypothesis is that Bug69 and Bug27 may be involved in the transport of Qa. To experimentally validate this conjecture, we first obtained Bug69 and Bug27 in recombinant form and purified them to homogeneity (Fig. 3A). We then employed a fluorescence-based thermal unfolding assay [20] to test the NAD precursors Qa, Na, and Nm as potential ligands (Fig. 3B). We included the pyridine picolinic acid (Pa) and tricarboxylic citric acid (Ca) as control molecules. Ca was previously co-crystallized with Bug27 in an open state, thereby being identified as a non-bona fide ligand of Bug27 [17]. An initial screen at a fixed concentration of 1 mM showed that, among the tested compounds, only Qa significantly affected the thermal stability of the proteins (Δ Tm), with a > 50-fold preference over Na in Bug69 and about 20-fold in Bug27. On the other hand, Ca induced modest shifts, around 10 and 5 times less than Qa, in Bug69 and Bug27, respectively (Fig. 3C). Both proteins feature a conserved Trp alongside varying quantities of tyrosines and phenylalanines, hinting at the potential utility of fluorescence spectroscopy in accurately evaluating quinolinate ligand binding. A comparison of the fluorescence emission spectra of Bug27 and Bug69 in the absence and presence of quinolinate showed a significant fluorescence quenching (Fig. 3D,E). However, this effect differed in magnitude for each protein, indicating slightly distinct changes in the tryptophan environment upon ligand binding. Tryptophan emission peaked at 330 nm (excited at 295 nm) for Bug27 and at 335 nm for Bug69, with the peak emission wavelength remaining unchanged upon quinolinate binding. Based on quinolinateinduced fluorescence change, the quantitative analysis revealed that the dissociation constants (K_D) for quinolinate binding to Bug27 and Bug69 were 136 ± 28 nm and 70 ± 22 nm, respectively (Fig. 3F,G). These results suggest that Bug69 and Bug27 proteins are high-affinity quinolinate binders. The affinity of Qa for Bug27 was about three times higher than that of Na, Nm, and Ca, as previously calculated with the same approach [17]. Hence, using two independent biophysical assays, we experimentally supported our bioinformatics prediction that Qa could be the natural ligand of both Bug69 and Bug27.



In silico binding analysis

Following the *in vitro* binding study, we next performed molecular docking to get insights into the binding selectivity of Oa. After removing the non-natural citrate ligand from the binding pocket, we used the available Bug27 (PDB: 2QPQ) structure as the template to probe Qa, Na, Nm, and citrate. In line with the in vitro screening and the bioinformatics predictions, Oa vielded the best scoring pose, as reflected by its most extensive interactions with the binding pocket over the other ligands (Fig. 4A-C). In detail, as shown in Fig. 4A, the carboxylic group in position 2 of Qa is firmly anchored with two hydrogen bonds to the main chain amides of Gly21 and Thr22 and with a third H-bond to Thr22 side chain, while the carboxylate in position 3 forms an H-bond with Cys165 (main chain). The pyridine ring is held in place by an H-bond between the ring nitrogen and the Asn228 side chain amide and is sandwiched between Phe12 and Thr183. Additional residues, namely Ser74, Asn74, and Gln142, indirectly contribute to the ligand-pocket H-bonding Fig. 3. In vitro characterization of Bordetella pertussis Bug69 and Bug27. (A) SDS/PAGE of purified Bug69 and Bug27. M. protein molecular marker: CE, crude extracts (CE), (B) Chemical structures of pyridine NAD precursors and other carboxylic compounds used for the in vitro binding screening. (C) Histograms of the $\Delta T_{\rm m}$ (°C) of Bug69 and Bug27 in the presence of 1 mm concentration of the tested compound. The data represent the mean + SD of three independent experiments. (D, E) Fluorescence emission spectra of Bug27 and Bug69 in the presence and absence of saturating guinolinate (4.5 µM). The excitation and emission wavelengths were 295 and 328 nm, respectively. (F, G) The amplitude of normalized fluorescence as a function of quinolinate concentration for Bug27 (left) and Bug69 (right). The solid line represents a fit to the equation provided in the methods section. Data represent the mean of three independent experiments.

network, while Cys136 further stabilizes the pyridine ring with a pi-alkyl interaction (Fig. 4A). Next, we ran a structure-based multiple sequence alignment with all members of the NAD-related cluster previously identified with the SSN analysis. As a result, the identified ligand binding residues are universally conserved within the cluster, supporting their evolutionary selection for Qa binding in these Bug subfamilies (Fig. 4D).

Along this line of thought, the Qa binding residues within the Bug69/27 family differ from their structural counterparts in other Bugs with known structures and ligands [34–39] (Fig. 5).

Discussion

Among the three known multicomponent transport systems that rely on the initial substrate recognition by a periplasmic binding protein, the Tripartite Tricarboxylate Transporter, prototyped by the TctABC citrate transporter system [40], is the most recent and the least characterized. Strikingly, the TTT substrate

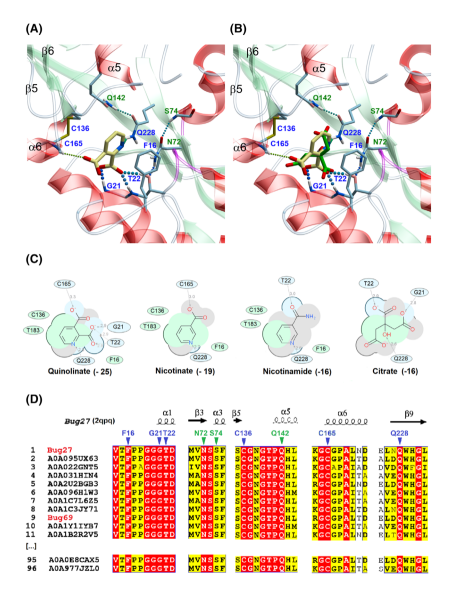


Fig. 4. *In silico* binding analysis of Bug27. (A) Close-up view of *Bordetella pertussis* Bug27 (PDB: 2qpq) binding site in ribbon representation with docked Qa (in yellow sticks). Secondary structure elements are color-coded as follows: light green for β-sheets, red for α-helices, and white for loops or turns. Binding residues in direct contact with the ligand (with H-bonds or hydrophobic interactions) are displayed in sticks with blue labels. Additional residues that indirectly contribute to the ligand stabilization (e.g., by forming an H-bonding network) are displayed with green labels. H-bonds are shown as colored dotted lines, while π -π interactions as black dotted lines. (B) Docked Qa (yellow) overlaid with co-crystallized citrate molecule (green) from PDB 2qpq. (C) 2D interaction diagram between tested ligands and receptor. Green and blue shading represents hydrophobic and H-bonding contacts, respectively. Dashed arrows represent hydrogen bonds. The docking score is in brackets. The ICM software docking Score is a Generalized Born Surface Area (GBSA)/Molecular Mechanics (MM)-type scoring function augmented with a directional hydrogen bonding term [46]. (D) Partial views of the structure-based sequence alignment of the NAD-related Bug27/69 TctC subfamily (Blue cluster in Fig. 2C). The alignment was performed using 96 representative sequences out of the 253 by applying an 80% sequence identity cutoff. Numbering and secondary structure elements derive from the structure of *B. pertussis* Bug27 (PDB: 2qpq). Conserved residues putatively involved in the quinolinate binding site are marked by triangles with the same color coding as panel A.

binding proteins (TctC homologs) are significantly over-represented in some bacterial pathogens, underscoring the extensive functional diversification within this family. In addition, most *tctC* genes lack readily detectable membrane counterparts in genomic proximity, hence their assignment as "orphan" proteins. In the pathogen *B. pertussis*, only two out of 76 TctC homologs (collectively renamed as Bug proteins) have

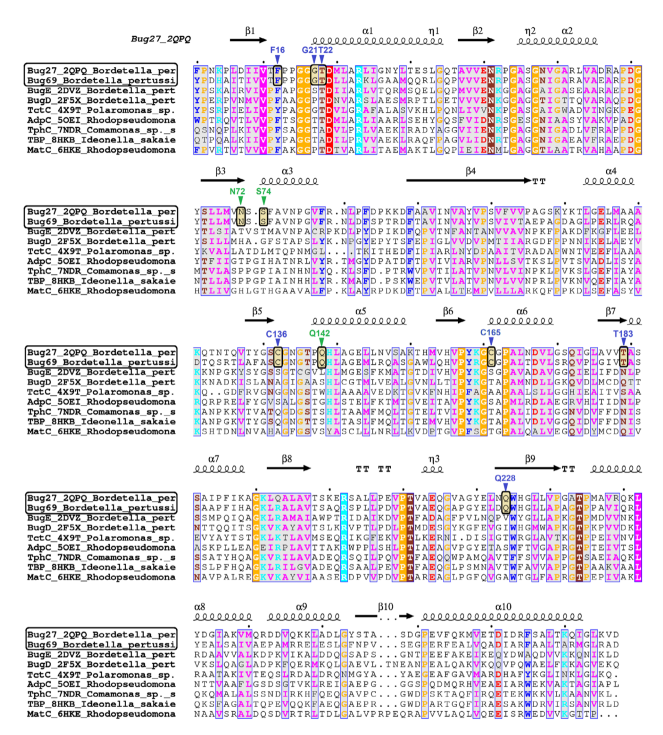


Fig. 5. Structure-based sequence alignment of TctC/Bug proteins with known ligand and structure. Secondary structure elements from Bug27 (PDB: 2qpq) are represented by spirals (α and 3₁₀ helices) and arrows (β strands). The 3₁₀ helix is labeled η . Partially conserved residues (numbered by Bug27 structure) are shown on a gray background, and universally conserved residues are highlighted by a solid background color. Triangles indicate residues identified in the quinolinic acid-binding pocket, as also shown in Fig. 4. Highlighted boxes mark the unique conservation of such residues among the Bug27-Bug69 pair. Ligands (in brackets) of TctC proteins used are as follows: *Bordetella pertussis* BugE (glutamate); *B. pertussis* BugD (aspartate); *Polaromonas* TctC Bpro_3516 (unknown); *Rhodopseudomonas palustris* AdpC (oxoadipate); *Comamonas* sp. TphC (terephthalate); *Ideonella sakaiensis* TBP ISF6_0226 (terephthalate); *Rhodopseudomonas palustris* MatC Rpa3494 (malate).

undergone a comprehensive biochemical and structural characterization: BugD and BugE, which are highaffinity binders of aspartate and glutamate, respectively [34,35]. Among the orphan Bug proteins, Bug27 and Bug69 have been previously associated with a pyridine transport function, but the results were preliminary. Herrou et al. [17] crystallized Bug27 liganded with citrate and found substantially identical affinity constants for citrate, nicotinate, and nicotinamide with a protein fluorescence quenching method. The test of Na and Nm was justified by the upregulation, among other genes, of Bug27 when bacterial cells were grown in the presence of Na. The substrate promiscuity of Bug27 and the observation that the inactivation of bug27 did not affect Na uptake [17] prompted us to speculate that this PBP might prefer a different pyridine precursor. In parallel, Brickman et al. [16], studying the essential role of Bordetella NadC in the Qa salvage pathway for NAD biosynthesis, found that the gene bug69, which shares nearly 60% sequence identity with bug27, occurs in a gene cluster with nadC and its transcriptional regulator nadQ in several Bordetella species. This observation led to the assumption that Bug69 could detect Qa. Nevertheless, the genetic inactivation of Bug69 did not yield any growth defect on Qa [16], leaving the functional role of Bug69 an open question as well.

Through sequence similarity network analysis and a gene context survey across one thousand bacterial species, we identified a putative isofunctional group comprising orthologs of Bug27 and Bug69 from β -proteobacteria. These orthologs are strongly associated with *nadC* and *nadQ*, individually or in combination, along with some clustering with other NAD-related genes.

Thus, the most plausible hypothesis is that both Bug69 and Bug27 might participate in the transport of Qa. However, genome context analysis did not reveal a membrane domain encoded adjacent to or near Bug69 or Bug27. This is unsurprising, as Bordetella uptake genes are a disproportionately represented family of orphan SBPs, i.e., SBPs for which membrane transport proteins are unknown. Consequently, membrane domain partners could be shared by multiple Bugs and be encoded in distant genomic sites. Despite the strong genetic association with NAD-related genes, other functions unrelated to NAD metabolism cannot be ruled out for Bug27 and Bug27. One possibility is that Bug27 and Bug69 are signaling proteins specialized in sensing quinolinate. This scenario could be particularly relevant in inflammation and immune response. In Eukaryotes, Qa is formed via the aerobic degradation of tryptophan (known as the kynurenine pathway) [41]. Several studies have suggested that inflammation induced by bacterial infections can lead to dysregulation of the kynurenine pathway with the concomitant increase in quinolinic acid production [42].

Through thermal shift assay, we demonstrated that Bug69 and Bug27 bind Qa with at least a 40-fold preference over the other NAD precursors, Na and Nm. $K_{\rm D}$ values of about 0.1 μ M were determined by Tryptophan fluorescence assay. These results show that Oa is an in vitro high-affinity binder and the likely natural ligand of these orphan Bugs. Our results allow us to reconcile the lack of functional defect of Bug69 mutant when grown on Qa [16] due to the same function exerted by Bug27 and, possibly, other uncharacterized proteins that facilitate Qa import. We also speculate that the high concentration of Qa used in the growth medium (30 µM) might also contribute to the absence of the Bug69 mutant's defect. Indeed, a significant amount of Qa could permeate the Bordetella cells at such concentration, sustaining growth even without an import facilitator like Bug69. Similarly, Streptococcus pyogenes carries a similar NAD metabolism gene set as Bordetella, lacking de novo biosynthesis but retaining a functional NadC to salvage Qa from the medium [15]. This study demonstrated that S. pyogenes can grow on Qa as a single pyridine precursor with a minimal requirement of $\sim 10 \,\mu\text{M}$ without an apparent Qaspecific transporter. As in pathogenic streptococci, genome context analysis in Bordetella pertussis did not point to any apparent transmembrane component that could partner with Bug69 or Bug27 in Qa import. Thus, Qa can permeate the plasma membrane by diffusion or share an unidentified pyridine or dicarboxylate transporter encoded by the *B. pertussis* genome.

In conclusion, our combined experimental and bioinformatics findings suggest that *Bordetella pertussis* Bug69 and Bug27 function as high-affinity, standalone periplasmic binding proteins of quinolinic acid. This pyridine precursor, akin to Na or Nm, can fuel NAD salvage biosynthesis in Bordetellae (Fig. 1). Our bioinformatics predictions enable confident projection of isofunctional orthologs in many β -proteobacteria families (Fig. 2). Hopefully, this approach could be replicated to dissect the complex functional diversification of the TctC/Bug family of periplasmic binding proteins.

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Conflict of interest

The authors declare that they have no conflict of interest related to this study.

Data accessibility

The data supporting the findings of this study are available from the corresponding author, L. Sorci, at l. sorci@staff.univpm.it, upon request.

Author contributions

LS and NR conceived and designed the project; LS, GM, and FM performed the experiments; LS performed the bioinformatics analyses; LS and GM acquired the data; LS, GM, AA, and NR analyzed and interpreted the data; LS and NR wrote the paper.

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