

The *Salmonella enterica* serovar Typhimurium virulence factor STM3169 is a hexuronic acid binding protein component of a TRAP transporter

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Abstract

The intracellular pathogen *S*. Typhimurium is a leading cause of foodborne illness across the world and is known to rely on a range of virulence factors to colonize the human host and cause disease. The gene coding for one such factor, *stm3169*, was determined to be upregulated upon macrophage entry and its disruption reduces survival in the macrophage. In this study we characterize the STM3169 protein, which forms the substrate binding protein (SBP) of an uncharacterized tripartite ATP-independent periplasmic (TRAP) transporter. Genome context analysis of the genes encoding this system in related bacteria suggests a function in sugar acid transport. We demonstrate that purified STM3169 binds D-glucuronic acid with high affinity and specificity. *S*. Typhimurium LT2 can use this sugar acid as a sole carbon source and the genes for a probable catabolic pathway are present in the genome. As this gene was previously implicated in macrophage survival, it suggests a role for D-glucuronate as an important carbon source for *S*. Typhimurium in this environment.

INTRODUCTION

Non-typhoidal *Salmonella* is a leading cause of foodborne illness worldwide [1]. In particular, *Salmonella enterica* serovar Typhimurium (S. Typhimurium) is a common cause of gastroenteritis and can lead to inflammatory diarrhoea and bacteraemia [2, 3]. This intracellular pathogen is able to survive and replicate within both phagocytic and non-phagocytic cells. The intracellular environment of a macrophage phagosome is harsh, containing acid, bactericidal compounds and reactive oxygen species [4, 5]. However, this also provides a niche for adaptive enteropathogenic bacteria such as *S*. Typhimurium to colonize and replicate within, isolated from patrolling immune cells [6]. These bacteria rely on a global change in gene expression to adapt to the intracellular environment, employing a range of virulence factors to mediate host cell entry, survival and growth.

Haneda *et al.* investigated the shift in protein levels in *S*. Typhimurium strain SH100 upon macrophage entry and identified over 50 proteins whose increased accumulation

was dependent on the stringent response-related signalling molecule ppGpp [7]. The gene encoding the most differentially regulated protein, *stm3169*, was then disrupted on the *S*. Typhimurium chromosome, and the resulting strain had a significant growth defect inside macrophage-like RAW264.7 cells [7]; as such, STM3169 was suggested to be important for macrophage survival and virulence.

In this study we have characterized the function of this protein, which encodes a substrate binding protein of an uncharacterized tripartite ATP-independent periplasmic (TRAP) transporter. We provide a novel perspective into important cellular functions needed for macrophage survival.

METHODS

Cloning, expression and purification of STM3169

The gene *stm3169* was amplified from *S*. Typhimurium LT2 (ATCC 700720) and cloned into pETYSBLIC3C [8] by ligation-independent cloning. This plasmid was transformed into the expression host *Escherichia coli* BL21 (DE3). Protein

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Abbreviations: CV, column volumes; FT-ICR MS, Fourier-transform ion cyclotron resonance MS; MST, microscale thermophoresis; SBP, substrate binding protein; TRAP, tripartite ATP-independent periplasmic.

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expression was carried out in 1 litre of Luria Bertani (LB) medium supplemented with kanamycin at 50 µg ml⁻¹, with 0.4 mM IPTG added once the optical density (OD₆₀₀) reached 0.6. Cells were incubated at 37 °C at 120 r.p.m. overnight, then harvested by centrifugation at 7800 g for 25 min. The cell pellet was resuspended in wash buffer (50 mM KPi pH 7.8, 200 mM NaCl, 20% glycerol, 40 mM imidazole) containing 0.4 mM AEBSF protease inhibitor (Sigma Aldrich). The sample was sonicated at 70 W for 3 min, followed by centrifugation at 27000 g at 4 °C for 30 min. Ni²⁺ affinity chromatography was used to purify the His_c-tagged protein. The cell lysate was loaded onto a HisTrap column (GE Healthcare) and washed with the aforementioned wash buffer, then eluted with the elution buffer (50 mM KPi pH 7.8, 200 mM NaCl, 20% glycerol, 500 mM imidazole). The protein of interest was identified by SDS-PAGE. The purified protein was buffer exchanged into 50 mM KPi pH 7.8 and 200 mM NaCl using the HiTrap Desalting column (GE Healthcare). Protein concentration was measured by determining the absorbance at 280 nm and using the Beer-Lambert law.

Protein renaturation

A fraction of the desalted protein was then loaded onto a 5 ml HisTrap column (GE Healthcare). The column was then washed with 30 column volumes (CV) of denaturation buffer (50 mM KPi pH 7.8, 200 mM NaCl, 20% glycerol, 40 mM imidazole, 2 M guanidine hydrochloride) to unfold the bound protein. A gradient wash across 12 CV was performed from the denaturation buffer to the previously mentioned guanidine hydrochloride-free wash buffer to refold the protein. A further 8 CV wash using the wash buffer was performed before elution using the elution buffer as before. Circular dichroism was used to check correct refolding of the renatured protein. CD spectra were determined using a J810 spectropolarimeter. Protein samples were dialysed into 20 mM KPi pH 7.8 buffer and diluted to a concentration of 0.2 mg ml⁻¹. The spectra were determined in a 1 mm pathlength quartz cuvette between 260 and 180 nm at 100 nm min⁻¹ with 1 nm pitch. Prior to any downstream experiments, the protein was desalted into 50 mM KPi pH 7.8 and 200 mM NaCl. The concentration of the refolded protein was measured by determining the absorbance at 280 nm and using the Beer-Lambert law and predicted extinction coefficient from ProtParam (Expasy).

Thermal shift assay

The Protein Thermal Shift Dye kit (Applied Biosystems) was used to carry out the thermal shift assay. A final concentration of 5μ M protein in 50 mM KPi, 20 mM NaCl and each ligand at final concentrations of 1 nM, 1μ M and 1 mM was added to the thermal shift dye in triplicate. The experiments were performed using a StepOne Real-Time PCR system (Applied Biosystems) with a temperature ramp of 1.5 °C every 2 min from 25 to 99 °C. The melting temperatures (T_m) were calculated by determining the inflection point of the plot of the normalized fluorescence measurements against temperature.

Intrinsic fluorescence spectroscopy

All intrinsic fluorescence measurements were performed using a FluoroMax 4 fluorescence spectrometer (Horiba Jobin-Yvon). STM3169 in 50 mM KPi pH 7.8 plus 200 mM NaCl buffer was used at a concentration of $0.5 \,\mu$ M. The protein was excited at 297 nm with slit widths of 3.5 nm and emission was monitored at 330 nm. To test for potential ligands, various sugar acids were sequentially added at concentrations of $100 \,\mu$ M.

To quantitatively assess binding of specific ligands, protein was titrated with increasing concentrations of ligand in order to elucidate the $K_{\rm D}$ of ligand binding. Cumulative fluorescence was then plotted in Prism 5 (GraphPad) and the $K_{\rm D}$ was calculated from the hyperbolic fit of the binding curve.

Microscale thermophoresis

The fluorescent dye NT-647-NHS was used at 30 μ M to label 15 μ M of STM3169. Any unreacted dye was removed using a HiTrap Desalting column (GE Healthcare). Protein and dye concentrations were measured by absorbance at 280 and 650 nm, respectively. The concentration of ligand was varied from 200 μ M to 5 nM in a 16-step dilution, then mixed with labelled protein in capillaries for data acquisition using the Monolith NT.115 (NanoTemper). Outliers were detected using the NanoTemper analysis software.

Mass spectrometry by FT-ICR-MS

The natively folded STM3169 containing an N-terminal tag (MGSSHHHHHHSSGLEVLFEGPA) attached to the mature peptide was used. A sample of this protein was dialysed into 35 mM ammonium acetate pH 5 and diluted to a final protein concentration of 100 µM. Fourier-transform ion cyclotron resonance MS (FT-ICR-MS) was performed using a solariX FT-MS device (Bruker Daltonics) with a 9.4-T superconducting magnet. Analysis was performed in denaturing conditions where the protein was diluted to $1 \,\mu\text{M}$ in 50% (v/v) acetonitrile containing 0.1% (v/v) formic acid before introduction via a TriVersa NanoMate (Advion BioSciences) source in positive-ion mode. The applied voltage was adjusted to 1.4–1.7 kV to achieve a stable ion current. A 120 °C nitrogen dry gas was supplied at 1.3 l min⁻¹ to aid desolvation. Instrument control and data acquisition used Compass 1.4 (Bruker Daltonics). Spectra were generated by the accumulation of 20 scans with 0.1 s ion cooling time and 0.4s scan time with 8 million data points recorded. To improve resolution and reduce space-charge effects, the 42+ protein charge states were isolated in the hexapole before scanning in the FT-ICR cell. Spectra from isolated charge states were deconvoluted to monoisotopic protein mass using version 2.0 of the SNAP averaging algorithm (C 4.9384, N 1.3577, O 1.4773, S 0.0417 and H 7.7583%), in DataAnalysis version 4.0 (Bruker Daltonics).

Homology modelling of STM3169

The predicted structure of the mature STM3169 was prepared by automated homology modelling on SWISS-MODEL (swissmodel.expasy.org) [9–12]. The structure of Bamb_6123 from *Burkholderia ambifaria* AMMD (PDB ID: 4N15) was used as a template for homology modelling of the mature peptide of STM3169. The QMEAN Z-score was predicted to be -0.69, which suggests that the model of STM3169 is consistent with other experimentally determined structures of a similar size. The derived GMQE (global model quality estimation) for the model was determined to be 0.74, indicating high reliability.

S. Typhimurium LT2 growth experiments

A stationary phase culture of *S*. Typhimurium LT2 (ATCC 700720) grown in Luria Bertani (LB) media was centrifuged and washed with M9 salts [13] to remove any residual LB. These were inoculated into 96-well microplates with an initial OD_{600} of 0.01 in M9 media (48 mM Na₂HPO₄, 22 mM KH₂PO₄, 19 mM NH₄Cl, 9 mM NaCl, 2 mM MgSO₄) with 5 mM carbon source (D-glucose, D-galactose, sodium D-glucuronate, sodium D-galactonate, L-arabinose; Sigma Aldrich). Triplicate cultures were grown in the microplates at 37 °C, and shaken in an Epoch2 microplate spectrophotometer (BioTek). OD₆₀₀ measurements were taken every hour for 15 h.

RESULTS AND DISCUSSION

The *stm3169* gene from *S*. Typhimurium LT2 is encoded within a typical operon for a TRAP transporter, in that it also includes genes encoding the two required membrane subunits (*stm3170* and *stm3171*) [14] and so is probably part of a functioning transporter (Fig. 1). STM3169 is the substrate binding protein (SBP) component that initially binds the substrate in

the periplasm; this component is usually expressed at high levels relative to the membrane domains, consistent with the high abundance of the STM3169 protein in the Haneda *et al.* study [7, 14].

TRAP transporters form a large family of diverse SBPdependent secondary transporters in bacteria, the functions of which are, largely, to move organic acids into bacterial cells [15–17]. A hallmark sign of this substrate preference is a conserved arginine residue in the SBP subunit that coordinates the carboxylate group of the acid ligand [18, 19]. This residue is conserved in STM3169 (Arg170), suggesting a typical organic acid-type ligand. Beyond predicting that the ligand contains a carboxylate group, the binding sites of different TRAP transporter SBPs are highly variable, allowing them to accommodate ligands of many different sizes and shapes, often using complex water networks [17, 20].

In S. Typhimurium LT2, the *stm3169-stm3171* TRAP transporter gene cluster is not found in a region containing any other associated genes that give any clue to its substrate and hence cellular function (Fig. S1a, available in the online version of this article). However, genome analysis of *stm3169* orthologues from related gamma-proteobacteria revealed a range of different cluster architectures that strongly suggest a role in sugar acid import and catabolism for these strains (Fig. 1). A common feature of these clusters is the presence of genes coding for proteins predicted to be involved in hexuronate catabolism. These proteins, UxuA, UxuB and UxuC, have been characterized in *Erwinia chrysanthemi* where they are responsible for the utilization of D-glucuronate following uptake by the hexuronate MFS transporter ExuT [21]. The



Fig. 1. Homologues of STM3169 may be encoded from operons containing hexuronate metabolic genes. In *S.* Typhimurium LT2, *stm3169* is found on an operon with *stm3170* and *stm3171*, the two putative membrane components of a TRAP transporter. This operon is not surrounded by other genes that might be involved in uptake or sugar metabolism. Some genes neighbouring *stm3169* homologues were predicted to code for the hexuronate catabolism enzymes UxuAB and a glucuronate isomerase, UxuC/UxaC. Genes are represented by coloured bars, identifying its encoded protein's putative function. Homologues were identified in other strains using MicrobesOnline.





presence of homologues of UxuABC (STM3135–STM3137) with ExuT (STM3134) in *S*. Typhimurium (Fig. S1b) suggests its ability to catabolize this sugar. Additionally, this pattern of co-occurrence is found in other genomes represented in MicrobesOnline [22], suggesting that this TRAP transporter might be involved in uptake of hexuronic acids.

To test this hypothesis, we expressed and purified recombinant STM3169 from *E. coli* using nickel-affinity chromatography (Fig S2a). A portion of the purified protein was denatured using guanidine hydrochloride and refolded, removing any endogenous ligand potentially carried over during the purification, a common phenomenon with these SBPs [23, 24]. Analysis of the protein by circular dichroism suggested minimal differences in the folding when comparing the native and refolded protein, suggesting successful protein refolding (Fig. S2b). The protein was analysed by MS to have a mass of 35741.96 Da, which matches the predicted monoisotopic mass of the protein (Fig. S2c). An additional signal at mass 35840.4 was observed, which, being 98 Da greater than the predicted signal, is probably a single phosphate adduct on the protein, an artefact we have observed with other SBPs [25].

To screen potential substrates for this TRAP transporter we used an established fluorophore-based protein thermal shift assay [17] using a range of monosaccharides and their related acid forms. Analysis of the natively purified protein showed no significant changes in protein thermal stability in the presence of any of the compounds tested: D-glucuronate, D-galacturonate, D-gluconate, D-galactonate, D-galactose and the negative control, L-arabinose (data not shown). However, for the refolded and presumably ligand-free protein, increases in protein thermal stability were observed in the presence of two sugar derivatives, D-galactonate and D-glucuronate (Fig. 2a). These data suggest that the purified protein from *E. coli* came with pre-bound ligand and also suggested ligands for continuation of *in vitro* binding experiments.

To further assess binding we used fluorescence spectroscopy, measuring the change in intrinsic protein fluorescence upon ligand binding. Using protein excitation at 295 nm, the same set of sugar acids were added sequentially to refolded STM3169, detecting fluorescence in time acquisition mode. Whilst addition of D-galactonate had little effect on fluorescence, D-glucuronate caused an ~10% quench in fluorescence (Fig. 2b). This discrepancy may be attributed to the nature of protein thermal shift experiments as non-specific interactions could give rise to a shift in thermal stability.

We proceeded to determine the binding affinity of STM3169 for D-glucuronate by titration and measuring the resulting changes in intrinsic fluorescence (Fig. 2c). The experimental data suggested D-glucuronate was binding to STM3169 with a binding constant ($K_{\rm D}$) of 0.17±0.077 µM. As a complementary biophysical methodology, we examined this interaction using microscale thermophoresis (MST), measuring a ligand-dependent change in thermophoresis of dyed protein molecules to indicate conformational changes that



Fig. 3. The putative binding site of STM3169 resembles that of characterized D-glucuronate binding proteins. Residues which were experimentally shown to contribute to ligand binding and their various homologous residues were identified. The carbon atoms and residue names of the various homologous residues were coloured by individual proteins. The D-glucuronate binding protein Apre_1383 from *Anaerococcus prevotii* PC1 (4N91, grey), the D-glucuronate and D-galacturonate binding protein Bpro_3107 from *Polaromonas* sp. JS666 (4MHF, orange) are shown. STM3169 from *S.* Typhimurium LT2 (light blue) was modelled using the crystal structure of Bamb_6123 (4N15) on SWISS-MODEL (swissmodel.expasy.org). Carbon atoms of the ligands, the alpha and beta anomers of D-glucuronate, are represented in light green. Nitrogen and oxygen atoms in the various structures are in blue and red, respectively. Residue names denoted in black are conserved between all structures analysed.

accompany ligand binding. This technique also suggested D-glucuronate was binding STM3169, and through repeated 16-step titrations a K_D of $0.25\pm0.048\,\mu$ M was estimated (Fig. 2d). No clear signal suggesting a conformational change upon D-galactonate binding was observed (data not shown). Together, these data are consistent with D-glucuronate being a ligand for STM3169, with affinities in the range determined for other high-affinity TRAP transporters such as the SBP of the sialic acid transporter of *Haemophilus influenzae*, SiaP, with a reported K_D of ~0.1 μ M [26].

A recent study by Vetting et al. [17] screened 158 SBPs against a large number of potential ligands to classify these proteins into isofunctional clusters. A number of these SBPs were then characterized structurally, some of which were successfully co-crystallized with D-glucuronate. We sought to compare these crystal structures against a model of STM3169 prepared on SWISS-MODEL (swissmodel.expasy.org) [9-12] to supplement our in vitro findings. These proteins were superposed using the secondary-structure matching (SSM) algorithm [27] on CCP4MG [28] and the measured root mean square distances of the alignments were found to be between 1 Å and 1.56 Å, suggesting highly similar overall structures. Analysis of the aligned binding sites (Fig. 3) initially revealed two conserved arginines (Arg150, Arg170 in STM3169) and an asparagine (Asn210 in STM3169), consistent with known D-glucuronate binding proteins [18]. As predicted, Arg170 in the model of STM3169 aligns with the residues responsible for coordinating the carboxyl group of the ligand in the D-glucuronate bound structures of Apre_1383 and Bpro_3107. The two other conserved residues, Arg150 and Asn210, have been shown to further coordinate the ligand by forming multiple hydrogen interactions. STM3169 residues Asn35, Gln36, Glu74 and Asp237 are fully conserved in the D-glucuronate-specific binding protein Apre_1383 and its corresponding residues form more hydrogen bonds with D-glucuronate. Bpro_3107, a D-glucuronate and D-galacturonate binding protein, possesses alternative residues also capable of coordinating the ligand. This intricate network of multiple predicted hydrogen bonds is likely to give rise to the high binding affinity for D-glucuronate shown though our *in vitro* assays.

Through in vitro studies and homology modelling, we have characterized a high-affinity substrate binding component of the TRAP transporter with apparent specificity for D-glucuronate. As S. Typhimurium also has homologous genes coding for proteins involved in D-glucuronate catabolism, it should be able to import D-glucuronate and utilize it as a carbon source. We confirmed this, an observation first observed in 1969 from the Ames group [29] (Fig. 4a); furthermore, we demonstrated the utilization of a selection of other sugars used in this study. From these data we propose a model (Fig. 4b) whereby the STM3169 protein forms part of a functioning TRAP transporter that functions alongside the putative UhpC/ExuT hexuronate transporter (STM3134) that is divergently transcribed from the uxuABC genes (stm3135-stm3137). After transport, the bacterium would use the UxuABC proteins and the enzymes of the Entner-Doudoroff pathway to convert D-glucuronate into central metabolites. The high-affinity nature of the TRAP transporter, compared to the lower affinity major facilitator superfamily protein ExuT, might suggest that D-glucuronate is present in the macrophage as a carbon source but at low micromolar concentrations, which would explain the significant increase in protein levels in the macrophage to ensure its efficient uptake [7].

This ability to metabolize sugar acids is key for the colonization of intracellular niches, where other substrates for glycolysis may not be present [16]. The landmark study of Eriksson et al. [30], which provided the first comparison of S. Typhimurium transcriptional changes upon moving to the intravacuolar environment of a macrophage, revealed that the organism appeared to be using the related sugar acid D-gluconate and the Entner-Douderoff pathway during growth [30]. In fact, the UhpT/ExuT-like transporter STM3134 was identified as being specifically upregulated as well as *dgoK* and *dgoA*, which encode enzymes that interconvert the UxuABC reaction products into pyruvate and glyceraldehyde-3-phosphate, suggesting a broader role for sugar acid molecules as metabolites. While STM3134 may be responsible for the transport of multiple hexuronate carbon sources, we have demonstrated that STM3169-STM3171 is a selective high-affinity transport system for D-glucuronate.

Despite the lack of evidence to suggest that the STM3169– STM3171 TRAP transporter is able to transport D-galactonate, our defined growth media assay (Fig. 4a) has hinted towards



Fig. 4. D-Glucuronate serves as a carbon source for *S*. Typhimurium. (a) *S*. Typhimurium LT2 was grown in minimal media using D-glucose, D-glucuronate, D-galactose, D-galactonate and L-arabinose as the sole carbon source. Data points are the average of three independent replicates and error bars indicate ±sD. (b) Proposed model for the uptake of D-glucuronate. D-Glucuronate can potentially be transported by the STM3169–STM3171 TRAP transporter system or the putative hexuronate transporter STM3134. Once inside the cell, UxuABC converts D-glucuronate to 2-keto-3-deoxygluconate (KDG)_ and, via the Entner–Doudoroff pathway, the latter is broken down into glycerol-3-phosphate and pyruvate for use. Coloured bars correspond to the various representations on Fig. 1.

another transporter capable of efficient D-galactonate uptake. This and other sugars found in the macrophage could be taken up by other high-affinity transporters, including further members of the TRAP family. There is previous evidence suggesting the two other members of the TRAP family in *S*. Typhimurium LT2, STM3671–STM3673 and STM4052–STM4054, have transport functions which could include carbon sources in the macrophage. The SBP of STM3671–STM3673 is 88% identical to the previously characterized *E. coli* 2,3-diketo-L-gulonate binding protein, YiaO [31–33], whilst STM4052–STM4054 is thought to be coregulated with proteins of the rhamnose utilization pathway by the activator RhaS (STM4048) through promoter analysis by RegPrecise [34].

Finally, we have shown evidence of function for the longidentified virulence protein STM3169 and suggested a role for its associated transport system in cell metabolism. Our discovery adds to our understanding of how pathogenic bacteria such as *S*. Typhimurium are able to survive within hostile host intracellular environments. Given its key role in virulence as observed by Haneda *et al.* [7], this knowledge could potentially aid in the design of competitive inhibitors.

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Author contributions

R.H., C.B.N. and A.L. performed the experiments and analysis. A.M. and A.A. generated the strain of $E.\ coli$ recombinantly expressing STM3169.

C.B.N. and G.H.T. conceptualized the experiments. R.H. and G.H.T. were involved in supervision. The manuscript was prepared by R.H., C.B.N. and G.H.T.

Conflicts of interest

The authors declare that there are no conflicts of interest.

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