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Expression and characterization of a histidine-rich protein, Hpn: potential for Ni\textsuperscript{2+} storage in Helicobacter pylori

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Hpn is a small cytoplasmic protein found in Helicobacter pylori, which binds Ni\textsuperscript{2+} ions with moderate affinity. Consisting of 60 amino acids, the protein is rich in histidine (28 residues, 46.7%), as well as glutamate, glycine and serine residues (in total 31.7%), and contains short repeating motifs. In the present study, we report the detailed biophysical characterization of the multimeric status and Ni\textsuperscript{2+}-binding properties of purified recombinant Hpn under physiologically relevant conditions. The protein exists as an equilibration of multimeric forms in solution, with 20-mers (approx. 136 kDa) being the predominant species. Using equilibrium dialysis, ICP-MS (inductively coupled plasma MS) and UV/visible spectroscopy, Hpn was found to bind five Ni\textsuperscript{2+} ions per monomer at pH 7.4, with a dissociation constant \(K_d\) of 7.1 \(\mu\)M. Importantly, Ni\textsuperscript{2+} binding to Hpn is reversible: metal is released either in the presence of a chelating ligand such as EDTA, or at a slightly acidic pH (pH for half dissociation, \(pH_{1/2}\) ~ 6.3). Ni\textsuperscript{2+} binding induces conformational changes within the protein, increasing \(\beta\)-sheet and reducing \(\alpha\)-helical content, from 22% to 37%, and 20% to 10% respectively. Growth curves of Escherichia coli BL21(DE3) both with and without the \(hpn\) gene performed under Ni\textsuperscript{2+} pressure clearly implied a role for Hpn to protect the cells from higher concentrations of external metal ions. Similarly, the accumulation of Ni\textsuperscript{2+} in these cells expressing Hpn from a plasmid was approx. 4-fold higher than in uninduced controls or control cultures that lacked the plasmid. Similarly, levels of Ni\textsuperscript{2+} in wild-type \(H.\ pylori\) 26695 cells were higher than those in \(H.\ pylori\) \(hpn\)-deletion mutant strains. Hpn may potentially serve multiple roles inside the bacterium: storage of Ni\textsuperscript{2+} ions in a ‘reservoir’; donation of Ni\textsuperscript{2+} to other proteins; and detoxification via sequestration of excess Ni\textsuperscript{2+}.

Key words: Helicobacter pylori, histidine-rich protein, Hpn, nickel, storage.

INTRODUCTION

Helicobacter pylori is a Gram-negative bacterium that colonizes the gastric mucosa of humans [1], and has been shown to be the causative agent of type B gastritis and peptic ulcers [2–4]. The development of gastric carcinoma and MALT (mucosa-associated lymphoid tissue) lymphoma are also strongly associated with chronic \(H.\ pylori\) infection [5,6]. To colonize and establish infection, \(H.\ pylori\) must produce an assortment of factors to generate a neutral pH in order to survive the extremely acidic environment in the stomach. One of them is a membrane-bound [Ni–Fe] hydrogen-uptake hydrogenase, which permits respiratory-based energy production for the bacteria in the mucosa [7,8]. Another is urease, a dinuclear Ni\textsuperscript{2+}–containing metalloenzyme that accounts for up to 6% of the soluble cellular proteins [9–11], which catalyses the hydrolysis of urea to yield ammonia and carbamate to neutralize the gastric acid, critical for \(H.\ pylori\)’s colonization and survival at acidic pH [12].

A constant supply of Ni\textsuperscript{2+} ions into \(H.\ pylori\) is therefore required for the synthesis and activity of these Ni\textsuperscript{2+}-containing metalloenzymes. To date, members of two general types of Ni\textsuperscript{2+}-specific, membrane-integrated uptake systems have been identified in \(H.\ pylori\) [13]: (i) the multiple-component ABC (ATP-binding cassette) transporters, which are believed to be a four-gene operon designated as \(abc\)ABC [14]; and (ii) the nickel–cobalt transporter family, comprising the putatively monomeric \(NixA\) protein [15].

However, when excess Ni\textsuperscript{2+} ions accumulate, they inhibit growth and exhibit toxic effects as a result of interference with normal protein–metal binding and catalysis, as well as the generation of reactive oxygen species [16–18]. The toxicity of metal ions is minimized through the strict control of their intracellular concentrations, which is generally achieved through two processes: regulation of ion transport, i.e. import and efflux from the cell, and sequestration, through the formation of tight complexes with metal-storage proteins [19,20]. As in other bacteria, \(H.\ pylori\) has to strike a delicate balance between the import of Ni\textsuperscript{2+} ions, their efficient intracellular storage and delivery to Ni\textsuperscript{2+}-dependent metalloenzymes when required.

The completion of the \(H.\ pylori\) genome sequence has revealed the presence of several putative ion-binding proteins and membrane transporters that are involved in the transport and cytoplasmic accumulation of bivalent cations [21]. As mentioned above, integral membrane proteins, such as the \(NixA\) and the \(abc\)ABC transporters, scavenge Ni\textsuperscript{2+} ions from the surrounding milieu and import them into the cell. In addition, two proteins with a high affinity for Ni\textsuperscript{2+} ions have been characterized in \(H.\ pylori\): the HspA heat-shock protein, a bacterial GroES homologue with a unique histidine-rich C-terminal domain [22], and an unusual protein called Hpn, which is abundant in the cell cytoplasm, accounting for approx. 2% of all protein synthesized [23]. As described previously [23], Hpn consists of 60 amino acids, including 28 histidine residues (Figure 1). The majority of the histidine residues are located within the central part of the protein (from residue 11 to residue 33) and include two separate stretches of six and seven consecutive histidine residues. There are two internal short repeats of Glu–Glu–Gly–Cys–Cys, at positions 38–42 and 51–55, four sets of paired histidine residues (one near the

Abbreviations used: ABC, ATP-binding cassette; Amp, ampicillin; DTT, dithiothreitol; ICP-MS, inductively coupled plasma MS; IPTG, isopropyl \(\beta\)-D-thiogalactoside; LB, Luria–Bertani; LMCT, ligand-to-metal charge transfer; MALDI, matrix-assisted laser-desorption ionization; MT, metallothionein; TTBS, Tris-buffered saline with Tween 20; UV/vis, UV/visible.

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N-terminus, three towards the C-terminus) and also an Xaa-Xaa-His motif at the N-terminus. All these sequence features indicate that this protein would bind metal ions strongly. Mutated strains of *H. pylori* lacking the *hpn* gene are four times more sensitive to ranitidine bismuth citrate, a metal-containing drug widely used to treat *H. pylori* infections, than the wild-type [24,25]. Recently, the *H. pylori* Ni^{2+}-uptake regulatory protein NikR was found to up-regulate not only the expression of the urease subunit genes, but also the expression of the *nixA* and *hpn* genes in the presence of excess Ni^{2+} [20]. This suggested that *H. pylori* may respond in a co-ordinated manner to the presence of extracellular Ni^{2+}, to allow the optimal import of Ni^{2+} ions (up-regulation of the *nixA* gene), optimal incorporation of Ni^{2+} into other proteins, as well as optimal storage of the free Ni^{2+} ions (up-regulation of the *hpn* gene) to minimize any toxic effects due to their accumulation [20].

In the present paper, we describe the cloning, expression, purification and characterization of the Hpn protein from *H. pylori*. Most notably, our results indicate that this protein would bind metal ions strongly. Mutated strains of *H. pylori* lacking the *hpn* gene would be four times more sensitive to ranitidine bismuth citrate, a metal-containing drug widely used to treat *H. pylori* infections, than the wild-type [24,25].

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions**

*H. pylori* 26695 was a gift from Department of Brucella, the University of Hong Kong, and was cultured on Brucella agar (Difco) plates supplemented with 10% defibrinated sheep blood at 37°C under microaerophilic conditions (5% CO₂, 4% O₂ and 91% N₂), maintained by CampyPak Plus (BBL). Plasmid DNA was maintained in *Escherichia coli* DH10B. *E. coli* BL21(DE3) (Stratagene) was used to express the *hpn* gene. Amp (ampicillin) (100 μg/ml) and IPTG (isopropyl β-D-thiogalactoside) (0.5 mM) were added to LB (Luria–Bertani) broth (USB) when indicated. Chromosomal *hpn*-deletion mutants were constructed as described previously by electroporating plasmid pPHN2KO (kindly supplied by A. G. Plaut, GRASP Digestive Disease Research Center, Tufts-New England Medical Center, Boston, MA, U.S.A.) into *H. pylori* 26695 cells in 10% (v/v) glycerol (Bio-Rad gene pulser at 12.5 KVCm, 25 μF, 600 Ω) [26]. Kanamycin-resistant colonies were isolated, and cell lysates of cultures grown from individual clones were analysed for Hpn production by Western blot analysis, to confirm gene deletion.

**DNA techniques**

Standard DNA manipulations were performed as described by Maniatis et al. [27], or as recommended by the manufacturers. Plasmid DNA was isolated using the QIAprep Spin mini kit (Qiagen). DNA fragments were purified from 1.0% agarose gels [1 × TAE (Tris/acetate/EDTA)] with the QIAquick gel extraction kit (Qiagen). PCR was performed on a PerkinElmer 2400 thermal cycler using the Expand High Fidelity PCR System (Roche). Oligonucleotide primers were synthesized by Proligo (Singapore). The *hpn* gene (HP1427) was amplified from unpurified *H. pylori* 26695 genomic DNA (from a picked colony) by PCR using primer 1 (5′-TACTCCATGGCTCATATGGCACACC-ATGAAGAACAGCA-3′) and primer 2 (5′-TATACTCGAGTT-ACCGTGATGCCTGGCAGCACCAAC-3′), which introduce NdeI and XhoI restriction sites (underlined) at the 5′- and 3′-ends of the PCR product respectively. After restriction digestion, the PCR products were ligated into the expression vector pET-32a (Novagen), pre-cut with the same enzymes and gel-purified, to generate the plasmid pET-Hpn. The plasmid was sequenced using ABI Prism BigDye (Applied Biosystems) and subsequently transformed into *E. coli* BL21(DE3) as described previously [27] for protein expression.

**Expression and purification of Hpn**

A saturated overnight culture of pET-hpn/*E. coli* BL21(DE3) was expanded (1:100) in LB medium supplemented with 100 μg/ml Amp, 0.5 mM NiSO₄, and 0.5% glucose, and was grown at 37°C until a *D₆₀₀* of 0.7 was reached. Protein expression was induced by the addition of IPTG to 0.5 mM, and the cultures were then incubated for a further 4 h at 37°C, before chilling to 4°C. Cells were harvested by centrifugation at 5000 g for 20 min at 4°C and were resuspended in 10 ml of ice-cold Buffer A (20 mM sodium phosphate buffer, pH 7.4, 500 mM NaCl, 100 mM imidazole and 1 mM PMSF) per litre of cell culture. Bacteria were disrupted by sonication in the presence of 1% (v/v) Triton X-100. The lysate was centrifuged at 30000 g for 30 min, and the supernatant was filtered through a 0.45-μm-pore-size cellulose acetate acetate syringe filter (Iwaki Glass) before loading on to a NiSO₄-impregnated HiTrap Chelating HP column (5 ml; Amersham Biosciences) that had been pre-equilibrated with 10 column vol. of Buffer A. After washing with 10 column vol. of Buffer A, Hpn protein was eluted with Buffer B (20 mM sodium phosphate buffer, pH 7.4, 500 mM NaCl and 500 mM imidazole). The purest fractions, as determined by SDS/PAGE (15% gel), were pooled, concentrated (Centricron YM-3; Millipore) and then purified further by gel-filtration chromatography on Superdex 75 resin (10/300 column; Amersham Biosciences) in buffer appropriate for subsequent use.

**Protein analysis**

Protein concentrations were determined by BCA (bicinchoninic acid) assay (Bio-Rad) using BSA as the standard. SDS/PAGE was performed according to the method of Maniatis et al. [27], and proteins were visualized with Coomassie Blue R-250.

Immunoblot analysis was used to determine the amount of Hpn present in the various purified Hpn fractions, using a standard curve constructed from Hpn solutions of known concentrations. Fractions were resolved on SDS/15% polyacrylamide gels then transferred on to PVDF membranes (Hybond-P; Amersham Biosciences). After unoccupied sites were blocked with 5% (v/v) non-fat dried milk in TTBS [Tris-buffered saline with Tween 20: 10 mM Tris/HCl, 150 mM NaCl, 0.01% (v/v) Tween 20], membranes were incubated with His·Tag® monoclonal antibody (1:1000 dilution) (Novagen) as recommended by the manufacturers. The membranes were then washed three times with TTBS, incubated with anti-mouse IgG conjugated to horseradish peroxidase, and developed using the ECL® (enhanced...
Characterization of a histidine-rich protein, Hpn

chemical blots (Amersham Biosciences). Image acquisition and analysis were performed on an ImageScanner (Amersham Biosciences) using ImageMaster 2D Elite software (Amersham Biosciences).

After resolution on an SDS/15% polyacrylamide gel and Coomassie Blue staining, the band corresponding to Hpn was excised and digested in-gel with chymotrypsin (Sigma), with overnight incubation at 37°C. Peptide fragments were then analysed by MALDI (matrix-assisted laser-desorption ionization)-MS.

To confirm the N-terminal sequence of the recombinant Hpn, the purified protein was subjected to N-terminal analysis by Edman degradation. Aliquots of Hpn (∼5 μg) was subjected to SDS/PAGE, transferred on to a PVDF membrane, and stained with 0.1% Coomassie Blue R-250, before excision and analysis with an automated HPG1000A liquid-phase protein sequencer (Hewlett-Packard).

The free thiolate content of Hpn was determined by the method of Ellman [28]. Briefly, Hpn protein was added to a 1 mM DTNB [5,5′-dithiobis-(2-nitrobenzoic acid)] solution in 20 mM Hepes, pH 7.4, and 100 mM NaCl, to final concentration of 10 μM. After incubation for 2 h at room temperature (25°C), the amount of generated p-nitrothiolate was determined using the molar absorption coefficient (ε412) of 13600 M⁻¹·cm⁻¹ [28].

Determination of the multimerization state of Hpn

To examine the effects of imidazole and DTT (dithiothreitol), as well as Ni²⁺ binding, on the multimerization state of Hpn, proteins (50 μM) were injected into a Superose 12 (10/300 column) (Amersham Biosciences) gel-filtration column, using running buffers containing 20 mM Hepes, pH 7.4, and 500 mM NaCl supplemented with 100 or 500 mM imidazole, 5 mM DTT or 250 μM Ni²⁺ (as NiSO₄) when needed. The proteins were equilibrated in the respective running buffer for at least 1 h before injection into the column. The column was calibrated with vitamin B₁₂ (1.355 kDa), lysozyme (14.3 kDa), β-lactoglobulin (35 kDa), BSA (66 kDa), human transferrin (80 kDa) and IgG (150 kDa). Blue Dextrin was used to determine the void volume. Molecular masses were determined by plotting log molecular masses of the standards against the partition coefficient (Kᵥ), where Kᵥ = (Vₑ - Vᵥ)/Vᵥ, where Vₑ represents elution volume, Vᵥ is the void volume, and Vₚ is the total column volume. The aggregation state of recombinant Hpn protein was also determined by non-denaturing PAGE, according to the method of Coligan et al. [29], with high-molecular-mass native markers (Amersham Biosciences) and stained using Coomassie Blue R-250.

UV/vis (visible) spectroscopy

Unless otherwise indicated, UV/vis absorption measurements were performed on samples of 50 μM proteins in 20 mM Hepes, pH 7.4, and 100 mM NaCl, using a Varian Cary 3E UV/vis spectrophotometer. Spectra were recorded over the wavelength range 230–650 nm in cuvettes with a pathlength of 1 cm.

CD spectroscopy

CD measurements were measured on a Jasco J-720 spectro-polarimeter calibrated with a 0.06% (+)-10-camphorsulfonic acid solution (Sigma–Aldrich), using a 0.1-cm-pathlength quartz cell. CD data were obtained for 5 μM Hpn in Tris/Cl buffer, pH 7.4, in both the absence and the presence of 1–5 molar equivalents of Ni²⁺ (as NiSO₄). Instrument optics and sample chamber were continually flushed with 3 litres/min of dry N₂ gas. Instrument settings used were: scan range, 190–250 nm; scan rate, 20 nm/min; wavelength step, 0.25 nm; sensitivity, 5 millidegrees; response time, 8 s. The final spectrum, representing an average of six scans, was corrected for the corresponding buffer and smoothed using adjacent averaging or an FFT (fast Fourier transform) filter. Quantitative estimations of the secondary-structure contents were made using the CDPro software package [30].

Influence of Ni²⁺ and Hpn on the growth of E. coli

(i) E. coli BL21(DE3) cultures, both with and without the plasmid pET-hpn (inoculated from individual colonies), were grown in LB medium (containing 100 μg/ml Amp when required) at 37°C. Stationary-phase overnight cultures (1 ml) were added to 50 ml of fresh medium supplemented with 500 μM NiSO₄, and were incubated for 24 h at 37°C, with IPTG added where applicable (to a final concentration of 500 μM) when the D₆₀₀ was approx. 0.6. Cell densities (determined by D₆₀₀) were measured after 0, 2, 4, 8, 16 and 24 h.

(ii) E. coli BL21(DE3) overnight cultures (as above, 10 μl) were used to inoculate 1 ml of fresh medium containing 0, 50, 100, 200, 500, 600, 700 or 800 μM NiSO₄ (as indicated). Cultures were grown for 18 h at 37°C, with IPTG added where applicable (to 500 μM), before the final cell density (determined by D₆₀₀) was measured.

Ni²⁺ accumulation in E. coli

E. coli BL21(DE3) overnight cultures (as above, 10 μl) were added to 1 ml of fresh medium supplemented with 500 μM NiSO₄. Cultures were grown for 18 h at 37°C, with IPTG (1 M) added when required (to a final concentration of 0.5 mM) when the D₆₀₀ reached approx. 0.6. Cells were pelleted at 4000 g for 10 min and washed three times with PBS, and the cellular Ni²⁺ content was determined by ICP-MS (inductively coupled plasma MS) (Agilent 7500) with 207Tl as an internal reference.

Ni²⁺ accumulation in H. pylori

Wild-type or Δhpn mutant H. pylori 26695 cells from fresh plates were resuspended in 10 ml of Brucella broth supplemented with 5% foetal bovine serum and 0.2 mM NiSO₄, and were grown for 48 h at 37°C. Cells were pelleted at 4000 g for 10 min and washed three times with PBS. The cellular Ni²⁺ content was then determined by ICP-MS similarly. All of the accumulation experiments were carried out with three replicates and an average value was used.

Ni²⁺ binding to Hpn

Binding of Ni²⁺ to Hpn was determined by equilibrium dialysis. The dialysis tubes (molecular-mass cut-off of 1 kDa), filled with 200 μl of protein suspension (5 μM), were placed into 1000 ml of 20 mM Hepes and 100 mM NaCl, pH 7.4, supplemented with a series of concentrations of Ni²⁺ from 0 to 100 μM, for overnight incubation at 4°C. The Ni²⁺ concentrations inside and outside the dialysis tubes were determined by ICP-MS. Each assay was repeated in triplicate. The results were subjected to Hill analysis for the determination of the stoichiometry and the binding affinity.

Alternatively, 15 molar equivalents of Ni²⁺ (as NiSO₄) were added to 5 μM protein in 20 mM Hepes at pH 7.4. After more than 1 h, the mixture was loaded on to a PD-10 desalting column (Amersham Biosciences) to remove excess Ni²⁺ ions. The fraction containing the Hpn–Ni²⁺ complex was collected, and the Ni²⁺ concentration was determined by ICP-MS.
Table 1  Purification of expressed Hpn

<table>
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<th>Fraction</th>
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<th>Hpn (mg)</th>
<th>Hpn in fraction (%)</th>
<th>Fold purification</th>
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<td>0.05</td>
<td>1</td>
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<tr>
<td>Crude supernatant</td>
<td>3279</td>
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<td>1.6</td>
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<td>2.3</td>
<td>85</td>
<td>1700</td>
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<tr>
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<td>1.4</td>
<td>&gt; 97*</td>
<td>1940</td>
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</tbody>
</table>

* As determined from a Coomassie Blue-stained gel.

RESULTS

Expression and purification of Hpn

The hpn gene (HP1427) was amplified by PCR from genomic H. pylori DNA and inserted into pET32a, via the NdeI/XhoI restriction sites, for high-level expression in E. coli. No fusion tag was included as we predicted that the separate stretches of six and seven consecutive histidine residues within the Hpn protein would result in strong binding to Ni2+-charged iminodiacetate-derivatized agarose (HiTrap chelating Sepharose). DNA sequencing revealed that the cloned hpn gene contained several (silent) mutations compared with the published sequence (GenBank® accession number U26361), but the translated product remained unchanged. It was found that adding IPTG to 0.5 mM at a D600 of 0.6–0.8, followed by incubation at 37 °C for 4 h, was optimal for the expression of soluble protein from pET-Hpn in E. coli BL21(DE3). Under these conditions, generally 2–3 mg of soluble Hpn protein was expressed per litre of cell culture. Hpn bound strongly to the Ni-charged chelating Sepharose resin in the presence of 100 mM imidazole, and could be eluted by 500 mM imidazole.

To confirm the identity of the purified recombinant Hpn protein, we performed Western blotting, N-terminal protein sequence analysis and MALDI–TOF (time-of-flight) mass spectrometric analysis. Based on the presence of six and seven consecutive histidine residues in the Hpn protein sequence, we predicted that it would bind with the His·Tag® monoclonal antibody with good affinity. Western blotting experiments confirmed this prediction, with a single intense band appearing on the PVDF membrane (Figure 2A), corresponding to the migratory position of the Hpn protein on a 15% polyacrylamide gel (as a weak band; results not shown).

To perform N-terminal sequencing, Hpn was purified by HiTrap Chelating HP to approx. 95–97% pure by Coomassie Blue staining (Table 1). Upon storage for several days at 4 °C in gel-filtration buffer (which lacked imidazole), the Hpn protein progressively aggregated and precipitated. This could be prevented by the addition of imidazole (500 mM), which increased protein stabilities in solutions to over 1 month (as determined by SDS/PAGE).

To confirm the identity of the purified recombinant Hpn protein, we performed Western blotting, N-terminal protein sequence analysis and MALDI–TOF (time-of-flight) mass spectrometric analysis. Based on the presence of six and seven consecutive histidine residues in the Hpn protein sequence, we predicted that it would bind with the His·Tag® monoclonal antibody with good affinity. Western blotting experiments confirmed this prediction, with a single intense band appearing on the PVDF membrane (Figure 2A), corresponding to the migratory position of the Hpn protein on a 15% polyacrylamide gel (as a weak band; results not shown).

To perform N-terminal sequencing, Hpn was purified by HiTrap Chelating HP to approx. 95% homogeneity, resolved further by SDS/PAGE, and then electroblotted on to a PVDF membrane to isolate homogeneous protein for N-terminal sequencing. The first seven amino acids were determined to be Ala-His-His-Glu-Glu-Gln-His. This revealed that the N-terminal methionine residue had been removed co- or post-translationally in E. coli. MALDI–MS spectrometry of the Hpn digested in-gel with chymotrypsin revealed a major peak at m/z 2773.183 (Figure 2D). This value agrees very well with the mass of m/z 2773.178 for the peptide AHHEEQHGHHHHHHTHHHHHY of Hpn as predicted by analysis with MS-digest software (http://prospector.ucsf.edu/).

![Image](354x462 to 525x724)

Figure 2  Analysis of expressed Hpn protein

(A) Western blotting with His·Tag® antibody. The arrow indicates the position of the recombinant Hpn protein. (B) Purified Hpn subjected to SDS/20% PAGE. The left-hand lane contains the protein markers (sizes are given in kDa), and the right-hand lane contains purified Hpn protein, which migrates to approx. 7 kDa. (C) Purified Hpn subjected to SDS/15% PAGE. The left-hand lane contains the protein markers (sizes are given in kDa), and the right-hand lane contains purified Hpn protein, which migrates to approx. 16 kDa. (D) MALDI-MS spectrum of Hpn showing the major peak at m/z 2773.183 corresponding to the N-terminal fragment of recombinant Hpn.

Analysis using Ellman’s reagent revealed that there were 3.7 thiolate groups per monomer, indicating that four cysteines in Hpn were in reduced form.

Molecular size

As was found previously [23], on standard SDS/10–15% polyacrylamide gels, Hpn migrated with an apparent molecular mass of approx. 16 kDa (Figure 2C), which is considerably larger than its predicted molecular mass of approx. 7 kDa. This is consistent with the properties of other histidine-rich proteins, which also exhibit retarded migration on SDS/polyacrylamide gels [32]. However, when SDS/20% polyacrylamide gels were used, Hpn migrated with an apparent molecular mass of approx. 7 kDa (Figure 2B), which corresponds closely to its predicted monomeric mass (7077 Da).

The native molecular mass of the protein in the presence of imidazole, DTT and Ni2+ was estimated by gel-filtration chromatography. As shown in the elution profile (Figure 3A), in the presence of 100 mM imidazole, a large fraction of the purified recombinant Hpn formed a high-molecular-mass aggregate of > 500 kDa (eluted at approx. 8.1 ml). There were six other major species observable, corresponding to molecular masses of 55, 34, 26, 20, 14 and 7 kDa, suggesting that Hpn was present in a range of multimeric forms. However, when DTT was added to the Hpn solution to a final concentration of 5 mM, before loading on to the Superose 12 column, the relative amount of the high-molecular-mass aggregate (eluted at approx. 8.1 ml) decreased significantly (Figure 3B), and the majority of the protein eluted
Figure 3 Analysis of Hpn multimeric state using gel-filtration chromatography

Elution profile (monitoring at 280 nm) of Hpn after gel filtration on Superose 12 (10/300) column, using running buffers containing 20 mM Hepes, pH 7.4, and 500 mM NaCl, supplemented with (A) 100 mM imidazole, (B) 100 mM imidazole plus 5 mM DTT, or (C) 500 mM imidazole plus 5 mM DTT. Proteins were equilibrated in the respective running buffer for at least 1 h immediately before analysis.

with an apparent molecular mass of 136 kDa (eluted at approx. 10.4 ml), with two very minor peaks at 55 and 26 kDa. When the eluted fraction corresponding to 136 kDa was re-injected into the gel-filtration column under the same conditions, a similar elution profile was observed (results not shown), suggesting that an equilibrium exists between multimeric forms. When the amount of imidazole was increased to 500 mM in the presence of 5 mM DTT, this fraction (approx. 136 kDa) become the dominant species with little high-molecular-mass aggregate (approx. 8.1 ml of eluate) (Figure 3C). When analysed on a SDS/15% polyacrylamide gel, all of the separately collected fractions migrated with an similar molecular mass of approx. 16 kDa (as in Figure 2C), showing that they are the same protein with different multimerization states.

Ni\textsuperscript{2+}–Hpn interaction

As shown in Figure 3, Hpn was present mainly as a 136 kDa multimer in buffers containing 500 mM imidazole and 5 mM DTT; otherwise, it will aggregate to over 500 kDa (Supplementary Figure S1 at http://www.BiochemJ.org/bj/393/bj3930285add.htm). It is impossible to investigate the Ni\textsuperscript{2+} binding in the presence of high concentrations of imidazole and DTT because of their competitive metal-binding capabilities. Therefore we studied the ‘effective’ Ni\textsuperscript{2+}–binding mode under physiologically relevant conditions (without imidazole). The choice of these experimental conditions was supported further by the findings that Hpn (2% of the total proteins of \textit{H. pylori}) was absent in the two-dimensional gels of \textit{H. pylori} cellular protein fractions (R. Ge, Q.-Y. He, J.-D. Huang and H. Sun, unpublished work), indicating that Hpn was unlikely to be present as a monomer inside the \textit{H. pylori} cells.

The binding of Ni\textsuperscript{2+} with apo-Hpn was studied by ICP-MS. It was found that freshly prepared Hpn contained fewer than one Ni\textsuperscript{2+} ion per ten Hpn molecules (results not shown). To a solution of Hpn (50 \mu M, in 20 mM Hepes buffer at pH 7.4), 15 molar equivalents of Ni\textsuperscript{2+} (as NiSO\textsubscript{4}) were added, then the excess Ni\textsuperscript{2+} was removed by desalting on a PD-10 column. The Hpn-bound Ni\textsuperscript{2+} was quantified by comparing the amount of Ni\textsuperscript{2+} present in the protein fraction with a standard curve (results not shown). This revealed that 5.1 ± 0.2 (n = 3, mean ± S.D.) Ni\textsuperscript{2+} ions were present per Hpn monomer. In order to examine whether free cysteine residues in Hpn were involved in the Ni\textsuperscript{2+} binding, the Ni\textsuperscript{2+}-bound Hpn was subjected to the method of Ellman [28], and the number of free thiolate groups was determined to be 0.4, indicating that almost all the four cysteine residues took part in the Ni\textsuperscript{2+} binding.

UV/vis spectroscopy was used to characterize further the Ni\textsuperscript{2+}–Hpn interaction. Complexation of Ni\textsuperscript{2+} to proteins normally leads to the production of new absorptions in the visible region due to LMCT (ligand-to-metal charge transfer) transitions [33]. Consequently, these new absorption peaks can be readily used to monitor protein–metal binding. The UV/vis absorption spectrum of a 50 \mu M Hpn solution was recorded (Figure 4A). After the sequential addition of different molar equivalents of Ni\textsuperscript{2+} to the purified protein solution, two shoulder peaks gradually appeared, centred at approx. 286 and 335 nm, indicative of Ni\textsuperscript{2+} binding (Figure 4A). The band at 335 nm was attributed to a ligand...
[e.g. N(His)] → Ni^{2+} LMCT transition [34]. A plot of ΔΔθ_{286} against the molar ratio (r) of Ni^{2+} (as NiSO_{4}) to the apo-protein is also shown in Supplementary Figure S2 (http://www.BiochemJ.org/bj/393/bj3930285add.htm). It can be seen that with the increase of r, the absorption at 286 nm (and 335 nm) increases in intensity and reaches a plateau at r ≈ 4.8 ± 0.2, which suggests that five Ni^{2+} ions bind strongly per Hpn monomer.

We investigated how the intensity of the absorption at 335 nm altered as the pH of a Ni^{2+}-saturated Hpn solution was systematically varied. As can be seen in the inset of Figure 4(A), there were no significant changes in the absorption at 335 nm at pH values greater than approx. 6.8. This indicated that the amount of Ni^{2+} bound to the protein remained relatively stable above this pH value. However, upon lowering the pH of the solution from approx. pH 6.8 to 5.8 (by the careful addition of 1 M HCl), the intensity of the absorption at 335 nm decreased markedly. At pH values lower than approx. 5.0, the absorption at 335 nm remained essentially unchanged. The pH for half dissociation of Ni^{2+} from the protein complex (pH_{1/2}) was determined to be 6.3 ± 0.1.

A competition binding assay between Hpn and EDTA for free Ni^{2+} was performed analogously. Upon the addition of a large excess of EDTA (120 molar equivalents) to a Ni^{2+}-saturated Hpn solution at pH 6.8, the absorption at 335 nm (characteristic of an LMCT) decreased exponentially with a half-life of 41 ± 1 min (Figure 4B). However, this peak was restored to original levels upon the addition of excess Ni^{2+} ions, demonstrating that Ni^{2+} binding is reversible (results not shown).

To determine the Ni^{2+}-binding affinity and capacity of the protein, the pure protein was dialysed against 20 mM Hepes buffer (pH 7.4) with different amount of Ni^{2+}. As shown in Figure 5, at the concentration used (5 μM), the binding of Ni^{2+} to Hpn reached saturation at 40 μM, and the protein was able to bind to 4.8 ± 0.1 Ni^{2+} ions per monomer in a slightly co-operative fashion (Hill coefficient of approx. 1.7). The dissociation constant (K_{d}) was calculated to be 7.1 μM.

The CD spectrum of Hpn in 20 mM Tris/HCl at pH 7.4 (Figure 6) is characterized by a negative peak with a maximum at approx. 217 nm and a positive peak with a maximum at 197 nm, attributed to the presence of a mixture of α-helix, β-sheet, turn and unordered form. Upon the addition of different amounts of molar equivalents of Ni^{2+} ions (as NiSO_{4}) to the apo-protein solution, the negative peak at 217 nm increased gradually, indicative of a conformational change within the protein induced by the metals. Quantitative analysis using CDPro [30] indicated that α-helix content was reduced from approx. 20% to 10%, whereas the β-structure increased from 22% to 37% after addition of 5 molar equivalents of Ni^{2+}.

Figure 5 Binding of Ni^{2+} to recombinant Hpn as determined by equilibrium dialysis

Hpn (5 μM, 200 μl) was dialysed overnight at 4°C against 1000 ml of 20 mM Hepes, pH 7.4, and 100 mM NaCl containing 0, 2, 5, 10, 20, 30, 40, 60 or 100 μM NiSO_{4}. Hpn-bound Ni^{2+} was determined by ICP-MS (subtracting Ni^{2+} concentrations inside and outside the dialysis tubing). Shown is a Hill plot of the molar ratio (r) of bound Ni^{2+} to Hpn protein against concentration of NiSO_{4} in the dialysis buffer. Each point represents the average value of three independent experiments.

Figure 6 Binding of Ni^{2+} to recombinant Hpn as determined by CD

CD data were obtained for 5 μM Hpn in Tris/HCl buffer, pH 7.4, in both the absence and the presence of 1–5 molar equivalents of Ni^{2+} (as NiSO_{4}). The arrow indicates the increases in ellipticity values (approx. 217 nm) as the protein was titrated with 0–5 molar equivalents of Ni^{2+}.

Figure 7 Effects of Ni^{2+} on growth of E. coli and Ni^{2+} accumulation in E. coli and in H. pylori

(A) Growth curves of freshly inoculated cultures of E. coli BL21(DE3) both with and without the hpn gene (pET-hpn) in LB medium containing Ni^{2+} (as NiSO_{4}, 500 μM). (B) The influence of Hpn expression on the growth of E. coli BL21(DE3) cells cultured in LB medium supplemented with NiSO_{4} at 0, 50, 100, 200, 300 or 500 μM. ▲ + pET-hpn, with IPTG induction (0.5 mM); ▲ + pET-hpn, no IPTG induction; ■ − pET-hpn, with IPTG induction (0.5 mM); ■ − pET-hpn, no IPTG induction. (C) E. coli BL21(DE3) cells were cultured in LB medium supplemented with 500 μM NiSO_{4} with or without the plasmid pET-Hpn, and with or without IPTG induction of Hpn expression. (D) The wild-type and Hpn-deleted H. pylori 26695 cells were cultured in fresh Brucella broth supplemented with 5% fetal bovine serum and 200 μM NiSO_{4}. The Ni^{2+} content associated with the cell pellet was determined using ICP-MS.

Effect of Ni^{2+} on the growth of E. coli

E. coli BL21(DE3) cells with and without the plasmid pET-hpn were grown in LB medium supplemented with different amounts of NiSO_{4}. The growth at 37°C in the presence of 500 μM Ni^{2+}, as a representative, was followed by the measurement of D_{600} values (Figure 7A). The D_{600} values for the E. coli BL21(DE3) cells with hpn were about twice those of cells without hpn after 8 h. In addition, the D_{600} values for the E. coli BL21(DE3)
cells with hpn were slightly higher upon addition of IPTG than those without (Figure 7B). In contrast with the very similar \( D_{600} \) values for the *E. coli* BL21(DE3) cells with hpn grown at \([\text{Ni}^{2+}]\) ≤ 500 \( \mu \)M, the \( D_{600} \) values for the *E. coli* BL21(DE3) cells without hpn fell significantly at \([\text{Ni}^{2+}] \geq 200 \, \mu \text{M} \). More importantly, the difference in \( D_{600} \) values (overnight culture) for *E. coli* BL21(DE3) cells harbouring hpn became more evident with \( \text{Ni}^{2+} \) concentrations increased to 500 \( \mu \text{M} \) or over and without IPTG, i.e. the \( D_{600} \) values were higher for those cells with IPTG than for those without.

**Cellular \( \text{Ni}^{2+} \) accumulation**

To comply with the conditions used in the expression of Hpn, the cells treated with 0.5 \( \text{mM} \) \( \text{Ni}^{2+} \) were pelleted and washed three times with PBS, and the cellular \( \text{Ni}^{2+} \) content was determined by ICP-MS. In *E. coli* BL21(DE3) harbouring pET-hpn where Hpn expression had been induced with IPTG, the \( \text{Ni}^{2+} \) ion content was found to be approx. 4-fold higher than in un-induced cells, or in cells that did not contain pET-hpn (Figure 7C). The cellular accumulation of \( \text{Ni}^{2+} \) was then investigated in cultures of *H. pylori* 26695 which contained a wild-type copy of the gene, and in constructed strains where the *hpn* gene had been deleted. The deletion of the *hpn* gene was confirmed by PCR (Supplementary Figure S3 at http://www.BiochemJ.org/bj/393/bj3930285add.htm), and by the disappearance of the Hpn protein from cell lysates, using Western blot analysis, probing with the His\(^{\text{Tag}}\) monoclonal antibody. After 48 h of culture in medium containing 0.2 \( \text{mM} \) \( \text{NiSO}_{4} \), ICP-MS analysis indicated that intracellular levels of \( \text{Ni}^{2+} \) were 1.5-fold higher in the wild-type strain than in the \( \Delta hpn \) strain (Figure 7D). In addition, the accumulation of \( \text{Ni}^{2+} \) in *H. pylori* 26695 is at least 2-fold greater than that in *E. coli*, on a gram-for-gram basis (wet cell mass).

**DISCUSSION**

The transition metal \( \text{Ni}^{2+} \) is essential for anaerobic metabolism in micro-organisms, including the medically important Gram-negative bacterium *H. pylori*. However, too much ‘free’ or un-bound \( \text{Ni}^{2+} \) can be toxic and therefore intracellular concentrations must be regulated tightly [10,13]. Initial studies have indicated that the small cytoplasmic Hpn protein may play a pivotal role in \( \text{Ni}^{2+} \) homeostasis, binding or transport [23], and Hpn was named to emphasize its origins in *H. pylori* and its affinity for \( \text{Ni}^{2+} \) [23]. Although the specific amino acids essential for metal binding remain to be established, several conserved histidine, glutamate and aspartate residues stand out as potential metal ligands. All four cysteine residues of Hpn were found to be involved in \( \text{Ni}^{2+} \) binding. It is possible that the two neighbouring double cysteines seem to be possible to take part in the coordination with the same \( \text{Ni}^{2+} \) ion. The motif Asp-Xaa-His-His-Xaa-Xaa-Glu (or DXHHXHXE) is a potential metal-binding site, and is conserved in approx. 300 proteins listed in the SwissProt database ([35]; http://www.expasy.org/sprot/). A significant proportion of these proteins are histidine-rich and have been implicated in metal binding, such as UreE [36], IGPD (imidazole glycerol-phosphate dehydratase) [37], HspA and HypB [38,46]. Another conserved pentapeptide motif: His-Glu-Xaa-Xaa-His (or HEXHX) appears regularly in protein sequences, especially in metalloproteases. Roughly half of all metalloprotease characterized to date contain this motif, which has been implicated in metal binding in X-ray crystal structures [39]. Low-stringency homologous sequence searching using various BLAST programs [40] failed to find any meaningful protein matches to Hpn, the only ‘hits’ containing ubiquitious short histidine-rich motifs (results not shown). Consequently, the structural fold and organization of Hpn cannot presently be inferred by homology.

The *hpn* gene was cloned into a pET32 vector, and was successfully expressed (untagged) in a predominantly soluble form in *E. coli*. Approx. 77% of the amino acid residues in Hpn are predicted to be > 16% solvent-exposed using Predict Protein ([41]; http://cubic.bioc.columbia.edu/predictprotein/), indicating that the protein should be highly hydrophilic. Denaturing conditions were not required for Hpn binding to \( \text{Ni}^{2+} \)-impregnated Sepharose resin, suggesting that enough contiguous histidine residues are solvent-exposed under the conditions used to be readily available either for chelation with the immobilized metal ions, or possibly for protein–protein interactions. This is congruent with the predicted hydrophilicity of the histidine-rich regions by PredictProtein (results not shown).

After purifying milligram quantities of recombinant Hpn, we were able to investigate in detail its multimeric form, conformation and Ni-binding properties in *vitro*. On native gels, Hpn migrated as a single band with an apparent molecular mass of > 500 \( \text{kDa} \) (Supplementary Figure S1 at http://www.BiochemJ.org/bj/393/bj3930285add.htm). However, it eluted as a mixture of species after gel-filtration chromatography on a Superose 12 column in a buffer containing 20 \( \text{mM} \) Heps, pH 7.4, 500 \( \text{mM} \) NaCl and 100 \( \text{mM} \) imidazole buffer (Figure 3A). Upon addition of 5 \( \text{mM} \) DTT to the running buffer, the relative amounts of the high-molecular-mass aggregate dramatically decreased, indicative of the presence of an intermolecular oxidation. The protein fractions with apparent molecular masses of approx. 55 and approx. 136 \( \text{kDa} \) correspond to a ‘dimer’ and ‘pentamer’ of tetramers respectively. It is possible that the 20 (-2)-mer Hpn multimer may be the physiologically relevant form of the protein in *vitro*. In the presence of higher amounts of imidazole, the relative amount of the high-molecular-mass aggregate decreased further, indicating that imidazole plays a role in the prevention of protein precipitation and restoring the preferred multimerization state.

A number of other histidine-rich proteins have been reported to exist as multimers (Supplementary Table S1 at http://www. BiochemJ.org/bj/393/bj3930285add.htm) [32,37,42–46]. Based on these data, it seems that the higher the percentage of histidine in a metal-binding protein, the greater the propensity of the protein to form higher-order multimers. One explanation for this is that histidine residues located on different protein monomers might facilitate protein multimerization in some way, possibly through their tendency to align or ‘stack’ on top of one another. It is conceivable that the disappearance of the low-molecular-mass Hpn band (as determined by SDS/PAGE) after storage for several days in buffers lacking imidazole may be due to the formation of a high-molecular-mass aggregate that is large enough to precipitate. Conversely, when present in concentrations of no less than 500 \( \text{mM} \), imidazole in solution may locate between the protein monomers, inhibiting intermolecular histidine stacking, consequently stabilizing the protein. However, we cannot exclude the possibility that other biophysical processes are behind multimer formation/disassociation.

We have found that Hpn can bind up to five \( \text{Ni}^{2+} \) ions per monomer under physiologically relevant conditions (i.e. at pH 7.4). More importantly, this binding is reversible, as revealed by both pH titration experiments and competition binding experiments with EDTA (Figure 4). This result raises the intriguing possibility that Hpn releases bound \( \text{Ni}^{2+} \) ions when the environmental pH decreases to approx. 6.8, a physiological pH value. The moderate affinity (\( K_{a} \) of approx. 7.1 \( \mu \text{M} \)) of Hpn for \( \text{Ni}^{2+} \) is comparable with that of HypB (\( K_{a} \) of approx. 2.3 \( \mu \text{M} \)), a protein that has been suggested to store \( \text{Ni}^{2+} \) in its N-terminal histidine-rich region (16 histidine residues out of 54 residues) in

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**Bradyrhizobium japonicum** [46]. In this manner, *H. pylori* may buffer itself against a potentially damaging fall in the environmental pH. As shown in a previous report [23], there appears to be no direct interaction between Hpn and urease, since *hpn* gene deletion has little affect on the urease activity. It is possible that Hpn may donate Ni^{2+} to a chaperone protein (e.g. HspA) with higher affinity, which would transport the Ni^{2+} ions to other proteins and ultimately to a Ni^{2+}-containing enzyme [21]. The presence of other functions of the protein is also possible and needs to be studied further.

Interestingly, Ni^{2+} release from the protein appeared to occur in a single process, although one may have expected five micro-steps (generating micro pk_{s} values). This may be due to the high kinetic liability of Ni^{2+} binding. A similar kinetic profile has been observed for Zn^{2+} (and Cd^{2+}) release from MT (metallothionein; Zn_{4}−MT), i.e. no more than two steps were observed [47]. A facile Ni^{2+}−to-ligand exchange mechanism would facilitate intraprotein metal transfer, possibly assisting rapid metal trafficking to and from other proteins or enzymes.

The appearance of absorption peaks at 286 and 335 nm upon the addition of Ni^{2+} to apo-Hpn indicates that the Ni^{2+} is probably co-ordinated to nitrogen (and thiolate) centres, the majority presumably from the imidazole moieties of histidine residues. Similar absorptions have been observed in other Ni^{2+}-binding proteins such as NikiR, a protein known to bind to Ni^{2+} via three histidine nitrogens and one cysteine [33,48]. These UV/vis data also indicated that Ni^{2+} co-ordinates to ligands in either a 4-coordinated diamagnetic square-planar or a distorted square-planar geometry [10,34], consistent with silent EPR signals of the Ni^{2+}-saturated Hpn (results not shown), which suggests that the Ni^{2+} ions may be in low-spin states. Further studies are required to firmly establish the co-ordination geometry of Ni^{2+}-bound Hpn.

Similar to the situation in other metal-binding proteins, CD spectroscopy revealed that the binding of Ni^{2+} ions induced a protein conformational change in Hpn, with the α-helical content reduced from approx. 20% to 10%, and β-sheet content increased from approx. 22% to 37% (Figure 6). We are currently investigating the biological and biophysical significance of this change in protein structure.

In *E. coli* BL21(DE3) cells where the expression of the plasmid-based *hpn* gene had been induced, clearly Ni^{2+} had less influence on the cell growth than in those cells without *hpn*, based on the ΔOD values (Figures 7A and 7B). The higher ΔOD values for *E. coli* BL21(DE3) cells with the *hpn* gene in the presence of IPTG than those in the absence of IPTG indicated that Hpn may have a protective role against excess Ni^{2+} (and probably other heavy metals) in the environment. A similar pattern was also observed in *H. pylori* previously [25]. These data are in agreement with Ni^{2+} accumulation by ICP-MS. Ni^{2+} accumulation from the extracellular milieu was approx. 4-fold higher than in uninduced cultures, or in cells that did not contain the plasmid (Figure 7C). An analogous Ni^{2+}-accumulation pattern was observed in *H. pylori* cultures under similar conditions. As the transmembrane metal transport machineries in *E. coli* and *H. pylori* are quite distinct, this suggests that Hpn may promote the gradual intracellular accumulation of Ni^{2+} through a ‘passive’ (equilibrium-driven) sequestration mechanism. The slight difference in Ni^{2+} accumulation between *H. pylori* and *H. pylori* with the *hpn* gene deleted also suggests that some other proteins, such as Hpn-like protein, play a similar role in cells.

In view of its short length (60 amino acids) and extremely high content of potential metal-binding residues, Hpn shares some similarities with MT, a zinc-storage protein rich in cysteine residues [47]. MT has been known to play a fundamental role in the metabolism of zinc and copper under various physiological conditions and its binding capability to metals is comparable with that of Hpn towards Ni^{2+}. Moreover, MT also appears to play a role in sequestering toxic metals such as Cd^{2+} and Hg^{2+}. It will be of great interest to see whether future work reveals any structural or functional similarities between these two important metal-binding proteins. With its large multimeric quaternary structure, possibly based around an arrangement of tetramers, Hpn may be regarded as a ‘ferritin-like’ Ni^{2+} protein, which regulates levels of Ni^{2+} within the bacterium.

In conclusion, the 60-amino-acid exceptionally histidine-rich Hpn protein from *H. pylori* binds tightly and reversibly to Ni^{2+} ions in a pH-dependent manner. Hpn binds approx. five Ni^{2+} ions per monomer with a K_{D} of 7.1 μM, and adopts a variety of multimeric structures in solution, with the 20 (± 2)-mer form predominating under mildly reducing conditions. The growth and accumulation experiments performed under Ni^{2+}-pressure clearly imply a role for Hpn to protect the cells from higher concentrations of external metal ions. Given its high intracellular concentration in *H. pylori* (approx. 2 % of total protein), our data are consistent with Hpn functioning as an intracellular ‘reservoir’ of Ni^{2+} ions, playing a key role in bacterial Ni^{2+} homeostasis.

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Characterization of a histidine-rich protein, Hpn 293


31 Reference deleted
