Identification of *rcnA* (*yohM*), a Nickel and Cobalt Resistance Gene in *Escherichia coli*

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We report here on the isolation and primary characterization of the *yohM* gene of *Escherichia coli*. We show that *yohM* encodes a membrane-bound polypeptide conferring increased nickel and cobalt resistance in *E. coli*. *yohM* was specifically induced by nickel or cobalt but not by cadmium, zinc, or copper. Mutation of *yohM* increased the accumulation of nickel inside the cell, whereas cells harboring *yohM* in multicopy displayed reduced intracellular nickel content. Our data support the hypothesis that YohM is the first described efflux system for nickel and cobalt in *E. coli*. We propose *rcnA* (resistance to cobalt and nickel) as the new denomination of *yohM*.

Nickel and cobalt are both required as trace elements in prokaryotes to fulfill a variety of metabolic functions, but high intracellular concentrations of these transition metals are toxic. One of the strategies evolved by bacteria to prevent damage is to export excess metal by efflux systems. Plasmid-borne determinants responsible for nickel and/or cobalt resistance have been described for the heavy-metal-resistant bacterium *Ralstonia metallidurans* (11, 15), among which are members of the resistance-nodulation-cell division superfamily: the best-characterized CzcCBA (cobalt-zinc-cadmium) three-component cation antipporter (14) and the homologous CnrCBA (cobalt-nickel resistance) (10) and NccCBA (nickel-cobalt-cadmium resistance) (18) efflux systems. Moreover, cobalt can be extruded from the cytoplasm by the cation diffusion facilitator CzcD of *R. metallidurans* at the expense of the proton motive force or a potassium gradient (15). Cobalt may also be a substrate of Zn-CpfX-type ATPases, as in *Helicobacter pylori* (8). There is no evidence for the transport of nickel by one of these two modes of efflux. Instead, this metal can be transported outside the cytoplasm by NreB from *R. metallidurans* (7) or NrsD from *Synechocystis* sp. strain PCC 6803 (6), which are members of the major facilitator superfamily and which each exhibit 12 putative transmembrane helices and a histidine-rich carboxy terminus contributing to nickel resistance.

In *Escherichia coli*, anaerobic hydrogenase isoenzymes and urease (in ureolytic strains) require incorporation of nickel to become active (12). Complex assembly processes involve accessory proteins, namely, HypB, implicated in nickel insertion into hydrogenase, and UreE, which delivers nickel to urease. HypB and UreE are well conserved among bacteria apart from a terminal histidine-rich stretch whose function would be nickel storage and which is absent in *E. coli* proteins (3, 5). In order to gain insights into nickel trafficking and, more precisely, to find proteins that would be involved in nickel resistance, we searched the *E. coli* genome database with a query based on a consensus alignment of the UreE and HypB histidine-rich variants. The best returned hit was *yohM*, whose product bears a histidine-rich domain in its center. The aim of the present work is to demonstrate the implication of *yohM* in nickel and cobalt trafficking in *E. coli*.

**Inactivation of *yohM* confers sensitivity to nickel and cobalt.** The *yohM* gene was identified because its product contains a remarkable histidine-rich loop (see below and Fig. 4, top). The *yohM* gene is surrounded upstream by *yohL*, which is divergently transcribed, and downstream by *yohN* (Fig. 1). The whole region was amplified from MC4100 chromosomal DNA by using the youp and yodwn primers, and the resulting 2,003-bp BamHI-EcoRI fragment was cloned into a pUC18 vector (19), resulting in plasmid pAR123. A *yohM* insertion mutant, in which a *uidA*-Kan cassette derived from pU1DK11 (2) was inserted into *yohM* at the NsiI site, was then constructed. The transcriptional *yohM-uidA* fusion was recombined back to the chromosome of *recBC sbcBC* strain JC7623 (17) and was further moved into the wild-type (wt) strain MC4100 (laboratory collection) via P1 phage transduction to obtain strain ARY023. To perform complementation studies, *yohM* alone was cloned. For that purpose, a DNA fragment amplified with the youp and yoMdwn primers was digested by PstI and HindIII and then introduced into pUC18, resulting in pAR020.

To assess whether *yohM* could be responsible for some metal resistance in *E. coli* and to determine the nature of the metals to which *yohM* would be sensitive, a plate sensitivity assay was carried out. Wild-type strain MC4100 was sensitive to all of the tested metals except manganese (Table 1). Among them, only nickel and cobalt promoted increased growth inhibition for the *yohM* mutant ARY023, since the zone of inhibition was increased by 38% for nickel and 30% for cobalt in comparison with wt strain MC4100. Furthermore, when expressed in *trans* from the multicopy plasmid pAR020, *yohM* conferred a marked enhancement of nickel and cobalt resistance to the host mutant strain (two- to threefold). The presence of plasmid-borne *yohM* did not affect the response to the other tested metals. Thus, *yohM* is shown to be a nickel and cobalt resistance gene in *E. coli*.
Determination of nickel and cobalt MICs. The toxic effect of nickel and cobalt on the wt and mutant strains was further assayed by measuring the final optical density at 600 nm (OD$_{600}$) of a 12-h culture in minimal medium supplemented with NiCl$_2$ or CoCl$_2$. In agreement with the results of the metal sensitivity plate assay, the yohM strain displayed higher nickel and cobalt sensitivities (Fig. 2). More precisely, growth arrest occurred at 4 μM NiCl$_2$ and 30 μM CoCl$_2$ for ARY023 (yohM) compared with 10 μM NiCl$_2$ and 50 μM CoCl$_2$ for MC4100 (wt). Interestingly, in the mutant transformed with pAR020 (yohM$^-$), the wild-type resistance levels were not only recovered but greatly enhanced, as in this case, ARY023/pAR020 was able to resist nickel or cobalt concentrations 100-fold higher. This result might be explained by two parameters, the high number of copies of vector pUC18 harboring yohM and the participation of the pUC18 lac promoter in yohM transcription, given that in the recombinant plasmid, yohM and Plac are in the same orientation.

The yohM gene is induced by nickel and cobalt. To analyze the metal-dependent expression of yohM, the transcriptional yohM-uidA fusion carried by ARY023 was used. After 6 h of growth in rich medium in the absence or in the presence of 0.5 mM CuSO$_4$, ZnCl$_2$, CdCl$_2$, CoCl$_2$, or NiCl$_2$, β-glucuronidase activity was assayed. In the absence of added metals, there was almost no expression of the yohM-uidA fusion, as an activity of

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<tr>
<th>Metal</th>
<th>MC4100 (wt)</th>
<th>ARY023 (yohM)</th>
<th>ARY023/pAR020 (yohM$^-$)</th>
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<tbody>
<tr>
<td>Ni</td>
<td>16</td>
<td>22</td>
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<tr>
<td>Co</td>
<td>17</td>
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<td>Cu</td>
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<td>0</td>
</tr>
<tr>
<td>Cd</td>
<td>14</td>
<td>14.5</td>
<td>14</td>
</tr>
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*Cells (10$^8$) were spread on N medium glucose agar plates (16). The zone of inhibition (in millimeters) around a filter disk containing 20 μl of a 5 mM solution of either NiCl$_2$, CoCl$_2$, CuSO$_4$, ZnSO$_4$, MnSO$_4$, or CdCl$_2$ was measured after a 24-h aerobic incubation at 37°C. Similar results were obtained after 24 h at 37°C under anaerobic conditions (GasPak jar).
yohM-induced by nickel or cobalt and yohM is solely expressed when these metals are present, strongly suggesting that the function of yohM is to detoxify the cell with regard to nickel or cobalt.

**yohM encodes a nickel-cobalt efflux system in E. coli.** The preceding results have clearly shown that yohM confers increased resistance to nickel or cobalt in *E. coli*. This suggests that YohM can function as an efflux system. In such a case, the concentration of cytosolic metal ions should be higher in the sensitive cells (yohM than in the resistant cells (wt). Alternatively, resistance could be the result of a storage mechanism, i.e., binding of the metal by the histidine-rich loop, and this would result in an increased concentration of metal ions in the resistant cells. The in vivo activity of YohM was monitored by using a 63Ni uptake assay as described previously (13). In favor of the first hypothesis, the yohM mutant accumulated nearly twice the level of nickel as the wild type (Fig. 3, filled symbols). This finding was strongly reinforced by the measurements recorded from the sensitive strain complemented by yohM (open symbols). YohM was specifically expressed in the presence of 5 μM 63NiCl2 (filled symbols) or 5 μM 65NiCl2 and 50 μM CoCl2 (open symbols). Aliquots (100 μl) were filtered at the indicated times. The intracellular concentration of 63Ni per milligram of bacterial dry weight was determined.

**A striking feature is the existence of the histidine-rich loop in various proteins from different species of alpha, beta, and gamma proteobacteria, cyanobacteria, and archaea.** Indeed, the curve obtained for the wild-type strain in the competition assay reflected an increased intracellular nickel accumulation, which reached a level similar to that found in the yohM mutant (Fig. 3, open symbols). The latter displayed no significant change between the two assays. Likewise, for the yohM mutant harboring the plasmid-borne yohM gene, the curves obtained in the presence and absence of cobalt are nearly superimposable, indicating the huge efflux capacity of the overexpressed system.

**YohM is a histidine-rich membrane protein.** YohM is predicted to encode a 274-amino-acid protein with a molecular mass of 30,419 Da and to possess a remarkable histidine-rich region (amino acids 121 to 146) containing 17 histidines, 3 aspartates, and 3 glutamates out of 26 residues (Fig. 4, top panel). Using yohMup and yohMdown primers, yohM was cloned into the expression vector pET30 at the NdeI and HindIII sites, resulting in plasmid pAR02T. YohM was specifically labeled with 35S-methionine in vivo by use of the T7 RNA polymerase system and analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). One major band with an estimated mass of 32 kDa was observed in the crude extracts, in close agreement with the calculated mass (Fig. 4, bottom panel, lanes 1 and 4). The separation of crude extract into membrane and soluble fractions allowed us to assign 35S-Met-labeled YohM to the membrane fraction (Fig. 4, bottom panel, lanes 1 and 4). Topology prediction, using the TM-Pred program (9), predicted YohM to be an inner membrane protein encompassing six transmembrane domains (Fig. 4, top panel). However, the proper orientation of the protein remains to be elucidated, as in silico analysis provided no clear indication of the orientation of the histidine-rich loop, which could be either periplasmic or cytoplasmic. When used as a probe for a BLAST query against the nonredundant database, YohM showed highest similarities with uncharacterized putative proteins from different species of alpha, beta, and gamma proteobacteria, cyanobacteria, and archaea (Fig. 4, top panel). A striking feature is the existence of the histidine-rich loop in all of them, as the BLAST default settings exclude this form of repeated residues from the query because they are considered...
low-complexity segments; these sequences were not retrieved because they possess a histidine stretch. When looking for conserved regions which could serve as signatures of the YohM family, two motifs were defined (Fig. 4, top panel). Considering the residues known to be putative nickel or cobalt ligands and located outside the histidine-rich region, nine of them can be highlighted because of their strict or strong conservation. With regard to the YohM sequence, these residues are H27, H33, H63, H121, H123, H153, H157, D160, and C187. Interestingly, all of these residues are present in one or the other signature motif.

From a functional point of view, YohM resembles NrsD from *Synechocystis* sp. strain PCC 6803 (6) and NreB from *R. metallidurans* (7). Indeed, all are membrane-bound proteins which possess a histidine-rich domain. They belong to the major facilitator superfamily and are strongly suggested to be responsible for nickel resistance by an efflux mechanism. However, in contrast to NreB and NrsD, which are predicted to contain 12 transmembrane helices and histidine-rich C termini, from *Synechocystis* sp. strain PCC 6803 (6) and NreB from *R. metallidurans* (7). Indeed, all are membrane-bound proteins which possess a histidine-rich domain. They belong to the major facilitator superfamily and are strongly suggested to be responsible for nickel resistance by an efflux mechanism. However, in contrast to NreB and NrsD, which are predicted to contain 12 transmembrane helices and histidine-rich C termini,
YohM is supposed to contain 6 transmembrane segments and a histidine-rich domain located in the center of the polypeptide. Moreover, YohM transports cobalt in addition to nickel, which is the sole metal transported by NreB and NrsD. YohM and similar proteins can be very partially aligned with members of the nickel cobalt transporter family (NiCoT), which are nickel uptake permeases (4) (data not shown). Nevertheless, YohM does not harbor the NiCoT signature present in the second transmembrane helix of these eight-helix permeases; YohM thus does not belong to the NiCoT family.

In conclusion, YohM seems to be the first reported member of a new class of nickel and cobalt exporters, and we propose to assign it the new designation *rnnA*, corresponding to resistance to cobalt and nickel.

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REFERENCES