The Iron Transport Protein NRAMP2 Is an Integral Membrane Glycoprotein That Colocalizes with Transferrin in Recycling Endosomes

By Samantha Gruenheid, François Canonne-Hergaux, Susan Gauthier, David J. Hackam, Sergio Grinstein, and Philippe Gros

From the *Department of Biochemistry and Center for Host Resistance, McGill University, Montreal, Quebec, Canada H3G 1Y6; and ‡Division of Cell Biology, The Hospital for Sick Children, Toronto, Ontario, Canada M5G 1X8

Summary

The natural resistance associated macrophage protein (N ramp) gene family is composed of two members in mammals, N ramp1 and N ramp2. N ramp1 is expressed primarily in macrophages and mutations at this locus cause susceptibility to infectious diseases. N ramp2 has a much broader range of tissue expression and mutations at N ramp2 result in iron deficiency, indicating a role for N ramp2 in iron metabolism. To get further insight into the function and mechanism of action of N ramp proteins, we have generated isoform specific anti-N ramp1 and anti-N ramp2 antisera. Immunoblotting experiments indicate that N ramp2 is present in a number of cell types, including hemopoietic precursors, and is coexpressed with N ramp1 in primary macrophages and macrophage cell lines. N ramp2 is expressed as a 90–100-kD integral membrane protein extensively modified by glycosylation (>40% of molecular mass). Subcellular localization studies by immunofluorescence and confocal microscopy indicate distinct and nonoverlapping localization for N ramp1 and N ramp2. N ramp1 is expressed in the lysosomal compartment, whereas N ramp2 is not detectable in the lysosomes but is expressed primarily in recycling endosomes and also, to a lower extent, at the plasma membrane, colocalizing with transferrin. These findings suggest that N ramp2 plays a key role in the metabolism of transferrin-bound iron by transporting free Fe²⁺ across the endosomal membrane and into the cytoplasm.

Key words: iron • anemia • transport • infection • macrophage

Naturally occurring (1) or experimentally induced (2) mutations at the N ramp1 (natural resistance associated macrophage protein 1) locus in vivo impair macrophage function and cause susceptibility to infection by intracellular pathogens such as Salmonella, Leishmania, and M. tuberculosis in mice. In humans, polymorphic variants at N ramp1 are associated with increased susceptibility to tuberculosis and leprosy (3, 4). Studies in vitro in explanted cell populations have indicated that mutations at N ramp1 affect the ability of the macrophage to restrict the intracellular replication of antigenically unrelated microorganisms. We cloned the N ramp1 gene (5) and showed that its mRNA is expressed abundantly in macrophages (6) and in neutrophils (7) and is inducible in macrophages by exposure to cytokines and bacterial endotoxin (6). Predicted amino acid sequence analysis indicates that N ramp1 has many characteristics of an integral membrane transport protein including 12 putative transmembrane (TM) domains, several predicted N-linked glycosylation sites, and a sequence signature previously identified in a number of eukaryotic and prokaryotic transport proteins (5). In macrophages, direct biochemical studies have shown that N ramp1 is a membrane phosphoglycoprotein of apparent mass 90–110 kD (8), which is expressed in the Lamp1-positive lysosomal compartment (9). Moreover, studies in phagosomes containing either latex beads or intact bacteria have shown that upon phagocytosis, N ramp1 is recruited to the membrane of the phagosome, where it remains during its maturation to phagolysosome (9). These findings suggest that N ramp1 may affect resistance to infection by modulating the intravesicular milieu of the bacterial phagosome.

We have identified a second N ramp gene in mammals, N ramp2, which encodes a protein highly similar to N ramp1 (78% identity over the hydrophobic core) (10). As opposed to the phagocyte-specific expression of N ramp1, N ramp2 mRNA expression has been detected in most tissues and
cell types analyzed (10–12). Recently, it was shown that the N ramp gene is mutated (G185R) in two animal models of iron deficiency, the mk mouse (13) and the Belgrade rat (14). The mk mouse displays deficiency in intestinal iron uptake and microcytic anemia (15, 16). The Belgrade rat also shows a defect in intestinal iron absorption (17). Moreover, studies in oocytes have shown that N ramp can transport a number of divalent cations such as Fe, Zn, and Mn in a pH-dependent, electrogenic fashion associated with the symport of a single proton (12). In addition, transient overexpression of the wild type but not G185R N ramp2 in HEK293T cells results in a robust stimulation of cellular 55Fe uptake (15). Taken together, these results indicate that N ramp2 is the transferrin-independent system responsible for dietary iron absorption in the intestine. However, the ubiquitous expression of N ramp2 mRNA suggests that it may be involved in iron metabolism in other tissues as well. As opposed to N ramp1, where the cellular and subcellular localization of the protein have been established, the lack of isoform-specific, anti-N ramp2 antibodies has precluded the identification of the cell type and of the subcellular compartment expressing this protein. Such information is critical to elucidate the role of the N ramp2 protein in cellular iron metabolism. In particular, the demonstration of H+-driven, Fe2+ transport activity of N ramp2, as well as its expression in a wide variety of tissues, make it a likely candidate not only for transferrin-independent iron absorption in the intestine but also for the transferrin-independent uptake of iron in peripheral tissues. It is well established that acidification of the endosomal compartment causes Fe3+ release from transferrin and that reductases then convert the Fe3+ to Fe2+, but the mechanism of transport of Fe2+ across the endosomal membrane has not yet been elucidated.

Materials and Methods

Immunogens. For the production of isoform-specific polyclonal antisera directed against N ramp2, rabbits were immunized with fusion proteins containing glutathione S-transferase (GST) fused to a peptide segment derived from the amino terminal region of N ramp2 (residues 1–71; for amino acid numbering see reference 10). This peptide is in a region of the protein which is not conserved in other N ramp family members, including N ramp1 (18). The GST–N ramp2 fusion protein was constructed in the plasmid vector pGEX (Pharmacia) as follows. The N ramp2 sequence was amplified by PCR using oligonucleotides N F2 (5'- AAAATCTATGGTGTGGATCC-3') and N R (5'-CTGAA

TTGACAAGGCCAGT-3') (nucleotides 1–268), and the full-length N ramp2 cDNA as template. The PCR product was digested with BglII and EcoRI and the resulting overhangs were paired using the Klenow fragment of DNA polymerase I (Pharmacia) before digestion with BglII. The digested PCR product was ligated into BglII- and Smal-digested pQE40 plasmid vector (Qiagen). The in-frame his-DHFR–N ramp2 fusion protein construct was transformed into E. coli strain M15(pREP4) for expression (Qiagen). Purification was performed on Ni-NTA agarose according to experimental conditions suggested by the manufacturer (Qiagen). The polyclonal antiserum directed against the GST fusion protein was purified against the his-DHFR fusion protein by a preparative immunoblotting procedure (20). The anti-N ramp1 polyclonal antiserum (8) was affinity purified against the corresponding N ramp1–GST fusion protein by the same protocol.

Cell Culture. The mouse monocyte-macrophage cell lines R AW 264.7 and J 774a, the mouse Sertoli cell line T M 4, and the mouse kidney line mCD3 were obtained from the American Type Culture Collection (ATCC). They were cultured in media and under conditions recommended by the ATCC. WEHI 3B (myelomonocyte), WEHI 231 (B lymphocyte), Bi 141 (T lymphocyte), and 70Z/3 (pre-B cell) cells were cultured as described previously (21). Chinese hamster ovary (CHO) cells LR 73 (22) were grown in α-MEM supplemented with 10% fetal calf serum, 2 mM L-glutamine, 50 U/ml penicillin and 50 μg/ml streptomycin. All media and media supplements were purchased from GIBCO BR L. Mouse macrophages were obtained by peritoneal lavage, as previously described (8). We have previously described the production and characterization of R AW macrophages expressing a transfected wild-type N ramp1 fused to a c-myc epitope (9). The c-myc-tagged N ramp2 expression plasmid was constructed by excising the c-myc-tagged N ramp2 cDNA from plasmid pBluescript (23) using SpeI and EcoRI sites from the polylinker, followed by cloning into the mammalian expression plasmid pCB6 (24). For expression in CHO cells, the same insert was cloned into the expression vector pMT2 (25). CHO cells were transfected by electroporation as described previously (9). Clones of stable transfecants were selected in geneticin (G418, 1 mg crude/ml final; GIBCO BRL) for 10–14 d, and under conditions recommended by the ATCC. They were cultured in media and under conditions recommended by the ATCC. WEHI 3B (myelomonocyte), WEHI 231 (B lymphocyte), Bi 141 (T lymphocyte), and 70Z/3 (pre-B cell) cells were cultured as described previously (21). Chinese hamster ovary (CHO) cells LR 73 (22) were grown in α-MEM supplemented with 10% fetal calf serum, 2 mM L-glutamine, 50 U/ml penicillin and 50 μg/ml streptomycin. All media and media supplements were purchased from GIBCO BR L. Mouse macrophages were obtained by peritoneal lavage, as previously described (8). We have previously described the production and characterization of R AW macrophages expressing a transfected wild-type N ramp1 fused to a c-myc epitope (9). The c-myc-tagged N ramp2 expression plasmid was constructed by excising the c-myc-tagged N ramp2 cDNA from plasmid pBluescript (23) using SpeI and EcoRI sites from the polylinker, followed by cloning into the mammalian expression plasmid pCB6 (24). For expression in CHO cells, the same insert was cloned into the expression vector pMT2 (25). CHO cells were transfected by electroporation as described previously (9). Clones of stable transfecants were selected in geneticin (G418, 1 mg crude/ml final; GIBCO BR L) for 10–14 d, and expanded and individually, and tested for protein expression by immunofluorescence using the anti-c-myc tag monoclonal antibody 9E10 (Babco).

Immunoblotting and Immunoprecipitation. Crude membrane fractions from the various cells were prepared as described previously (27). Protein concentration of the membrane fraction was determined by the Bradford assay (BioRad). Proteins were separated on SDS-polyacrylamide gels and transferred by electroblotting to nitrocellulose membranes. For experiments where the membrane was to be stripped and reprobed, a polyvinylidene fluoride membrane was used (Westman; Schleicher and Schuell) to reduce protein loss from the membrane during stripping. Equal loading and transfer of proteins was verified by staining the blots with Ponceau S (Sigma Chemical Co.). The blots were blocked in T BST (10 mM Tris/C, pH 8, 150 mM NaCl, 0.05% Tween 20, pH 8) plus 5% skim milk powder for 1 h at room temperature. Primary antibodies used were as follows: affinity purified rabbit anti-mouse N ramp2 (1:100 dilution); affinity purified rabbit anti-
Blots were prehybridized in a solution containing 10% dextran onto denaturing agarose gels containing 0.66 M formaldehyde.

denatured in a formamide–formaldehyde mixture and loaded by addition of ddH₂O was added in place of PNGase F.

DNA (200 μg/ml) at 65°C for 2–16 h. Hybridization was for 24 h at 65°C in the same buffer containing the radiolabeled probe (10⁶ cpm/ml of hybridization buffer; specific activity 10⁴ cpm/μg DNA). Blots were washed under conditions of increasing stringency up to 0.1× SSC and 0.1% SDS at 65°C and then exposed to Kodak X R film with two intensifying screens at −70°C for 18 h to 7 d at −80°C.

Phagosome Fractionation. Phagosomes were isolated from J774a cells by a modification of a method described previously (9). 10 subconfluent 150 mm dishes of each cell line were fed with a 1:200 dilution of blue-dyed latex beads (0.8 μm; Sigma Chemical Co.) in culture medium for 1 h at 37°C in 5% CO₂. The cells were then washed in PBS and harvested in the presence of protease inhibitors (1 μg/ml leupeptin, 1 μg/ml aprotonin, 1 μg/ml pepstatin, and 100 μg/ml PM SF; all Boehringer Mannheim) and recovered by centrifugation (2,000 g, 5 min). The cell pellets were washed and resuspended in homogenization buffer (8.5% sucrose, 3 mM imidazole, pH 7.4) and homogenized by passage through a 22G needle until 90% of the cells were broken, as monitored by light microscopy. Nuclei and unbroken cells were pelleted and the supernatant loaded onto a sucrose step gradient as follows: the supernatant was brought up to 40% sucrose by addition of 62% sucrose and loaded on top of a 1-ml 62% sucrose cushion. Layers of 2 ml of 35, 25 and finally 10% sucrose were assembled and labeled using PhotoShop, Metamorph, and Freehand software. To label the lysosomal compartment, cells were incubated with 1 mg/ml lysine-fixable FITC–dextran (Molecular Probes) in growth medium for 4 h at 37°C in 5% CO₂. After washing, cells were incubated an additional 30 min to chase the dextran from the early endosomal to the lysosomal compartments. For identification of the early and recycling endosomal compartment, cells were incubated in serum-free medium containing 50 μg/ml FITC–transferrin (Molecular Probes) for 30 min at 37°C in 5% CO₂. Phagosomes were formed by incubating the cells with 3 μm latex beads (Sigma Chemical Co.) diluted 1:200 in complete culture medium for 15 min at 37°C in 5% CO₂. After treatments to identify the specific subcellular compartments, cells were fixed in 4% paraformaldehyde and processed for immunofluorescence.

Results

To generate an isomform-specific anti-N ramp2 antisera, a protein segment derived from the amino terminus of N ramp2 was selected based on its predicted antigenicity and its sequence divergence from N ramp1 (28% identity). A GST–N ramp2 fusion protein containing the amino terminal N ramp2 segment was produced and used for immunization, and the anti-N ramp2 fraction was further isolated from the immune serum by affinity purification against a second immobilized N ramp2–DH FR fusion partner. The antisera was tested for specificity by immunoblotting crude membrane fractions as well as by immunoprecipitation of [35S]methionine-labeled cell lysates from transfected CHO cell clones expressing either c-myc-tagged N ramp1 or c-myc-tagged N ramp2 proteins (Fig. 1). The anti-c-myc monoclonal antibody (9E10 [30]) specifically recognized a species of apparent molecular mass 90–100 kD in the N ramp2-transfected CHO cells, and a protein of 85–95 kD in the N ramp1-transfected cells that were absent from extracts of untransfected CHO controls (Fig. 1, A and B, bottom). These protein species migrated as broad bands in SDS–acylamide gels. In membrane preparations from N ramp2-transfected CHO cells (CHON2), the affinity-
purified anti-Nramp2 antiserum recognized a single protein species of apparent molecular mass 90-100 kD (Fig. 1 A, top). Immunoblotting analysis of the same set of membrane fractions with anti-Nramp1 antiserum revealed a single protein species of 85-95 kD in membranes from the Nramp1-transfected CHO cells (Fig. 1 A, middle). These immunoreactive bands were absent from untransfected CHO cells and from transfected cells expressing the other Nramp isoform. The electrophoretic mobility characteristics of the Nramp1 and Nramp2 detected by the respective polyclonal antiserum were very similar to those of the species detected by the anti-c-myc antibody in the same cell extracts (Fig. 1 A, bottom). The reactivity and isoform specificity of the antibodies were confirmed by immunoprecipitation studies of [35S]methionine metabolically labeled cell extracts from CHO cells or from N r a m p 1 and Nramp2 CHO transfec-

Figure 1. After affinity purification, anti-Nramp1 and anti-
Nramp2 antisera were tested for specificity and reactivity by immu-

nobl otting against crude membrane fractions (A) as well as by immunoprecipitation of [35S]methionine-labeled cell lysates (B) from CHO cells or from the same cells transfected with a c-myc-tagged (tag sequence EQKLISEEDL) Nramp1 (CHON1) or a c-myc-tagged Nramp2 (CHON2). Protein ex-

Experimental Design: Membrane Fractions and Immuno- blot Analysis

Membrane fractions were prepared from different cultured cell lines as well as from thioglycolate-induced primary mouse macrophages (Mø) and is expressed most exclusively in mononuclear phagocytes, N r a m p 2 mRNA is expressed in most tissues (10-12). W e asked whether the two N r a m p proteins would display an overlapping or mutually exclusive expression pattern. N o rth-ern blot analysis of total cellular RNA from a panel of mu-
rine hematological cell lines revealed a readily detectable

level of N r a m p 2 mRNA expression in the macrophage lines RAW 264.7 and J774a as well as in Friend virus-
transformed erythroleukemia (MEL) cells (Fig. 2 A, top; exposure time 1 wk). A much lower level of expression of N r a m p 2 was found in other cell lines: WEHI 231 (B lymphocyte), WEHI 3B (melanoma monocyte), 70/2 (pre-B lymphocyte), and B1 141 (T lymphocyte). In comparison, N r a m p 1 mRNA expression was restricted to the macrophage cell lines RAW 264.7 and J774a and is expressed at levels ~50-fold higher than that of N r a m p 2 (Fig. 2 A, middle; exposure time 24 h). Thus, N r a m p 1 and N r a m p 2 mRNA are coexpressed in macrophages.

To analyze N r a m p 2 protein expression in macrophages, membrane fractions were prepared from thioglycolate-

induced primary mouse macrophages, from J774a and RAW 264.7 cultured macrophages. For comparison, membranes were also prepared from cell lines derived from tissues previously shown to express a high level of N r a m p 2 mRNA: the Sertoli cell line T M 4 and the kidney inner medullary collecting duct line mM C D -3 (17). M embranes were also prepared from control and Nramp2-transfected CHO cells, as well as from two cell lines expressing low levels of N r a m p 2 mRNA (Fig. 2 A, WEHI 231 and WEHI 3B). The membranes were analyzed by immunoblotting with the anti-Nramp2 antibody (Fig. 2, B and C). The antibody detected a major heterogeneous immunoreactive protein species of broad electrophoretic mobility with an apparent molecular mass of 80-90 kD in all cells tested, with the exception of WEHI 231 cells and untransfected CHO cells. The protein was most abundant in TM 4, RAW 264.7, J774a and MEL cells. The protein was also detected in the membranes prepared from primary mouse macrophages and mM C D -3 cells, although at a lower level. WEHI 3B membranes showed the lowest level of N r a m p 2 expression,
whereas WEHI 231 and untransfected CHO cells were negative for N ramp2 expression. In the positive membrane samples, the electrophoretic mobility and heterogeneity of the immunoreactive species varied, possibly due to different posttranslational modification of the protein in these cell types. Thus, N ramp2 is expressed in a wide variety of tissues, including macrophages, and macrophages coexpress N ramp1 and N ramp2.

The apparent mass of endogenous N ramp2 estimated by SDS-PAGE is considerably greater than the 62.3 kD molecular mass predicted by the primary amino acid sequence of the cDNA. Together with the broadness of the immunoreactive band, this anomalous mobility suggests that N ramp2 may be posttranslationally modified by glycosylation. To test this hypothesis, membrane fractions from J774a cells and N ramp2-transfected CHO cells were treated with endoglycosidases followed by electrophoresis and immunoblotting. N ramp2 was resistant to digestion with Endo H (Fig. 2 C, lanes 1 and 2), which specifically cleaves high mannose and some hybrid N-linked oligosaccharides from glycoproteins. In contrast, PNGase F, which hydrolyzes high mannose, hybrid, and complex oligosaccharides, converted the 82-kD N ramp2 species into smaller forms of approximate apparent molecular masses of 50–55 kD (Fig. 2 C, lane 4). PNGase treatment of membranes from the N ramp2 CHO transfectants also resulted in a shift of the apparent molecular mass of the protein from 85 to ~56 kD (Fig. 2 C, lanes 6 and 7). Therefore, N ramp2 is posttranslationally modified extensively by complex N-linked glycosylation.

To gain insight into the subcellular localization of N ramp2, we first performed immunofluorescence studies on CHO and RAW 264.7 transfected cells, using an antibody directed against the c-myc epitope attached to the carboxy terminus of the transfected N ramp2 protein. The high levels of transfected protein in these cell lines facilitated nonambiguous localization of N ramp2, without possible limitation associated with low levels of expression of the endogenous protein. We have previously observed that the c-myc-tagged N ramp2 protein is functional in both CHO and RAW macrophage backgrounds and carries out active Fe²⁺ transport in these cells (Govoni, G., and P. Gros, unpublished results). We initially tested whether N ramp2 localizes to the late endosomal/lysosomal compartment, as found earlier for N ramp1 (9). To label the lysosomal compartment, cells were cultured in the presence of FITC-conjugated dextran, followed by a chase period of 30 min to remove the dextran from the early endosomal compartments, before fixation and immunostaining with the anti-c-myc antibody. In N ramp1-transfected CHO cells, there was clear colocalization of the anti-c-myc staining (Fig. 3 A) and the dextran-loaded late endosomal/lysosomal compartment (Fig. 3 B). N ramp1-stained vesicles negative for FITC-dextran were also detected. In contrast, in N ramp2-transfected CHO cells, anti-c-myc staining revealed an intracellular network of finer punctate vesicles distributed throughout the cytoplasm (Fig. 3 C). This staining does not appear to colocalize with the FITC-dextran (Fig. 3 D). Similar results were obtained in parallel experiments using the c-myc-N ramp1 and c-myc-N ramp2-transfected RAW cells (data not shown), indicating that N ramp2 is not expressed in the lysosomal compartment. Thus, N ramp1 and N ramp2 clearly appear to have distinct, nonoverlapping subcellular sites of expression.

Since N ramp2 is implicated in cellular iron uptake, it appears logical that N ramp2 be present at the plasma membrane and/or in recycling endosomes. To label these compartments, CHO (Fig. 4) and RAW transfected cells (Fig. 5) were cultured in the presence of FITC-conjugated transferrin before fixation and immunostaining with the anti-c-myc antibody. Analysis by confocal microscopy indicated that, as expected, transferrin (green) stained both the plasma membrane (ring-like staining at the edge of the cells) and the recycling endosomes (subcellular punctate staining) (Fig. 4 B). A very similar and overlapping pattern was observed for N ramp2, as revealed by the anti-c-myc antibody (red, Fig. 4 A). Superimposition of the two images (Fig. 4 C) clearly identifies colocalization (yellow) of the two signals. Certain cells stained with FITC-transferrin but...
were negative for the c-myc staining, suggesting that although these cells are positive for the pSV2neo plasmid and are resistant to G418, they failed to express c-myc-tagged Nramp2. Such cells provide an internal control for the specificity of the anti-c-myc staining and for the overlapping staining with FITC–transferrin. Similarly, when untransfected CHO cells were identically processed and examined, the cells were not stained with the anti-c-myc antibody (Fig. 4 D), but displayed a normal pattern of staining with FITC–transferrin (Fig. 4 E). Finally, when the lysosomes of the Nramp2-transfected cells were stained with FITC–dextran (Fig. 4 H) and with the anti-c-myc antibody for Nramp2 (Fig. 4 G), no significant overlap between the two signals was detected (Fig. 4 I). A similar colocalization of Nramp2 and FITC–transferrin was noted in parallel experiments with RAW macrophages expressing c-myc-Nramp2 (Fig. 5). These results confirm that Nramp1 and Nramp2 have nonoverlapping, subcellular localization, and that Nramp2 colocalizes with transferrin in the early recycling endosomal compartment.

To confirm the endosomal localization of Nramp2 determined in transfected CHO and RAW cells, we performed immunofluorescence in nontransfected cell lines that tested positive (MEL, TM4) or negative (WEHI 231) for Nramp2 expression by immunoblotting (Fig. 2). Immunofluorescence was performed with the anti-Nramp2 polyclonal antisera and FITC–transferrin, and results are shown in Fig. 6. In MEL (Fig. 6, A and B) and TM4 (C and D) cells, the endogenous Nramp2 protein (B and D) showed very similar staining pattern to that generated by FITC–transferrin (A and C), similar to that seen in transfected cells (Figs. 4 and 5). Finally, in agreement with the absence of Nramp2 expression in WEHI 231 cells noted by immunoblotting (Fig. 2 B), no Nramp2 staining was observed in WEHI231 cells (Fig. 6 F), although the endosomal compartment of these cells could readily be labeled by FITC–transferrin (Fig. 6 E). Together, these results verify data obtained in transfected CHO and RAW cells.

The localization of Nramp2 to the plasma membrane and endosomal network raised the possibility that like Nramp1, Nramp2 may become associated with phagosomal membranes after phagocytosis. Phagosomes are initially

Figure 4. Colocalization of Nramp2 and transferrin in early endosomes was determined by double immunofluorescence and confocal microscopy in either CHO control cells (D–F) or CHO cells transfected with a c-myc-tagged Nramp2 protein (A–C and G–I). Cells were cultured in the presence of FITC–conjugated transferrin (A–F) or FITC–conjugated dextran (G–I) before fixation and immunostaining with the anti-c-myc tag antibody and a rhodamine-conjugated secondary antibody (A, D, and G). The slides were then examined by confocal microscopy, and the FITC (green; B, E, and H) and rhodamine (red; A, D, and G) images were overlaid (yellow identifies colocalization; C, F, and I). In the Nramp2-transfected CHO cells, the Nramp2 staining (A and G) showed extensive overlap with the distribution of FITC–transferrin (A + B = C) but not with lysosomal FITC–dextran (G + H = I). FITC–transferrin staining was observed in the untransfected CHO cells (E), but the cells were negative for anti-c-myc staining (D).
derived from the plasma membrane and are known to sequentially interact and acquire proteins from both early and late endosomes before their final fusion with lysosomes (29). We have shown that Nramp1 is acquired during the phagosomal maturation process, using the model system of latex bead–containing phagosomes (9). Latex bead–containing phagosomes are ideal for microscopic examination and can also be purified from cell homogenates by flotation on sucrose gradients. To determine whether Nramp2 can associate with the phagosomal membrane, J774a cells were fed latex beads for 1 h at 37°C, and the phagosomal fraction was isolated from cell homogenates by fractionation on sucrose gradient. Equal amounts of the purified phagosomal fraction and of a crude total membrane fraction were separated by SDS-PAGE, and the relative amount of endogenous Nramp2 in each sample was determined by immunoblotting. As shown in Fig. 7 A (left), Nramp2 was significantly enriched in purified phagosomes as compared to the crude membrane preparation. In these experiments, the Lamp1 protein (marker of the phagolysosome, center) and the transferrin receptor (marker of the plasma membrane, right) were used as controls. As expected, significant enrichment of Lamp1 was seen in the phagosomal fraction whereas the transferrin receptor was not enriched in phagosomes even though it is readily detectable in crude membrane fractions. These results suggest that Nramp2 becomes associated with the phagosome during its maturation to phagolysosome. Possible association of Nramp2 with latex beads phagosomes was further analyzed in J774a cells by double immunofluorescence and confocal microscopy. J774a cells were fed latex beads and then fixed and processed by double immunofluorescence using anti-Nramp2 and anti-Lamp1 antibodies (Fig. 7 B). In J774a cells, the Nramp2 signal obtained with our antibody was weak, which limited the analysis. Nevertheless, in several of the sections analyzed a portion of the Nramp2 signal could clearly be seen...
at the periphery of the bead, suggesting association with the phagosome. However, this signal was much weaker than that obtained using the anti-Lamp1 antibody. Thus, results from immunoblotting and immunofluorescence suggest the possibility that a portion of the endosomal N ramp2 protein becomes associated with the phagosome during phagolysosome maturation.

Discussion

A large body of biochemical data supports the proposal that N ramp2 functions as a transporter for several divalent cations, including Fe2+. N ramp2 is mutated in the mk mouse and in the Belgrade rat, with both animals exhibiting a severe microcytic hypochromic anemia and a severe defect in iron absorption by intestinal cells. However, in vitro studies have shown that iron acquisition is also decreased in the peripheral cells and tissues of these animals and that the anemia cannot be corrected by direct iron injections, suggesting a second block of iron entry into peripheral tissues. Thus, physiological consequences of N ramp2 mutations in vivo strongly suggest that N ramp2 is not only involved in iron uptake at the level of the intestinal enterocyte but also participates in iron acquisition in other cell types as well. In peripheral tissues, cellular iron uptake is through the transferrin cycle (for review see reference 38). Diferric transferrin binds to the transferrin receptor and is internalized, and acidification of the internalized vesicles results in release of iron from transferrin followed by alkalinization of the vesicles and recycling of the receptor to the cell surface. Iron escapes the acidified endosomal compartment to reach the cytoplasm, where it can be captured by mitochondria for heme biosynthesis and incorporation into heme-containing proteins, stored in the cytoplasm in the form of ferritin, and/or used directly for synthesis of nonheme-containing proteins (e.g., ribonucleotide reductase). The mechanism by which iron is extruded from the acidified endosome to enter the cytoplasm is unknown and has been a matter of considerable debate.

In the current study, we have raised isoform specific anti-N ramp2 antiserum and have used it to verify a number of structural and biochemical features of N ramp2 predicted from the primary amino acid sequence deduced from the cDNA. These analyses have shown that N ramp2 is an integral membrane protein which is extensively modified by N-linked glycosylation. As opposed to N ramp1, which is macrophage-specific, N ramp2 protein was found ubiquitously expressed in a majority of cell lines analyzed. The current study has also clearly established that N ramp2 and N ramp1 localize to distinct subcellular compartments. Whereas N ramp1 colocalizes with FITC–dextran in the lysosomal compartment, N ramp2 is not detectable in this compartment but rather shows clear colocalization with FITC–transferrin both at the plasma membrane and in recycling endosomes. The demonstration of N ramp2 expression in several peripheral tissues, the colocalization of N ramp2 and transferrin in plasma membrane and recycling endosomes, the iron transport properties of the N ramp2 protein, and the effect of N ramp2 mutations on iron metabolism in peripheral tissues are strong evidence that N ramp2 is responsible for transporting Fe2+ into the cytoplasm after acidification of the transferrin-positive endosome (Fig. 8). Interestingly, this acidification would simultaneously provide a gating mechanism for iron transport by N ramp2 and for release from transferrin (Fig. 8). Indeed, the pH dependence of iron transport by N ramp2 has been demonstrated in several systems, including X enus oocytes (12), transfected HEK 293T cells (14), and CHO cells (G ovoni, G., and P. Gros, unpublished results). Likewise, release of iron from transferrin and its subsequent release from endosomes is dependent on endosome acidification, which can be inhibited by bafilomycin (40) and concanamycin (41), specific inhibitors of the vacuolar H+-ATPase, but is insensitive to the Na+, K+-ATPase inhibitor ouabain (41, 42). This suggests a critical role for the vacuolar H+-ATPase in this process. Indeed, the association of vacuolar H+-ATPase with transferrin-positive endosomal vesicles has been demonstrated by immunohistochemical means in LLC–porcine kidney epithelial 1 cells (43).
The demonstration that Nramp1 and Nramp2 are coexpressed in the same cell type with distinct subcellular localizations suggests possible functional parallels between Nramp1 and Nramp2 (Fig. 8). Both the yeast Smf1 and the mammalian Nramp2 proteins can transport Mn\(^{2+}\), with the latter also transporting Fe\(^{2+}\) and other divalent cations. Nramp2 and Smf1 share approximately 40% sequence identity within the conserved hydrophobic core (18). As the mammalian Nramp1 and Nramp2 proteins share almost 80% sequence identity within their hydrophobic cores, it is likely that Nramp1 is involved in the transport of divalent cations as well. The removal of such metabolically essential ions from the phagosomal space would provide an attractive explanation for the observed pleiotropic effect of Nramp1 mutations in vivo on the replicative potential of internalized microbes that inhabit the phagosomal space in macrophages. Additionally, the observation that a portion of Nramp2 associates with latex bead phagosomes in J774a cells suggests that Nramp2 may also play a role in depleting the phagosomal space of divalent cations necessary for microbial survival.

Despite considerable efforts, transport studies in CHO and RAW cells transfected and overexpressing Nramp1 protein have so far failed to demonstrate an Nramp1-mediated transport of either \(^{54}\text{Mn}\) or \(^{55}\text{Fe}\) transport in these cells (Govoni, G., and P. Gros, unpublished results). The distinct, nonoverlapping distribution of Nramp1 and Nramp2 reported here provides an explanation for this apparent lack of transport activity associated with Nramp1. While the two proteins may have the same transport potential, the observed targeting of Nramp2 to the plasma membrane and recycling endosome compartment would result in a net increase in cellular accumulation of extracellularly added, radiolabeled ligand under acidic pH transport assay conditions. On the other hand, the restricted expression of Nramp1 to the lysosomal compartment would not cause a similar increased cellular uptake of a ligand presented in the extracellular milieu, although it could act on such ligand if present in the phagolysosomal space. Therefore, it is tempting to speculate that both Nramp1 and Nramp2 have similar transport function but act at different, nonoverlapping, intracellular sites (Fig. 8). If Nramp1 and Nramp2 do indeed transport the same substrates, it is also tempting to speculate that vesicular acidification via the vacuolar H\(^+\)-ATPase may provide a key common gating mechanism for the activation of both transporters, through fusion with vacuolar H\(^+\)-ATPase-positive vesicles (Fig. 8). Possible similarities and differences in the mechanism of action and regulation of Nramp1 and Nramp2 in macrophages are currently being investigated.
References


