Function of human Rh based on structure of RhCG at 2.1 Å

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The five human Rh (Rhesus) family proteins fall into two functionally distinct groups: the ammonia transporting Rh glycoproteins (RhAG, RhBG, and RhCG) and the nontransporting (nonglycosylated) Rh proteins (RhD and RhCE). Erythroid RhD is recognized for its key role in blood group incompatibility (Rh+) vs. (Rh−) (1, 2) whereby Rh− individuals (lacking RhD antigen) can develop an immune response when exposed to Rh+ blood. Along with RhD, RhAG and RhCE are specific components of the erythrocyte Rh complex with incompletely defined stoichiometry (3, 4). RhBG and RhCG are found in a variety of tissues including brain, liver, gut, and kidney. Proteins of the Rh family have 20–27% identity to the “methyl-ammonia permease” (MEPs) in yeast and to the “ammonia transporters” (Amts) in prokaryotes, plants, and some invertebrates (5). RhAG, RhBG, and RhCG have been shown to transport ammonia (6, 7). RhCG is responsible for ammonia secretion across the membranes of the epithelial cells of the renal collecting duct (8, 9). RhCG knockout mice show abnormal acidification of the blood due to impaired urinary removal of NH\textsubscript{4}\textsuperscript{+} emphasizing the importance of renal transport for human physiology.

Aqueous ammonia (NH\textsubscript{3}) and ammonium ion (NH\textsubscript{4}\textsuperscript{+}) are in equilibrium governed by a pK\textsubscript{a} of 9.25. Thus at physiological pH, NH\textsubscript{3} is the preponderance of evidence suggested that the Amts and their homologs transport the charged species, NH\textsubscript{4}\textsuperscript{+}, however, the atomic structure of AmtB and conductivity measurements (10, 11) showed that the prokaryotic Amts serve as channels for passage of uncharged NH\textsubscript{3} across the membrane, later confirmed by subsequent transport measurements (12, 13), as had been postulated previously (14). NH\textsubscript{3} is formed by reprotonation of NH\textsubscript{4}\textsuperscript{+} in solution.

Here, we report the atomic resolution structure of a eukaryotic Rh glycoprotein, human RhCG, expressed in modified HEK293s cells (15). The structural differs from its prokaryotic homologs in having an additional transmembrane helix and in other Rh-specific structural features that further elaborate the NH\textsubscript{3} transport mechanism vital to human renal function. The high sequence similarity of RhCG to the erythroid Rh proteins allows construction of reliable models of their structures that inform the stoichiometry and assembly of the erythrocyte complex.

Results and Discussion

RhCG Forms Trimmers of 12 Crossing Transmembrane Channels.

The atomic structure of human RhCG at 2.1 Å resolution contains an additional N-terminal transmembrane helix, termed M0 relative to Amt (10–12) or the prokaryotic NeRh (16, 17). The 12 transmembrane \(\alpha\)-helices (M0–M11) form a right-handed helical bundle that surrounds an apparently empty channel (Fig. 1). As in bacterial Amt proteins, helices M1–M5 and M6–M10 are related by an in-plane quasi-2-fold symmetry with respect to the membrane plane. Helices M0 and M11 break this quasi-symmetry and are linked by loops to the 10-helix core. These loops vary in sequence and length between the human Rh family (Fig. S1). In RhD and RhCE, the loops are relatively short and are resistant to proteolytic cleavage. In the human Rh glycoproteins, RhAG, RhBG, and RhCG, the loops (with the exception of M10–M11 in RhAG) are longer in sequence and are susceptible to proteolytic cleavage (4). These differences suggest that the loops are functionally differentiated.

The RhCG M0 region is conserved across higher eukaryotes (Fig. S2). However, in the prokaryotic Amts (10–12) and NeRh (16, 17), a shorter stretch of residues constitute a cleaved signal sequence. M0 lies against the lipid-accessible surface of the trimer and at the interface between subunits. A single glutamine (Q19), halfway along the M0 helix in the center of the lipid bilayer, is hydrogen bonded to glutamine (Q101) on M2 (Fig. 2). In all members of the Rh family that have M0, a glutamine or glu-tamate are conserved at positions equivalent to 19 and 101 (Fig. S2). These polar residues in the center of a transmembrane region would otherwise be exposed to lipid. Therefore, M0 is pinned to M2 by this conserved midmembrane interaction.

Each of the human Rh glycoproteins contains a single conserved N-glycosylation site (N48 in RhCG) on the loop between M0 and M1. For our construct, expressed in a human endothelial kidney (HEK) cell line engineered to produce a short homogenous core oligosaccharide (Man\textsubscript{3}GlcNAc\textsubscript{2}), termed GloT- (15) (SI Methods), N48 would be homogeneously glycosylated. The

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The authors declare no conflict of interest.

Data deposition: Atomic coordinates and structure factors for the human RhCG crystal structure are deposited in the Protein Data Bank under accession code 3HID. Comparative models deposited as the “HumanRhesus” model set into MODBASE (18). Correspondence and requests for materials should be addressed to stroud@msg.ucsf.edu.

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mass of 55.1 ± 0.1 kDa measured by MALDI mass spectroscopy is consistent with this predicted mass of 55.163 kDa for the singly glycosylated species.

In RhCG, the M0–M1 loop (~25 amino acids) contains several hydrophobic residues (A36, A38, W40, W41, and L49) that would be shielded from solvent by the normal branched polysaccharide. Similar hydrophobic character is seen in the M0–M1 loop of the other human Rh glycoproteins. The nonglycosylated RhD/CE have a shorter (~11 amino acids) M0–M1 loop. The glycosylation may serve to cover the hydrophobic sequence and protect the longer loops from proteolytic degradation.

**Mechanism of NH₃ Permeation.** Comparison of RhCG to the prokaryotic Amt (10–12) and NeRh (16, 17) structures confirms distinct conserved family specific features including acidic residues that line the apertures, an external aperture gated by phenylalanines, a largely hydrophobic lumen of the channel, and twin coplanar histidines in the center of the channel proposed to facilitate passage of the substrate. Because of these conserved features, the RhCG channel aperture is expected to attract NH₄⁺, whereas the lumen excludes NH₄⁺ and facilitates passage of neutral NH₃. The central pore of the channel is hydrophobic (Fig. 3), and most of residues are conserved among the human Rh glycoproteins (Fig. S1) favoring passage of uncharged NH₃ as in AmtB (10). This implies deprotonation of NH₄⁺ within the vestibule.

Conserved twin histidines (H185 and H344), required for optimal NH₃ permeation (21), are hydrogen bonded to each other via their NH–N6 and intrude into the center of the channel (Fig. 3, blue box). These coplanar and antiparallel histidines are conserved underlying their importance in the mechanism of transport. Despite these histidines, the hydrophobic lumen contains no ordered water molecules. The absence of any ordered water further supports the exclusion of NH₄⁺, which would require a replacement for its hydration shell, from the channel.

At the extracellular vestibule, Amt channels have a characteristic tryptophan proposed to aid in recruiting NH₃ through a π-cation interaction (10). RhCG and NeRh (16, 17) lack this tryptophan, however acidic residues E166 in the extracellular vestibule and D218, D278, and E329 in the intracellular vestibule
may serve to attract NH$_3^+$ (Fig. 3, labeled with *). Transport of NH$_3^+$ by the Rh glycoproteins has been shown to be pH dependent (7, 23). Because the hydrophobicity of the channel lumen is phylogenetically well conserved, pH dependence may be conferred at the vestibules. In the kidney, the environment of the extracellular vestibule in RhCG differs between the apical and basal surface of the epithelium of the renal intercalated cell. The basal surface exposes the extracellular vestibule of RhCG to the kidney interstitium with a pH of ~7.3, thereby resulting in a net extracellular charge of ~2.6 (~2.3 in the vestibule) for each RhCG monomer, assuming unaltered side chain pK$_a$'s. The apical surface exposes the extracellular vestibule to the urinary space with a pH of ~5.5, suggesting protonation of five histidines (39, 46, 111, 247, and 374) for a net extracellular charge of +3.1 (+0.4 in the vestibule) assuming unaltered side chain pK$_a$'s. Thus the extracellular vestibule in the urinary space has reduced affinity for NH$_3^+$, perhaps accounting for the pH dependence of the delivery of NH$_3^+$ to the urinary space via RhCG (24).

Another acidic residue (D177) near the vestibule has a key structural role rather than substrate recruitment (Fig. 3 D177 ▼). RhCG D177 is hydrogen bonded to the backbone amides of residues 180–183 in the extracellular vestibule. These interactions satisfy the available bonds and occlude the acidic side chain from solvent. The structures of Amt, NeRh, and RhCG show that this position plays a key structural role in capping the M5 helix (10). Similarly, at the intracellular vestibule, quasi-symmetric D336 acts as an N-terminal cap to the M10 helix (Fig. 3 D336 ▽). The D160A mutation in AmtB (25), and the homologous D177N mutation in RhCG (26) result in loss of transport of the orthogonal radio-labeled substrate, C14-methyl ammonia, most probably, we argue, due to structural disruption.

The extracellular pore in the Amt subfamily is gated by two residues 180 and 181 in the 180-183 loop, serving as a cap to the M5 helix (10). In contrast to Amt, the outer phenylalanine (F130) does not obstruct the pore of RhCG or NeRh (Fig. 3, orange box) (16, 17). This conformation may be favored due to the nearby D129, a buried, charged residue that is hydrogen bonded to the hydroxyl of Y254 of an adjacent monomer within the trimeric assembly and the HN6 of H67. Each of these residues is conserved in the human Rh glycoproteins. The limited mobility of D129 may restrict F130, thereby locking the gate in the open conformation. Small voids on each side of the F235 side chain allow for freedom of motion necessary to open the “Phe gate.” Mutation of F235 to valine reduced NH$_3^+$ transport, supporting a role in passage (27).

An RhCG feature not present in Amts is a pocket that extends from the cytosolic aperture to the lateral exterior surface in contact with the lipid hydrocarbon chains that we term the “shunt” (Fig. 3, green box). Amino acids that line the shunt are similar among human Rh glycoproteins but conserved in RhD and RhCE, which do not function as NH$_3^+$ channels. The equivalent region in the Amt structures contained a large void but no opening to the cytosol or membrane. The shunt is also seen in the structure of NeRh, suggesting that it is common to the Rh subfamily. The cytosolic aperture of this shunt has an acidic residue E329, which may play a role in recruiting charged NH$_3^+$.

The extracellular pore in the Amt subfamily is gated by two residues 180 and 181 in the 180-183 loop, serving as a cap to the M5 helix (10). In contrast to Amt, the outer phenylalanine (F235) acts as an N-terminal cap to the M10 helix (Fig. 3 D235 ▼). Similarly, at the intracellular vestibule, quasi-symmetric D336 was observed, consistent with a lack of nonspecific protein-mediated permeability and no leakage of NH$_3^+$.

RhCG Transports Ammonia but Not Water. The permeation of NH$_3^+$ was measured using recombinant RhCG-containing proteoliposomes, where the influx of NH$_3^+$, but not NH$_2^-$, induces a rise in internal pH assayed by the pH-sensitive fluorescence of 5-carboxy-fluorescein (CF) located inside. (Fig. 4A) (10, 28, 29). Relative rates were assayed by stopped-flow external addition of 5 mM NH$_4$Cl. In the absence of protein, liposomes alone exhibited a rate of 11.1 ± 0.6 s$^{-1}$ (n = 6). With RhCG, the rate was 14.3 ± 1.9 s$^{-1}$ (n = 6). Doubling the RhCG increased the rate to 23.4 ± 2.4 s$^{-1}$, demonstrating that RhCG transports NH$_3^+$. Substitution of 5 mM NaCl for 5 mM NH$_4$Cl made no detectable change in fluorescence, demonstrating that changes in CF fluorescence were due to NH$_3^+$ influx rather than to water efflux resulting from any change in osmolarity. When the Escherichia coli acetate transporter, YaaH, was reconstituted with liposomes (as control for the reconstitution procedure), no significant change in the control rate at 10.4 ± 1.7 s$^{-1}$ (n = 6) was observed, consistent with a lack of nonspecific protein-mediated permeability and no leakage of NH$_3^+$ due to the reconstitution procedure. These results indicate that RhCG is an efficient channel for NH$_3^+$ and NH$_4^+$ conductance is not precluded by the liposome assays, nor the different indications from electrophysiological recordings (6, 7, 30) resolved, however the structure of RhCG seems to imply exclusion of NH$_3^+$ from the hydrophobic lumen.

Water permeability of RhCG was assessed by measuring conduction of proteoliposomes upon osmotic challenge, as assayed by concentration-dependent self-quenching of CF (29) (Fig. 4B). For protein-free liposomes, the fluorescence change on stopped-flow mixing with 275 mM sucrose indicated that water efflux was 4.0 ± 0.1 s$^{-1}$ (n = 6) versus 3.9 ± 0.1 s$^{-1}$ (n = 6) for RhCG proteoliposomes, indicating no additional water permeability in RhCG proteoliposomes. Doubling the protein concentration resulted in an increase to 4.6 ± 0.1 s$^{-1}$ and YaaH gave 5.7 ± 0.1 s$^{-1}$. This can be compared with reconstitutions of the water channel AqpZ into proteoliposomes under similar conditions which gave ~150 s$^{-1}$ (10). Thus, RhCG does not transport water and the liposomes were well sealed. This conclusion is critical to kidney physiology, in which the high solute concentration of the urinary space must be maintained.

![Fig. 3.](https://example.com/fig3.png)

**Fig. 3.** RhCG NH$_3^+$ channel and shunt. Partially transparent surface generated by HOLLOW (22) of the channel, voids, and shunt. Select side chains that compose this surface are displayed as sticks. Water molecules within the channel surface displayed as red spheres. All atoms colored by atom type with carbons in white except for the shunt with carbons in green.
Modeling the Erythrocyte Rh Complex. Early identification of the erythrocyte Rh complex (3) suggested a composition of RhAG: RhD: RhCE with ∼2:1:1 stoichiometry, in tetrameric assemblies (4). The specific arrangement of proteins can now be addressed by protein modeling. Among the human Rh glycoproteins, 35% of residues and 46% of transmembrane positions are conserved (Fig. S1). Thus, the homotrimeric RhCG currently provides a better template for modeling the erythroid complex and the binary interfaces between them than any other known structure.

To probe the stoichiometry and configuration of the subunits in the erythrocyte complex, models of all possible trimeric arrangements of the RhAG, RhD, and RhCE subunits were built based on the RhCG trimer template (31). In the RhCG trimer, four exposed polar regions of each subunit are buried by a complementary region on the opposing subunit: P1(Y32, N252), P2(Y60, Q64, D65, S234), P3(H67, D129, Y254) (Fig. 5), and P4(S87, D432, W436). The polarity of these amino acid residues is preserved in the erythrocytic proteins (Fig. S1), and is distinct from the Amt and MEP proteins.

The modeled assemblies were used to test the various possible homo and heterologeric arrangements in the erythrocyte Rh complex. Conservation of the subunit interfaces argues that the human erythrocyte Rh complex is trimeric, just as in the RhCG trimer. Computational assessment of all homomeric and heteromeric trimer models shows various scores that are less favorable than in the parent RhCG structure (Table S1). However, the scores among the models are most favorable for homotrimers of RhAG, followed by trimers that replace one RhAG by RhD, or RhCE. They are least favorable for homotrimers of RhD, or RhCE, suggesting that RhAG is more compatible with the template trimeric configuration than RhD and RhCE (Table S1). Although the assessment scores cannot be used to distinguish between existing and nonexistent configurations, they suggest a stochastic subunit arrangement within trimers with the overall experimentally observed ratio of ∼2:1:1 for RhAG: RhD: RhCE (4).

Functional Insights Derived from Models. RhD and RhCE do not transport NH₃. However, NH₃ permeation cannot be measured directly because these two proteins can only be expressed in the presence of RhAG and NH₃ passage measured in two different experimental methods can be attributed to RhAG alone (24, 32, 33). In the RhD/CE protein models, the positions occupied by conserved histidines within the central region of the channel are replaced. The outer histidine (H1185) found in the Rh glycoproteins is replaced by tyrosine and the inner histidine (H344) by phenylalanine. These substitutions do not fully obstruct the lumen of the channel, however the histidines (in AmtB) have been shown to be essential; mutants in which the histidines were replaced by phenylalanine and alanine lost NH₃ conduction (21). The histidine C-H hydrogen-bonding potential may be a key to freezing the tautomeric form of NH₃ as suggested based on electron density difference maps between high and low NH₃ concentration in crystal structures (10).

Debate continues as to a possible role for the Rh erythroid complex as the long-sought-after conductor of CO₂ in the red blood cell. Such a role was suggested based on reduced growth of green algae following sRNAi down regulation of a native Rh homolog (34, 35). Support for this proposal was augmented by a feature of the prokaryotic NcRh structure (16), where a CO₂ binding site was identified in a cleft near the cytosolic aperture of the channel. However, RhCG shows that this site is not present in the human Rh glycoproteins. RhCG Q283 replaces an alanine at the equivalent position in NcRh. This glutamate side chain occupies the proposed CO₂ site and it is conserved in all the human Rh glycoproteins. Therefore this site is not available for CO₂ transport in human Rh.

Nevertheless, RhAG and the erythrocyte aquaporin AQP1 do allow passage of both NH₃ and the less hydrophilic CO₂ measured based on changes in surface pH in Xenopus oocytes (36). However, RhAG demonstrated 4-fold greater preference for passage of NH₃ versus CO₂ than seen in AQP1. Furthermore, AQP1 is far more abundant than RhAG in the erythrocyte, making it the

Fig. 4. Channel NH₃ transport in proteoliposomes. (A) Relative vesicular pH change indicated by CF fluorescence for RhCG at protein to lipid ratio 1:90 by weight (open blue triangles) and 1:45 (closed blue triangles) demonstrated protein dependent NH₃ permeability relative to empty liposomes (black squares) and a control protein YaaH (red diamonds). (B) Proteoliposomes from the same sample preparations after addition of 275 mM sucrose at t = 0 were monitored for transport of water resulting in decreased vesicular volume, increased CF concentration, and subsequent self-quenching. Data displayed for the exponential fluorescence decay after lag time to reach self-quenching concentration and exponential fit (black lines).

Fig. 5. Phenylalanine gate and polar cluster P3. The partially transparent channel surface and nearby void surfaces are displayed to reveal the waters with observed electron density in the molecular structure. The side chains of Phe gate and P3 residues are shown as sticks colored by atom type. The P3 cluster for each monomer (structure model in white) involves Y254 from the adjacent monomer (labeled with * and in blue).
more probable primary pathway for CO₂. However, the potential remains for some Rh mediated CO₂ passage through the central channel in the same manner proposed for NH₃.

Renal Rh Glycoprotein Ammonia Transport. NH₄⁺ secretion in the kidneys is critical to systemic acid-base homeostasis (37–39). In humans, inability to appropriately dispose of excess H⁺ in urine results in distal renal tubular acidosis that depresses arterial blood pH (< 7.35), and in extreme cases results in kidney stones, rickets, and renal failure (40, 41). Amongst many epithelial cell types, RhCG and RhBG are found in the cells of the collecting tubule and collecting duct acid-secreting intercalated cells as well as in the majority of the cells of the distal collecting tubule (7, 8, 42–46).

Growing consensus supports maintenance of acid/base equilibrium and H⁺/NH₄⁺ excretion by the following model (Fig. 6). Whereas renal RhBG is found predominately on the basal epithelial surface (in contact with the kidney interstitium), RhCG is abundant in both the basal and apical surfaces (in contact with the urinary space). Based on studies of the RhCG knockout mouse, NH₃ can pass through the epithelial cell membrane, however RhCG dramatically increases the permeability as evidenced by recovery rates under acid load (8, 9). The diffusion of NH₃ through RhCG in the distal portions of the nephron epithelium appears to be driven from the interstitium, by progressively lowered pH to the epithelial cell cytosol, and into the lumen.

The pH balance is maintained by the basolateral HCO₃⁻/Cl⁻ exchanger (49–51), the luminal V-type H⁺-ATPase, and luminal H⁺/K⁺-ATPase. The secretion of protons, mostly from H⁺-ATPases, and to a lesser extent from H⁺/K⁺-ATPase acidifies urine (48, 52). The more acidic environment of the urine shifts the equilibrium [NH₄⁺]/[NH₃] ~100:1 at pH 7.25 toward 10⁴⁺:1 at pH 5.25 in urine. This equilibrium favors concentration of NH₄⁺ in the urinary space, thereby lowering the [NH₃] and so providing the NH₃ gradient to drive concentration. This also allows for the accumulation of high [NH₃⁺], resulting in net excretion of NH₄⁺. Urinary concentrations of NH₄⁺ of greater than 200 mM can exist, while across the cell membrane NH₄⁺ remains ~8 mM (7, 38). Consequently, long-term maintenance of blood pH is dependent on NH₃ passage by RhCG in the kidney.

Methods

RhCG was expressed in HEK293S cells (15) and purified using FLAG affinity purification and size-exclusion chromatography. Crystals were grown in vapor diffusion and diffraction data were collected from cryoprotected crystals at beamline 8.3.1 of the Advanced Light Source (Lawrence Berkeley National Laboratories). The structure was determined by molecular replacement in Phaser (53) using NeRh (3B9W) as search model. Model building and refinement were calculated in CCP4 (54) and REFMAC (55), ARP/wARP (56), and Coot (57) (Table 1). Protoproline permeability of NH₃ was indicated by pH-dependent response of CF by stopped-flow cytometry. Comparative models were generated using MODELLER-9v6 (31) and assessed using Z-DOPE (58), a normalized atomic distance-dependent statistical potential based on known protein structures. Detailed methods are provided in SI Text.

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Data

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Numbers in parentheses correspond to the high-resolution shell for data refinement (2.18–2.10 Å).

References

6. Bakouh N, et al. (2004) NH3 is involved in the NH₃-transport by the following model (Fig. 6) and Patricia Greene for assistance in editing the manuscript. This research was supported by National Institutes of Health Grants R01 GM24485, GM73210, and GM74929.


