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**Graphical abstract**

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Enterotoxigenic *Escherichia coli*–blood group A interactions intensify diarrheal severity

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Enterotoxigenic *Escherichia coli* (ETEC) infections are highly prevalent in developing countries, where clinical presentations range from asymptomatic colonization to severe cholera-like illness. The molecular basis for these varied presentations, which may involve strain-specific virulence features as well as host factors, has not been elucidated. We demonstrate that, when challenged with ETEC strain H10407, originally isolated from a case of cholera-like illness, blood group A human volunteers developed severe diarrhea more frequently than individuals from other blood groups. Interestingly, a diverse population of ETEC strains, including H10407, secrete the EtpA adhesin molecule. As many bacterial adhesins also agglutinate red blood cells, we combined the use of glycan arrays, biolayer interferometry, and noncanonical amino acid labeling with hemagglutination studies to demonstrate that EtpA is a dominant ETEC blood group A–specific lectin/hemagglutinin. Importantly, we have also shown that EtpA interacts specifically with glycans expressed on intestinal epithelial cells from blood group A individuals and that EtpA–mediated bacterial–host interactions accelerate bacterial adhesion and effective delivery of both the heat-labile and heat-stable toxins of ETEC. Collectively, these data provide additional insight into the complex molecular basis of severe ETEC diarrheal illness that may inform rational design of vaccines to protect those at highest risk.

**Introduction**

Enterotoxigenic *Escherichia coli* (ETEC) is a major cause of infectious diarrhea in the developing world (1, 2). These organisms are estimated to be responsible for millions of infections and hundreds of thousands of deaths, particularly of young children (3). Likewise, these pathogens are a principal cause of diarrhea in travelers to endemic areas and an occasional cause of illness in developed countries (4–7). Despite significant declines in deaths from acute diarrhea (8), enteric infections caused by ETEC and other pathogens remain inextricably associated with malnutrition, stunted growth, and impaired intellectual development (9), poorly understood sequelae that appear to result from gut dysfunction that follows repeated episodes of diarrhea in childhood (10). The ultimate impact of these illnesses is further amplified by poor responses to vaccines, increased mortality from other infections, and emerging evidence that early gut dysfunction may be reflected in the subsequent development of obesity and its attendant metabolic complications (11, 12). Consequently, these infections continue to exact an extraordinary toll on global health.

The ETEC pathovar is defined by the production of plasmid-encoded heat-labile (LT) and/or heat-stable (ST) enterotoxins (13). Effective delivery of these toxins to cognate receptors on the surface of intestinal epithelial cells is ultimately required to drive net salt and water efflux into the lumen of the small intestine, which results in diarrhea. Although ETEC was originally identified in individuals suffering from severe cholera-like diarrhea (14), the manifestations of infection may range from asymptomatic colonization to fulminant illness clinically indistinguishable from that caused by *Vibrio cholerae* (15–17). The molecular basis for these varied clinical presentations is presently unclear.

In the classic paradigm for ETEC virulence, these organisms adhere to small intestinal epithelia via plasmid-encoded fimbrial colonization factors to effect toxin delivery. However, recent studies suggest that the molecular pathogenesis of this diverse group of pathogens may involve a repertoire of additional virulence features (18–20), including a variety of different adhesins to interact with the intestinal epithelium (21).

Because of their impact on global health, these pathogens are a principal target for vaccine development (22). Although there is
ness following H10407 challenge tends to be significantly more severe than that associated with other ETEC strains (42, 43), the severity of diarrhea can vary substantially among volunteers within the same cohort, implicating the involvement of one or more host factors in disease outcome.

Here, we demonstrate that, when challenged with H10407, naive human volunteers with blood group A are at substantially increased risk for severe diarrhea relative to other blood groups. In addition, we found that the EtpA adhesin is a lectin that selectively promotes interaction of ETEC with human small intestinal epithelial cells expressing blood group A glycans. These EtpA-dependent ETEC-host interactions potentially provide a molecular basis for selection and an accelerated course of illness in blood group A–positive hosts.

Results

Blood group A volunteers challenged with ETEC have more severe disease. Because pathogen interactions with human blood group antigens can influence the outcome of infection by other gastrointestinal pathogens (44, 45), the severity of diarrhea can vary substantially among volunteers within the same cohort, implicating the involvement of one or more host factors in disease outcome.

Here, we demonstrate that, when challenged with H10407, naive human volunteers with blood group A are at substantially increased risk for severe diarrhea relative to other blood groups. In addition, we found that the EtpA adhesin is a lectin that selectively promotes interaction of ETEC with human small intestinal epithelial cells expressing blood group A glycans. These EtpA-dependent ETEC-host interactions potentially provide a molecular basis for selection and an accelerated course of illness in blood group A–positive hosts.
challenged with 1 to 2 × 10^7 colony-forming units of ETEC H10407 and was ETEC-naïve prior to experimental challenge. All subjects were treated with antibiotics upon meeting early treatment criteria (Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/JCI97659DS1) or at 120 hours after inoculation to clear their infections.

Interestingly, we found that development of severe diarrheal illness was significantly (P = 0.015) accelerated in individuals with blood group A relative to those in non-A blood groups (Figure 1A), with volunteers in the A blood group requiring early initiation of antibiotic therapy more frequently (72%) than those in blood group B or O (43%). Despite earlier antibiotic administration, maximal diarrheal volumes over 24 hours, total diarrheal volume prior to treatment, and the frequency of diarrheal episodes tended to be higher in A versus non-A individuals (Supplemental Figure 1, A–D). Notably, individuals with blood groups B or O were considerably more likely to remain free of diarrhea or have mild diarrhea (44%) following challenge than volunteers expressing the A blood group (19%) (Table 1). Conversely, of the 106 subjects, 81% of those expressing the A blood group antigen (type A or AB) had moderate-to-severe diarrhea (MSD) compared with 56% of individuals with non-A (B or O) blood types (P = 0.017), and the overall relative risk of MSD among individuals with the A blood group compared with the non-A blood groups was 1.44 (95% CI = 1.1–1.9, P = 0.009) (Figure 1B and Supplemental Table 2). These data therefore suggest that, upon challenge with H10407, blood group A volunteers (Supplemental Tables 3 and 4) are markedly more likely to manifest the severe, cholera-like spectrum of illness originally associated with this isolate.

**Table 1. Diarrheal illness and severity by blood group**

<table>
<thead>
<tr>
<th>Diarrheal severity</th>
<th>O or B</th>
<th>A or AB</th>
<th>Total subjects (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None to mild</td>
<td>35 (44)</td>
<td>5 (19)</td>
<td>40 (38)</td>
</tr>
<tr>
<td>MSD</td>
<td>44 (55)</td>
<td>22 (81)</td>
<td>66 (62)</td>
</tr>
<tr>
<td>Total subjects</td>
<td>79</td>
<td>27</td>
<td>106</td>
</tr>
</tbody>
</table>

\( ^a \text{Pearson } \chi^2 = 5.694, 1 \text{ df, } P = 0.017. \)

EtpA interacts specifically with blood group A glycans. Pathogens use adhesins possessing glycocalyx-specific lectin activity (46) to engage blood group antigens expressed on mucosal surfaces, including the gastrointestinal tract (47). The apical surface of intestinal epithelial cells is heavily decorated with a variety of glycoproteins that form the glyocalyx (48), and glycosylation patterns of these proteins are in part determined by blood group (47, 49). Because earlier studies had suggested that EtpA, a secreted adhesion molecule discovered in H10407 (25), engages the surface of gastrointestinal epithelia, we conducted glycan-array screening to further elucidate the nature of EtpA-mediated bacterial adhesion. Using recombinant EtpA (rEtpA) to probe arrays containing more than 400 unique glycan species, we found that rEtpA interacted almost exclusively with glycans related to blood group A. In contrast, we observed little or no binding to other glycans, including those of blood group B or blood group O (Figure 2A and Supplemental Data Set 1). Next, in biolayer interferometry experiments, we found that EtpA interacted with blood group A–related glycans with high affinity when the glycan was presented as a multivalent trisaccharide conjugated to polyacrylamide (PAA): GalNACα1-3 (Fucα1,2)Galβ1-PAA, where Gal indicates galactose; kD, ~9 × 10^-8 M (Figure 2B). Similarly, we demonstrated that EtpA can engage N-acetylgalactosamine (GalNAc) (kD, ~1.6 × 10^-8 M), the terminal sugar in blood group A, with higher affinity than Gal (kD, ~1.9 × 10^-7), the terminal sugar of blood group B (Figure 2C). Collectively, these studies suggest that the EtpA adhesin is a blood group A–specific lectin.

Many bacterial adhesins possess the ability to agglutinate RBCs. Included among these adhesins is filamentous hemagglutinin from Bordetella pertussis (50), a component of acellular pertussis vaccines (51), which, like EtpA, belongs to the 2-partner family of bacterial exoproteins (25). We found that EtpA could decorate the surface of human RBCs belonging to blood group A1 (Figure 3A) and, to a lesser extent, cells from those individuals from blood group A2 who exhibit lower levels of the glycosyltransferase enzyme activity required for addition of terminal GalNAc residues that determine the A blood group (52); however, only a small percentage of blood group B or O erythrocytes had detectable EtpA on the surface. Similarly, in RBC pull-down assays, more rEtpA remained associated with erythrocytes from the A1 and A2 blood groups (Figure 3B) than those from the B or O blood groups. In addition, incubation of EtpA with RBCs from blood group A resulted in agglutination, but yielded poor agglutination activity with other RBCs (Figure 3C). In addition, pretreatment of target A1 cells with α-N-acetylgalactosaminidase completely abrogated the hemagglutinin activity of EtpA (Figure 3D) as did addition of excess GalNAc, but not GlcNAc or Gal, the terminal sugar of blood group B (Supplemental Figure 2). We also demonstrated that latex microparticles coated with EtpA specifically agglutinated A1 RBCs (Figure 3E). EtpA therefore belongs to a large family of bacterial adhesins that possess lectin/hemagglutinin activity (46), and this protein preferentially targets GalNAc residues presented in the context of the A blood group.

EtpA is a predominant blood group A–binding partner in ETEC. Next, to determine whether EtpA is the predominant blood group A–binding partner in ETEC, we examined interaction of A blood group glycans with subcellular protein fractions isolated from H10407. Blood group A glycans interacted only with a high molecular weight species (in excess of 150 kD) in culture supernatants, with no detectable interaction with preparations from the inner or outer membranes (Figure 4A). Because the EtpA adhesin is the only known ETEC-specific virulence molecule in the ETEC secretome of this size (53), we next compared binding of blood group A glycans to secreted proteins from the WT H10407 strain and isogenic etpA mutants. Binding occurred only in supernatants prepared from WT ETEC or the complemented *etpA* mutant strain, but not the *etpA* isogenic deletion mutant (Figure 4, B–D). Interestingly, blood group A erythrocytes have been traditionally used to screen for the presence of CFA/I fimbriae in ETEC (54). However, we found that CFA/I-mediated nonspecific hemagglutination of RBCs regardless of blood type, a result that is consistent with the idea that CFA/I interactions are mediated via sialic acid (55) (Supplemental Figure 3). To further confirm the role of EtpA in mediating interaction with the A blood group, we next tagged the ETEC proteome using the
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Glycans on the surface of intestinal enterocytes might therefore direct EtpA-mediated interactions required for effective bacterial adhesion and toxin delivery. Using HT-29 intestinal cells derived from a blood group A–positive host, we found that EtpA bound to the surface of these cells and colocalized with blood group A glycans (Figure 5A). Conversely, rEtpA bound very poorly to the surface of HT-29A–/– cells, an HT-29–derived line that was CRISPR-Cas9 engineered to eliminate the gene encoding the glycosyltransferase responsible for the addition of terminal GalNac residues to the core H (blood group O) antigen (45). Likewise, we found that EtpA from ANL-tagged WT ETEC interacted preferentially with HT-29 parental cells compared with the mutant HT-29A–/– cells (Figure 5B). Interestingly, EtpA-expressing ETEC localized to regions of blood group A expression on the surface of HT-29 cells (Figure 5C), and EtpA-expressing ETEC adhered noncanonical amino acid azidonorleucine (ANL) (56) and incubated tagged supernatant and outer membrane protein (OMP) fractions with erythrocytes. Again, we found that a single band from tagged WT supernatant, corresponding to EtpA, bound appreciably only to A1 erythrocytes (Figure 4E). However, testing of ANL-labeled OMPs purified from H10407 failed to detect any specific interactions of ETEC OMPs with RBCs of any major blood group (Supplemental Figure 4), further suggesting that the secreted EtpA molecule is the dominant blood group A–specific lectin expressed on the surface of ETEC. We also demonstrated that EtpA expression by a geographically diverse group of ETEC conferred the ability to bind blood group A, suggesting that these findings are not limited to the H10407 prototype strain (Supplemental Figure 5, A–D).

**EtpA directs ETEC interaction with blood group A on host intestinal epithelial cells.** We questioned whether blood group A–related glycans on the surface of intestinal enterocytes might therefore direct EtpA-mediated interactions required for effective bacterial adhesion and toxin delivery. Using HT-29 intestinal cells derived from a blood group A–positive host, we found that EtpA bound to the surface of these cells and colocalized with blood group A glycans (Figure 5A). Conversely, rEtpA bound very poorly to the surface of HT-29A–/– cells, an HT-29–derived line that was CRISPR-Cas9 engineered to eliminate the gene encoding the glycosyltransferase responsible for the addition of terminal GalNac residues to the core H (blood group O) antigen (45). Likewise, we found that EtpA from ANL-tagged WT ETEC interacted preferentially with HT-29 parental cells compared with the mutant HT-29A–/– cells (Figure 5B). Interestingly, EtpA-expressing ETEC localized to regions of blood group A expression on the surface of HT-29 cells (Figure 5C), and EtpA-expressing ETEC adhered noncanonical amino acid azidonorleucine (ANL) (56) and incubated tagged supernatant and outer membrane protein (OMP) fractions with erythrocytes. Again, we found that a single band from tagged WT supernatant, corresponding to EtpA, bound appreciably only to A1 erythrocytes (Figure 4E). However, testing of ANL-labeled OMPs purified from H10407 failed to detect any specific interactions of ETEC OMPs with RBCs of any major blood group (Supplemental Figure 4), further suggesting that the secreted EtpA molecule is the dominant blood group A–specific lectin expressed on the surface of ETEC. We also demonstrated that EtpA expression by a geographically diverse group of ETEC conferred the ability to bind blood group A, suggesting that these findings are not limited to the H10407 prototype strain (Supplemental Figure 5, A–D).
more readily to parental cells expressing blood group A (Figure 5, D and E, and Supplemental Figure 6A). Importantly, we found that optimal delivery of LT toxin required both EtpA (Supplemental Figure 6B) and the presence of the A blood group antigen (Figure 5F). Similarly, we noted that addition of exogenous GalNAc impaired both EtpA-mediated adhesion and toxin delivery to HT29 parental cells (Supplemental Figure 7, A and B), suggesting that this terminal glycan on blood group A is important for EtpA pathogen–epithelial cell interactions. Notably, while purified LT was significantly less potent than cholera toxin (CT) in activating cAMP in target cells, direct toxin-mediated activation of this cyclic nucleotide was actually enhanced in cells lacking the A blood group (Supplemental Figure 8), recapitulating recent observations with CT (45). Collectively, these data suggest that the presence of blood group A effectively promotes bacterial adhesion and ETEC-host interactions that culminate in enhanced toxin delivery.

HT-29 cells are derived from a metastatic colorectal carcinoma, and glycan distribution can vary significantly with location within the gastrointestinal tract (57). In addition, ETEC are thought to preferentially target the small intestine, where the bulk of fluid secretion occurs following colonization, rather than the colon.

We sought, therefore, to confirm these interactions using small intestinal enteroids derived from human stem cells obtained from individuals belonging to each of the major ABO blood groups. Using ileal enteroids, we found that, while EtpA bound very poorly to cells expressing either B or O blood groups (Figure 6, A and B). We also found EtpA-expressing bacteria within areas of blood group A expression on the epithelial surface of polarized enteroids (Figure 6C), and interestingly, we also observed apparent rearrangement or “footprints” of the A blood group at sites of bacterial attachment, suggesting an intimate interaction of ETEC with these cell-surface glycans (Figure 6D).

Finally, we found that ETEC adhered much more readily to polarized intestinal monolayers derived from individuals with blood group A compared with other blood groups (Figure 6E) and that adhesion of WT ETEC to A blood group enteroids was significantly accelerated relative to that of bacteria lacking etpA (Figure 6F), culminating in enhanced toxin delivery by the EtpA-expressing WT (Figure 6, G and H). EtpA therefore appears to be a dominant pathovar-specific lectin uniquely capable of preferentially engaging A blood group glycans expressed on small intestinal epithelial surfaces to accelerate effective ETEC pathogen-host interactions.

Discussion

Blood group–dependent microbial-host interactions can dictate clinical outcomes of infection with important gastrointestinal pathogens. In the case of *Helicobacter pylori*, the BabA adhesin facilitates attachment to the gastric mucosa by engaging ABO (60) and Le^b^, blood group antigens potentiating the delivery of other secreted virulence factors that lead to development of gastric ulcers and cancer (44, 60, 61). Similarly, the clear epidemiologic

**Figure 3. EtpA is a blood group A–specific hemagglutinin.**

(A) Binding of rEtpA to the surface of group A1 and A2 RBC ghosts relative to B and O RBCs is shown in FACS data and a graph. (B) RBC pull-down of rEtpA with different blood group ghost erythrocytes. Shown in the immunoblot is rEtpA-myc-his identified by anti-myc antibodies. (C) EtpA agglutinates A1 RBCs, but exhibits minimal or no hemagglutination activity with RBCs from those with A2, B, or O blood groups. The column on the right includes anti–A blood group antibodies as a positive control. (D) EtpA-mediated hemagglutination is dependent on terminal GalNAc residues. A1 RBCs pretreated with N-acetylgalactosaminidase (top row) fail to agglutinate in the presence of rEtpA. Shown (row 2, left) are positive controls (no enzyme pretreatment) and blood group O–negative controls (bottom rows). (E) EtpA-coated latex microspheres (top row) agglutinate RBCs from blood group A, but not B or O. Bottom row shows BSA-coated particles as controls. Each of the images are representative of 3 experimental replicates. Original magnification, ×40.
associations between severe *V. cholerae* infections and the O blood group (62–66) are reflected in recent observations that CT preferentially activates enterocytes of blood group O lineage (45). Both CT and the closely related LT toxin of enterotoxigenic *E. coli* appear to share in a differential binding of blood group antigens that favors intoxication of blood group O enterocytes (67, 68). Interestingly, because of the molecular similarity between CT and LT, some earlier field studies examined ETEC outcomes relative to blood group with the implicit assumption that LT-producing ETEC would lead to more severe disease following infection of O blood group hosts; however, these studies yielded conflicting results (69, 70).

In contrast, from the human volunteer ETEC challenge studies, a surprising, but clear, relationship between the A blood group and disease severity began to emerge. The ETEC controlled human infection model has a number of potential advantages over prior field studies (3, 62, 69, 70) in defining a molecular basis for disease. First, the present studies were confined to ETEC-naive adults challenged with a defined inoculum of a single strain (H10407) isolated from an individual with severe cholera-like illness (41). The controlled human infection model may therefore permitted us to unmask somewhat subtle molecular associations that might have otherwise been hidden by confounding variables common to field studies of enteric infections in developing countries, including malnutrition, breast feeding, altered microbiota, and coinfection with other enteric pathogens as well as the genetic variability inherent in the infecting ETEC strains.

The studies presented here provide an expanded view of ETEC pathogen-host interactions beyond a classical paradigm in which these organisms engage the intestinal surface via fimbrial colonization factors and deliver LT and/or ST enterotoxins. The present studies of ETEC paint a more complex picture in which A blood group expression potentially facilitates EtpA-mediated lectin-specific bacterial interactions with intestinal epithelia that culminate in effective toxin delivery and symptomatic diarrheal illness.

Our studies suggest that the many ETEC strains that produce the EtpA adhesin (27, 29) may be particularly well equipped to preferentially infect hosts who express the A blood group antigen on mucosal surfaces. When infected with H10407, an EtpA-producing strain, immunologically naive subjects with blood group A appear to be at significantly increased risk for diarrheal illness, and the ensuing diarrhea tends to be more severe relative to the other major blood groups. Intriguingly, the ETEC H10407 strain, isolated from a case of severe cholera-like diarrheal illness more than 40 years ago, is typically among the most virulent strains to be tested in human-challenge studies (43, 71).

It is likely however, that while EtpA leads to enhanced virulence in blood group A hosts, other putative virulence factors identified in H10407, including the mucin-degrading EatA protease (19), collectively augment virulence by accelerating toxin delivery that culminates in severe diarrhea.

Several additional features of our studies deserve comment. We were not able to examine “secretor status,” i.e., expression
Because approximately 20% of the population are nonsecretors due to defects in FUT2 (72), it is likely that some of the A blood group–positive individuals did not express the A antigen on their gastrointestinal epithelia. Theoretically, this might account for some individuals in the A blood group who failed to become ill following challenge. Conversely, while the A blood group is a risk factor in this model, the development of severe diarrheal illness among non-A volunteers demonstrates that expression of A glycans is not an absolute requirement for disease.

*V. cholerae* infections provide an important analogous paradigm for severity of illness related to blood group. While severe cholera is significantly associated with the O blood group, it is not exclusively observed in blood group O individuals (66). We should also note that the ETEC-challenge studies were done with a relatively high inoculum of bacteria (~10^7) designed to assure a reasonable attack rate (73), a feature that may not be reflected in naturally occurring infections in young children where the A blood group also appears to be a risk for development of symptomatic infections (3). It is also worth noting that volunteers in these studies were treated with antibiotics to clear the infection once they met the definition of severe diarrheal illness, a feature that might also have affected the ultimate outcome.

The secreted EtpA lectin likely acts in concert (21) with at least 2 additional glycan-binding tip adhesins of ETEC fimbriae with different substrate specificities. These include FimH of the chromosomally encoded type 1 pili, which binds to mannosylated surface proteins on epithelial cells (74), and the CfaE adhesin located at the tips of canonical plasmid-encoded CFA/I pili, which putatively binds to sialylated glycoproteins (55, 75). The latter is consistent with the observation that ETEC expressing CFA/I mediates mannose-resistant (54) and neuraminidase-sensitive (75, 76) hemagglutination in all major blood groups and across species to include chicken and bovine erythrocytes. Therefore, it appears that EtpA is the only blood group A–specific lectin identified in ETEC to date. Nevertheless, we cannot rule out the possibility that our studies failed to identify another blood group A lectin that was not efficiently expressed under laboratory culture conditions.
cification of EtpA as blood group A lectin could focus antigen development and inform vaccine design to specifically protect individuals that are at highest risk for severe cholera-like diarrheal illness.

Methods

**ETEC controlled human infection volunteer challenge studies.** Samples for blood group analysis were obtained from a convenience sample of 106 human volunteers from 4 prior independent controlled human infection model ETEC challenge studies conducted at the Johns Hopkins Center for Immunization Research (ClinicalTrials.gov NCT01739231, NCT01060748, NCT00844493, and NCT01922856). All volunteers were experimentally infected with 1 to 2 × 10⁷ colony-forming units of the EtpA-expressing ETEC strain H10407. Subjects in these studies

Figure 6. EtpA and EtpA-expressing ETEC interact preferentially with the surface of blood group A enteroids. (A) Confocal imaging of EtpA binding to surface of polarized small intestinal enteroid monolayers from a blood group A subject. Differential interference contrast (DIC) image at bottom shows architecture of polarized enteroids on Transwell filter. Original magnification, ×63. (B) Kinetic ELISA quantification of biotinylated EtpA binding to surface of small intestinal enteroid cells from blood groups A, B, and O (n = 4 technical replicates representative of 3 separate experiments). *P < 0.05, **P < 0.01, ANOVA, Kruskal-Wallis. (C) Association of EtpA-expressing ETEC H10407 with A blood group on surface of small intestinal cells. Images represent velocity-processed confocal laser-scanning microscopy (CLSM) data of ETEC H10407 (red) to blood group A (green) and nuclei (blue). Lower panel shows polarized orientation of cells with bacteria adherent to the apical surfaces of enterocytes. Original magnification, ×63. (D) Top panel: CLSM of ETEC (green) and A blood group (red) on jejunal enteroids. Bottom panel: A blood group “footprints” (arrows) at sites of bacterial attachment. Data shown in C and D are representative of 3 experimental replicates. Original magnification, ×60. (E) Bacterial density on surface of small intestinal enteroids determined by CLSM quantification of H10407 (serotype O78). Quantitation based on imaging 10 fields per blood group, at ×20 magnification. *P = 0.039, **P < 0.0001, ANOVA, Kruskal-Wallis. (F) Adhesion of WT H10407 and etpA mutant bacteria to blood group A small intestinal enteroids (n = 5 technical replicates; representative of 3 biological replicates). Bars represent geometric mean values. (G) Production of cAMP by blood group A target enteroids following infection with WT (H10407) or the etpA mutant. Data represent total of 5 technical replicates from 2 separate experiments. (H) Production of cAMP by blood group A small intestinal enteroids following infection with WT (n = 6) or the etpA mutant (n = 4). Data are representative of 3 independent biological replicates. Inset graph indicates relative (mean ± SD, n = 6 technical replicates) production of heat stable toxin (ST) by the WT (blue bars) vs. the mutant (gray bars). **P < 0.01, Mann-Whitney U test, 2 tailed nonparametric testing.

Interestingly, findings presented here are in keeping with earlier birth-cohort studies of children under the age of 2 years in Bangladesh who also demonstrated an association between the A blood group and symptomatic ETEC infection (3). Therefore, while our recent studies have potentially important clinical and vaccinology implications, further study of the relationship among blood group, disease severity, and antigen expression could guide and inform use of these antigens in vaccines.

These additional molecular details regarding the nature of ETEC pathogen-host interactions could promote an improved understanding of the virulence elements and host determinants related to the most severe forms of illness that are clinically indistinguishable from severe cholera (15–17). Consequently, identification of EtpA as blood group A lectin could focus antigen development and inform vaccine design to specifically protect individuals that are at highest risk for severe cholera-like diarrheal illness.
were treated with antibiotics once they met previously defined end-point criteria (summarized in Supplemental Table 1) or at 120 hours after inoculation to clear their infections. Additional details of the of the study design are included in the Supplemental Methods.

Glycan-array screening. Glycan arrays containing 411 separate features were fabricated as previously reported, except for the addition of a washable fluorescent dye, the free acid of DyLight 649 (0.7 μg/ml, Thermo Scientific), to the print buffer as an indicator of successful liquid deposition and spot morphology. The array format and assay have been described previously (77) along with analysis of reproducibility (78) and validation with numerous antibodies and lectins (79, 80). Slides were blocked overnight at 4°C with 3% BSA (w/v; 200 μl/well) in PBS and then washed 6 times with PBST (PBS with 0.05% Tween 20; 200 μl/well). Polystyrene-tagged rEtpA-6His was diluted to final concentrations of 20 and 200 μg/ml in PBST buffer containing 3% BSA (MilliporeSigma) and 1% HSA (MilliporeSigma), and then 50 μl of each sample was added into 2 different wells on different slides. Incubation buffer alone served as the negative control for samples analyzed in duplicate. After incubation with rEtpA for 2 hours at 37°C in an orbital shaker (100 rpm), arrays were washed and then probed with mouse monoclonal IgG anti-His, antibody (Qiagen, 24660) diluted 1:500 to a final concentration of 0.4 μg/ml (1 hour at 37°C; 100 rpm), followed by DyLight 549-conjugated affinity purified goat anti-mouse IgG (Jackson ImmunoResearch, 115-505-071; 1 hour at 37°C; 100 rpm). After washing 7 times with PBST (200 μl/well), slides were immersed in wash buffer for 5 minutes and centrifuged at 200 g for 5 minutes. Slides were scanned at 10 μm resolution with a GenePix 4000B Microarray Scanner (Molecular Devices Corp.) with excitation/emission wavelengths of 532/575 nm and analyzed with GenePix Pro 6.0 software, as previously reported (77). Heatmaps were assembled in R version 3.2.2 (The R Foundation for Statistical Computing), using the gplots and RColorBrewer packages made available through the Comprehensive R Archive Network (CRAN; http://cran.wustl.edu/). Additional details of the glycan-array screening can be found in Supplemental Methods.

Bacterial strains and culture conditions. ETEC strain H10407 was provided from GMP stock produced at the Walter Reed Army Institute of Research (Silver Spring, Maryland, USA). The etpA mutant H10407 (jf1668) and complemented (jf1697) strains were generated in previous studies. ETEC strains were routinely grown in Luria broth (LB) or LB agar (2% agar) at 37°C from glycerol stocks preserved at -80°C. Bacterial strains and plasmids used in this study are listed in Supplemental Table 5. Strains 100137, 200145, 30051, 400643, 500632, 601142, and 700241 were provided by David Rasko (Institute for Genome Sciences, University of Maryland School of Medicine, Baltimore, Maryland, USA). Strain 12005000131-1 was provided by Elizabeth Cebelinski (Molecular Epidemiology Section of the Infectious Diseases Laboratory, Minnesota Department of Health, St. Paul, Minnesota, USA).

Protein expression and purification. The 2-partner secreted EtpA glycoprotein was purified from culture supernatants of E. coli Top10 coexpressing plasmids pJL017 and pJL030 (81). Briefly, E. coli Top10 carrying pJL017 and pJL030 (Supplemental Table 5) was grown in LB with ampicillin (100 μg/ml) and chloramphenicol (15 μg/ml) at 37°C, in a shaking incubator at 230 rpm, until OD 600 nm reached approximately 0.5–0.6. Recombinant protein expression was then induced with 0.0002% arabinose for 6 hours at 37°C. Culture supernatant was concentrated using a 100 kD Molecular Weight Cutoff (MWCO) filter (Millipore), and rEtpA-myc-6His protein was purified using immobilized metal affinity chromatography as described previously (81).

ETEC subcellular fractionation and far-Western analysis. ETEC H10407 was grown overnight in 10 ml cultures of LB at 37°C as described above, diluted into 500 ml cultures the next morning, and grown for 3 hours at 37°C. Cells were harvested by centrifugation at 5000 g for 10 minutes at 4°C and the pellet resuspended in 8 ml 50 mM Tris buffer pH 7.8 containing 1 mM EDTA and 1 mg DNAse. Subcellular fractions were then prepared as previously described (82). Briefly, cells were disrupted using a French press, followed by the addition of 2 mM MgCl2 to E. coli lysate and centrifugation at 5000 g for 10 minutes at 4°C to remove large debris/intact cells. Supernatant was collected, supplemented with PMSF to a final concentration of 1 mM, and ultracentrifuged at 100,000 g for 1 hour at 4°C. Supernatant from this step was saved as a cytosolic fraction. The remaining cell pellet was resuspended in ice-cold 10 mM HEPES pH 7.4, layered over a gradient of 69%, 52.8%, and 26.4% sucrose in HEPES pH 7.4 and ultracentrifuged at 113,000 g for 16 hours at 4°C. The visible upper layer containing inner membranes and the lower high-density layer with outer membranes were collected. Membrane proteins were reconstituted in ice-cold HEPES pH 7.4 and centrifuged for 1 hour at 100,000 g, 4°C. Finally, the protein pellets from the inner and outer membranes were resuspended in 0.25 and 0.5 ml dH2O, respectively.

Far-Western blotting was used to test blood group A glycan interaction with ETEC proteins. Briefly, approximately 10 μg of ETEC proteins were resolved by SDS-PAGE on NuPAGE 4%-20% Tris-Glycine Gel (Life Technologies) and transferred to nitrocellulose membranes. Membranes were blocked with 5% BSA in TBS-T (Tris[hydroxymethyl]aminomethane–buffered saline, 0.01% Tween-20) for 1 hour and then incubated with a final concentration of 5 μg/ml of biotinylated multivalent blood group A trisaccharide (Glycotech; 01-032) overnight at 4°C. Biotinylated glycans were detected using streptavidin–HRP (SA-HRP) conjugate (1:20,000), and EtpA was detected by immunoblotting using anti-EtpA rabbit polyclonal antibody.

Hemagglutination studies. We examined the ability of EtpA to agglutinate human erythrocytes belonging to blood groups A1, A2, B, and O (Immucor, catalog 0002338). Briefly, 2-fold dilutions of rEtpA in PBS in a final volume of 10 μl were mixed with 40 μl of approximately 1% cell suspensions (in PBS) of RBCs expressing the A1, A2, B, and O antigens in 96-well round-bottom plates, then incubated for 30 minutes at 4°C. Plates were then imaged to show agglutinated versus nonagglutinated (settled) RBCs. In parallel experiments, recombinant polystyrene-tagged EtpA was immobilized onto cobalt-impregnated super-paramagnetic polysacrylate beads (Dynabeads TALON, Thermo Fisher, catalog 101.01D) that were then incubated with RBCs.

To examine the specificity of EtpA-induced agglutination of RBCs, we used a protocol similar to that previously described by Evans (54). Briefly, bacteria were grown overnight on CYE agar (1% casein acids; BD), 0.15% yeast extract (BD),
and the extracts adjusted to 2 mM MgCl₂ and 100 μM of DNase I (MilliporeSigma) was added to each tube. RBCs were resuspended in 20 μl ice-cold lysis buffer (50 mM Tris, 1 mM EDTA, pH 7.8) and lysed by sonication (4 × 20-second bursts). Approximately 1 mg of DNase I was added to each tube.

For FACS analysis, 15 μg biotinylated EtpA or BSA was added to 190 μl of RBC ghosts and rotated at RT for 1.5 hours. Cells were washed twice in PBS and pelleted by spinning at 25,000 g for 5 minutes, rotated for 30 minutes with a 1:200 dilution of SA-conjugate nanocrystals (Qdot 525, QI0141MP, Thermo Fisher), and then washed twice in PBS and resuspended in 200 μl of PBS. Fluorescence was measured using FlowJo software (BD Biosciences), and data were analyzed using FlowJo software, v10.

Molecular cloning of mutant Met-RS. To permit noncanonical amino acid labeling of ETEC proteins with azidonorlucine (Anl), a mutant form of the methionyl-tRNA synthetase (NLL-MetRS) gene from pAM1 (56) was cloned (In-Fusion, Clontech) into the NcoI and EcoRI sites of pBAD/Myc-His B (Supplemental Table 5) using primers jf091715.1 and jf091715.2 (Supplemental Table 6). The resulting plasmid pBAD-MetRS was transformed into WT H10407 or the jf1668 etpA mutant to generate jf3422 and jf3904, respectively.

Protein interaction studies using noncanonical amino acid labeling of ETEC. Proteomes of strains jf3422 and jf3904 bearing the MetRS expression plasmid were labeled with growth with azidonorlucine as previously described. Briefly, strains were grown overnight in LB containing 100 μg/ml ampicillin, 0.2% arabinose, and 1 mM azidonorlucine (HAA1625, Iris Biotech GmbH). Cell-free supernatants were concentrated 10-fold using 10 kD MWCO centrifugal concentrators and dialyzed against PBS prior to use.

OMPs were prepared from ANL-tagged bacteria grown in the same fashion using an established protocol (83). Briefly, labeled bacteria were resuspended in 20 ml ice-cold lysis buffer (50 mM Tris, 1 mM EDTA, pH 7.8) and lysed by sonication (4 × 20-second bursts). Approximately 1 mg of DNase I (MilliporeSigma) was added to each tube and the extracts adjusted to 2 mM MgCl₂ and 100 μM PMSF. Intact cells and large debris were removed by low-speed centrifugation at 3,000 g for 10 minutes, and membranes were pelleted from clarified lysates by ultracentrifugation for 1 hour at 100,000 g. Membrane pellets were then resuspended in 2 ml 10 mM HEPES pH 7.4 and layered onto discontinuous sucrose gradients consisting of a top layer of 0.77 M sucrose, a middle layer of 1.44 M sucrose, and a bottom layer of 2.02 M sucrose. After centrifugation at 113,000 g for 16 hours at 4°C, the OMPs were recovered from the interface between the 1.44 and 2.02 M layers. OMP pellets were washed by resuspension in 14 ml 10 mM HEPES pH 7.4, collected by centrifugation at 100,000 g for 1 hour, and resuspended in 0.5 ml PBS.

Erythrocyte membranes were prepared using the method previously described by Steck et al. (84). Briefly, suspensions of RBCs were sedimented at 2,300 g for 10 minutes at 4°C, the pellet recovered, and then resuspended in 5 volumes of cold PBS, pH 8.0. After centrifugation, packed RBCs were hemolyzed by mixing with 5 mM sodium phosphate, pH 8.0, and incubating on ice for 1 minute. The resulting membranes were then harvested by centrifugation at 22,000 g for 10 minutes and the supernatant removed by aspiration.

HT-29 membranes or membranes from the HT-29A⁻ line were prepared as follows. Cells grown in monolayers on a 725 flask were washed with PBS and the supernatant decanted. After addition of 4 ml fresh PBS, cells were recovered by scraping and the suspension centrifuged at 5 minutes at 4,200 g. Cell pellets were resuspended in 1 ml of ice-cold lysis buffer (5 mM PO4, pH 8, 0.5 mg/ml DNase I, containing Pierce Protease Inhibitor; Thermo Fisher, 88665) and transferred to a microfuge tube spun for 5 minutes at 3,400 g. Supernant was removed and replaced with 1 ml fresh lysis buffer. Cells were then incubated on ice for 30 minutes, flash-frozen in an ethanol-dry ice bath, and thawed at 37°C. After repeating the freeze-thaw cycles twice, the extract was passed through a 26-gauge needle 4 to 5 times to shear DNA and spun at 25,000 g for 10 minutes at 4°C. The resulting translucent membrane pellets were then resuspended in PBS.

Erythrocyte ghosts or HT-29 membrane preparations were incubated with ANL-labeled bacterial proteins at 4°C overnight, pelleted by centrifugation, washed in 1 ml of ice-cold PBS, then resuspended and boiled in click buffer (50 mM HEPES, 0.5% [w/v] SDS, pH 7.5). ANL-containing proteins were labeled for 1 hour at 37°C in the dark with 5 μM tetramethylrhodamine-alkyne (TAMRA-alkyne) (Molecular Probes) in the presence of 25 mM ascorbate, 1.5 mM CuSO₄, and 2.5 mM Tris (3-hydroxypropyltriazolymethyl)amine. Labeled proteins were separated by SDS-PAGE in the dark and the gel scanned on a Typhoon FLA 9500 phosphorimager (GE Healthcare Life Sciences) using excitation at 532 nm and a 575 nm long-pass filter. Positive control TAMRA-labeled rEtpA was produced by reacting EtpA with 120 μM azidobutyric acid NHS ester (Lumiprobe). The unreacted label was then removed by dialysis, and the azide-labeled EtpA reacted with TAMRA-alkyne, as before.

Bio-layer interferometry. Bio-layer interferometry was used to determine affinity of EtpA-blood group binding using Octet Red96 (Pall ForteBio Corp.). Briefly, biotinylated blood group A antigens conjugated to PAA (either as the disaccharide GalNAcβ3Galβ-PAA-biotin; GlycoTech, catalog 01-017; or the trisaccharide GalNAcα3[Fucol,2]Galβ-PAA-biotin, GlycoTech 01-032) or related target sugars (α-Gal-PAA-biotin, GlycoTech 01-003; GalNAcα3GalNAcβ-PAA-biotin, GlycoTech 01-026; αGalNAc-PAA-biotin, GlycoTech 08-010) were immobilized onto the SA biosensors (18-5019; Pall ForteBio Corp.). Those were then incubated with 2-fold serial dilutions (2000–500 nM) of purified rEtpA or control BSA (2000 nM) prepared in 1× PBS as analyte. Affinity constant (kD) values were calculated using Octet software version 8.1.

Growth and differentiation of human enteroids from different blood groups. Enteroid experiments were conducted at either John Hopkins University School of Medicine or at Washington University School of Medicine, and detailed experimental protocols for both institutions are included in the Supplemental Methods. In brief, biopsy samples were obtained from adult patients undergoing routine endoscopy at each institution. Small intestinal epithelial (enteroid) cell lines were then generated from dissociated crypts as previously described (85–87), stored, and then accessed from the Washington University Digestive Diseases Research Core Center Biospecimens Core or the respective aligned program at John Hopkins University School of Medicine.
Purified cell lines of each of the described ABO types were thawed and resuspended in Matrigel (BD Biosciences) and then propagated at 37°C, 5% CO₂ with advanced DEM/F12 media (Invitrogen) supplemented with Wnt3a, R-spondin 1, and Noggin (WRN). Protocols for cellular differentiation and growth of polarized epithelial monolayers were performed as described in detail in the Supplemental Methods. Blood typing by PCR and immunofluorescence microscopy was performed as previously described.

**ol-13-N-acetylgalactosaminyltransferase-deficient cell line.** Cells lacking ol-13-N-acetylgalactosaminyltransferase (GI 5573627; GenBank D82843.2) were constructed from HT-29 cells (ATCC HTB-38) by CRISPR/Cas9 genome editing in the Genome Engineering and iPSC Center at Washington University (http://geic.wustl.edu) as previously reported (43). Both parental HT-29 cells and the resulting HT-29A- line were propagated in McCoy’s-5A medium (Gibco, Life Technologies) supplemented with 10% FBS.

**ETEC adhesion and toxin delivery assays.** To examine delivery of the LT toxin, cultures of HT-29 or HT-29A- were differentiated by growing cultures at confluence for 1 week prior to use in 96-well plates. Where indicated, phosphodiesterase (PDE) inhibitors vardeparin (Sigma, PHR1503), and rolipram (MilliporeSigma, R6520) were added in the figure legends.

**Confocal laser scanning microscopy.** HT-29 cells were seeded onto glass coverslips pretreated with poly-L-lysine in 24-well tissue culture plates. Enteroids grown on Transwell filters were processed by removing the filter from the well and then treated as below. Cells were blocked with 1% BSA in PBS and incubated with biotinylated EtpA (50 μg/ml final concentration) for 2 hours at 37°C or overnight at 4°C. After washing with PBS 3 times, EtpA was detected using Qdot 605 SA conjugate (Life Technologies). A blood group antigen was detected using mouse monoclonal antibody (IgM; Santa Cruz Biotechnology Inc., sc-69951) against human blood group A, followed by Alexa Fluor 647-conjugated goat anti-mouse IgM heavy chain (Molecular Probes, A21238). CellMask deep red plasma membrane stain (Thermo Fisher Scientific, C10046) (1:2,000) and DAPI (1:6,000) were used to stain cellular membranes and nuclei, respectively. ETEC H10407 (serotype O78:H11) was visualized with polyclonal rabbit antibodies against O78 obtained from the Penn State University E. coli Reference Center, followed by detection with goat anti-rabbit IgG (H&L) cross-absorbed Alexa Fluor (488 or 594) secondary antibody conjugates (Invitrogen, A11070, A11072).

Confocal microscopy images were captured using a Zeiss LSM 510 Meta Confocal Laser Scanning Microscope. Image files were saved in LSM format and processed using Fiji (89), Image J v 2.0.0 (NIH), and Volocity 3D analysis software (version 6.3, PerkinElmer).

**Blood group typing.** Coded, deidentified samples forwarded to Washington University were used in a blinded fashion to type for the major ABO blood group antigens. Plasma or serum was used in reverse immunoagglutination for antibodies directed at the ABO antigens, which was performed using standard methods. Briefly 2 μl RBCs (Immucor) from each blood group were incubated with 4 μl of patient sera or plasma at RT for 5 minutes in a 96-well plate, then examined microscopically for agglutination. For available saliva, DNA was extracted from 50 μl of sample (Qiagen 69504), and genotyping by PCR was performed as previously described (90). Salivary genotyping results were confirmed by dot immunoblotting. Briefly, 0.5 μl saliva diluted 1:4 in PBS was dotted onto nitrocellulose membranes that were then incubated in a solution of H₂O₂ (3%) for 5 minutes. Membranes were then incubated with 1:20 dilutions of anti-A, anti-B, or anti-H antibodies (MilliporeSigma) for 30 minutes, washed 3 times in PBS with 0.05% Tween, and probed with goat anti-mouse IgM HRP-conjugate (1:5,000 dilution; Santa Cruz Biotechnology Inc., sc-2064) and developed using the ECL Western Blotting Substrate (Thermo Fisher).

ABO determinations were then matched to diarrheal outcomes in which the severity of illness was dichotomized into none (no or mild disease) and MSD using established criteria (71).

**Statistics.** We used log-rank (Mantel-Cox) test to compare time-to-treatment outcomes, with p < 0.05 considered significant. Meta-analysis (pooled fixed effects) was used to compare the relative risk of moderate to severe diarrheal illness. Pearson χ² testing was used to compare diarrheal illness and severity by blood group. Mann-Whitney U testing was used to compare nonparametric data. More than 2 groups of data were analyzed using Kruskal-Wallis testing for multiple comparisons of nonparametric data.

**Study approval.** Institutional Review Board approval was obtained at each participating study site (Washington University, the Naval Medical Research Center, and Johns Hopkins University) prior to sample interrogation and analysis. Written, informed consent was received from all participants prior to inclusion in the studies.

**Author contributions**

JMF, MAC, and MD were involved in study design, data curation, manuscript preparation, and project administration. TJV, AP, and PK conducted cloning of MetRS and ANL-labeling experiments. SC, ALB, DAS, BD, CDH, RLG, CKP, MGP, and MSR designed and conducted ETEC controlled human infection model studies in volunteers and processed the data and samples. WSW and JCG conducted glycan-array experiments. PK FMK, JFA, and BT were involved in protein interaction, hemagglutination, microscopy, adhesion, and toxin delivery studies. AS and MK conducted flow cytometry experiments. MAC, SS, JFA, MD, MK, and BT acquired, maintained, and propagated small intestinal stem cell banks or
intestinal epithelial cells. PK and TJV were responsible for recombinant protein production.

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