Molecular Basis of Age-dependent Gastric Inactivation of Rhesus Rotavirus in the Mouse

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Abstract
Rotavirus requires specific proteolytic activation by trypsin for efficient replication in tissue culture. To observe the nature of intestinal proteolytic activation of rotavirus in vivo, metabolically labeled rhesus rotavirus (RRV) grown in the presence of trypsin inhibitors was administered to adult and 10-d-old suckling mice by gavage. In the adult stomach, vp4 was cleaved in a manner distinct from in vitro trypsin cleavage. In the suckling stomach, RRV vp4 remains largely uncleaved. The alternative cleavage in the adult stomach was associated with a profound decrease in viral infectivity. vp4 from RRV recovered from the suckling small intestinal lumen was cleaved in a pattern similar or identical to in vitro trypsin-activated virus with bands comigrating with vp5* and vp8*. In contrast, vp4 was not observed in any recognizable form in RRV recovered from adult intestines. Comparison of infectivity of virus recovered from suckling and adult intestines revealed a 10,000-fold decrease in titer in the virus recovered from the adult intestine. In vitro digestions of RRV revealed that pepsin digestion can cleave RRV vp4 and markedly enhance acid-induced loss of rotavirus infectivity. Subsequent digestion with chymotrypsin removes most of the pepsin cleavage products of vp4. Virus injected directly into jejunal loops of adult mice and virus administered orally to adult mice pretreated with antacid drugs retained infectivity. These studies indicate the development of gastric acid and peptic secretion may be an important host defense factor in rotavirus gastroenteritis. (J. Clin. Invest. 1992. 89:1741–1745.)

Key words: viral enteritis • protease • pathogenesis • host restriction

Introduction
Rotaviruses are an important cause of severe infant gastroenteritis in humans as well as in a variety of other mammals. Symptomatic rotavirus infection usually occurs in young animals; and in some species, such as the mouse, there appears to be an absolute age restriction. In natural infection the virus is transmitted by the fecal–oral route. Thus virions must pass through the stomach en route to their target cells on the villi of the small intestine. During this gastrointestinal luminal phase, infecting virus particles are exposed to a variety of harsh conditions, including various proteases, bile salts, and gastric acid. As with other viruses that replicate in the gastrointestinal tract, rotavirus has evolved not only to survive these harsh conditions but also to depend on proteolytic activation with trypsin for efficient growth.

Rotaviruses are members of the Reoviridae and consist of 11 double-stranded RNA segments within a bilayered protein capsid. The outer capsid is comprised of two major proteins, vp7 and vp4. vp4 is cleaved by trypsin in vitro, and presumably in vivo, into smaller peptides vp5* and vp8*, which remain associated with the virion (1). This proteolytic cleavage is critical for efficient viral replication both in tissue culture (2, 3) and in vivo (4). Although it has long been presumed that intestinal luminal trypsin activates rotavirus in vivo, this has not been directly demonstrated. In this report, vp4 of rhesus rotavirus (RRV) is shown to be cleaved to products closely resembling vp5* and vp8* in the infant mouse intestine. In contrast, in the adult mouse, vp4 is first cleaved in the stomach, probably by pepsin, with a resultant striking loss of infectivity. In the adult intestine, further proteolysis, perhaps by chymotrypsin, results in a loss of most of the peptic cleavage product.

Methods

Animals. Rotavirus-free 8–10-d-old suckling and 6–8-wk-old adult CD1 mice were obtained from Charles River Breeding Laboratories (Portage, MI).

Virus. RRV has been previously described (5). 35S-labeled virus was grown on MA104 cells and purified by fluorocarbon extraction and CsCl gradient centrifugation as previously described (6), with the modification of adding 0.5 ml 1% aprotinin (Sigma Chemical Co., St. Louis, MO) to each 150-cm² flask when the 35S methionine was added. This allowed us to prepare virus that was largely uncleaved by trypsin.

Oral in vivo prophylaxis experiments. Animals were administered 10–100,000 cpn (1–5 × 10⁶ peroxidase focus units [pfu]) of purified 35S-labeled RRV in 50 μl of TNC buffer (10 mM Tris HCl, 100 mM NaCl, 0.1 mM CaCl₂) by gavage with stainless steel feeding needles and killed by cervical dislocation 30–60 min later. In some experiments adult mice were pretreated with 5 mg cimetidine orally 1–2 h before inoculation with labeled RRV. The stomach and small intestine were removed separately and flushed with 2 ml ice-cold TNC buffer. The washes were clarified by centrifugation in a microfuge at 10,000 g for 5 min at 4°C, and aliquots were taken for scintillation counting and titration on MA104 cells by peroxidase focus counting as previously described (7). Virus (recovered or inoculum) was not trypsin activated before the titrations. The remainder of the washes were pelleted at 100,000 g for 1 h and analyzed by SDS–PAGE (12% acrylamide) and fluorography. In most experiments, ~ 30% (range 15–36%) of the administered cpn were recovered in the clarified washes after the low speed centrifugation. Another 20–25% of administered cpn were recovered with eluted intestinal cells, with most of the remaining administered cpn appearing in the low speed pellet. Intestinal epithelial cells were eluted by incubating the washed intestines everted in Ca/Mg-free

1. Abbreviations used in this paper: pfu, peroxidase focus units; RRV, rhesus rotavirus; TNC, buffer consisting of 10 mM Tris HCl, 100 mM NaCl and 0.1 mM CaCl₂.
Hanks' buffered salt solution containing 2 mM EDTA for 30 min. The eluted enterocytes were washed four times in cold PBS and aliquots were boiled in Laemmli sample buffer before SDS–PAGE and fluorography. Intestinal and gastric pH was estimated with pH paper on undiluted luminal secretions. The accuracy of the pH paper was verified by comparing results obtained with the paper to those obtained on a pH meter (pH; Beckman Instruments, Inc., Fullerton, CA) on prepared buffers. The pH paper yielded results within 0.2 pH of those obtained with the meter.

**Jejunal loop experiments.** Mice were anesthetized and 2–3-cm jejunal loops were formed by suture ligation, with care to preserve the mesenteric blood supply. The loops were inoculated with 20 μl radiolabeled RRV (20,000–40,000 cpn, 10⁶ pfu) in TNC via a 30-gauge needle. After 1 h, mice were killed while still anesthetized, and the virus was recovered by flushing and centrifugation as described above.

**In vitro proteolysis.** All proteases were from Sigma Chemical Co. Metabolically labeled virus was suspended in either TNC buffer (pH 7.4) or 1× standard saline citrate buffer at the pH indicated. Enzymes were added at a final concentration of 10 μg/ml. For sequential digestions, the pH was adjusted to 7.0 after the initial pepsin digestion by addition of 1 M NaOH before the addition of the second enzyme. After incubation at 37°C for the indicated times, aliquots for titration of infectivity were removed, and the reactions were stopped by addition of Laemmli sample buffer and immediate boiling for 3 min.

**Results**

**Intraluminal digestion of RRV.** To examine the gastrointestinal luminal processing of rotavirus, purified radiolabeled RRV was administered to adult and suckling mice by gastric gavage. 1 h later luminal virus was recovered by flushing the intestines and stomachs with ice-cold TNC buffer. The washes were concentrated by centrifugation and analyzed by SDS–PAGE and fluorography. Fig. 1 depicts the results of a typical experiment. In virus recovered from the stomach of a suckling mouse (Fig. 1, lane SS), vp4 was present largely in its uncleaved 86.5-kD form. In virus recovered from the adult stomach (Fig. 1, lane AS) vp4 was apparently cleaved with a resulting 60-kD band and a faint ~ 20-kD band. Lane SI depicts virus recovered from the intestine of a suckling mouse in which vp4 is cleaved to 60- and 28-kD products similar or identical to vp5* and vp8* in in vitro trypsin-digested virus (lane T). In contrast, virus recovered from adult intestine (lane AI) was largely devoid of vp4 or its cleavage products.

To ensure that the protein profile of luminal virus reflected that of virus that had attached to its target cells, enterocytes were isolated from the small intestines of mice inoculated with radiolabeled RRV, washed, and analyzed by SDS–PAGE. Again, virus associated with suckling mouse enterocytes (Fig. 2, lane SI) demonstrated a profile identical to in vitro trypsin-activated virus with cleavage of vp4 to vp5* and vp8* (lane T), whereas virus associated with adult mouse enterocytes had a profile lacking identifiable vp4, vp5*, and vp8* (Fig. 2, lane AI). A single band of ~ 20 kD is seen in lane AI of Fig. 2 and lane AS in Fig. 1 which may be a fragment of vp4.

**Infectivity of recovered virus.** To determine whether the altered cleavage of vp4 in the adult mouse gut was associated with changes in infectivity, virus recovered from gastric and intestinal washes was titered on MA104 cells in 96-well plates by a peroxidase focus assay as previously described (7). The results, normalized for cpm recovered (Fig. 3), demonstrated a dramatic fall in titer (~ 10,000-fold) in virus recovered from adult mouse stomach and intestine relative to both inoculum and virus recovered from suckling animals.

**Gastric secretions are responsible for the loss of RRV infectivity in the adult mouse.** In preliminary experiments, we observed that the gastric pH in adult and suckling mice differed significantly (pH 2.9 vs. 3.5, respectively). We hypothesized that developmentally determined changes in gastric secretions might account for the dramatic fall in rotavirus titer and the change in vp4 cleavage observed in the adult mouse gastrointestinal lumen. To test this hypothesis, labeled virus was injected into isolated jejunal loops of adult mice, bypassing the gastric lumen. The recovered virus had a normal trypsin-activated protein profile by SDS–PAGE and fluorography (data not shown) and was highly infectious in the peroxidase focus assay (Fig. 4). In fact, the recovered virus was approximately fivefold more infectious than the inoculum, consistent with trypsin-mediated activation of the virus.

![Figure 1. Comparison of 35S-labeled RRV recovered from washes of stomachs or small intestines of adult or suckling mice 1 h after intragastric inoculation. Lane C, inoculum RRV; T, RRV treated in vitro with 10 μg/ml trypsin at 37°C for 1 h; Ch, RRV treated in vitro with 10 μg/ml chymotrypsin at 37°C for 1 h; AS, adult stomach; SS, suckling stomach; AI, adult intestine; SI, suckling intestine. Structural viral proteins are identified. The photograph is of a montage of two exposures of the same autoradiograph to optimize visualization of the structural proteins.](image1)

![Figure 2. Enterocyte-associated radiolabeled RRV recovered from the small intestines of suckling (S) or adult mice (A) 1 h after inoculation. Lane C, inoculum RRV; T, RRV treated in vitro with 10 μg/ml trypsin at 37°C for 1 h; Ch, RRV treated in vitro with 10 μg/ml chymotrypsin at 37°C for 1 h. Enterocytes were eluted with 2 mM EDTA from everted intestines and washed four times in PBS before SDS–PAGE and fluorography.](image2)
Data presented by Eisolated (lane PC) resulted from the Trypsin treatment of vp4 and the pH extensive destruction of virus proteases were secreted from the small intestine. Figure 1000000 from the stomach or intestines of adult or suckling mice 1 h postinoculation with purified radiolabeled RRV (10^6 pfu, 100,000 cpm). Virus in intestinal and gastric washes was titrated by peroxidase focus method on MA104 cells and normalized for cpm recovered. Data presented are means of three animals in each group.

Further evidence for a major role for gastric secretions in the loss of rotavirus infectivity within the adult mouse gastrointestinal tract was obtained by pretreating adult mice with the H2 receptor antagonist, cimetidine. This treatment resulted in an increase in gastric pH from ~3.0–3.7 as determined by pH paper (data not shown). Fig. 5 shows that, in adult mice pretreated with cimetidine (lane T), most of the virus recovered from the small intestine had 60- and 28-kD bands, consistent with the products of trypsin-cleaved vp4, vp5*, and vp6* (poorly seen). Virus recovered from control animals (lane C) had no recognizable vp4 or vp4 cleavage products. Results of peroxidase focus assays shown in Fig. 6 confirmed that most of the loss of viral titers could be prevented by decreasing gastric acid secretion with this drug. Similar results were obtained with the proton pump antagonist, omeprazole (data not shown).

In vitro proteolysis. In vitro digestions with several enteric proteases were performed in attempts to duplicate the effects on virus structure and infectivity observed in vivo. Initially, pepsin at various pH values was used to examine its effects on the structural viral proteins. Fig. 7 shows the results of such an experiment. At pH values <2.5, pepsin digestion resulted in extensive destruction of virtually all of the structural proteins. As the pH approached 3.0, pepsin digestion resulted in loss of the vp4 band and the appearance of an ~60-kD band.

In other experiments, digestions with selected proteases either individually or sequentially were performed (Fig. 8). In this experiment, trypsin, pepsin, and chymotrypsin (lanes T, P, and CH respectively) digestions were all found to greatly reduce the amount of vp4 in the virions with the simultaneous generation of a new band of ~60 kD. In the chymotrypsin digestion there was also the appearance of an ~52-kD band. Trypsin treatment of pepsin-digested virus (lane PT) resulted in the appearance of yet another band of 80 kD, possibly a vp2 cleavage product. Pepsin followed by chymotrypsin treatments (lane PC) resulted in a substantial reduction of the 60-kD pepsin cleavage product, whereas pepsin–trypsin treatment (lane PT) did not.

The infectivity of RRV after in vitro treatment with trypsin, chymotrypsin, pepsin, and pH 3.5 buffer for 1 h is shown in Fig. 9. As noted previously, trypsin treatment of rotavirus resulted in enhanced infectivity, whereas chymotrypsin treatment had little effect on viral titers. Low pH treatment reduced titers ~100-fold from baselines, whereas combined low pH and pepsin treatment resulted in almost a complete loss of infectivity. In other experiments (data not shown) trypsin-activated RRV infectivity was likewise dramatically reduced by acid/pepsin treatment.

Discussion

Prior studies in both tissue culture systems and in mice have indirectly suggested the importance of intestinal luminal protease activity in the in vivo replication of rotavirus. In this study we sought to examine directly the luminal proteolytic processing of rotavirus in vivo and to compare such processing in adult and suckling animals. We chose to use the simian rotavirus RRV for these studies because it grows efficiently, can be metabolically labeled to high specific activity, and is able to infect suckling mice and cause diarrheal disease. In our experiments we observed striking differences between suckling and adult mouse enteric luminal processing of rotavirus as well as striking differences in the viability of virus recovered from the gut. These differences appear to be largely due to differences in the adult and suckling mouse gastric secretion.

Figure 3. Infectivity of RRV recovered from the stomach or intestines of adult or suckling mice 1 h postinoculation with purified radiolabeled RRV (10^6 pfu, 100,000 cpm). Virus in intestinal and gastric washes was titrated by peroxidase focus method on MA104 cells and normalized for cpm recovered. Data presented are means of three animals in each group.

Figure 5. Effects of cimetidine pre-treatment on radiolabeled RRV recovered from the small intestine of adult mice. Mice were given 5 mg of cimetidine (T) or saline (C) 1 h before intragastric inoculation with radiolabeled RRV.

Figure 4. Infectivity of virus recovered from isolated jejunal loops of adult mice. Closed jejunal loops in anesthetized mice were injected with purified radiolabeled RRV (40,000 cpm, 10^6 pfu) and recovered by washing after 1 h. Infectivity was determined by peroxidase focus assay and normalized for cpm recovered. Data presented are means of two experiments.

Figure 6. Infectivity by peroxidase focus titration of RRV recovered from the small intestines of cimetidine- or saline (control)-treated adult mice 1 h after inoculation with radiolabeled RRV (5 × 10^6 pfu, 10,000 cpm). Data represent the means of three animals per group. a, Stomach; b, intestine.
Figure 7. Digestion of radiolabeled RRV with 10 μg/ml pepsin at indicated pH values. +, pepsin digestion; −, pH buffer control.

In our in vivo experiments, we observed a striking loss of infectivity as RRV traversed the adult mouse stomach. This was associated with a decrease in vp4 content and the appearance of bands of ~ 60 and 20 kD (Fig. 1, lane AS). Although the 60-kD band comigrates vp5* (Fig. 1, lane T), it is unlikely to be identical to the product of trypsin digestion of vp4 for several reasons. First, AS recovered or pepsin-treated virus is not infectious in contrast to trypsin-treated virus (Figs. 3 and 9). Second, the 60-kD band in pepsin-treated virus is susceptible to chymotrypsin digestion (Fig. 8, lane PC), whereas vp5 is not (Fig. 8, lane TC). Finally, acid proteases such as pepsin cleave peptides at sites distinct from those utilized by trypsin.

In the intestinal lumen the 60-kD band disappeared. We speculate that the initial cleavage in the adult stomach may be due to pepsin and/or other acid-resistant proteases, as demonstrated in in vitro experiments (Fig. 8, lane P). The subsequent loss of the 60-kD band in the intestine is probably a result of intestinal luminal proteases such as chymotrypsin, as seen in our in vitro studies (Fig. 8, lane PC). Such a loss of a 60-kD fragment of vp4 from SA11 has been previously observed with high dose chymotrypsin treatment (1). The cleavage of vp4 by pepsin in vitro and in vivo correlates with the observed loss of RRV infectivity. It is tempting to speculate that changes in vp4, which contains neutralizing epitopes, mediates hemagglutination, and probably mediates cell attachment and penetration, could lead to a loss of infectivity. The lack of viral inactivation in adult intestinal loops (Fig. 4) or in cimetidine-treated animals (Fig. 6) argues against inactivation of RRV in the adult intestinal lumen.

Several previous studies have examined the interaction of rotavirus with various proteases. Early work demonstrated enhancement of rotavirus growth in the presence of trypsin and demonstrated that the effect was on the viral particle rather than on host cells (8). Trypsin activation of rotavirus is associated with the cleavage of the outer capsid protein vp4 into vp5* and vp8* (1). We observed an identical cleavage of intact vp4 in the murine small intestine (Figs. 1 and 5). Chymotrypsin, another intestinal luminal protease, fails to activate rotavirus and cleaves vp4 into a fragment of similar electrophoretic mobility to vp5* and several fragments of lower molecular weight than vp8* (1, 9, 10).

The mechanism by which trypsin cleavage of vp4 mediates enhanced infectivity is not definitely known, although some studies have proposed that it facilitates viral penetration of target cell membranes (7, 11). Experiments using MAbs specific for the vp8* product of vp4 have suggested that vp8* may be involved in cell attachment (12).

Although it has long been hypothesized that the effects of trypsin on rotavirus growth in tissue culture parallel events in the intestinal lumen, little direct evidence has been produced to date. Protease inhibitor studies indirectly demonstrated the importance of intestinal luminal proteolysis for rotavirus replication in vivo (4). By methods previously used to study the intestinal luminal activation of reovirus (13, 14), we have directly demonstrated the cleavage of rotavirus vp4 into vp5* and vp8* in the suckling mouse intestine. Similar cleavage and activation of infectivity were observed in the adult mouse intestine only if gastric secretions were physically bypassed or pharmacologically suppressed.

Several previous workers have examined the inactivation of rotaviruses by acid and gastric secretions (15–17). Although methodologies differ, most have shown rotaviruses to be relatively stable at pH > 4 and with inactivation half-lives of 10–100 min at pH 3. We observed a comparable drop in infectivity of RRV in vitro at pH 3.5, which was greatly enhanced in the presence of pepsin (Fig. 9). Pepsin has previously been shown...
to inactivate serotype 1 reovirus in association with the removal of $\alpha 3$ (13). At pH values of < 3, we found that pepsin digestion resulted in proteolysis of the entire particle (Fig. 7). Indeed, we have found that, at pH values < 2.5, RRV rapidly loses infectivity and hemagglutination titers whether or not pepsin is present (data not shown). These data are consistent with previous morphological studies in which collapse of the rotavirus outer capsid was observed at low pH values (18). The nonfasting gastric pH in our sucking mice was ~ 3.5, whereas in the adult mouse stomach it was ~ 3.0. These values may not reflect the microenvironment of rotavirus particles in the stomach, where foodstuffs may locally buffer the gastric pH.

The differences noted between the effects of gastric secretions from adult and sucking mice on RRV infectivity are consistent with previously observed differences in acid and pepsin secretion in the developing rodent (19). Rodents generally have very low acid and pepsin secretion until the time of weaning. It has been proposed that this reduced secretion facilitates the enteric absorption of functional maternal antibodies from breast milk.

It seems doubtful that the enhanced gastric acid/pepsin secretion noted in the adult mice is the sole determinant of the strict host age restriction for symptomatic rotavirus disease in the murine model. If rotavirus replication were highly efficient in the adult mouse small intestine, one would expect disease to occur even if only a few infectious virions succeeded in surviving the gastric pH barrier. We were able to detect small amounts of infectious virus in the small intestines of untreated adult mice (Fig. 3), in preliminary studies using cimetidine-treated, seronegative adult mice inoculated with $10^7$ pfu of RRV, we have observed more frequent excretion of rotavirus measured by ELISA of fecal suspensions in treated (4/6) versus control mice (1/4) but no evidence of diarrhea in any mouse. Thus, other factors affecting the efficiency of rotavirus replication in the intestine, such as a relative paucity of viral receptors on adult enterocytes (20, 21), or affecting the expression of overt disease, such as increased intestinal absorptive reserve in adult mice, must also be important in host age restriction of rotavirus diarrhea.

Although humans do increase their capacity to secrete both acid and pepsin as they mature, the most significant increases occur during the first three months of life (22), before the peak age for rotavirus disease. The reduced gastric acid observed in human neonates might account for the persistence of low virulence strains of rotavirus in some nurseries (23).

It seems likely that gastric inactivation of rotavirus virions is an important host defense mechanism in natural infections. A study of the minimal infective dose of rotavirus strain OSU in piglets detected gastric secretion factor(s) that reduced the infectivity of the inoculum and were both pH and concentration dependent (15). Human vaccine studies using RRV and RIT 4237 rotavirus strains demonstrated that a higher proportion of infants responded to the vaccine if they were pretreated with bicarbonate buffer (24, 25). Greatly reduced gastric acid and pepsin secretion have been observed in malnourished and even well-nourished children in developing countries (26, 27). This reduced gastric acid barrier may well predispose these children to rotavirus as well as other enteric infections.

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References


